IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

REQUEST FOR FILING CONTINUATION / DIVISIONAL APPLICATION UNDER 37 C.F.R. § 1.53(b)

Commissioner for Patents P.O. Box 1450 Alexandria, VA 22313-1450

Customer Number 21839

Sir:

Π NON-PUBLICATION IS REQUESTED. A CERTIFICATION IS BELOW.

This is a request for filing a 🛛 continuation 🗍 divisional application under 37 C.F.R. § 1.53(b) of the following pending application, Application No. 10/551,205, filed November 14, 2006 entitled ORAL FORMULATIONS OF CLADRIBINE

by the following named inventor(s):

Nicholas S. BODOR Yogesh DANDIKER

- The entire disclosure of the prior application from which a copy of the oath or declaration is supplied herewith is considered as being part of the disclosure of the accompanying application and is hereby incorporated by reference therein.
- Applicant(s) hereby requests that the above-captioned application NOT BE PUBLISHED under 35 U.S.C. § 122(b) and 37 C.F.R. § 1.211. The undersigned hereby certifies that the invention disclosed in this application has not and will not be the subject of an application filed in another country, or under a multilateral international agreement, that requires publication at eighteen (18) months after filing.
- This application is being filed by less than all the inventors named in the prior application. In \square accordance with 37 C.F.R. § 1.63(d)(2), the Commissioner is requested to delete the name(s) of the following person or persons who are not inventors of the invention being claimed in this application.
- This application is being filed by more than all the inventors named in the prior application. In accordance with 37 C.F.R. § 1.63(d)(5), a new oath or declaration is enclosed, and the Commissioner is requested to add the name(s) of the following person or persons who are inventors of the invention being claimed in this application.
- \boxtimes Applicant(s) suggests Figure 1 for inclusion on the front page of the patent application publication and patent.
- \boxtimes Applicant(s) requests that the published application include the following assignment information: Ares Trading, S.A., Aubonne, Switzerland.
- \square Small entity status is claimed.
- Enclosed is a copy of the prior Application No. 1. \square as originally filed on , including copies of the specification, claims, drawings and the executed oath or declaration as filed.
- \boxtimes Enclosed is a revised prior application and a copy of the prior executed oath or declaration as 2. filed. No new matter has been added to the revised application.

Buchanan Ingersoll & Rooney PC

Attorneys & Government Relations Professionals



Hopewell EX1058 Hopewell v. Merck IPR2023-00480

3. \boxtimes The filing fee has been calculated as follows
and in accordance with the enclosed **Preliminary Amendment:**

					F	EES
Basic Application Filing Fo	ee (1011)				\$	330
Examination Fee (1311)						220
Search Fee (1111)						540
	No. of Claims		Extra Claims	Rate		
Total Claims	64	Minus 20=	44	x \$ 52 (1202)		2288
Independent Claims	4	Minus 3=	1	x \$ 220 (1201)		220
If multiple dependent claims are presented, add 390				\$	0	
App. Size Fee (app + dwgs. exceeding 100 sheets) - \$270 for each 50 sheets (1081)			\$	0		
Total Application Fee			\$	3598		
Small Entity Status claimed - subtract 50% of Total Application Fee				0		
Add Assignment Recording Fee of \$ 40 if Assignment document is enclosed			\$	0		
TOTAL APPLICATION FEE DUE			\$	3598		

- 4. This application is being filed without a filing fee.
- 5. to Deposit Account No. 02-4800 for the fee due. Charge _____
- 6, \boxtimes Charge <u>\$ 3598</u> to credit card.
- 7. \boxtimes The Director is hereby authorized to charge any appropriate fees under 37 C.F.R. §§ 1.16, 1.17 and 1.21 that may be required by this paper, and to credit any overpayment, to Deposit Account No. 02-4800.
- \boxtimes 8. New drawings are enclosed.
- Applicant(s) claims priority under 35 U.S.C. §§ 119 of the following application(s): 9.

Country		Appl. No.	Filing Date MM-DD-YYYY
			MM-DD-YYYY
	The certified copy of the priority app	blication:	

The certified copy of the priority application:

is enclosed. Continuation/Divisional Patent Application Transmittal Letter Application No. <u>unassigned</u> Attorney Docket No. <u>0056192-000067</u>

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			was filed on, filed on, filed on, filed on, and acknowledged by the Examiner on, in Paper No		
			was filed in the International Bureau and acknowledged by the Examiner on		
			has not yet been filed.		
10.		A Prelir	ninary Amendment is enclosed.		
11.	\boxtimes	An Info enclose	rmation Disclosure Statement, Form Substitute PTO-1449 and (2) documents are ed.		
12.		A Gene enclose	A General Authorization of Payment of Fees and Petitions for Extensions of Time is enclosed.		
13.	\boxtimes	An Application Data Sheet.			
14.		Also enclosed is			
15.	\boxtimes	The power of attorney in the prior application is to Customer Number 21839.			
	a.	—	The power appears in the original papers in the prior application.		
	b.		Since the power does not appear in the original papers, a copy of the power in the prior application is enclosed.		
	C.	\boxtimes	Address all future communications to: (may only be completed by applicant, attorney, or agent of record)		
			Buchanan Ingersoll & Rooney pc Customer Number 21839		
Date:	<u>January</u>	<u>7, 2011</u>	By Mary Kitherine Downeuter Mary Katherine Baumeister Registration No. 26254		

Customer No. 21839 703 836 6620

inventor(s)
assignee of complete interest
attorney or agent of record
filed under 37 C.F.R.§ 1.34(a)

APPLICATION DATA SHEET

January 7, 2011

Application Information

Application Number::

Filing Date::

Application Type::

Subject Matter::

Utility

Regular

Suggested Classification::

Suggested Group Art Unit::

CD-ROM or CD-R?::

Number of CD Disks::

Number of Copies of CDs::

Sequence Submission?::

Computer Readable Form (CRF)?::

Number of Copies of CRF::

Title::

ORAL FORMULATIONS OF CLADRIBINE Attorney Docket Number:: 0056192-000067 Request for Early Publication?:: No Request for Non-Publication?:: No Suggested Drawing Figure:: 1 1 Total Drawing Sheets:: Small Entity?:: No

Latin Name::

Variety Denomination Name::

Petition Included?::NoPetition Type::Licensed US Govt. Agency::Contract or Grant Numbers::Secrecy Order in Parent Appl.?::No

Applicant Information

Applicant Authority Type::	Inventor
Primary Citizenship Country::	United States
Status::	Full Capacity
Given Name::	Nicholas
Middle Name::	S.
Family Name::	BODOR
Name Suffix::	
City of Residence::	Bal Harbour
State or Province of Residence::	Florida
Country of Residence::	United States
Street of Mailing Address::	10225 Collins Avenue Unit 1002/1004
City of Mailing Address::	Bal Harbour
State or Province of Mailing Address::	Florida

Country of Mailing Address::	United States
Postal or Zip Code of Mailing Address::	33154
Applicant Authority Type::	Inventor
Primary Citizenship Country::	United Kingdom
Status::	Full Capacity
Given Name::	Yogesh
Middle Name::	
Family Name::	DANDIKER
Name Suffix::	
City of Residence::	Toronto
State or Province of Residence::	
Country of Residence::	Canada
Street of Mailing Address::	57 Fenn Avenue
City of Mailing Address::	Toronto
State or Province of Mailing Address::	
Country of Mailing Address::	Canada
Postal or Zip Code of Mailing Address::	M2L 1M9

Correspondence Information

Correspondence Customer Number:: 21839		
Phone Number::	(703) 836-6620	
Fax Number:	(703) 836-2021	

Representative Information

Representative Customer Number:: 21839

Domestic Priority Information

Application::	Continuity Type::	Parent Application::	Parent Filing Date::
This Application	Continuation of	10/551,205	November 14, 2006
10/551,205	National Stage of	PCT/US2004/009387	March 26, 2004
PCT/US2004/009387	Claims benefit under 35 U.S.C. §119(e) of	60/458,922	March 28, 2003
PCT/US2004/009387	Claims benefit under 35 U.S.C. §119(e) of	60/484,756	July 2, 2003
PCT/US2004/009387	Claims benefit under 35 U.S.C. §119(e) of	60/541,247	Feb 4, 2004

Foreign Priority Information

Country::

Application Number::

Filing Date::

Claimed::

Priority

Assignee Information

Assignee Name::

Street of Mailing Address::

City of Mailing Address::

State or Province of Mailing Address::

Country of Mailing Address::

Postal or Zip Code of Mailing Address::

Switzerland CH-1170

Aubonne

Respectfully submitted,

BUCHANAN INGERSOLL & ROONEY PC

ARES TRADING S.A.

Zone Industrielle d l'Ouriettaz

Date: January 7, 2011

inter By:

Mary Katherine Baumeister Registration No. 26254

Customer No. 21839 703 836 6620

COMBINED DECLARATION AND POWER OF ATTORNEY FOR UTILITY OR DESIGN PATENT APPLICATION

As a below named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name:

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled:

ORAL FORMULATIONS OF CLADRIBINE

the specification of which (check only one item below):

- is attached hereto.

was filed as United States Patent application Number _ and was amended on on (if applicable).

was filed as PCT International application Number March 26, PCT/US2004/009387 _____ on and was amended on 2004 ____ (if applicable).

I hereby state that I have reviewed and understand the contents of the above-identified specification, including the claims, as amended by any amendment referred to above.

I acknowledge the duty to disclose to the Office all information known to me to be material to patentability as defined in Title 37, Code of Federal Regulations, § 1.56.

I hereby claim foreign priority benefits under Title 35, United States Code, §§ 119 (a)-(d), 172 or 365(a) of any foreign application(s) for patent or inventor's certificate or of any international (PCT) application(s) designating at least one country other than the United States of America listed below and have also identified below any foreign application(s) for patent or inventor's certificate or any PCT international (PCT) application(s) designating at least one country other than the United States of America filed by me on the same subject matter having a filing date before that of the application(s) of which priority is claimed:

PRIOR FOREIGN/PCT APPLICATION(S) AND ANY PRIORITY CLAIMS UNDER 35 U.S.C. §§119(a)-(d), 172 or 365(a):				
COUNTRY (if PCT, indicate "PCT")	APPLICATION NUMBER	DATE OF FILING (MM/DD/YYYY)	PRIORITY UNDER 3 §§119, 172 Yes	CLAIMED 35 U.S.C. OR 365(a) No

I hereby appoint the attorneys and agents associated with the following PTO Customer Number of Buchanan Ingersoll PC (including attorneys from Burns, Doane, Swecker & Mathis) to prosecute said application and to transact all business in the Patent and Trademark Office connected therewith and to file, prosecute and transact all business in connection with international applications directed to said invention:

· Customer Number 21839

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

FULL NAME OF SOLE OR FIRST INVENTOR	Nicholas S. Bodor
Signature	hilder S. Morr
Date	11-14-05
Residence (City, State, Country)	Bal Harbour, Florida, US
Citizenship	United States
Mailing Address	10225 Collins Avenue, Unit 1002/1004
City, State, ZIP, Country	Bal Harbour, Florida 33154, US
FULL NAME SECOND INVENTOR, IF ANY	Yogesh Dandiker
Signature	theme
Date	1/26/08.
Residence (City, State, Country)	TORONITO, CAMADA
Citizenship	United Kingdom
Mailing Address	57 FENN AVENUE
City, State, ZIP, Country	TOROLITO CAMADA, M2L 1Mg
FULL NAME OF THIRD INVENTOR, IF ANY	
Signature	
Date	
Residence (City, State, Country)	
Citizenship	
Mailing Address	
City, State, ZIP, Country	

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ORAL FORMULATIONS OF CLADRIBINE

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CROSS-REFERENCE TO EARLIER APPLICATIONS

This application is a continuation of prior copending US Application No. 10/551,205 filed November 14, 2006, now allowed, which is the US national stage of International Application No. PCT/US2004/009387, filed March 26, 2004, which claims benefit under 35 U.S.C. § 119(e) of United States Provisional Application No. 60/458,922, filed March 28, 2003; of United States Provisional Application No. 60/484,756, filed July 2, 2003; and of United States Provisional Application No. 60/541,247, filed February 4, 2004, all of said applications being hereby incorporated by reference herein in their entireties and relied upon.

BACKGROUND OF THE INVENTION

Cladribine, which is an acid-labile drug, has the chemical structure as set forth below:



It is also known as 2-chloro-2'-deoxyadenosine or 2-CdA. Cladribine exists as a white, nonhydroscopic, crystalline powder, consisting of individual crystals and of crystalline aggregates.

Cladribine is an antimetabolite which has use in the treatment of lymphoproliferative disorders. It has been used to treat experimental leukemias such as L1210 and clinically for hairy cell leukemia and chronic lymphocytic leukemia as well as Waldenstrom's macroglobulinaemia. It has

also been used as an immunosuppressive agent and as a modality for the treatment of a variety of autoimmune conditions including rheumatoid arthritis, inflammatory bowel disease (e.g., Crohn's disease, ulcerative colitis) and multiple sclerosis (see e.g., J. Liliemark, Clin. Parmacokinet, 32(2): 120-131, 1997). It has also been investigated, either experimentally or clinically in, for example, lymphomas, Langerhan's cell histiocytosis, lupus erythematosus, chronic plaque psoriasis, Sezary syndrome, Bing-Neel syndrome, recurrent glioma, and solid tumors.

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Oral delivery of drugs is often preferred to parenteral delivery for a 10 variety of reasons, foremost patient compliance, or for cost or therapeutic considerations. Patient compliance is enhanced insofar as oral dosage forms alleviate repeated health care provider visits, or the discomfort of injections or prolonged infusion times associated with some active drugs. At a time of escalating health care costs, the reduced costs associated with oral 15 administration versus parenteral administration costs gain importance. The cost of parenteral administration is much higher due to the requirement that a health care professional administer the cladribine in the health care provider setting, which also includes all attendant costs associated with such administration. Furthermore, in certain instances, therapeutic considerations such as the need for a slow release of cladribine over a prolonged period of time may be practically met only by oral or transmucosal delivery.

However, to date the oral delivery of cladribine has been plaqued by low bioavailability (see, e.g., J. Liliemark et al., J. Clin. Oncol., 10(10): 1514-1518, 1992), and suboptimal interpatient variation (see, e.g., J. Liliemark, Clin. Pharmacokinet, 32 (2): 120-131, 1997). See also, A. Tarasuik, et al. reporting poor absorption and pH dependent lability (Arch. Immunol. et Therapiae Exper., 42: 13-15, 1994).

Cyclodextrins are cyclic oligosaccharides composed of cyclic α -(1 \rightarrow 4) linked D-glucopyranose units. Cyclodextrins with six to eight units have

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been named α -, β - and γ -cyclodextrin, respectively. The number of units determines the size of the cone-shaped cavity which characterizes cyclodextrins and into which drugs may be included to form stable complexes. A number of derivatives of α -, β - and γ -cyclodextrin are known in which one or more hydroxyl groups is/are replaced with ether groups or other radicals. These compounds are thus known complexing agents and have been previously used in the pharmaceutical field to form inclusion complexes with water-insoluble drugs and to thus solubilize them in aqueous media.

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Recently, Schultz et al., in U.S. Patent No. 6,194,395 B1, have described complexing and solubilizing cladribine with cyclodextrin. The Schultz et al. patent primarily addresses the problems inherent in previously described aqueous formulations of cladribine, particularly for subcutaneous and intramuscular injection. Schultz et al. have found that cladribine is not only significantly more soluble in aqueous media when formulated with cyclodextrin, but also is more stable against acid-catalyzed hydrolysis when combined with cyclodextrin. The latter finding is taught to be of particular benefit in the formulation of solid oral dosage forms, where the compound would normally undergo hydrolysis in the acid pH of the stomach contents. Schultz et al. do not appear to have described any actual work in connection with solid oral dosage forms. In fact, they describe only one method of preparing the solid dosage form, which is a melt extrusion process, in which the cladribine and cyclodextrin are mixed with other optional additives and then heated until melting occurs. Furthermore, the broad dosage ranges of 1 mg to 15 mg of cladribine and 100 mg to 500 mg of cyclodextrin listed in the patent suggest no criticality to the particular amount of cyclodextrin to be present with a given amount of cladribine in a solid oral dosage form. Indeed, these dosage ranges include many combinations which may be suitable as mixtures but not for complex formation. For example, a ratio of 1

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mg of cladribine to 500 mg of cyclodextrin contains too much cyclodextrin, so

that the drug would not readily leave the complex and achieve its therapeutic function. On the other hand, 15 mg of cladribine and only 100 mg of cyclodextrin would not be enough to complex that amount of cladribine.

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The Schultz *et al.* patent does suggest improving the stability of cladribine in oral dosage forms by combining/complexing it with cyclodextrin, but does not suggest improving the drug's oral bioavailability by such means; in fact, the patent does not describe or suggest a method for enhancing or maximizing the bioavailability of cladribine from a solid oral dosage form of cladribine and cyclodextrin, or a composition specially designed to do so.

Many workers have studied the solubility of specific drugs in water containing various concentrations of selected cyclodextrins in order to demonstrate that increasing concentrations of cyclodextrins increase the solubility of the drugs at selected temperatures and pH levels, as for example reported in the Schultz *et al.* patent. Phase solubility studies have also been performed by various workers in order to elucidate the nature of the complex formation, for example, whether the cyclodextrin and drug form a 1:1 complex or a 1:2 complex; see, for example, Harada *et al.* U.S. Patent No. 4,497,803, relating to inclusion complexes of lankacidin-group antibiotics with cyclodextrin, and Shinoda *et al.* U.S. Patent No. 4,478,995, relating to a complex of an acid addition salt of (2'-benzyloxycarbonyl)phenyl trans-4guanidinomethylcyclohexanecarboxylate with a cyclodextrin.

While Schultz *et al.* teach that a cladribine-cyclodextrin complex improves the water solubility and acid stability of cladribine, the art does not suggest how to maximize or enhance the benefits of the complexation in terms of bioavailability and interpatient variation when the complex is to be administered in a solid oral dosage form.

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SUMMARY OF THE INVENTION

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It has now been found that amorphous cyclodextrins can be combined with cladribine to form a particularly advantageous product which can be incorporated into a solid oral dosage form. This product is a complex cladribine-cyclodextrin complex, and the solid oral dosage form containing it improves oral bioavailability and/or achieves lower interpatient and/or intrapatient variation of the drug.

The present invention provides a complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous
inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex, and a pharmaceutical composition comprising said complex, formulated into a solid oral dosage form. Thus, the cyclodextrin itself is amorphous, the inclusion complex with cladribine is amorphous (and is preferably saturated with cladribine) and the free cladribine which forms the non-inclusion complex is amorphous.

The invention also provides a method for increasing or enhancing the oral bioavailability of cladribine comprising orally administering to a subject in need thereof, a pharmaceutical composition comprising a complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex, formulated into a solid oral dosage form which maximizes the amount of cladribine in the inclusion and non-inclusion complexes.

The invention further provides for treatment of conditions responsive to administration of cladribine in mammals by administering thereto the composition of the invention. Use of cladribine in the preparation of the pharmaceutical compositions of the invention for administration to treat

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cladribine-responsive conditions and for enhancing the oral bioavailability of cladribine is also provided.

Still further, the invention provides a process for the preparation of a complex cladribine-cyclodextrin complex which comprises the steps of:

(i) combining cladribine and an amorphous cyclodextrin in water at a temperature of from about 40 to about 80°C and maintaining said temperature for a period of from about 6 to about 24 hours;

(ii) cooling the resultant aqueous solution to room temperature; and

(iii) lyophilizing the cooled solution to afford an amorphous product.

In yet a further aspect the invention provides a pharmaceutical composition obtainable by a process comprising the steps of:

(i) combining cladribine and an amorphous cyclodextrin in water at a temperature of from about 40 to about 80°C and maintaining said temperature for a period of from about 6 to about 24 hours;

(ii) cooling the resultant aqueous solution to room temperature;

(iii) lyophilizing the cooled solution to afford an amorphous product;

and

(iv) formulating the amorphous product into a solid oral dosage form.

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BRIEF DESCRIPTION OF THE DRAWING

A more complete appreciation of the invention and its many attendant advantages will be readily understood by reference to the following detailed description and the accompanying drawing, wherein the sole Figure is a graphical representation of the results of a phase solubility study where various molar concentrations of hydroxypropyl- β -cyclodextrin (HP β CD) are plotted against various cladribine molar concentrations, with (•) representing

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the data points obtained for complexation under conditions specified in EXAMPLE 2 below.

DETAILED DESCRIPTION OF THE INVENTION

Throughout the instant specification and claims, the following definitions and general statements are applicable.

The patents, published applications, and scientific literature referred to herein establish the knowledge of those with skill in the art and are hereby incorporated by reference in their entirety to the same extent as if each was specifically and individually indicated to be incorporated by reference. Any conflict between any reference cited herein and the specific teachings of this specification shall be resolved in favor of the latter. Likewise, any conflict between an art-understood definition of a word or phrase and a definition of the word or phrase as specifically taught in this specification shall be resolved in favor of the latter.

The term "inclusion complex" as used herein refers to a complex of cladribine with the selected cyclodextrin wherein the hydrophobic portion of the cladribine molecule (the nitrogen-containing ring system) is inserted into the hydrophobic cavity of the cyclodextrin molecule. This is often referred to simply as a cyclodextrin complex of the drug.

The term "non-inclusion complex" refers to a complex which is not an inclusion complex; rather than the hydrophobic portion of cladribine being inserted in the cyclodextrin cavity, the non-inclusion complex is formed primarily by hydrogen-bonding of the hydroxyls and amino group on "free" cladribine, (*i.e.* cladribine not in the inclusion complex) to the hydroxyls on the exterior of the cyclodextrin torus (*e.g.* in the case of hydroxypropyl- β -cyclodextrin, hydroxypropyl and hydroxyl groups on the glucose rings). This is a more loosely-held association than an inclusion complex.

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As used herein, whether in a transitional phrase or in the body of a claim, the terms "comprise(s)" and "comprising" are to be interpreted as having an open-ended meaning. That is, the terms are to be interpreted synonymously with the phrases "having at least" or "including at least". When used in the context of a process, the term "comprising" means that the process includes at least the recited steps, but may include additional steps. When used in the context of a composition, the term "comprising" means that the composition includes at least the recited features or components, but may also include additional features or components.

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10 The terms "consists essentially of" or "consisting essentially of" have a partially closed meaning, that is, they do not permit inclusion of steps or features or components which would substantially change the essential characteristics of a process or composition; for example, steps or features or components which would significantly interfere with the desired properties of 15 the compositions described herein, *i.e.*, the process or composition is limited to the specified steps or materials and those which do not materially affect the basic and novel characteristics of the invention. The basic and novel features herein are the provision of a complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous

inclusion complex of cladribine with an amorphous cyclodextrin and (b) 20 amorphous free cladribine associated with amorphous cyclodextrin as a noninclusion complex, formulated into a solid oral dosage form, so as to provide improved bioavailability and/or lower interpatient and/or intrapatient variation following administration. Essential to the invention is the combination of the

amorphous nature of the starting cyclodextrin, and the level of water 25 solubility exhibited by cladribine (about 5 mg/ml at room temperature), and consequently its capability for hydrogen bonding, which can be taken advantage of under particular conditions described hereinafter, and which afford a special amorphous mixture uniquely well-suited for optimizing the 30 oral bioavailability of cladribine.

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The terms "consists of" and "consists" are closed terminology and allow only for the inclusion of the recited steps or features or components.

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As used herein, the singular forms "a," "an" and "the" specifically also encompass the plural forms of the terms to which they refer, unless the content clearly dictates otherwise.

The term "about" is used herein to means approximately, in the region of, roughly, or around. When the term "about" is used in conjunction with a numerical range, it modifies that range by extending the boundaries above and below the numerical values set forth. In general, the term "about" or "approximately" is used herein to modify a numerical value above and below the stated value by a variance of 20%.

The term "amorphous" is used herein to refer to a noncrystalline solid. The cyclodextrins encompassed herein themselves are amorphous because they are each composed of a multitude of individual isomers, and their complexes with cladribine are also amorphous. Further, conditions for complexation can be selected (elevated temperature and prolonged complexation times, as described hereinafter) so that a supersaturated cladribine solution will be formed. When cooled, because of the amorphous nature of the complex and the cyclodextrin, some excess free cladribine does not precipitate but rather is trapped in amorphous form in intimate admixture with the (preferably saturated) amorphous cladribine-cyclodextrin

inclusion complex. This excess cladribine forms a loosely-held association, or non-inclusion complex, with the cyclodextrin through hydrogen bonding. This, then, further increases the amount of cladribine in the product; this

additional cladribine, because it is amorphous and also because it is in intimate admixture with the amorphous inclusion complex, is expected to be somewhat protected from degradation by stomach acid (although it may not be as protected as the cladribine which is in the form of the inclusion complex).

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The term "saturated" when used in conjunction with a complex of cladribine in amorphous cyclodextrin means that the complex is saturated with cladribine, that is, the complex contains the maximum amount of cladribine which can be complexed (by means of both inclusion and noninclusion complexes) with a given amount of cyclodextrin under the conditions of complexation used. A phase solubility study can be used to provide this information, as described in more detail hereinafter. (Conditions for the complexation are also described in more detail below.) Alternatively, a saturated complex may be arrived at empirically by simply adding cladribine to an aqueous solution of the selected cyclodextrin until no more cladribine goes into solution; ultimately, excess cladribine, if any, is removed (by filtration or centrifugation) and the solution lyophilized to provide the dry saturated complex.

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The expression "substantially', as in "substantially free" means within 20% of the exact calculated amount, preferably within 10%, most preferably within 5%.

The term "interpatient variability" refers to variation among patients to which a drug is administered. The term "intrapatient variability" refers to variation experienced by a single patient when dosed at different times.

As used herein, the recitation of a numerical range for a variable is intended to convey that the invention may be practiced with the variable equal to any of the values within that range. Thus, for a variable which is inherently discrete, the variable can be equal to any integer value of the numerical range, including the end-points of the range. Similarly, for a

variable which is inherently continuous, the variable can be equal to any real value of the numerical range, including the end-points of the range. As an example, a variable which is described as having values between 0 and 2, can be 0, 1 or 2 for variables which are inherently discrete, and can be 0.0,

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0.1, 0.01, 0.001, or any other real value for variables which are inherently continuous.

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In the specification and claims, the singular forms include plural referents unless the context clearly dictates otherwise. As used herein, unless specifically indicated otherwise, the word "or" is used in the "inclusive" sense of "and/or" and not the "exclusive" sense of "either/or."

Technical and scientific terms used herein have the meaning commonly understood by one of skill in the art to which the present invention pertains, unless otherwise defined. Reference is made herein to various methodologies and materials known to those of skill in the art. Standard reference works setting forth the general principles of pharmacology include Goodman and Gilman's *The Pharmacological Basis of Therapeutics*, 10th Ed., McGraw Hill Companies Inc., New York (2001).

Reference is made hereinafter in detail to specific embodiments of the
invention. While the invention will be described in conjunction with these specific embodiments, it will be understood that it is not intended to limit the invention to such specific embodiments. On the contrary, it is intended to cover alternatives, modifications, and equivalents as may be included within the spirit and scope of the invention as defined by the appended claims. In
the following description, numerous specific details are set forth in order to provided a thorough understanding of the present invention. The present invention may be practiced without some or all of these specific details. In other instances, well-known process operations have not been described in detail, in order not to unnecessarily obscure the present invention.

25 There is provided by the present invention compositions, as well as methods of making and of using pharmaceutical compositions, useful to achieve desirable pharmacokinetic properties. Such compositions stem from the discovery that solutions of cyclodextrin and cladribine in which cladribine is in a high thermodynamic state, when presented to the gastric mucosa

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through which they are absorbed are associated with improved cladribine absorption, as reflected by higher bioavailability and/or lower interpatient variation.

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It is postulated, without wishing to so limit the invention, that upon dissolution (*e.g.*, by contact with a fluid, such as a bodily fluid), dry compositions according to the invention form a locally saturated cladribine solution in which cladribine is in the state of highest thermodynamic activity (HTA), thus favoring absorption. Cladribine has a fairly low, although not insignificant, intrinsic aqueous solubility; it is in fact somewhat water soluble. The free cladribine formed from dissociation of the inclusion and noninclusion complexes in a saturated aqueous solution seeks a more stable activity level by being absorbed through the gastric mucosa.

In view of the foregoing, it is apparent that to produce optimal pharmaceutical compositions, in a solid oral dosage form, these dosage forms should be formulated to release a localized saturated cladribine solution, upon contact of the solid dosage forms with body fluid at the mucosa, in which cladribine is in its HTA state. To provide such a localized saturated solution *in vivo*, it is important to first identify the optimal ratio of cladribine to amorphous cyclodextrin, which ratio is referred to herein as the HTA ratio, to be used in the solid dosage form.

The HTA ratio is empirically determined and is identified as the ratio of cladribine to amorphous cyclodextrin which corresponds to the maximum amount of cladribine that can be complexed with a given amount of the cyclodextrin. The HTA ratio may be determined using an empirical method such as a phase solubility study to determine the saturation concentration of cladribine that can be solubilized with different concentrations of amorphous cyclodextrin solutions. Hence, the method identifies the concentrations at which a saturated cladribine-cyclodextrin complex is formed. It is noted that the molar ratio represented by a point on the phase solubility graph shows

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how many moles of amorphous cyclodextrin are the minimum needed to maintain the drug in the complex, under given conditions; this may then be converted to a weight ratio. For example, if a phase solubility diagram shows that 9 moles of a given cyclodextrin are needed to maintain the cladribine in a saturated complex, then multiplying the number of moles of cladribine by its molecular weight and multiplying the number of moles of the selected cyclodextrin by its molecular weight, one can arrive at the ratio of the products as an appropriate optimized weight ratio. A phase solubility study also provides information about the nature of the cladribine-

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cyclodextrin inclusion complex formed, for example whether the inclusion complex is a 1:1 complex (1 molecule of drug complexed with 1 molecule of cyclodextrin) or a 1:2 complex (1 molecule of drug complexed with 2 molecules of cyclodextrin).

In accordance with the present invention, one can start using either the selected amorphous cyclodextrin, such as hydroxypropyl-β-cyclodextrin (HPBCD) or hydroxypropyl-y-cyclodextrin, or cladribine as the fixed variable to which an excess of the other is added to identify various solubility data points (indicating saturated cladribine-cyclodextrin complexes) and draw the resultant line. Typically, cladribine is added to an aqueous solution having a known concentration of amorphous cyclextrin under conditions empirically found to promote complex formation. Generally, the complexation is conducted with heating, for example at about 45 to about 60°C for a significant period of time, e.g., at least 6-9 hours; it is believed that even better results can be obtained by heating at up to about 80°C for up to 24 hours. Excess precipitated cladribine is then removed and the cladribine

25 concentration is subsequently measured. This concentration represents the amount of cladribine solubilized for a given amorphous cyclodextrin concentration. This process is repeated for a different known concentration of cyclodextrin until several data points are obtained. Each data point represents the concentration of the cladribine dissolved in a known

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concentration of the selected amorphous cyclodextrin. The data points are then plotted to show the concentration of cladribine against the various cyclodextrin concentrations used. The graph is a phase solubility diagram which can be used to determine the amount of cladribine for any specific concentration of cyclodextrin used to form the solution under a given set of complexation conditions. It will be appreciated that the aqueous solubility of cladribine is about 5 mg/ml at room temperature and would be higher at elevated temperature. Consequently, the data points correspond to the amount of cladribine dissolved in aqueous HPBCD or other amorphous

-14-

10 cyclodextrin under the selected conditions; when later lyophilized, the solution yields a complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex. If equilibrium conditions are reached during the complexation, the amorphous cladribine-cyclodextrin complex will be saturated with cladribine.

One of skill in the art will appreciate that concentrations at which saturated complexes of cladribine with amorphous cyclodextrins are formed (and thus HTA ratios as well) may be identified by a variety of alternative methodologies. Accordingly, any method known in the field suitable to identify these concentrations is within the scope of the invention.

It has been discovered that desirable pharmacological properties (improved bioavailability and/or coefficient of variation as compared to traditional approaches) are associated with mixtures of inclusion complexes and non-inclusion complexes of cladribine and cyclodextrin.

Using intrinsically amorphous cyclodextrins, for example hydroxypropyl-β-cyclodextrin, hydroxypropyl-γ-cyclodextrin, randomly methylated cyclodextrins, and the like, with cladribine, which is a somewhat water soluble compound (capable of H-bonding through its free hydroxyl and

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amino groups), the cladribine provides increased solubility in solutions of these cyclodextrins. Not only is there increased water solubility but also Hbonded association of the cladribine with the cyclodextrin, separately from the actual inclusion complexed material.

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One of skill in the art will appreciate that the phase solubility diagram for each given starting concentration ratio represents the starting point of one's investigation on the basis of which variables (reactants' concentrations, temperature and time) may be altered to promote inclusion complex and non-inclusion complex associations favoring a higher or lower proportion of either type of association in the final product. Departure from the ratio of cladribine to cyclodextrin, the temperature and/or the dilution empirically found to promote equilibrium towards complex formation is then analyzed to promote the formation of mixtures of inclusion complexes and non-inclusion complexes of cladribine and cyclodextrin in various proportions according to the invention.

Thus, for example, by starting with more diluted cyclodextrin (*i.e.*, larger water volumes than that used for solubility plot analysis) logically will accommodate more cladribine in solution sequestering more of the same from complex formation. Upon evaporation, some of the solubilized cladribine will tend to associate with cyclodextrin in a non-inclusion complex fashion. By altering the initial dilution, one may shift equilibrium towards inclusion complex or non-inclusion complex formation. Similarly, by increasing complexation temperature, the water solubility of cladribine may be increased while decreasing the stability of inclusion complexes, thus promoting non-inclusion complexes. Thus, by altering complexation temperature, one may shift equilibrium towards inclusion complex or noninclusion complex formation. Finally, complexation time may be altered to favor the formation of mixtures of inclusion complexes and non-inclusion complexes of cladribine and cyclodextrin according to the invention.

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As exemplified hereinafter, it is possible to maximize the cladribine in solid amorphous mixtures, by forcing additional cladribine into solution (using more dilute solutions of cyclodextrin, higher temperatures and longer complexation times, as indicated above). When the solution is cooled off, the extensively amorphous nature of these cyclodextrins does not allow crystallization of an excess amount of cladribine beyond that which forms an inclusion complex with the cyclodextrin; and upon freezedrying/lyophilization, one obtains an amorphous mixture of cladribinecyclodextrin inclusion complex (which is amorphous) and amorphous free cladribine, loosely associated with uncomplexed cyclodextrin (and even with complexed cyclodextrin) by hydrogen-bonding, that is, the non-inclusion complex.

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As shown in the EXAMPLES, this may be done by maximizing solubilization by elevating the temperature (for example, to about 50° to 80°C), and stirring for many hours (up to 24 hours) before freeze-drying. The weight/weight ratios obtained were about 1:14 and 1:11. The apparent optimum weight/weight ratio under these exemplified conditions is the higher of these, or about 1:14 of cladribine: cyclodextrin. If too much excess caldribine is added to the complexation medium, then crystallization of some of the cladribine takes place, which would in turn result in some crystalline cladribine in the product; this undesired excess cladribine is not in solution and is not H-bonded to the amorphous cyclodextrin and lowers the weight ratio. Therefore, it is desirable to carefully control the amount of excess cladribine beyond that which will form the inclusion complex to only the amount which will dissolve in the solution. The desired amorphous mixture of amorphous inclusion complex and amorphous free cladribine can be termed a "complex cladribine-cyclodextrin complex," which includes both inclusion and non-inclusion/H-bonded complexes. The inclusion complex is a complex of cladribine inserted into the hydrophobic cavity of the selected amorphous cyclodextrin, while the non-inclusion/H-bonded complex is

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amorphous free cladribine loosely hydrogen-bonded to the cyclodextrin. It is estimated that about two-thirds (60 to 70%) of the cladribine will be in the non-inclusion complex, with the remaining one third (30 to 40%) being in the inclusion complex when the product is obtained as exemplified hereinbelow (17% HPβCD solution, 45 to 50°C complexation temperature for about 9 hours); by increasing the percentage of cyclodextrin used and/or manipulating the temperature, products can be readily obtained in which a much greater proportion of the amorphous mixture is in the form of the inclusion complex. In the case of a representative amorphous cyclodextrin, hydroxypropyl- β -cyclodextrin (HP β CD) a cladribine:cyclodextrin weight ratio of from about 1:10 to about 1:16 is appropriate for the exemplified conditions; the ratio is expected to be the same for hydroxypropyl- γ -

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cyclodextrin under those conditions. The material obtained is characterized by rapid dissolution of the cladribine in aqueous media.

Freeze-drying, also known as lyophilization, comprises three basic stages: first a freezing stage, then a primary drying stage and finally a secondary drying stage. EXAMPLE 2 below provides details of lyophilization as conducted on the batches described therein. This procedure can be further optimized by following the principles described by Xiaolin (Charlie) Tang and Michael J. Pikal in *Pharmaceutical Research*, Vol. 21, No. 2, February 2004, 191-200, incorporated by reference herein in its entirety and relied upon.

The above-described method requires amorphous cyclodextrins rather than originally crystalline cyclodextrins which have relatively low water solubilities, such as α -, β - or γ -cyclodextrin, 2,6-dimethyl- β -cyclodextrin and the like, because these cyclodextrins would allow crystallization of cladribine in excess of that forming an inclusion complex and therefore would not afford the desired amorphous mixture. The method also would not be useful if cladribine were highly hydrophobic/lipophilic, because in such a situation the drug would not have intrinsic aqueous solubility/H-bonding capability and

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could not provide the unique mixture obtained herein. However, in point of fact, cladribine has an aqueous solubility of 5 mg/ml at room temperature. thus a significant amount of the drug will be simply soluble in the water phase especially at higher than room temperature; also, as in the case of HPBCD, for example, some of the cladribine will be associated by hydrogenbonding to the 2-hydroxypropyl and free glucose-OH groups in the cyclodextrin via the two hydroxy functions found in the deoxyadenosine moiety of the cladribine.

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The cyclodextrins within the scope of this invention are amorphous 10 derivatives of the natural cyclodextrins α -, β - or y-cyclodextrin wherein one or more of the hydroxy groups are substituted, for example, by alkyl, hydroxyalkyl, carboxyalkyl, alkylcarbonyl, carboxyalkoxyalkyl, alkylcarbonyloxyalkyl, alkoxycarbonylalkyl or hydroxy-(mono or polyalkoxy)alkyl groups; and wherein each alkyl or alkylene moiety 15 preferably contains up to six carbons. Although commonly referred to as a single entity, an amorphous cyclodextrin is actually a mixture of many different entities, since the substituent groups can be located on various hydroxyls of the basic cyclodextrin structure. This in turn results in the amorphous nature of these cyclodextrins, which is indeed well-known.

20 Moreover, these cyclodextrins can be obtained in varying degrees of substitution, for example from 1 to 14, preferably from 4 to 7; the degree of substitution is the approximate average number of substituent groups on the cyclodextrin molecule, for example, the approximate number of hydroxypropyl groups in the case of the hydroxpropyl-ß-cyclodextrin

25 molecule, and all such variations are within the ambit of this invention. Substituted amorphous cyclodextrins which can be used in the invention include polyethers, for example, as described in U.S. Patent No. 3,459,731. Further examples of substituted cyclodextrins include ethers wherein the hydrogen of one or more cyclodextrin hydroxy groups is replaced by

C1-6alkyl, hydroxy-C1-6alkyl, carboxy-C1-6alkyl or C1-6alkyloxycarbonyl-

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C₁₋₆alkyl groups or mixed ethers thereof. In particular, such substituted cyclodextrins are ethers wherein the hydrogen of one or more cyclodextrin hydroxy groups is replaced by C1-3alkyl, hydroxy-C2-4alkyl or carboxy-C₁₋₂alkyl or more particularly by methyl, ethyl, hydroxyethyl, hydroxypropyl, hydroxybutyl, carboxymethyl or carboxyethyl. The term "C1-6alkyl" is meant to include straight and branched saturated hydrocarbon radicals, having from 1 to 6 carbon atoms such as methyl, ethyl, 1-methylethyl, 1,1-dimethylethyl, propyl, 2-methylpropyl, butyl, pentyl, hexyl and the like. Other cyclodextrins contemplated for use herein included glucosyl-β-cyclodextrin and maltosyl-βcyclodextrin. Of particular utility in the present invention are randomly methylated ß-cyclodextrin and polyethers such as hydroxypropyl-ßcyclodextrin, hydroxyethyl-β-cyclodextrin, hydroxypropyl-y-cyclodextrin, and

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cyclodextrin sulfobutyl ether. In addition to simple cyclodextrins, branched cyclodextrins and cyclodextrin polymers may also be used. Other cyclodextrins are described, for example, in Chemical and Pharmaceutical Bulletin 28: 1552-1558 (1980); Yakugyo Jiho No. 6452 (28 March 1983); Angew. Chem. Int. Ed. Engl. 19: 344-362 (1980); U.S. Patent Nos. 3,459,731 and 4,535,152; European Patent Nos, EP 0 149 197A and EP 0 197 571A; PCT International Patent Publication No. VVO90/12035; and UK Patent Publication GB 2,189,245.

hydroxyethyl-γ-cyclodextrin, as well as sulfobutyl ethers, especially β-

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References describing cyclodextrins for use in the compositions according to the present invention, and/or which provide a guide for the preparation, purification and analysis of cyclodextrins include the following: Cyclodextrin Technology by Jozsef Szejtli, Kluwer Academic Publishers (1988) in the chapter Cyclodextrins in Pharmaceuticals; Cyclodextrin Chemistry by M. L. Bender et al., Springer-Verlag, Berlin (1978); Advances in Carbohydrate Chemistry, Vol. 12, Ed. By M. L. Wolfrom, Academic Press, New York in the chapter "The Schardinger Dextrins" by Dexter French, pp. 189-260; Cyclodextrins and their Inclusion Complexes by J. Szejtli,

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Adakemiai Kiado, Budapest, Hungary (1982); I. Tabushi, *Acc. Chem. Research*, 1982, 15, pp. 66-72; W. Sanger, *Angewandte Chemie*, 92, p. 343-361 (1981); A. P. Croft *et al.*, *Tetrahedron*, 39, pp. 1417-1474 (1983); Irie *et al. Pharmaceutical Research*, 5, pp. 713-716 (1988); Pitha *et al.*, *Int. J. Pharm.* 29, 73 (1986); U.S. Patent Nos. 4,659,696 and 4,383,992; German Patent Nos. DE 3,118,218 and DE-3,317,064; and European Patent No. EP 0 094 157A. Patents describing hydroxyalkylated derivative of β - and γ cyclodextrin include Pitha U.S. Patent Nos. 4,596,795 and 4,727,064, Müller U.S. Patent Nos. 4,764,604 and 4,870,060 and Müller *et al.* U.S. Patent No. 6,407,079.

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Amorphous cyclodextrins of particular interest for complexation with cladribine include: hydroxyalkyl, *e.g.* hydroxyethyl or hydroxypropyl, derivatives of β - and γ -cyclodextrin; carboxyalkyl, *e.g.* carboxymethyl or carboxyethyl, derivatives of β - or γ -cyclodextrin; β -cyclodextrin sulfobutyl ether; and randomly methylated β -cyclodextrin. 2-Hydroxypropyl- β -cyclodextrin (HP β CD), 2-hydroxypropyl- γ -cyclodextrin (HP γ CD), randomly methylated β -cyclodextrin sulfobutyl ether, carboxymethyl- β -cyclodextrin (CM β CD) and carboxymethyl- γ -cyclodextrin (CM γ CD) are of special interest, especially hydroxypropyl- β -cyclodextrin and hydroxypropyl- γ -cyclodextrin.

Compositions of an amorphous mixture of amorphous free cladribine and an amorphous, preferably saturated, cladribine-cyclodextrin inclusion complex for use in the present invention can be prepared under conditions favoring complex formation in a liquid environment as described and as exemplified herein. The resultant liquid preparations can be subsequently converted to a dry form suitable for administration as a solid oral or transmucosal dosage form.

One of skill will appreciate that a variety of approaches are available in the field to prepare compositions as described herein. One available

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method exemplified herein includes the steps of mixing the cladribine in an aqueous solution of an amorphous cyclodextrin, separating un-dissolved cladribine (e.g., by filtering or centrifugation), and (yophilizing or freezedrying the saturated solution to form a solid amorphous mixture.

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Pharmaceutical compositions according to the invention may optionally include one or more excipients or other pharmaceutically inert components. One of the advantages of the invention, however, is that cladribine drug forms as described herein can be prepared with the minimal amount of excipients necessary for shaping and producing the particular form, such as a tablet or patch. Excipients may be chosen from those that do not interfere with cladribine, with cyclodextrin or with complex formation.

Dosage forms are optionally formulated in a pharmaceutically acceptable vehicle with any of the well-known pharmaceutically acceptable carriers, diluents, binders, lubricants, disintegrants, scavengers, flavoring 15 agents, coloring agents, and excipients (see Handbook of Pharmaceutical Excipients, Marcel Dekker Inc., New York and Basel (1998); Lachman et al. Eds., The Theory and Practice of Industrial Pharmacy, 3rd Ed., (1986); Lieberman et al., Eds. Pharmaceutical Dosage Forms, Marcel Dekker Inc., New York and Basel (1989); and The Handbook of Pharmaceutical Excipients, 3rd Ed., American Pharmaceutical Association and Pharmaceutical Press, 2000); see also Remington's Pharmaceutical Sciences, 18th Ed., Gennaro, Mack Publishing Co., Easton, PA (1990) and Remington: The Science and Practice of Pharmacy, Lippincott, Williams & Wilkins, (1995)). A simple solid oral dosage form consists of the amorphous mixture of amorphous free cladribine and amorphous cladribine-cyclodextrin complex (preferably saturated) as described above, *i.e.* the complex cladribine-cyclodextrin complex, compressed with a small amount (e.g.

about 1% by weight) of a suitable binder or lubricant such as magnesium stearate.

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In certain instances, oral absorption may be further facilitated by the addition of various excipients and additives to increase solubility or to enhance penetration, such as by the modification of the microenvirionment.

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The methods and pharmaceutical compositions described herein offer novel therapeutic modalities for the treatment of patients in need of treatment with cladribine. As shown herein, the invention addresses the problems of poor bioavailability traditionally associated with oral cladribine.

The compositions of the invention are particularly suitable as modalities for the treatment of any cladribine-responsive disease. Several disease states responsive to cladribine are well-documented in the literature (see *infra*). For any target disease state, an effective amount of the complex cladribine-cyclodextrin comples, *i.e.* the amorphous mixture of the optimized amorphous saturated cladribine-amorphous cyclodextrin complex with amorphous free cladribine as described above is used (*e.g.*, an amount affective for the treatment of multiple sclerosis, rheumatoid arthritis, or leukemia).

The term "therapeutically effective amount" or "effective amount" is used to denote treatments at dosages effective to achieve the therapeutic result sought. Therapeutically effective dosages described in the literature include those for hairy cell leukemia (0.09 mg/kg/day for 7 days), for multiple sclerosis (from about 0.04 to about 1.0 mg/kg/day (see U.S. Patent No. 5,506,214)); for other diseases, see also U.S. Patent Nos. 5,106,837 (autohemolytic anemia); 5,310,732 (inflammatory bowel disease); 5,401,724 (rheumatoid arthritis); 5,424,296 (malignant astrocytoma); 5,510,336 (histiocytosis); 5,401,724 (chronic myelogenous leukemia); and 6,239,118 (atherosclerosis).

Further, various dosage amounts and dosing regimens have been reported in the literature for use in the treatment of multiple sclerosis; see, for example: Romine et al., *Proceedings of the Association of American*

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Physicians, Vol. 111, No. 1, 35-44 (1999); Selby et al., The Canadian Journal of Neurological Sciences, 25, 295-299 (1998); Tortorella et al., Current Opinion in Investigational Drugs, 2 (12), 1751-1756 (2001); Rice et al., Neurology, 54, 1145-1155 (2000); and Karlsson et al., British Journal of Haematology, 116, 538-548 (2002); all of which are incorporated by reference herein in their entireties and relied upon.

Moreover, the route of administration for which the therapeutically effective dosages are taught in the literature should be taken into consideration. While the instant compositions optimize the bioavailability of cladribine following oral administration, it will be appreciated that even optimal bioavailability from oral dosage forms is not expected to approach bioavailability obtain after intravenous administration, particularly at early time points. Thus, it is often appropriate to increase a dosage suggested for intravenous administration to arrive at a suitable dosage for incorporation

- 15 into a solid oral dosage form. At the present time, it is envisioned that, for the treatment of multiple sclerosis, 10 mg of cladribine in the instant complex cladribine-cyclodextrin complex in the instant solid dosage form would be administered once per day for a period of five to seven days in the first month, repeated for another period of five to seven days in the second
- 20 month, followed by ten months of no treatment. Alternatively the patient would be treated with 10 mg of cladribine in the instant complex cladribinecyclodextrin complex in the instant dosage form once per day for a period of five to seven days per month for a total of six months, followed by eighteen months of no treatment. For further dosing information, see also U.S.
- Provisional Patent Application No. _____ [IVAX0021-P-USA/Attornev 25 Docket No. 033935-011], and U.S. Provisional Patent Application No. [IVAX0022-P-USA/Attorney Docket No. 033935-012], both entitled "Cladribine Regimen for Treating Multiple Sclerosis", both filed on March 25, 2004 and incorporated by reference herein in their entireties.

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Furthermore, one of skill will appreciate that the therapeutically effective amount of cladribine administered herein may be lowered or increased by fine tuning and/or by administering cladribine according to the invention with another active ingredient. The invention therefore provides a method to tailor the administration/treatment to the particular exigencies specific to a given mammal. Therapeutically effective amounts may be easily determined, for example, empirically by starting at relatively low amounts and by step-wise increments with concurrent evaluation of beneficial effect.

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10 As noted in the preceding paragraph, administration of cladribine in accord with this invention may be accompanied by administration of one or more additional active ingredients for treating the cladribine-responsive condition. The additional active ingredient will be administered by a route of administration and in dosing amounts and frequencies appropriate for each 15 additional active ingredient and the condition being treated. For example, in the treatment of multiple sclerosis, other useful drugs include interferon beta (Rebif[®], Betaseron[®]/Betaferon[®], Avonex[®]), identical to the naturally occurring protein found in the human body; glatiramer acetate (Copaxone[®]), a random chain (polymer) of the amino acids glutamic acid, lysine, alanine 20 and tyrosine; natalizumab (Antegren®), a monoclonal antibody; alemtuzumab (Campath-1H[®]), a humanized anti-CD52 monoclonal antibody; 4aminopyridine (also known as 4-AP and Fampridine), a drug that blocks the potassium channels in neurons; and amantadine, an anti-viral agent which improves muscle control and reduces muscle stiffness and is used to 25 alleviate the symptoms of fatigue in multiple sclerosis, a purpose for which pemoline (Cylert[®]) and L-Carnitine (a herbal product) may also be useful. In the treatment of hairy cell leukemia, additional active ingredients may include interferon alpha, pentostatin, fludarabine, rituximab (an anti-CD 20

monoclonal antibody) and the anti-CD22 recombinant immunotoxin BL 22; other additional active ingredients may be appropriate in other types of

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leukemias. In the treatment of rheumatoid arthritis, there are many other active ingredients which may be selected. These include NSAIDS (nonsteroidal anti-inflammatory drugs), which are of three types: salicylates such as aspirin, traditional NSAIDS such as ibuprofen and indomethacin, and COX-2 inhibitors such as celecoxib (Celebrex[®]), rofecoxib (Vioxx[®]), meloxicam (Mobic[®]), valdecoxib (Bextra[®]), lumiracoxib (Prexige[®]) and etoricoxib (Arcoxia®). Other drugs useful in treating rheumatoid arthritis which may be used in conjunction with the present invention include DMARDS, glucocorticoids, biological response modifiers and non-NSAID analgesics. DMARDS are disease-modifying anti-rheumatic drugs which include methotrexate, plaquenil, leflunomide (Arava[®]), sulfasalazine, gold, penicillamide, cyclosporine, methyl cyclophosamide and azathioprine. Glucocorticoids include dexamethasone, prednisolone, triamcinolone and many others. Biological response modifiers (which restore the diseasefighting ability of the immune system), include etanercept (Enrel®), a tumornecrosis factor inhibitor, infliximab (Remicade[®]), which is also an anti-TNF drug, anakinra (Kineret[®]), a selective IL-1 blocker, and Humira[®], a human monoclonal antibody which is another anti-TNF drug. The non-NSAID

hydrocodone, oxycodone and propoxyphene. Generally speaking, those drugs which work by a mechanism different from that of cladribine are particularly useful for concomitant therapy with the cladribine composition described herein. Those drugs which are effective by the oral route of administration and which are compatible with the instant cladribine
 complexes in a single dosage form may be incorporated into the instant dosage forms; otherwise, they should of course be separately administered

analgesics include acetaminophen as well as narcotic analgesics such as

in amounts, frequencies and via administration routes suitable to them. As used herein, "treating" means reducing, preventing, hindering the development of, controlling, alleviating and/or reversing the symptoms in the

individual to which a compound of the invention has been administered, as

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compared to the symptoms of an individual not being treated according to the invention. A practitioner will appreciate that the complexes, compositions, dosage forms and methods described herein are to be used in concomitance with continuous clinical evaluations by a skilled practitioner (physician or veterinarian) to determine subsequent therapy. Such evaluation will aid and inform in evaluating whether to increase, reduce or continue a particular treatment dose, and/or to alter the mode of administration.

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The methods of the present invention are intended for use with any 10 subject/patient that may experience the benefits of the methods of the invention. Thus, in accordance with the invention, the terms "subjects" as well as "patients" include humans as well as non-human subjects, particularly domesticated animals.

Any suitable materials and/or methods known to those of skill can be 15 utilized in carrying out the present invention. However, preferred materials and methods are described. Materials, reagents and the like to which reference are made in the following description and examples are obtainable from commercial sources, unless otherwise noted.

The following examples are intended to further illustrate certain 20 preferred embodiments of the invention and are not limiting in nature. Those skilled in the art will recognize, or be able to ascertain, using no more than routine experimentation, numerous equivalents to the specific substances and procedures described herein.

EXAMPLES

EXAMPLE 1 PHASE SOLUBILITY STUDY

A phase solubility study can be carried out as follows. Excess cladribine is added to cyclodextrin solutions of various concentrations of

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hydroxypropyl- β -cyclodextrin (HP β CD) and allowed to complex as described in EXAMPLE 2 below. The excess, undissolved cladribine is removed by filtration. The amount of cladribine in the complexation solution is measured to obtain a data point. This process is repeated with different known concentrations of HP β CD until several data points are obtained. These data points are then plotted graphically, each data point representing the amount of cladribine that can be dissolved in water with a specific concentration of cyclodextrin. Points on the line generated by the data points represent ratios for the product. One of skill in the art will realize the same results will be generated if excess cyclodextrin is added to cladribine solutions of known concentration.

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The molar concentrations of cladribine to cyclodextrin obtained are plotted and presented graphically. A representative phase solubility diagram is shown in the Figure. The plotted lines for cladribine-HPβCD represent cladribine solubilization for the conditions tested, that is, the ratio of the concentration of cladribine to the concentration of cyclodextrin. The area above each of the plotted lines represents conditions where excess insoluble cladribine is present. The area below each of the plotted lines represents the conditions where cyclodextrin is in excess.

The plot for cladribine-HP β CD shown in the Figure is approximately linear; this is indicative of a 1:1 complex, in which one molecule of the drug is complexed with one molecule of cyclodextrin. The Figure also shows that additional cyclodextrin is needed to maintain the cladribine in the complex. For example, about 0.14 mole of HP β CD is needed to maintain about 0.049 mole of cladribine dissolved under the selected conditions, which will ultimately provide the amorphous mixture of the amorphous, preferably saturated, cladribine-HP β CD inclusion complex and amorphous free cladribine (as a non-inclusion complex). Under the conditions of EXAMPLE 2 below, a significant portion of the cladribine in the product can be expected to be not in the inclusion complex but rather in amorphous form loosely held

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in intimate admixture therewith by hydrogen bonding as a non-inclusion complex.

EXAMPLE 2

PREPARATION OF CLADRIBINE-CYCLODEXTRIN COMPLEX FOR HUMAN TRIALS

Cladribine is complexed with HPBCD by the following method.

In 825 mL of distilled water, 172.5 g of hydroxypropyl-β-cyclodextrin are dissolved (forming an approximately 17% solution), then cladribine is added and the mixture is stirred at about 45 to about 50°C for about nine

10 hours. Stirring is continued for an additional 6 to 9 hours at room temperature. Any undissolved cladribine is removed by filtration and the solution is cooled to room temperature. To form the amorphous mixture of amorphous cladribine-cyclodextrin complex and amorphous free cladribine, the aqueous cladribine-cyclodextrin solution is dried by lyophilization prior to 15 incorporation into solid oral tablets. The lyophilization procedure comprises a freezing stage of rapidly bringing the complexation solution to about -40°C to about -80°C (e.g., about -45°C) for approximately 2 to 4 hours (preferably about 3 to 4 hours), followed by a primary drying stage at about -25°C for approximately 80-90 hours, typically under low pressure, and a second drying stage at about 30°C for about 15-20 hours.

Product made by the foregoing general procedure can be analyzed by HPLC (utilizing a Hypersil ODS 3 micron column and an acetonitrile based mobile phase, with UV detection at 264 nm) to find the weight ratio of cladribine to cyclodextrin in the final product. Final product preparations can be further characterized by methods known in the art, including, for example by inspecting appearance, ascertaining the overall impurity content by HPLC, ascertaining the water content using a Karl Fischer titrator, determining the dissolution profile by a standard method, for example using USP<711>Apparatus II equipment and UV detection at 264 nm, inspecting

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the content uniformity and performing quantitative assay by HPLC analysis of the active ingredient.

Two batches of cladribine/cyclodextrin product, FD04 and FD05, were made by the foregoing general procedure as follows:

Purified water (825 mL) was pre-heated at 48°C (target range 45°C to

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 50° C) in a 1-liter glass vessel by immersion in a water bath. The heated water was stirred to achieve a controlled central vortex. 2-hydroxypropyl- β -cyclodextrin (172.50 g) was weighed and slowly added to the heated water over a period of 40 minutes. The resulting solution was stirred for a further 10 minutes to ensure complete dissolution of the cyclodextrin. Cladribine (12.00 g for FD04 and 18.75 g for FD05) was weighed and added to the stirred cyclodextrin solution, which turned cloudy before becoming clear. The resulting clear solution was maintained at 48°C and continually stirred for 9 hours. Stirring continued for a further 7 hours while the solution cooled to room temperature.

Use of a larger amount of cladribine in the preparation of FD05 was part of an attempt to optimize the procedure; however, it was found that the initial amount of cladribine in that case was too great and precipitation was observed at the end of the cooling step for batch FD05. The solution was filtered to remove the precipitate. Analysis of the resultant product revealed (assay value = 87.2%) that 16.35 g of cladribine had been incorporated into the cyclodextrin complex in the case of FD05. No filtration was required for batch FD04, indicating that the amounts used in the preparation of FD04 were more appropriate and that the FD05 procedure could be optimized by beginning with a smaller amount of cladribine (16.35 g rather than 18.75 g), thus avoiding the filtration step.

After cooling to room temperature and, in the case of FD05, filtering, the solutions were filled into 100 mL lyophilization vials (20 mL solutions per vial), the filled vials were partially stoppered and lyophilized. The

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lyophilization included freezing at -45°C for about 200 minutes, a primary drying phase at -25°C under a pressure of 100 mTorr for about 5,200 minutes and a secondary drying phase at 30°C for about 1,080 minutes as set forth below:

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Step	Process	Temperature	Pressure (mTorr)	Time (min)
1	Load	4°C		
2	Load Hold	4ºC	n/a	120
3	Ramp	-45°C	n/a	120
4	Freezing	-45°C	n/a	200
5	Ramp	-25°C	100	120
6	Primary drying	-25°C	100	5200
7	Ramp	30°C	50	240
8	Secondary drying	30ºC	50	1080
9	Finish	30°C	Vials closed und	er vacuum

TABLE I

The FD04 and FD05 batches of cladribine/cyclodextrin product made by the foregoing procedure were analyzed by HPLC (utilizing a Hypersil ODS 3 micron column and an acetonitrile based mobile phase with UV detection at 264 nm) and empirically found to have the following characteristics:

TAE	BLE	Ξ II	
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Lot No.	Cladribine: HPβCD w/w	Cladribine: HPβCD Weight Ratio
FD04	12.00g:172.50g	1:14.38
FD05	16.35g:172.50g	1:10.55

The products were analyzed by DSC thermograms and X-ray diffraction methods to determine any free crystalline cladribine in the lyophilized material. Importantly, the samples exhibited no transitions in the region of 210°C to 230°C, which is associated with the melting of crystalline cladribine. In both cases, no significant thermal activity was recorded in the range of 210°C to 230°C, suggesting that the complexes obtained at the end of the lyophilization do not have any significant amount of free crystalline cladribine, considering the sensitivity of the analytical method (up to 3% w/w). This conclusion was supported by the absence of peaks for crystalline cladribine from X-ray diffraction traces for both complexes FD04 and FD05.

The products are amorphous mixtures of amorphous cladribine-HP β CD inclusion complex and amorphous free cladribine hydrogen-bonded to the cyclodextrin as a non-inclusion complex. The cladribine:HP β CD weight ratios obtained were about 1:14 and 1:11.

Generally speaking, amorphous mixtures within the scope of the present invention have cladribine:HP β CD weight ratios of from about 1:10 to 1:16.

EXAMPLE 3

PREPARATION OF ORAL TABLETS

Tablets were manufactured using batches of amorphous mixtures FD04 and FD05 described in EXAMPLE 2 for use in a clinical study.

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Batch N0120 was manufactured using cladribine-2-HP β CD complex mixture DF05 to a batch size of 3,000 tablets and batch N0126 was manufactured using cladribine-HP β CD complex mixture FD04 to a batch size of 800 tablets. The master formulations for the two batches are shown in TABLE III. Batch N0120 represented 3.0 g tablets and Batch N0126 represented 10 mg tablets for clinical study.

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**************************************		mg/tablet	mg/tablet
Constituent	Lot Number	3.0 mg	10.0 mg
		Batch N0120	Batch N0126
Cladribine-HPβCD complex mix	FD05	30.60*	
Cladribine-HPβCD complex mix	FD04		153.75**
Sorbitol powder NF	1007403	68.4	44,25
Magnesium stearate NF	1006280	1.00	2.00
Total		100.00	200.00

TABLE III

*Equivalent to 3.0 mg cladribine per tablet. **Equivalent to 10.0 mg cladribine per tablet.

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The following table sets forth the method of manufacture of the Batch N0120 and N0126 tablets.

TABLE IV

1.	Pre-mix the magnesium stearate with an approximately equal quantity of sorbitol power.
2.	Pass the cladribine-HP β CD complex and the remainder of the sorbitol powder into a one-liter glass jar via a 40-mesh screen.
3.	Blend the contents for 10 minutes at 12 rpm.
4.	Pass the magnesium stearate/sorbitol powder pre-mix into the glass jar via the 40-mesh screen.
5.	Blend the final mixture for 5 minutes at 12 rpm.
6.	Compress into 3.0 mg and 10.0 mg tablets at a target compression weight of 100 mg and 200 mg, respectively.

Both the Batch N0120 3.0 mg tablets and the Batch N0126 10.0 mg tablets were round, with one side flat-beveled edged and the other side shallow convex. The Batch N0120 3.0 mg tablets had an average weight of 100 mg, a thickness of 2.7 mm, a friability of 0.2%, a hardness of 4 Kp and a disintegration time of 3 minutes. The Batch N0126 10.0 mg tablets had an average weight of 198 mg, a thickness of 4.2 mm, a friability of 1%, a hardness of 2.8 Kp and a disintegration time of 5 minutes 42 seconds.

The Batch N0120 3.0 mg and N0126 10.0 mg tablets were used in the clinical study summarized in EXAMPLE 5 below.

EXAMPLE 4

CLINICAL STUDY: RELATIVE BIOAVAILABILITY

The objective of this study was to assess the relative bioavailability of three oral cladribine formulations: (1) a cyclodextrin-based formulation according to the instant invention (Tablet 1: complex FD05, i.e. Batch No.
15 N0120 tablets described above); (2) a mucoadhesive formulation (Tablet 2: containing 3.0 mg cladribine, 10 mg of Carbopol 71G NF, 22.2 mg of dicalcium phosphate, 64.3 mg of lactose and 0.5 mg of magnesium stearate, Batch No. N0121); and (3) a hard-gel capsule (Capsule containing 3.0 mg cladribine, 5.0 mg Carbopol 974P, 91.3 mg Avicel PH101, 100.0 mg Avicel
20 PH102, 0.2 mg colloidal silicon dioxide and 0.5 mg magnesium stearate, Batch No. RD03030) in comparison with one fixed subcutaneous clardribine administration (reference formulation) in patients with MS (multiple sclerosis).

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This study was a 2 center, open-label, randomized, 4-way crossover single dose study using twelve patients with MS. Patients received randomly three different fixed oral doses (3.0 mg) and a fixed subcutaneous dose of 3.0 mg. The four treatment days were separated by a drug-free interval of at

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least 5 days. In each treatment period, blood samples were collected over a 24-hour period for evaluation of plasma cladribine.

The plasma concentration of cladribine was measured by a HPLC/MS/MS method. Using this method, the relationship between concentration versus peak area ratio was found to be linear within the range of 100 pg/ml to 50,000 pg/ml for cladribine. The limit of quantification was 100 pg/ml, Analysis of samples was carried out in 16 runs. No calibrator had to be excluded from fitting of the calibration curve and accuracy of each quality control sample met the GLP requirements.

10 576 clinical plasma samples were analyzed and concentration values of cladribine were determined. The results were compiled and are summarized in the tables below (Tables V and VI). In these tables, the following definitions are applicable: T_{max} is the time to reach maximum concentration in the plasma; T_{1/2} is the half-life of cladribine in the plasma; 15 C_{MAX} is the maximum concentration of cladribine in the plasma; AUC_{inf} is the area under the curve for the measured data from zero extrapolated to infinity; AUCt is the area under the curve for the measured data (from zero to the last time point); Geom Mean is the geometric mean; CV is the coefficient of variation (relative standard deviation); LL is the lower limit; UL is the upper limit.

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TABLE VSummary Statistics for Pharmacokinetic Parameters for Cladribine StudyObtained via Non-Compartmental Analyses. (n=12).

Pharmacokinetic Parameter	3.0 mg	y subcuta	aneous									
				3n	ng Tablet	1	3r	ng Tablet	2	3 г	3 mg Capsule	
	Geom	Mean	CV**	Geom	Mean	CV**	Geom	Mean	CV**	Geom	Mean	CV**
	Mean	± SD	(%)	Mean	± SD	(%)	Mean	± SD	(%)	Mean	±SD	(%)
T _{max} (hr)	N/A	.313 ±.113	36.2	N/A	.521 ±.167	32.1	N/A	1.25 ±.839	67.1	N/A	2.25 ±.622	27.7
T _½ (hr)	N/A	6.69 ±2.01	30.1	N/A	7.55 ±2.50	33.1	N/A	6.73 ±2.82	41.9	N/A	6.27 ±2.31	36.9
C _{max} (pg/ml)	23186	N/A	40.1	6597	N/A	24.7	5041	N/A	52.6	3818	N/A	36.8
AUC _{inf} (hr⋅pg/ml)	57254	N/A	44.4	24936	N/A	28.8	21676	N/A	42.7	22604	N/A	39.5
AUC _t (hr·pg/ml)	54725	N/A	43.8	23182	N/A	28.0	20063	N/A	42.1	20951	N/A	42.0

5 **CV=SD/mean for T_{max} and $T_{\frac{1}{2}}$ and CV% geometric mean for C_{max} , AUC_{inf} and AUC_t.

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TABLE VI

Ratios of Oral to Subcutaneous Pharmacokinetic Parameters and Corresponding Two-Sided 90% Confidence Intervals for Cladribine Study (n=12).

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Pharmacokinetic Parameter	3 mg Tablet 1		3mg Tablet 2		3mg Capsule	
	Ratio*	LL, UL	Ratio*	LL, UL	Ratio*	LL, UL
AUCinf	43.1	35.7, 52.1	38.4	31.8, 46.4	38.9	32.1, 47.0
AUCt	41.9	34.6, 50.8	37.2	30.7, 45.0	37.6	31.0, 45.5

*Ratios (dose normalized) and Corresponding 95% LL obtained via inverse transformation of log-transformed data.

EXAMPLE 5

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CLINICAL STUDY: DOSE RESPONSE AND ABSOLUTE BIOAVAILABILITY

The objective of this study was to assess the systemic availability of cladribine after oral administration in two different fixed oral doses, in comparison with one fixed intravenous administration (reference formulation) in patients with MS (multiple sclerosis), and to evaluate the safety and tolerability of cladribine in this population.

This study was a 3 center, open-label, randomized, 3-way crossover

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single dose study using twenty-six patients with MS. Patients received
randomly two different fixed oral doses (3.0 mg and 10.0 mg) and a fixed
intravenous dose of 3.0 mg (administered as a 1 hour infusion). The three
treatment days were separated by a drug-free interval of at least 5 days. In

each treatment period blood samples were collected over a 24-hour period for evaluation of plasma cladribine.

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The plasma concentrations of cladribine were measured by a HPLC/MS/MS method. Using this method the relationship between concentrations versus peak area ratios was found to be linear within the range of 100 pg/ml to 50,000 pg/ml for cladribine. The limit of quantification was 100 pg/ml. Analysis of samples was carried out in 16 runs. Except the first run (which had to be rejected because of equipment failure), all other runs could be accepted. No calibrator had to be excluded from fitting of the calibration curve and accuracy of each quality control sample met the GLP requirements.

858 clinical plasma samples were analyzed and concentration values of cladribine were determined. The results were compiled and are summarized in the tables below [TABLES VII through X]. In these tables, the following definitions are applicable: T_{max} is the time to reach maximum concentration in the plasma; $T_{1/2}$ is the half-life of cladribine in the plasma; C_{max} is the maximum concentration of cladribine in the plasma; AUC_{inf} is the area under the curve for the measured data from zero extrapolated to infinity; AUC_t is the area under the curve for the measured data (from zero to the last time point); Geom Mean is the geometric mean; CV is the coefficient of variation (relative standard deviation); LL is the lower limit; UL is the upper limit; σ^2 is the mean variance; σ_B^2 is the mean variance between subjects; σ_W^2 is the mean variance within subjects; CV_T is the total coefficient of variation; and CV_W is the coefficient of variation within subjects.

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TABLE VII

Summary Statistics for Pharmacokinetic Parameters for Cladribine Study Obtained via Non-Compartmental Analysis (n=26)

Pharmaco- kinetic Parameter	3.0 mg IV infusion				Or	al Adm	inistratio	on	
					3.0 mg			10.0 mg	
	Geom	Mean	CV***	Geom	Mean	CV**	Geom	Mean	CV**
	Mean	± SD	(%)	Mean	± SD	(%)	Mean	± SD	(%)
Tmax(hr)	N/A	.817	48.6	N/A	.548	54.8	N/A	.558	36.5
		±.397			±.300			±.204	
T1/2(hr)	N/A	6.50	19.5	N/A	5.85	20.2	N/A	5.60	13.3
		±1.27			±1.18			±0.75	
C _{max} (pg/ml)	21425	N/A	27.6	5608	N/A	49.5	21242	N/A	50.5
AUCinf	58528	N/A	24.0	20159	N/A	35.0	76690	N/A	30.3
(hr∙pg/ml)			_						
AUCt	56396	N/A	24.0	19166	N/A	36.9	74532	N/A	30.3
(hr·pg/ml)									

**CV=SD/mean for T_{max} and T¹/₂ and CV% geometric mean for C_{max}, AUC_{inf} and AUC_t.

TABLE VIII

Ratios of Oral to I.V. Pharmacokinetic Parameters and Corresponding Lower Limit (LL) for the one-sided 95% Confidence Interval for Cladribine Study (n=26)

Pharmacokinetic Parameter		Oral Adm	ninistration		
	3.0	mg	10.0 mg		
	Ratio*	LL	Ratio*	LL	
AUC _{inf}	34.5	31.7	39.1	35.9	
AUCt	34.0	31.2	39.4	36.1	

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*Ratios (dose normalized) and Corresponding 95% LL obtained via inverse transformation of log-transformed data.

TABLE IX

Ratios and Corresponding two-sided 90% Confidence Intervals for Cladribine Study (n=26)

Pharmacokinetic Parameter	·	10.0 mg/3.0 mg	
	Ratio*	LL	UL
C _{max}	112.6	95.1	1.33.3
AUC	113.3	104.2	123.3
AUCt	115.8	106.1	126.5

*Ratios (dose normalized) and Corresponding 90% CI obtained via inverse transformation of log-transformed data.

TABLE X

Variance components for Cladribine Study (n=26)

Source of variation	C _{max}	AUCinf	AUCt
Between $(\sigma_{\rm B}^{2})$.0380	.0487	.0492
With (σ_{W}^{2})	.1315	.0330	.0357
TOTAL $(\sigma_B^2 + \sigma_W^2)$.1695	.0816	.0849
CV _T (%)	43.0	29.2	29.8
CV _w (%)	37.5	18.3	19.1

Where PK parameters are dose-adjusted and $CV = \sqrt{\exp(\sigma^2) - 1}$

The foregoing is considered as illustrative only of the principles of the invention. Further, since numerous modifications and changes will readily occur to those skilled in the art, it is not desired to limit the invention to the exact construction and operation shown and described, and accordingly, all suitable modifications and equivalents thereof may be resorted to, falling within the scope of the invention claimed.

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WHAT IS CLAIMED IS:

1. A pharmaceutical composition comprising a complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex, formulated into a solid oral dosage form, said composition comprising no significant amount of free crystalline cladribine therein.

2. The pharmaceutical composition according to Claim 1, wherein the complex is saturated with cladribine.

3. The composition according to Claim 1, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin, hydroxypropyl-γ-cyclodextrin, randomly methylated β-cyclodextrin, carboxymethyl-β-cyclodextrin or sulfobutyl-β-cyclodextrin.

4. The composition according to Claim 1, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin or hydroxypropyl-γ-cyclodextrin.

5. The composition according to Claim 1, wherein the weight ratio of cladribine to amorphous cyclodextrin is from about 1:10 to about 1:16.

6. The composition according to Claim 5, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin or hydroxypropyl-γ-cyclodextrin.

7. The composition according to Claim 1, wherein the approximate molar ratio of cladribine to amorphous cyclodextrin corresponds to a point located on the curve of a phase solubility diagram for saturated complexes of cladribine in varying concentrations of the cyclodextrin.

8. The composition according to Claim 1, wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b).

9. A method for enhancing the oral bioavailability of cladribine comprising orally administering to a subject in need thereof a pharmaceutical composition comprising a complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex, formulated into a solid oral dosage form, said composition comprising no significant amount of free crystalline cladribine therein.

10. The method according to Claim 9, wherein the complex is saturated with cladribine.

11. The method according to Claim 9, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin, hydroxypropyl- γ -cyclodextrin, randomly methylated β -cyclodextrin, carboxymethyl- β -cyclodextrin or sulfobutyl- β -cyclodextrin.

12. The method according to Claim 9, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin or hydroxypropyl- γ -cyclodextrin.

13. The method according to Claim 9, wherein the weight ratio of cladribine to amorphous cyclodextrin is from about 1:10 to about 1:16.

14. The method according to Claim 13, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin or hydroxypropyl-γ-cyclodextrin.

15. The method according to Claim 9, wherein the approximate molar ratio of cladribine to amorphous cyclodextrin corresponds to a point located on the curve

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of a phase solubility diagram for saturated complexes of cladribine in varying concentrations of the cyclodextrin.

16. The method according to Claim 9, wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b).

17. A method for the treatment of symptoms of a cladribine-responsive condition in a subject suffering from said symptoms comprising orally administering to said subject a pharmaceutical composition comprising a complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex, formulated into a solid oral dosage form, said composition comprising no significant amount of free crystalline cladribine therein.

18. The method according to Claim 17, wherein the complex is saturated with cladribine.

19. The method according to Claim 17, wherein the cladribine-responsive condition is selected from the group consisting of multiple sclerosis, rheumatoid arthritis and leukemia.

20. The method according to Claim 19, wherein the cladribine-responsive condition is multiple sclerosis.

21. The method according to Claim 17, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin, hydroxypropyl-γ-cyclodextrin, randomly methylated β-cyclodextrin, carboxymethyl-β-cyclodextrin or sulfobutyl-β-cyclodextrin.

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22. The method according to Claim 17, wherein the weight ratio of cladribine to amorphous cyclodextrin is from about 1:10 to about 1:16.

23. The method according to Claim 17, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin or hydroxypropyl-γ-cyclodextrin.

24. The method according to Claim 17, wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b).

25. A complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex.

26. The complex according to Claim 25, saturated with cladribine.

27. The complex according to Claim 25, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin, hydroxypropyl-γ-cyclodextrin, randomly methylated β-cyclodextrin, carboxymethyl-β-cyclodextrin or sulfobutyl-β-cyclodextrin.

28. The complex according to Claim 25, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin or hydroxypropyl-γ-cyclodextrin.

29. The complex according to Claim 25, wherein the weight ratio of cladribine to amorphous cyclodextrin is from about 1:10 to about 1:16.

30. The complex according to Claim 29, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin or hydroxypropyl-γ-cyclodextrin.

31. The complex according to Claim 25, wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b).

32. A process for the preparation of a complex cladribine-cyclodextrin complex as claimed in Claim 25, which comprises the steps of:

 (i) combining cladribine and an amorphous cyclodextrin in water at a temperature of from about 40 to about 80°C and maintaining said temperature for a period of from about 6 to about 24 hours;

(ii) cooling the resultant aqueous solution to room temperature; and

(iii) lyophilizing the cooled solution to afford an amorphous product.

33. The process according to Claim 32, further comprising a filtration step following step (ii).

34. The process according to Claim 32, wherein step (i) is performed at a temperature of from about 45 to about 60°C.

35. The process according to Claim 32, wherein step (i) is performed at a temperature of from about 45 to about 50°C.

36. The process according to Claim 34, wherein step (i) is performed with stirring.

37. The process according to Claim 36, wherein step (i) is performed for a period of from about 6 to about 9 hours.

38. The process according to Claim 32, wherein step (ii) is performed for a period of from about 6 to about 9 hours.

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39. The process according to Claim 32, wherein step (iii) comprises an initial freezing stage in which the solution is cooled to from about -40 to about -80° C, and held at said temperature for a period of from about 2 to about 4 hours.

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40. The process according to Claim 39, wherein, in the initial freezing stage of step (iii), the solution is cooled to about -45°C.

41. The process according to Claim 34, wherein 12.00 parts by weight of cladribine and 172.50 parts by weight of hydroxypropyl- β -cyclodextrin are introduced in step (i), or wherein 16.35 parts by weight of cladribine and 172.50 parts by weight of hydroxypropyl- β -cyclodextrin are introduced in step (i).

42. The process according to Claim 41, wherein 825 parts by volume of water are introduced in step (i).

43. The process according to Claim 32, wherein the lyophilization step (iii) comprises:

(a) an initial freezing stage in which the complexation solution is brought to from about -40°C to about -80°C for approximately 2 to 4 hours;

(b) a primary drying stage at about -25°C for approximately 80 to 90 hours; and

(c) a secondary drying stage at about 30°C for approximately 15 to 20 hours.

44. The process according to Claim 43, wherein stage (a) of the lyophilization is conducted at about -45°C for approximately 3 to 4 hours.

45. The process according to Claim 43, wherein stage (b) of the lyophilization is conducted under a pressure of about 100 mTorr.

46. A pharmaceutical composition according to Claim 1, obtainable by a process comprising the steps of:

(i) combining cladribine and an amorphous cyclodextrin in water at a temperature of from about 40 to about 80°C and maintaining said temperature for a period of from about 6 to about 24 hours;

(ii) cooling the resultant aqueous solution to room temperature;

(iii) lyophilizing the cooled solution to afford an amorphous product; and

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(iv) formulating the amorphous product into a solid oral dosage form.

47. The pharmaceutical composition according to Claim 46, wherein the process further comprises a filtration step following step (i) or (ii).

48. The pharmaceutical composition according to Claim 46, wherein step (i) of the process is performed at a temperature of from about 45 to about 60°C.

49. The pharmaceutical composition according to Claim 46, wherein step (i) of the process is performed at a temperature of from about 45 to about 50°C.

50. The pharmaceutical composition according to Claim 48, wherein step (i) of the process is performed with stirring.

51. The pharmaceutical composition according to Claim 50, wherein step (i) of the process is performed for a period of from about 6 to about 9 hours.

52. The pharmaceutical composition according to Claim 46, wherein step (ii) of the process is performed for a period of from about 6 to about 9 hours.

53. The pharmaceutical composition according Claim 46, wherein step (iii) comprises an initial freezing stage in which the solution is cooled to from about -40 to about -80°C, and held at said temperature for a period of from about 2 to about 4 hours.

54. The pharmaceutical composition according to Claim 53, wherein, in the initial freezing stage of step (iii), the solution is cooled to about -45°C.

55. The pharmaceutical composition according to Claim 48, wherein 12.00 parts by weight of cladribine and 172.50 parts by weight of the hydroxypropyl-β-cyclodextrin are introduced in step (i) of the process, or wherein 16.35 parts by weight of cladribine and 172.50 parts by weight of the hydroxypropyl-β-cyclodextrin are introduced in step (i) of the process.

56. The pharmaceutical composition according to Claim 55, wherein 825 parts by volume of water are introduced in step (i) of the process.

57. The pharmaceutical composition according to Claim 46, wherein the lyophilization step (iii) of the process comprises:

(a) an initial freezing stage in which the complexation solution is brought to from about -40°C to about -80°C for approximately 2 to 4 hours;

(b) a primary drying stage at about -25°C for approximately 80 to 90 hours; and

(c) a secondary drying stage at about 30°C for approximately 15 to 20 hours.

58. The pharmaceutical composition according to Claim 57, wherein stage (a) of the lyophilization is conducted at about -45°C for approximately 3 to 4 hours.

59. The pharmaceutical composition according to Claim 57, wherein stage (b) of the lyophilization is conducted under a pressure of about 100 mTorr.

60. The pharmaceutical composition according to Claim 46, wherein the formulation step (iv) of the process comprises blending the complex with magnesium stearate and compressing into tablets.

61. The pharmaceutical composition according to Claim 60, wherein magnesium stearate is pre-mixed with sorbitol powder before blending with the complex.

62. The method according to Claim 17, wherein the administration of cladribine is accompanied by administration of one or more additional active ingredients for treating the cladribine-responsive condition.

63. The method according to Claim 62, wherein the cladribine-responsive condition is multiple sclerosis.

64. The method according to Claim 63, wherein one or more additional active ingredients for treating multiple sclerosis is/are selected from the group consisting of interferon beta, glatirimer acetate, natalizumab, alemtuzumab, 4-aminopyridine and amantadine.

ABSTRACT

Provided are compositions of cladribine and cyclodextrin which are especially suited for the oral administration of cladribine..



1/1

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re F	atent Application of	MAIL STOP AMENDMENT		
Nichol	as Bodor et al.	Group Art Unit:		
Applic	ation No.:) Examiner:		
Filed:	January 5, 2011) Confirmation No.:		
For:	ORAL FORMULATIONS OF CLADRIBINE	 Continuation of Application No. 10/551,205, filed November 14, 2006, now allowed 		

FIRST INFORMATION DISCLOSURE STATEMENT

)

Commissioner for Patents P.O. Box 1450 Alexandria, VA 22313-1450

Sir:

In accordance with the duty of disclosure as set forth in 37 C.F.R. § 1.56, the accompanying information is being submitted in accordance with 37 C.F.R. §§ 1.97 and 1.98. Applicants request the Examiner's consideration of the documents listed on the accompanying substitute Form PTO-1449.

All of the listed documents were previously made of record in prior Application No. 10/551,205, filed November 14, 2006, upon which Applicants rely for the benefits provided in 35 U.S.C. § 120. In accordance with 37 C.F.R. § 1.98, a copy of each of the listed documents, is not required. For the Examiner's convenience, however, further copies of the International Search Report (ISR) and International Preliminary Report on Patentability and Written Opinion issued during the international phase of the parent application are provided herewith.

This Statement, substitute Form PTO-1449 and other documents are being submitted within three (3) months of the filing or entry of the national stage of this application or before the first Office Action on the merits, whichever is later. Since these documents are being filed within the time period set forth in 37 C.F.R. § 1.97(b), no fee or statement is required.

The following remarks are offered with respect to the listed documents which are not in English:

<u>DE 31 18 218</u> is in German. Applicants do not have an English translation. However, an English abstract was provided in the parent; the abstract and the citation of the document in the instant specification serve as a brief statement of relevance. Applicants consider this a general state of the art reference.

<u>DE 33 17 064</u> is in German. Applicants do not have an English translation but provided an English abstract in the parent. The abstract and citation of the document in the specification serve as a brief statement of relevance. Applicants consider this a general state of the art reference.

<u>EP 0 194 197 B1</u> is in German, although the claims are also present in English and a full English translation was provided in parent Application No. 10/551,205, filed November 14, 2006. This document is cited in the specification and applicants consider it to be a general state of the art reference.

The non-patent literature document by Shen GAO was cited during prosecution of the corresponding Chinese application and is in the Chinese language. An English translation of the cited section (page 105, lines 25-29) was filed in the parent application. This reference was cited as relevant only to some of the process and product-by-process claims.

The Van Axel Castelli et al. article and the three non-patent literature documents listed immediately after it are not prior art. They were submitted during prosecution of the parent to support positions taken by applicants therein.

It is respectfully requested that an Examiner-initialed copy of the enclosed Substitute Form PTO-1449 be returned to the undersigned.

The Director is hereby authorized to charge any appropriate fees under 37 C.F.R. §§ 1.16, 1.17 and 1.21 that may be required by this paper, and to credit any overpayment, to Deposit Account No. 02-4800.

Respectfully submitted,

BUCHANAN INGERSOLL & ROONEY PC

Date: January 7, 2011

By:

Mary Katherine Baumeister Registration No. 26254

Customer No. 21839 703 836 6620 Substitute for form 1449/PTO & 1449B/PTO

FIRST INFORMATION DISCLOSURE STATEMENT BY APPLICANT

(use as many sheets as necessary)

Sheet 1 of 4

U.S. PATENT DOCUMENTS

Application Number

First Named Inventor

Attorney Docket No.

Examiner Name

Filing Date

Complete if Known

January 5, 2011

Nicholas S. Bodor

0056192-000067

Examiner Initials	Document Number- Kind Code	Publication Date MM-DD-YYYY	Name of Patentee or Applicant of Cited Document	Pages, Columns, Lines Where Relevant Passages or Figures Appear
	US-4,383,992	05-17-1983	Lipari	
	US-6,239,118 B1	05-29-2001	Shatz et al.	
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	US-4,659,696	04-21-1987	Hirai et al.	
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	US-4,497,803	02-05-1985	Harada et al.	
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	US-4,727,064	02-23-1988	Pitha	
	US-4,596,795	06-24-1986	Pitha	
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	US-4,535,152	08-13-1985	Szejtli et al.	
	US-6,174,873	01-16-2001	Wrenn, Jr. et al.	
	US-6,699,849	03-02-2004	Loftsson et al.	

FOREIGN PATENT DOCUMENTS											
	Foreign Patent Document						S	TATUS	1		
Examiner Initials	Country Code ¹ , Number, Kind Code	Publication Date (MM-DD-YYYY)	Name of Patentee or Applicant of Cited Document	Translation	Partial Translation	Eng. Lang. Summary	Search Report	IPER	Abstract	Cited Pg	in Spec. / . No(s).
	EP 0 197 571 A2	10-15-1986	Janssen Pharmaceutica N.V.							x	
	WO 90/12035A1	10-18-1990	Janssen Pharmaceutica N.V.							x	
	DE 31 18 218 A1	04-22-1982	Chinoin Gyogyszer Es Vegyeszet						x	x	
	DE 33 17 064 A1	11-15-1984	Consortium Elektrochem Ind						x	x	
	GB 2 189 245 A	10-21-1987	American Maize- Products Company							x	
	EP 0 149 197 B1	07-24-1985	Janssen Pharmaceutica N.V.	Х						x	
	EP 0 094 157 A1	11-16-1983	Janssen Pharmaceutica N.V.							x	
	WO 99/42111	08-26-1999	Cyclops, ehf								

Examiner Signature Date Considered

*EXAMINER: Initial if reference considered, whether or not citation is in conformance with M.P.E.P. § 609. Draw line through citation if not in conformance and not considered. Include copy of this form with next communication to Applicant.

Substitute for form 1449/PTO & 1449B/PTO

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FIRST INFORMATION DISCLOSURE STATEMENT BY APPLICANT

(use as many sheets as necessary)

Application Number	
Filing Date	January 5, 2011
First Named Inventor	Nicholas S. Bodor
Examiner Name	
Attorney Docket No.	0056192-000067

Sheet 2 of 4

FOREIGN PATENT DOCUMENTS

¹Enter Office that issued the document, by the two-letter code.

	NON-PATENT LITERATURE DOCUMENTS
Examiner Initials	Include name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of the item (book, magazine, journal, serial, symposium, catalog, etc.), date, page(s), volume-issue number(s), publisher, city and/or country where published.
	TARASIUK et al., "Stability of 2-Chloro-2'-Deoxyadenosine at Various pH and Temperature",
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	KARLSSON et al., "Oral cladribine for B-cell chronic lymphocytic leukaemia: report of a phase II trial
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	ALREPTION et al. "On the bioavailability of 2 ablars 2' desveradorsains (CdA)" Fur LOin
	Pharmacol., Vol. 44, pp. 579-582, 1993, Springer-Verlag, Germany
	AHN et al., "Chiral Recognition in Gas-Phase Cyclodextrin: Amino Acid Complexes-Is the Three
	Point Interaction Still Valid in the Gas Phase?", J Am Soc Mass Spectrom, Vol. 12, pp. 278-287,
L	2001, Elsevier Science, Inc., US

Examiner		Date	
Signature		Considered	
*EXAMINER	Initial if reference considered, whether or not citation is in co	nformance with M.P.E.I	P. § 609. Draw line through citation if not in

Substitute for form 1449/PTO & 1449B/PTO	Complete if Known		
FIRST	Application Number		
INFORMATION DISCLOSURE	Filing Date	January 5, 2011	
STATEMENT BY APPLICANT	First Named Inventor	Nicholas S. Bodor	
(use as many sheets as necessary)	Examiner Name		
	Attorney Docket No.	0056192-000067	

Sheet 3 of 4

NON-PATENT LITERATURE DOCUMENTS

Examiner Initials	Include name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of the item (book, magazine, journal, serial, symposium, catalog, etc.), date, page(s), volume-issue number(s), publisher, city and/or country where published.
	BAKTHIAR et al., "A study of the complexation between dimethyl-β-cyclodextrin and steroid hormones using electrospray ionization mass spectrometry". <i>Rapid Communications in Mass</i>
	Spectrometry, Vol. 11, pp. 1478-1481, 1997, John Wiley And Sons Ltd, England
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	n 1743-1753 1998 American Chemical Society US
	UEKAMA et al. "Cyclodextrin Drug Carrier Systems" Chem Rev. Vol. 98 pp. 2045-2076, 1998
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Examiner Signature		Date Considered			
*EXAMINER: Initial if reference considered, whether or not citation is in conformance with M.P.E.P. § 609. Draw line through citation if not in conformance and not considered. Include copy of this form with next communication to Applicant.					
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Substitute for form 1449/PTO & 1449B/PTO		Complete if Known	
FIRST	Application Number		
INFORMATION DISCLOSURE	Filing Date	January 5, 2011	
STATEMENT BY APPLICANT	First Named Inventor	Nicholas S. Bodor	-
(use as many sheets as necessary)	Examiner Name		
	Attorney Docket No.	0056192-000067	

Sheet 4 of 4

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<u>OTHER</u>
International Search Report dated October 12, 2004 for PCT/US2004/009387, filed March 26, 2004
PCT International Preliminary Report on Patentability and Written Opinion for International Application No. PCT/US2004/009387, International Filing date March 26, 2004.

PATENT COOPERATION TREATY

PCT

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY

(Chapter I of the Patent Cooperation Treaty)

(PCT Rule 44bis)

Applicant's or agent's file reference 033935-008	FOR FURTHER ACTION	See item 4 below		
International application No. PCT/US2004/009387	International filing date (day/month/year) 26 March 2004 (26.03.2004)	Priority date (<i>day/month/year</i>) 28 March 2003 (28.03.2003)]		
International Patent Classification (IPC) or national classification and IPC 7 A61K 9/20, 47/48, 31/52				
Applicant IVAX CORPORATION				

- 1. This international preliminary report on patentability (Chapter I) is issued by the International Bureau on behalf of the International Searching Authority under Rule 44 *bis*.1(a).
- 2. This REPORT consists of a total of 8 sheets, including this cover sheet.

In the attached sheets, any reference to the written opinion of the International Searching Authority should be read as a reference to the international preliminary report on patentability (Chapter I) instead.

3. This report contains indications relating to the following items:

Box No. I	Basis of the report
Box No. II	Priority
Box No. III	Non-establishment of opinion with regard to novelty, inventive step and industrial applicability
Box No. IV	Lack of unity of invention
Box No. V	Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement
Box No. VI	Certain documents cited
Box No. VII	Certain defects in the international application
Box No. VIII	Certain observations on the international application

4. The International Bureau will communicate this report to designated Offices in accordance with Rules 44*bis*.3(c) and 93*bis*.1 but not, except where the applicant makes an express request under Article 23(2), before the expiration of 30 months from the priority date (Rule 44*bis*.2).

	Date of issuance of this report 01 October 2005 (01.10.2005)		
The International Bureau of WIPO 34, chemin des Colombettes 1211 Geneva 20, Switzerland	Authorized officer Beate Giffo-Schmitt		
Facsimile No. +41 22 740 14 35	Telephone No. +41 22 338 87 20		

Form PCT/IB/373 (January 2004)

PATENT COOPERATION TREATY

To:					POT DEC 2004
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see form PCT/ISA/220		WRITTEN OPINION OF THE			
				()	PCT Rule 43 <i>bis.</i> 1)
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nteri PCT	national application N NUS2004/009387	Vo. 7	International filing date (26.03.2004	i day/month/year)	Priority date <i>(day/month/year)</i> 28.03.2003
nter	national Patent Class	sification (IPC) or	both national classification	and IPC	
61	K9/20, A61K47/4	8, A61K31/52			
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	This opinion co	ntains indicati	ons relating to the fol	lowing items:	
	Box No. I	Basis of the or	binion		
	Box No. II	Priority			
	Box No. III	Non-establish	ment of opinion with rea	ard to novelty, inventi	ve step and industrial applicability
		Lack of unity of	of invention	ard to novery, inventi-	to stop and matsing approaching
Box No. V Reasoned statement under Rule 4 applicability; citations and explana				s.1(a)(i) with regard to	novelty, inventive step or industrial
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		Certain defect	e in the international an	plication	
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	If a demand for i written opinion o the applicant cho International Bur will not be so cor	nternational pre f the Internation poses an Author eau under Rule nsidered.	Ilminary examination is al Preliminary Examinin ity other than this one to 66.1 <i>bis</i> (b) that written o	made, this opinion will ig Authority ("IPEA"). I o be the IPEA and the opinions of this Interna	I usually be considered to be a -lowever, this does not apply where chosen IPEA has notifed the ational Searching Authority
	If this opinion is, submit to the IPE months from the whichever expire	as provided abo A a written rep date of malling as later.	ove, considered to be a ly together, where appro of Form PCT/ISA/220 o	written opinion of the opriate, with amendme r before the expiration	IPEA, the applicant is invited to onts, before the expiration of three of 22 months from the priority date,
	For further option	ns, see Form Po	CT/ISA/220.		
	For further detail	s, see notes to	Form PCT/ISA/220.		
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3. Nam	e and mailing addres	ss of the ISA:		Authorized Officer	13 Minister Palaana
3. Nam	e and mailling addres	ss of the ISA: Patent Office Junich		Authorized Officer Toulacis, C	

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Box No. I Basis of the opinion

- 1. With regard to the **language**, this opinion has been established on the basis of the international application in the language in which it was field, unless otherwise indicated under this item.
 - This opinion has been established on the basis of a translation from the original language into the following language , which is the language of a translation furnished for the purposes of international search (under Rules 12.3 and 23.1(b)).
- 2. With regard to any **nucleotide and/or amino acid sequence** disclosed in the international application and necessary to the claimed invention, this opinion has been established on the basis of:
 - a. type of material:

- □ a sequence listing
- \Box table(s) related to the sequence listing
- b. format of material:
 - in written format
 - in computer readable form
- c. time of filing/furnishing:
 - □ contained in the international application as filed.
 - filed together with the international application in computer readable form.
 - furnished subsequently to this Authority for the purposes of search.
- 3. In addition, in the case that more than one version or copy of a sequence listing and/or table relating theretc has been filed or furnished, the required statements that the information in the subsequent or additional copies is identical to that in the application as filed or does not go beyond the application as filed, as appropriate, were furnished.
- 4. Additional comments:

WRITTEN OPINION OF THE INTERNATIONAL SEARCHING AUTHORITY

International application No. PCT/US2004/009387

Box No. II Priority

1. Image: The following document has not been furnished:

copy of the earlier application whose priority has been claimed (Rule 43*bis*.1 and 66.7(a)).

□ translation of the earlier application whose priority has been claimed (Rule 43*bis*.1 and 66.7(b)).

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Consequently it has not been possible to consider the validity of the priority claim. This opinion has nevertheless been established on the assumption that the relevant date is the claimed priority date.

2. This opinion has been established as if no priority had been claimed due to the fact that the priority claim has been found invalid (Rules 43*bis*.1 and 64.1). Thus for the purposes of this opinion, the international filing date indicated above is considered to be the relevant date.

3. Additional observations, if necessary:

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WRITTEN OPINION OF THE INTERNATIONAL SEARCHING AUTHORITY

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Box No. III Non-establishment of opinion with regard to novelty, inventive step and industrial applicability					
The	e questions whether the claimed vious), or to be industrially applic	l inve able	ntion appears to be novel, to involve an inventive step (to be non have not been examined in respect of:		
	the entire international applicat	tion,			
\boxtimes	claims Nos. 13-35 with regard to industrial applicability				
bed	cause:				
⊠	the said international application, or the said claims Nos. 13-35 relate to the following subject matter which does not require an international preliminary examination (<i>specify</i>):				
	see separate sheet		· .		
	the description, claims or drawings <i>(indicate particular elements below)</i> or said claims Nos. are so unclear that no meaningful opinion could be formed <i>(specify)</i> :				
	the claims, or said claims Nos. are so inadequately supported by the description that no meaningful opinion could be formed.				
\boxtimes	no international search report has been established for the whole application or for said claims Nos.				
	the nucleotide and/or amino acid sequence listing does not comply with the standard provided for in Annex C of the Administrative Instructions in that:				
	the written form		has not been furnished		
			does not comply with the standard		
	the computer readable form		has not been furnished		
			does not comply with the standard		
	the tables related to the nucleo not comply with the technical r	otide requir	and/or amino acid sequence listing, if in computer readable form only, dc ements provided for in Annex C- <i>bis</i> of the Administrative Instructions.		

□ See separate sheet for further details

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Box No. V Reasoned statement under Rule 43*bis*.1(a)(i) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

1. Statement

J

Novelty (N)	Yes: No:	Claims Claims	1-98
Inventive step (IS)	Yes: No:	Claims Claims	1-98
Industrial applicability (IA)	Yes: No:	Claims Claims	1-12, 36-98

- 2. Citations and explanations
- see separate sheet
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Claims 13-35 relate to subject-matter considered by this Authority to be covered by the provisions of Rule 67.1(iv) PCT. Consequently, no opinion will be formulated with respect to industrial applicability of the subject-matter of these claims (Article 34(4)(a)(I) PCT).

V

For the assessment of the present claims 13-35 on the question whether they are industrially applicable, no unified criteria exist in the PCT Contracting States. The patentability can also be dependent upon the formulation of the claims. The EPO, for example, does not recognize as industrially applicable the subject-matter of claims to the use of a compound in medical treatment, but may allow, however, claims to a known compound for first use in medical treatment and the use of such a compound for the manufacture of a medicament for a new medical treatment.

- (N) A cladribine-cyclodextrin complex which is an intimate <u>amorphous admixture of</u>: a) an <u>amorphous inclusion complex</u> of cladribine with an amorphous cyclodextrin and b) <u>amorphous cyclodextrin as non-inclusion complex</u> (claim 58), is not disclosed in the documents cited in the search report. The same applies to a pharmaceutical composition comprising said cladribinecyclodextrin complex (claim 1), to a method for enhancing the oral bioavailability of cladribine comprising oral administration of said complex (claim 13), to a method for the treatment according to claim 25 comprising oral administration of said complex, to the use of said cladribine-cyclodextrin complex in the formulation of a solid oral dosage form (claims 36 and 47), to a process for the preparation of said complex (claim 67), and to a pharmaceutical composition being obtainable by a process according to claim 67 (claim 82).
- (IS) The object of the present application is to provide an oral pharmaceutical composition showing improved <u>bioavailability</u> of cladribine and <u>interpatient</u> <u>variation</u>, <u>when administered orally</u> (description, page 4, lines 22-26). This object has been achieved, by formulating a cladribine-cyclodextrin complex as defined in claim 58, i.e. is an intimate amorphous admixture of a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and b) amorphous cyclodextrin as non-inclusion complex, into a solid oral dosage form (description; page 33, example 4; page 35, table V; page 36, table VI; example 5, tables VII-X).

WRITTEN OPINION OF THE INTERNATIONAL SEARCHING AUTHORITY (SEPARATE SHEET)

This is not suggested by document US-B-6 194 395 (D1), which only discloses as an oral dosage form of cladribine, a milled extrudate comprising 1-15 mg cladribine and 100-500 mg cyclodextrin without giving an exact ratio and being silent about amorphous forms (D1; column 6).

(IA) The industrial applicability of claims 1-12 and 36-98 is beyond any doubt.

Form PCT/Separate Sheet/237 (Sheet 2) (EPO-January 2004)

Electronic Patent Application Fee Transmittal						
Application Number:						
Filing Date:						
Title of Invention:	ORAL FORMULATIONS OF CLADRIBINE					
First Named Inventor/Applicant Name:	Nicholas S. Bodor					
Filer:	Mary Katherine Baumeister/Diana Francis					
Attorney Docket Number:	00	56192-000067				
Filed as Large Entity						
Utility under 35 USC 111(a) Filing Fees						
Description		Fee Code	Quantity	Amount	Sub-Total in USD(\$)	
Basic Filing:			<u> </u>			
Utility application filing		1011	1	330	330	
Utility Search Fee		1111	1	540	540	
Utility Examination Fee		1311	1	220	220	
Pages:						
Claims:						
Claims in excess of 20		1202	44	52	2288	
Independent claims in excess of 3		1201	1	220	220	
Miscellaneous-Filing:						

Description	Fee Code	Quantity	Amount	Sub-Total in USD(\$)
Petition:				
Patent-Appeals-and-Interference:				
Post-Allowance-and-Post-Issuance:				
Extension-of-Time:				
Miscellaneous:				
	Tot	al in USD	(\$)	3598

Electronic Acknowledgement Receipt				
EFS ID:	9185944			
Application Number:	12986310			
International Application Number:				
Confirmation Number:	6100			
Title of Invention:	ORAL FORMULATIONS OF CLADRIBINE			
First Named Inventor/Applicant Name:	Nicholas S. Bodor			
Customer Number:	21839			
Filer:	Mary Katherine Baumeister/Diana Francis			
Filer Authorized By:	Mary Katherine Baumeister			
Attorney Docket Number:	0056192-000067			
Receipt Date:	07-JAN-2011			
Filing Date:				
Time Stamp:	12:04:57			
Application Type:	Utility under 35 USC 111(a)			

Payment information:

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Payment Type	e	Credit Card	Credit Card						
Payment was	successfully received in RAM	\$3598	\$3598						
RAM confirma	ation Number	8455	8455						
Deposit Acco	unt								
Authorized U	ser								
File Listing:									
Document Number	Document Description	75 ile Name	File Size(Bytes)/ Message Digest	Multi Part /.zip	Pages (if appl.)				

1	Transmittal of New Application	005619267CONT.pdf	133899	no	3			
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Warnings:								
Information	:		1		1			
2	Application Data Sheet	005619267ADS.pdf	88641	no	5			
Warnings:								
Information								
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3	Oath or Declaration filed	0056192670ATH.pdf	111313	no	2			
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Warnings:								
Information	:		1					
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Multipart Description/PDF files in .zip description								
	Document Des	scription	Start	E	nd			
	Specificati	ion	1	39				
	Claims		40		48			
	Abstrac	t	49		49			
	Drawings-only black and v	white line drawings	50		50			
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5	Transmittal Letter	005619267IDS.pdf	96019	no	2			
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6	Information Disclosure Statement (IDS)	005619267Fm1449.pdf	392829	no				
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8	NPL Documents	IPER.pdf	330298	no	8	
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Warnings:						
Information:						
9	Fee Worksheet (PTO-875)	fee-info.pdf	37904	no	2	
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Information:			1			
		Total Files Size (in bytes)	37	16294		
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National Stage of an International Application under 35 U.S.C. 371 If a timely submission to enter the national stage of an international application is compliant with the conditions of 35 U.S.C. 371 and other applicable requirements a Form PCT/DO/EO/903 indicating acceptance of the application as a national stage submission under 35 U.S.C. 371 will be issued in addition to the Filing Receipt, in due course. <u>New International Application Filed with the USPTO as a Receiving Office</u> If a new international application is being filed and the international application includes the necessary components for an international filing date (see PCT Article 11 and MPEP 1810), a Notification of the International Application Number and of the International Filing Date (Form PCT/RO/105) will be issued in due course, subject to prescriptions concerning national security, and the date shown on this Acknowledgement Receipt will establish the international filing date of the application.						

PATENT APPLICATION FEE DETERMINATION RECORD Substitute for Form PTO-875								Application or Docket Number 12/986,310			
	APP		S FILE	D - PART I	lumn 2)		SMALL	ENTITY	OR	OTHEF SMALL	THAN ENTITY
	FOR	NUMBE	R FILE		ER EXTRA		RATE(\$)	FEE(\$)		RATE(\$)	FEE(\$)
BAS (37 C	SIC FEE FR 1.16(a), (b), or (c))	N	/A	1	N/A		N/A			N/A	330
SEA (37 C	RCH FEE FR 1.16(k), (i), or (m))	N	/A	1	N/A		N/A			N/A	540
EXA (37 C	MINATION FEE FR 1.16(0), (p), or (q))	N	/A	1	N/A		N/A			N/A	220
TO1 (37 C	AL CLAIMS FR 1.16(i))	64	minus	20= *	44				OR	× 52 =	2288
IND (37 C	EPENDENT CLAI FR 1.16(h))	MS 4	minus	3 = *	1				1	× 220 =	220
API FEI (37	APPLICATION SIZE FEE (37 CFR 1.16(s)) If the specification and drawings exceed 100 sheets of paper, the application size fee due is \$270 (\$135 for small entity) for each additional 50 sheets or fraction thereof. See 35 U.S.C. 41(a)(1)(G) and 37 CFR 1.16(s).							0.00			
MUI	_TIPLE DEPEND	ENT CLAIM PRE	SENT (3	7 CFR 1.16(j))					1		0.00
*lft	he difference in co	olumn 1 is less th	an zero,	enter "0" in colu	mn 2.		TOTAL			TOTAL	3598
	APPLICATION AS AMENDED - PART II OTHER THAN										
		CLAIMS		HIGHEST	(Column 3)		OWIALL		1		
NT A		REMAINING AFTER AMENDMENT		NUMBER PREVIOUSLY PAID FOR	PRESENT EXTRA		RATE(\$)	ADDITIONAL FEE(\$)		RATE(\$)	ADDITIONAL FEE(\$)
μ	Total (37 CFR 1.16(i))	*	Minus	**	=		X =		OR	x =	
END	Independent (37 CFR 1.16(h))	*	Minus	***	=		x =		OR	x =	
AM	Application Size Fe	ee (37 CFR 1.16(s))			•]		
	FIRST PRESENT	ATION OF MULTIPL	E DEPEN	IDENT CLAIM (37 (CFR 1.16(j))				OR		
							TOTAL ADD'L FEE		OR	TOTAL ADD'L FEE	
		(Column 1)		(Column 2)	(Column 3)						
NT B		CLAIMS REMAINING AFTER AMENDMENT		HIGHEST NUMBER PREVIOUSLY PAID FOR	PRESENT EXTRA		RATE(\$)	ADDITIONAL FEE(\$)		RATE(\$)	ADDITIONAL FEE(\$)
Σ	Total (37 CFR 1.16(i))	*	Minus	**	=		X =		OR	x =	
	Independent (37 CFR 1.16(h))	*	Minus	***	=		x =		OR	x =	
AM	Application Size Fe	ee (37 CFR 1.16(s))							1		
	FIRST PRESENT	TION OF MULTIPL	E DEPEN	IDENT CLAIM (37 (CFR 1.16(j))				OR		
						. 1	TOTAL ADD'L FEE		OR	TOTAL ADD'L FEE	
k Krik	 If the entry in cc If the "Highest N If the "Highest Nu The "Highest Num 	olumn 1 is less th Jumber Previous umber Previously I ber Previously Paid	an the er y Paid F Paid For" For" (Tota	ntry in column 2, or" IN THIS SPA IN THIS SPACE i al or Independent) is	write "0" in colu CE is less than s less than 3, er s the highest foun	umr n 20 nter nd in	n 3. I, enter "20". "3". the appropriate box	in column 1.	-		

	United State	<u>'s Patent</u>	and Tradem	UNITED STATES D United States Pater Address: COMMISSIONE PO. Box 1450 Alexandra, Virgini www.uspto.gov	EPARTMENT OF C at and Trademark C ER FOR PATENTS a 22313-1450	OMMERCE Office
APPLICATION NUMBER	FILING or 371(c) DATE	GRP ART UNIT	FIL FEE REC'D	ATTY.DOCKET.NO	TOT CLAIMS	IND CLAIMS
12/986,310	01/07/2011	1623	3598	0056192-000067	64	4
				CO	NFIRMATION	NO. 6100
21839				FILING RECE	EIPT	
BUCHANAN, INGERSOLL & ROONEY PC POST OFFICE BOX 1404 ALEXANDRIA, VA 22313-1404					00000045534863	

Date Mailed: 01/20/2011

Receipt is acknowledged of this non-provisional patent application. The application will be taken up for examination in due course. Applicant will be notified as to the results of the examination. Any correspondence concerning the application must include the following identification information: the U.S. APPLICATION NUMBER, FILING DATE, NAME OF APPLICANT, and TITLE OF INVENTION. Fees transmitted by check or draft are subject to collection. Please verify the accuracy of the data presented on this receipt. If an error is noted on this Filing Receipt, please submit a written request for a Filing Receipt Correction. Please provide a copy of this Filing Receipt with the changes noted thereon. If you received a "Notice to File Missing Parts" for this application, please submit any corrections to this Filing Receipt with your reply to the Notice. When the USPTO processes the reply to the Notice, the USPTO will generate another Filing Receipt incorporating the requested corrections

Applicant(s)

Nicholas S. Bodor, Bal Harbour, FL; Yogesh Dandiker, Toronto, CANADA;

Assignment For Published Patent Application

ARES TRADING S.A., Aubonne, SWITZERLAND **Power of Attorney:** The patent practitioners associated with Customer Number <u>21839</u>

Domestic Priority data as claimed by applicant

This application is a CON of 10/551,205 11/14/2006 which is a 371 of PCT/US2004/009387 03/26/2004 which claims benefit of 60/458,922 03/28/2003 and claims benefit of 60/484,756 07/02/2003 and claims benefit of 60/541,247 02/04/2004

Foreign Applications (You may be eligible to benefit from the **Patent Prosecution Highway** program at the USPTO. Please see <u>http://www.uspto.gov</u> for more information.)

If Required, Foreign Filing License Granted: 01/18/2011

The country code and number of your priority application, to be used for filing abroad under the Paris Convention, is **US 12/986,310**

Projected Publication Date: 04/28/2011

Non-Publication Request: No

Early Publication Request: No

Title

ORAL FORMULATIONS OF CLADRIBINE

Preliminary Class

514

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Almost every country has its own patent law, and a person desiring a patent in a particular country must make an application for patent in that country in accordance with its particular laws. Since the laws of many countries differ in various respects from the patent law of the United States, applicants are advised to seek guidance from specific foreign countries to ensure that patent rights are not lost prematurely.

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Applicants may wish to consult the USPTO booklet, "General Information Concerning Patents" (specifically, the section entitled "Treaties and Foreign Patents") for more information on timeframes and deadlines for filing foreign patent applications. The guide is available either by contacting the USPTO Contact Center at 800-786-9199, or it can be viewed on the USPTO website at http://www.uspto.gov/web/offices/pac/doc/general/index.html.

For information on preventing theft of your intellectual property (patents, trademarks and copyrights), you may wish to consult the U.S. Government website, http://www.stopfakes.gov. Part of a Department of Commerce initiative, this website includes self-help "toolkits" giving innovators guidance on how to protect intellectual property in specific countries such as China, Korea and Mexico. For questions regarding patent enforcement issues, applicants may call the U.S. Government hotline at 1-866-999-HALT (1-866-999-4158).

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page 2 of 3

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APPLICATION NUMBER	FILING OR 371(C) DATE	FIRST NAMED APPLICANT	ATTY. DOCKET NO./TITLE
12/986,310	01/07/2011	Nicholas S. Bodor	0056192-000067
			CONFIRMATION NO. 6100
21839		PUBLICA	TION NOTICE
BUCHANAN, INGERSOL POST OFFICE BOX 1404 ALEXANDRIA, VA 22313	L & ROONEY PC - -1404		OC000000047400536*

Title:ORAL FORMULATIONS OF CLADRIBINE

Publication No.US-2011-0097306-A1 Publication Date:04/28/2011

NOTICE OF PUBLICATION OF APPLICATION

The above-identified application will be electronically published as a patent application publication pursuant to 37 CFR 1.211, et seq. The patent application publication number and publication date are set forth above.

The publication may be accessed through the USPTO's publically available Searchable Databases via the Internet at www.uspto.gov. The direct link to access the publication is currently http://www.uspto.gov/patft/.

The publication process established by the Office does not provide for mailing a copy of the publication to applicant. A copy of the publication may be obtained from the Office upon payment of the appropriate fee set forth in 37 CFR 1.19(a)(1). Orders for copies of patent application publications are handled by the USPTO's Office of Public Records. The Office of Public Records can be reached by telephone at (703) 308-9726 or (800) 972-6382, by facsimile at (703) 305-8759, by mail addressed to the United States Patent and Trademark Office, Office of Public Records, Alexandria, VA 22313-1450 or via the Internet.

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Office of Data Managment, Application Assistance Unit (571) 272-4000, or (571) 272-4200, or 1-888-786-0101

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OR Practitioner(s) named	below (if more than ten patent	practitioners are to	be named, then a cu	ustomer number must	be used):
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Signature	Tulm	Via Ka	Huga	Date 10.04	.2012
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STATEMENT UNDER 37 CFR 3.73(b)	
Applicant/Patent Owner: Ares Trading S.A.	
Application No./Patent No.: 12/986,310 Filed/Issue Date: 01/0	7/2011
Titled: ORAL FORMULATIONS OF CLADRIBINE	
Ares Trading S.A. , a CORPORATION	
(Name of Assignee) (Type of Assignee, e.g., corporation, partr	nership, university, government agency, etc.
states that it is:	
1. X the assignee of the entire right, title, and interest in;	
2. an assignee of less than the entire right, title, and interest in (The extent (by percentage) of its ownership interest is%); or	
3. the assignee of an undivided interest in the entirety of (a complete assignment from a	one of the joint inventors was made)
the patent application/patent identified above, by virtue of either:	
A. An assignment from the inventor(s) of the patent application/patent identified above. the United States Patent and Trademark Office at Reel, Frame	The assignment was recorded in, or for which a
OR	
B. X A chain of title from the inventor(s), of the patent application/patent identified above,	to the current assignee as follows:
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As required by 37 CFR 3.73(b)(1)(i), the documentary evidence of the chain of title from or concurrently is being, submitted for recordation pursuant to 37 CFR 3.11.	the original owner to the assignee was,
[NOTE: A separate copy (<i>i.e.</i> , a true copy of the original assignment document(s)) must accordance with 37 CFR Part 3, to record the assignment in the records of the USPTO. <u>S</u>	be submitted to Assignment Division in See MPEP 302.08]
The undersigned (whose title is supplied below) is authorized to act on behalf of the assignee.	
/Mary Katherine Baumeister/	JULY 30, 2012
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Mary Katherine Baumeister, Reg. No. 26254	Agent for Applicant(s)
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EFS ID:	13371700			
Application Number:	12986310			
International Application Number:				
Confirmation Number:	6100			
Title of Invention:	ORAL FORMULATIONS OF CLADRIBINE			
First Named Inventor/Applicant Name:	Nicholas S. Bodor			
Customer Number:	21839			
Filer:	Martin A. Bruehs/Louie Malloy			
Filer Authorized By:	Martin A. Bruehs			
Attorney Docket Number:	0056192-000067			
Receipt Date:	30-JUL-2012			
Filing Date:	07-JAN-2011			
Time Stamp:	15:17:32			
Application Type:	Utility under 35 USC 111(a)			

Payment information:

Submitted with Payment no						
File Listing	g:					
Document Number	Document Description		File Name	File Size(Bytes)/ Message Digest	Multi Part /.zip	Pages (if appl.)
1	Power of Attorney		AresTradingSA pdf	72559	no	1
·	Tower of Actorney		Ales Hadings Apar	6fb92021a1041a66c651e9a9487e28c9768 4de7b		
Warnings:						
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Assignee showing of ownership per 37		0056192-0067.pdf	30069	no	1
	CFR 3.73(b).		3d732cb3cdfcc2c4058981d4bf267b0451e 28d0f		
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APPLICATION NUMBER	FILING OR 371(C) DATE	FIRST NAMED APPLICANT	ATTY. DOCKET NO./TITLE
12/986,310	01/07/2011	Nicholas S. Bodor	0056192-000067
13974 SNR DENTON US LLP P.O. BOX 061080 Chicago, IL 60606-1080			CONFIRMATION NO. 6100 EPTANCE LETTER

NOTICE OF ACCEPTANCE OF POWER OF ATTORNEY

This is in response to the Power of Attorney filed 07/30/2012.

The Power of Attorney in this application is accepted. Correspondence in this application will be mailed to the above address as provided by 37 CFR 1.33.

/snguyen/

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Application Number		12986310
Filing Date		2011-01-07
First Named Inventor	Nicho	las S. BODOR
Art Unit		1623
Examiner Name Jonat		han S. LAU
Attorney Docket Number		20009904-0067

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	1 5	99/62958	WO		A1	1999-12-09	Janssen Pharmace N.V.	utica	Corresponds to JP 2002-517521	
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	Application Number		12986310	
	Filing Date		2011-01-07	
INFORMATION DISCLOSURE	First Named Inventor Nichola		iolas S. BODOR	
(Not for submission under 37 CFR 1.99)	Art Unit		1623	
	Examiner Name	Jonat	han S. LAU	
	Attorney Docket Number		20009904-0067	

Examiner Initials*	Cite No	Inclu (bool publi	nclude name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of the item pook, magazine, journal, serial, symposium, catalog, etc), date, pages(s), volume-issue number(s), ublisher, city and/or country where published.					
	1 Drugs in Japan (Nihon lyakuhinn Shu), 2004 edition, JIHOU Inc. March 1, 2003, pp. 651-654, "Cladrivine" (especially, "Composition") - relevance explained in January 10, 2012, Japanese Official Action							
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INFORMATION DISCLOSURE	Application Number		12986310	
	Filing Date		2011-01-07	
	First Named Inventor Nichola		olas S. BODOR	
STATEMENT BY APPLICANT (Not for submission under 37 CER 1 99)	Art Unit		1623	
	Examiner Name	Jonat	han S. LAU	
	Attorney Docket Number		20009904-0067	

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Please see 37 CFR 1.97 and 1.98 to make the appropriate selection(s):

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The fee set forth in 37 CFR 1.17 (p) has been submitted herewith.

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Signature	/Mary Katherine Baumeister/	Date (YYYY-MM-DD)	2012-08-09
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Bibliographic data: WO9962958 (A1) --- 1999-12-09

ACYLATED ALKYLATED CYCLODEXTRIN DERIVATIVES AND THEIR USE AS CARRIERS FOR MEDICAMENTS

Inventor(s):	UEKAMA KANETO [JP]; HIRAYAMA FUMITOSHI [JP]; KONDO AKIRA [JP]; KAWAJI HIROSHI [JP]; OHTA MASAAKI [JP]; OKAMOTO YASUHIRO [JP] <u>*</u>					
Applicant(s):	JANSSEN PHARMACEUTICA NV [BE]; UEKAMA KANETO [JP]; HIRAYAMA FUMITOSHI [JP]; KONDO AKIRA [JP]; KAWAJI HIROSHI [JP]; OHTA MASAAKI [JP]; OKAMOTO YASUHIRO [JP] <u>*</u>					
	- international:	A61K47/48; C08B37/16; (IPC1- 7): A61K47/48; C08B37/16				
Classification:	- European:	<u>A61K47/48W18B; B82Y5/00;</u> <u>C08B37/00M2B</u>				
Application number:	WO1999JP02806 19990527					
Priority number(s):	JP19980164465 19980529					
Also published as:	JP2002517521 (A	N) EP1084149 (A1) AU3954799 (A)				

Abstract of WO9962958 (A1)

Cyclodextrin derivatives having at least one lower alkyl group and at least one C2-20 alkanoyl group in the molecule, are disclosed pharmaceutical preparations wherein the derivative and a medicament are in such a state that they are closely compounded, are also disclosed. The cyclodextrin derivative having lowered hemolytic activity and its use as a medicament carrier.

Last updated: 14-03-2012 Worldwide Database 5.7.38; 93p



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INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification ⁶ :		(11) International Publication Number: WO 99/62958				
C08B 37/16, A61K 47/48	AI	(43) International Publication Date: 9 December 1999 (09.12.99)				
(21) International Application Number: PCT/JP	Minami–isshiki, Nagaizumi–cho, Sunto–gun, Shizuoka 411–0932 (JP).					
(22) International Filing Date: 27 May 1999 ((30) Priority Data: 10/164465 29 May 1998 (29.05.98)	(74) Agents: ODAJIMA, Heikichi et al.; Odajima Patent Office, Nippon Jitensha Building, 9–15, Akasaka 1–chome, Mi- nato-ku, Tokyo 107–0052 (JP).					
 (71) Applicant (for all designated States except US): J. PHARMACEUTICA N.V. [BE/BE]; Turnhoutse B-2340 Beerse (BE). (72) Inventors; and (75) Inventors/Applicants (for US only): UEKAMA, [JP/JP]; 4-18, Nagaminehigashi 2-chome, Kumar Kumamoto 862-0938 (JP). HIRAYAMA, F [JP/JP]; 4-1-302, Koto 1-chome, Kumar Kumamoto 862-0909 (JP). KONDO, Akira Fuji Laboratory, Janssen-Kyowa Co., Ltd., Minami-isshiki, Nagaizumi-cho, Sunto-gun, 411-0932 (JP). KAWAJI, Hiroshi [JP/JP]; Fuji La Janssen-Kyowa Co., Ltd., 600-8, Minami-isshi gaizumi-cho, Sunto-gun, Shizuoka 411-0932 (JP). Masaaki [JP/JP]; Fuji Laboratory, Janssen-Kyowa Co., Ltd., 600-8, Minami-isshiki, Nagaizumi-cho, Su Shizuoka 411-0932 (JP). OKAMOTO, Yasuhiro Fuji Laboratory, Janssen-Kyowa Co., Ltd., Fuji Laboratory, Janssen-Kyowa Co., Ltd., 600-8, Minami-isshiki, Nagaizumi-cho, Su Shizuoka 411-0932 (JP). 	 (81) Designated States: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG). Published A, With international search report. 					

(54) Title: ACYLATED ALKYLATED CYCLODEXTRIN DERIVATIVES AND THEIR USE AS CARRIERS FOR MEDICAMENTS

(57) Abstract

Cyclodextrin derivatives having at least one lower alkyl group and at least one C2-20 alkanoyl group in the molecule, are disclosed pharmaceutical preparations wherein the derivative and a medicament are in such a state that they are closely compounded, are also disclosed. The cyclodextrin derivative having lowered hemolytic activity and its use as a medicament carrier.



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DESCRIPTION

ACYLATED ALKYLATED CYCLODEXTRIN DERIVATIVES AND THEIR USE AS CARRIERS FOR MEDICAMENTS

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Technical Field

This invention relates to acylated alkylated cyclodextrin derivatives, and a process for preparing the same and use of the same as carriers for medica-10 ments.

Background Art

Cyclodextrin (hereinafter also referred to as CyD) is an oligosaccharide wherein glucose residues are 15 cyclicly bound by α -1,4 bond and composed of 6, 7 or 8 glucose residues, and ones called α , β or γ -CyD are known. Further, so-called branched cyclodextrins (hereinafter also referred to as branched CyD) are also known wherein glucosyl group(s) or maltosyl group(s) is/are 20 α -1,6 bound to one or two of the glucose units of these

20 α-1,6 bound to one or two of the glucose units of these CyDs.

These CyDs and branched CyDs have high inclusion ability on certain chemical substances, and are utilized for various uses such as stabilization of

25 unstable substances, retention of volatile substances and solubilization of water-sparingly soluble or insoluble substances, in the pharmaceutical, food and cosmetic fields.

Further, in order to utilize the physicochemi-30 cal characteristics and inclusion ability of the above CyDs as polyfunctional medicament carriers, etc., various CyD derivatives are provided. However, in the course of these researches, the presence of several problems in physical properties, inclusion ability,

35 intracorporeal kinetics, economical efficiency, etc. has been pointed out, and it has come to be made clear that

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there is some limitation on their use. Particularly, use thereof as medicament carriers, namely dissolution aids (solubilizers) or tissular disorder-preventing agents for injections or various preparations to be applied to the mucosae (e.g., eye drops, suppositories, etc.), on which low hemolytic activity, low actions to give topical irritation and low actions to cause tissular disorder are required, has been extremely inconvenient.

On the other hand, α -CyD has a solubility in water as comparatively high as 14.5 g/100 mL (25°C) and its hemolytic activity and muscular irritation are lower than those of β -CyD, but there is a limitation on α -CyD that the guest compounds of inclusion are limited to

- 15 small molecules. Further, its price is 30 times as high as that of β -CyD and it has a disadvantage point also in an economical aspect. γ -CyD is the best among α -, β and γ -CyD on the aspect of safety such as hemolytic activity and actions to cause tissular disorder and has
- 20 inclusion ability equal to that of β -CyD, but its price is about 100 times as high as that of β -CyD and therefore it has not so been utilized from economical reason. Further, glucosylated or maltosylated branched CyDs rouse interest partially because their solubilities in
- 25 water are increased compared with the corresponding unbranched CyDs, but they are not always satisfactory in behavior as carriers for the above medicaments.

Thus, attempts have been made to improve the physical properties or functionality of β -CyD, which is 30 easy to obtain, by chemically modifying it. For example, there have been obtained thereby heptakis (2,6-di-0-methyl)- β -CyD (hereinafter referred to as DM- β -CyD) wherein the hydroxyl groups at the 2- and 6-positions of the glucose are methylated, heptakis (2,3,6-tri-0-

35 methyl)- β -CyD (hereinafter referred to as TM- β -CyD) wherein all the hydroxyl groups at the 2-, 3- and

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6-positions of the glucose are methylated, 2-hydroxypropyl- β -CyD (hereinafter referred to as HP- β -CyD) wherein a hydroxypropyl group is introduced in the hydroxyl group mainly at the 6-position of the glucose, etc. Szejtili et al. suggest that it is possible to utilize DM- β -CyD as a dissolution aid for injection (J. Incl. Phenom., 1(2), 135 (1983)).

However, this $DM-\beta-CyD$ is extremely easy to dissolve in water and has strong inclusion ability, but 10 has a problem that since its solubility and stability

constant strikinly decrease at the side of high temperatures and the dissociation of the medicament from the medicament inclusion composite becomes easy, the designation of sterilization conditions for the injection is

- 15 hindered. Moreover, $DM-\beta$ -CyD has a stronger hemolytic activity than β -CyD, and its action to cause tissular disorder at the time of intramusclular injection is also larger than β -CyD. This tendency is the same in TM- β -CyD, and TM- β -CyD shows intermediate values between
- 20 DM- β -CyD and β -CyD. On the other hand, as to HP- β -CyD, large improvement is made on the lowering of solubility and the lowering of stability constant at high temperatures, and actions to cause tissular disorder such as hemolytic activity and muscular irritation are also
- 25 considerably improved compared with β -CyD, but they are equal to those of α -CyD, and it is the state of things that HP- β -CyD is far inferior to γ -CyD which has the lowest hemolytic activity and muscular irritation among natural CyDs.
- 30 Uwagama et al. disclose a pharmaceutical preparation wherein 2-hydroxyethyl-CyD (hereinafter referred to as HE- β -CyD) wherein a 2-hydroxyethyl group is introduced or 2,3-dihydroxypropyl-CyD (hereinafter referred to as DHP- β -CyD) is used as a carrier for
- 35 medicaments utilizing its low hemolytic activity or action to inhibit hemolytic activity, low action to

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cause tissular disorder or action to prevent tissular disorder, low action to give topical irritation or action to lower topical irritation (Japanese Laid-open Patent Publication No. 61430/1989). However, the

5 hemolysis-inhibiting action of HE- β -CyD and DHP- β -CyD is almost equal to that of γ -CyD, and further improvement will be desired to provide them for clinical use.

Szejtili et al. propose (carboxy)alkyloxyalkyl derivatives of CyD and a pharmaceutical composition 10 comprising such a derivative and a medicament (WO92/ 14762). However, there is no specific description on whether they show a sufficient hemolytic activity-inhibiting action or not.

15 Disclosure of Invention

Under such a situation, it becomes very important to provide a low hemolytic medicament carrier. Namely, this is because if a low hemolytic medicament carrier, which makes it possible to administer sparingly

- 20 soluble medicaments parenterally, can be provided, it can be expected, also for such a medicament that it has been thought to be impossible to apply it, to maintain its high concentration in the blood, and it is thought to make great contribution to the field of pharmaco-
- 25 therapy. Therefore, the objects of the invention lie in providing CyD derivatives satisfying the above needs, and providing the actual use of such a CyD derivative as a carrier or delivery tool for sparingly soluble medicaments.
- 30 For solving the above problems, the present inventors have synthesized various CyD derivatives, and examined their hemolysis-inhibiting action. As a result, they found that CyDs having an acyl group and an alkyl group together in the molecule are CyD derivatives
- 35 having a hemolytic activity that is significantly lower even compared with HE- β -CyD and γ -CyD whose hemolytic

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activities have hitherto been recognized to be low. It was further recognized that these derivatives sufficiently retain the medicament inclusion ability of the corresponding CyDs.

Therefore, according to the invention are provided acylated alkylated CyDs useful as solubilizers, adsorbents or agents having inclusion ability.

The acylated alkylated CyD according to the invention is, specifically, an acylated alkylated cyclo-10 dextrin derivative represented by the formula (I)



wherein n is any of integers 6, 7 and 8, and the 1-position and the 4-position of the sugar residues at the both ends are mutually bound by a covalent bond,

 R^1 , R^2 and R^3 independently represent hydrogen atoms, lower alkyl groups or C_{2-20} alkanoyl groups, or in some case, represent glucosyl groups or maltosyl groups whose hydroxyl group(s) may be replaced with lower alkyloxy group(s) or C_{2-20} alkanyloxy group(s),

provided that any of R^1 , R^2 and R^3 of the number of total $3 \times n$ composed of each n are simultaneously at least one lower alkyl group and at least one acyl group, and the residual groups, when exist, are hydrogen atoms or the glucosyl groups or maltosyl groups of the number of up to at most 2. There is a case where such derivatives are

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provided as compounds wherein the degree of substitution of the acyl group and/or alkyl group is different or part thereof are epimerized, depending on starting materials, reaction conditions, etc. for preparing them, 5 or there is also a case where it is convenient to provide them as a form of a mixture. Moreover, the acylated alkylated CyD derivatives sufficiently meet the objects of the invention, even in the form of mixtures, and thus such mixtures are also provided by the inven-10 tion.

The acylated alkylated CyD derivatives or mixtures of two or more of the derivatives can efficiently be prepared by acylation reaction using corresponding partially alkylated CyD derivatives as starting 15 materials. Thus, such a process for preparing an acy-

lated alkylated CyD is also provided by the invention. The acylated alkylated CyD derivatives or mixtures of two or more of the derivatives, even if they are derived from β -CyD, not only show hemolytic activi-20 ties significantly lower compared with HE- β -CyD and

 γ -CyD which have been recognized to have low hemolytic activity, but sufficiently retain the inclusion ability on medicaments which parent β -CyD inherently has. Moreover, rabbit muscular irritation of the acylated 25 alkylated CyD derivatives is much weaker than that of DM- β -CyD.

Thus, according to the invention is provided the above acylated alkylated CyD derivatives or use of the derivatives as carriers or delivery tools for water

- 30 soluble, sparingly water soluble or water insoluble medicaments. As a specific embodiment of this use is provided a pharmaceutical preparation which comprises such an acylated alkylated cyclodextrin derivative or mixture of two or more of the derivatives and such a
- 35 medicament in such a state that they are closely compounded. A process for preparing such a pharmaceutical

preparation is also provided.

Brief Description of Drawings

Fig. 1 is the mass spectrum (matrix: metha-5 nol, glycerol and m-nitrobenzyl alcohol, which is the same hereinafter) of DMA-β-CyD obtained in Example 1.

Fig. 2 is the ¹H-NMR spectrum of DMA- β -CyD obtained in Example 1.

Fig. 3 is the ¹H-NMR spectrum of DMA4- β -CyD 10 obtained in Example 2.

Fig. 4 is the ¹H-NMR spectrum of butyrated $DM-\beta-CyD$ obtained in Example 3.

Fig. 5 is the ¹H-NMR spectrum of octanoylated $DM-\beta-CyD$ obtained in Example 4.

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Fig. 6 is a drawing showing the results of the hemolytic activity test on various CyD derivatives. In figure 6, the white square (\Box) , black triangle (\blacktriangle) , white triangle (\bigtriangleup) , white circle (\bigcirc) , black circle (\bigcirc) , white inverted triangle (\bigtriangledown) , and white diamond

- 20 (\diamond) represent DMA- β -CyD, DMA4- β -CyD, β -CyD, DM- β -CyD, TM- β -CyD, 2-HP- β -CyD with a degree of substitution (D.S.) of 4.8 and sulfobutyl ether β -CyD with a D.S. of 3.5, respectively.
- Fig. 7 is a graph showing the released amounts 25 of cholesterol from the intact erythrocytes at the time when various CyD derivatives are contacted with erythrocytes. The vertical axis represents the released amount (%) of cholesterol supposing that the amount of cholesterol in all the erythrocytes is 100 %.

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Detailed Description of the Invention

The "acylated alkylated" in the invention means such a state that an acyl group and an alkyl group exist simultaneously on one molecule. Therefore, in the 35 acylated alkylated CyD derivative in the invention, at

least one of the hydroxyl groups in the CyD molecule is

converted to an acyl ester, and at least one of the other hydroxyl groups is converted to an alkyl ether.

Surprisingly, such a CyD derivative which simultaneously has an acyl group and an alkyl group on the CyD molecule has a significantly lower hemolytic activity than the corresponding CyD, as stated above. However, in view of significantly lowering hemolytic activity, in the above formula (I), is preferred a CyD derivative wherein 50 % (e.g, 7 as to β -CyD) or more of R¹ and R³ of the number of total 2×n (e.g, 14 as to β -CyD) are lower alkyl groups, the residual R¹ and R³ and R² are at least one acyl group, and the residual R¹, R² and R³, when exist, are hydrogen atoms, or a mixture of two or more of the derivatives. As a further pre-

15 ferred one, there can be mentioned a CyD derivative wherein 50 % or more, particularly about 100% of R¹ and R³ of the number of total $2 \times n$ are lower alkyl groups, and 50 % or more, particularly about 100% of R² of the number of n are C₂₋₂₀ alkanoyl groups, or a mixture of 20 two or more of the derivatives.

When harmony between the inclusion ability of the medicament and economical efficiency is taken into account, preferred is one which corresponds to n = 7 in the formula (I), i.e. β -CyD, and does not have a gluco-

- 25 syl group or maltosyl group as a branched sugar residue. Thus, as a still further preferred acylated alkylated derivative, there cas be mentioned a β -CyD derivative wherein 7 or more of 14 R¹s and R³s are lower alkyl groups and 4 or more of 7 R²s are C₂₋₂₀ alkanoyl groups,
- 30 or a mixture of two or more of the derivatives. As the mixture, there can, for example, be mentioned a mixture of two or more of compounds selected from the group consisting of compounds wherein 7 to 14 of all the R^1 s and R^3 s are alkyl groups. In this occasion, the number
- 35 of R^2 which is a C_{2-20} alkanoyl group is the same or different between the two or more of compounds.

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However, most preferably, there can be mentioned a β -CyD derivative wherein all of R¹s and R³s are lower alkyl groups and all of R²s are C₂₋₂₀ alkanoyl groups, and a mixture wherein such derivatives are 5 mainly (i.e, exceeding 50 %) contained.

The lower alkyl groups include straight-chain or branched alkyl groups having 1 to 6 carbon atoms, but as preferred ones, there can be mentioned methyl, ethyl, n-propyl, isopropyl, n-butyl, sec-butyl groups, etc., 10 and further preferred among them is a methyl group.

As to the C_{2-20} alkanoyl groups, the alkyl part may be straight-chain or branched, and as preferred ones, there can be mentioned acetyl, n-propanoyl, n-butanoyl, n-pentanoyl (or valeryl), n-hexanoyl (or

15 caproyl), n-heptanoyl (or enanthoyl), n-octanoyl (or capryloyl), n-dodecanoyl (or lauroyl), n-tetradecanoyl (or myristoyl) and n-octadecanoyl (or stearoyl) groups, etc., and, above all, acetyl, n-propanoyl, n-butanoyl and n-hexanoyl are preferred, and further preferred 20 among them is an acetyl group.

Thus, as particularly preferred ones among acylated alkylated CyD derivatives or mixtures of two or more of the derivatives according to the invention, there can be mentioned heptakis (2,6-di-0-methyl-3-

- 25 acetyl)- β -CyD with a degree of substitution (D.S.) of 7 at the 3-position (hereinafter referred to as DMA- β -CyD, and the following abbreviations follow this) and mixtures mainly containing this DMA- β -CyD, or a mixture of acetylated DA- β -CyDs with a lower substitution (D.S.
- 30 3.5-6) at the 3-position.

Such an acylated alkylated CyD of the invention can be prepared by the following process, as another embodiment of the invention, which comprises reacting a partially alkylated CyD derivative repre-35 sented by the formula (II)

(II)



wherein n is any of integers 6, 7 and 8, and the 1-position and the 4-position of the sugar residues at the both ends are mutually bound by a covalent bond,

 $R^{\,4},\ R^{\,5}$ and $R^{\,6}$ independently represent hydrogen atoms, lower alkyl groups, glucosyl groups or maltosyl groups, provided that R^4 , R^5 and R^6 of the number of total $3 \times n$ composed of each n are, simultaneously, at least one lower alkyl group and at least one hydrogen atom, and the number of the glucosyl groups and maltosyl groups is at most 2,

20 or a mixture of the derivatives with an activated $\mathrm{C}_{2\,-\,2\,0}$ alkanoic acid in a polar solvent, if necessary in the presence of a condensing agent.

The partially alkylated CyD derivatives of the formula (II) themselves are known or available on the 25 market, but ones prepared according to preparation processes known per se can also be used. Further, as the activated C_{2-20} alkanoic acid, acid anhydrides or acid halides (chlorides, bromides) of alkanoic acids can be used, but preferably acid anhydrides can be men-

30 tioned. When an acid halide is used, it is desirable to make a basic organic amine such as triethylamine coexist as a hydrogen halide-capturing agent, but it is advantageous to use pyridine as a solvent and a hydrogen halide-capturing agent or condensing agent.

35 When an acid anhydride is used as an acylating reactant and pyridine is used as a solvent, the acyl

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groups of the desired number can be introduced into a compound of the formula (II) by using pyridine in an amount enough to dissolve the CyD reactant and the acid anhydride reactant and carrying out reaction, usually,

- 5 at a temperature around 80°C for several hours to 72 hours. The desired acylated alkylated CyD derivatives can be isolated and purified from the thus obtained reaction mixture using solvent extraction, various chromatographies, and recrystallization per se known,
- 10 but as stated above, they can also be separated, in a state of a mixture of two or more of the derivatives, from the reaction solvent and the unreacted reactant or the side reaction products. Usually, after the completion of the reaction, the reaction mixture is added
- 15 dropwise into ice water to decompose the excess acid anhydride, and the desired CyD derivative is extracted with chloroform. Sodium carbonate is added to the extract, and the mixture is desalted and subjected to separation and purification using silica gel columns,
- 20 and if necessary, subjected to recrystallization from an appropriate solvent. The desired CyD derivative can be obtained by concentrating the obtained substance to dryness. The structure of the obtained substance can be confirmed by mass spectrum, elementary analysis, etc.

As stated above, the thus obtained acylated alkylated CyD derivatives or mixtures of the derivatives of the invention have hemolytic activities and muscular irritation significantly lowered, compared with the previous CyDs, and have an action to solubilize water-

30 sparingly soluble or insoluble medicaments at room temperature, and are useful as carriers or delivery tools for such medicaments.

Therefore, as another embodiment of the invention, there is provided a pharmaceutical preparation

35 which comprises such an acylated alkylated CyD derivative or a mixture of two or more of the derivatives and
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such a medicament in such a state that they are closely compounded. The "state that they are closely compounded" means such a state that the CyD derivative and the medicament are homogeneously mixed or such a state that the medicament and the CyD derivative form an inclusion compound.

The preparation in such a state can be prepared by sufficiently kneading the CyD derivative and the medicament, in an aqueous solvent (including a mixed 10 solvent between methanol, ethanol, acetonitrile,

dimethylformamide or the like and water), in such a state that the CyD derivative and the medicament are suspended or dissolved, using a kneader or the like regularly used for the preparation of formulations.

15 The pharmaceutical preparation can be administered in an administration form such as parenteral administration, namely intravenous injection, intramuscular injection, subcutaneous injection or topical administration to the skin or mucosa., but administra-20 tion methods are not limited thereto, and it can also be administered by oral administration.

Medicaments or active ingredients applicable to the preparation according to the invention may be any medicaments including water-soluble or sparingly soluble 25 ones, so long as they meet the objects of the invention, but there can, generally, be mentioned water-sparingly soluble or insoluble medicaments, or unstable medicaments.

Further suitable active ingredients are those 30 which exert a local physiological effect, as well as those which exert a systemic effect, either after penetrating the mucosa or in the case of oral administration - after transport to the gastro-intestinal tract with saliva. The dosage forms prepared from the compositions

35 according to the present invention are particularly suitable for active ingredients which exert their activ-

ity during an extended period of time, i.e. drugs having a half-life of at least several hours. Examples thereof are: analgesic and anti-inflammatory drugs (NSAIDs, flurbiprofen, fentanyl, indomethacin, ketoprofen,

- 5 nabumetone, paracetamol, piroxicam, tramadol); antiarrhythmic drugs (procainamide, quinidine, verapamil); antibacterial and antiprotozoal agents (amoxicillin, ampicillin, benzathine penicillin, benzylpenicillin, cefaclor, cefadroxil, cefprozil, cefuroxime axetil,
- 10 cephalexin, chloramphenicol, chloroquine, ciprofloxacin, clarithromycin, clavulanic acid, clindamycin, doxyxycline, erythromycin, flucloxacillin, halofantrine, isoniazid, kanamycin, lincomycin, mefloquine, minocycline, nafcillin, neomycin, norfloxacin, ofloxacin,
- 15 oxacillin, phenoxymethyl-penicillin, pyrimethaminesulfadoxime, streptomycin); anti-coagulants (warfarin); antidepressants (amitriptyline, amoxapine, butriptyline, clomipramine, desipramine, dothiepin, doxepin, fluoxetine, gepirone, imipramine, lithium carbonate, mian-
- 20 serin, milnacipran, nortriptyline, paroxetine, sertraline; 3-[2-[3,4-dihydrobenzofuro[3,2-c]pyridin-2(1H)y1]-ethy1]-2-methy1-4H-pyrido[1,2-a]pyrimidin-4-one); anti-diabetic drugs (glibenclamide, metformin); antiepileptic drugs (carbamazepine, clonazepam, ethosux-
- 25 imide, phenobarbitone, phenytoin, primidone, topiramate, valpromide); antifungal agents (amphotericin, clotrimazole, econazole, fluconazole, flucytosine, griseofulvin, itraconazole, ketoconazole, miconazole nitrate, nystatin, terbinafine, voriconazole); antihistamines
- 30 (astemizole, cinnarizine, cyproheptadine, decarboethoxyloratadine, fexofenadine, lunarizine, levocabastine, loratadine, norastemizole, oxatomide, promethazine, terfenadine); anti-hypertensive drugs (captopril, enalapril, ketanserin, lisinopril,
- 35 minoxidil, prazosin, ramipril, reserpine, terazosin); anti-muscarinic agents (atropine sulphate, hyoscine);

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antivirals (acyclovir, AZT, ddC, ddI, ganciclovir, loviride, tivirapine, 3TC, delavirdine, indinavir, nelfinavir, ritonavir, saquinavir); antineoplastic agents and antimetabolites (adriamycine, cladribine, 5 cisplatin, dactinomycin, daunorubicin, doxorubicin, etoposide, irinotecan, mitomycin, mitoxantrone,

- tamoxifen, taxol, taxotere, topotecan, trimetrexate, vincristine, vinblastine); anti-migraine drugs (almotriptan, alniditan, eletriptan, frovatriptan,
- 10 naratriptan, rizatriptan, sumatriptan, zolmitriptan); anti-Parkinsonian drugs (bromocryptine mesylate, levodopa, selegiline); antipsychotic, hypnotic and sedating agents (alprazolam, buspirone, chlordiazepoxide, chlorpromazine, clozapine, diazepam, flupen-
- 15 thixol, fluphenazine, flurazepam, 9-hydroxyrisperidone, lorazepam, mazapertine, olanzapine, oxazepam, pimozide, pipamperone, piracetam, promazine, risperidone, selfotel, seroquel, sertindole, sulpiride, temazepam, thiothixene, triazolam, trifluperidol, ziprasidone, zol-
- 20 pidem); anti-stroke agents (lubeluzole, lubeluzole oxide, riluzole, aptiganel, eliprodil, remacemide); antitussive (dextromethorphan, laevodropropizine); beta-adrenoceptor blocking agents (atenolol, carvedilol, metoprolol, nebivolol, propanolol); cardiac inotropic
- 25 agents (amrinone, digitoxin, digoxin, milrinone); corticosteroids (beclomethasone dipropionate, betamethasone, budesonide, dexamethasone, hydrocortisone, methylprednisolone, prednisolone, prednisone, triamcinolone); disinfectants (chlorhexidine); diuretics (aceta-
- 30 zolamide, frusemide, hydrochlorothiazide, isosorbide); enzymes; essential oils (anethole, anise oil, caraway, cardamom, cassia oil, cineole, cinnamon oil, clove oil, coriander oil, dementholised mint oil, dill oil, eucalyptus oil, eugenol, ginger, lemon oil, mustard oil,
- 35 neroli oil, nutmeg oil, orange oil, peppermint, sage, spearmint, terpineol, thyme); gastro-intestinal agents

(cimetidine, cisapride, clebopride, diphenoxylate, domperidone, famotidine, lansoprazole, loperamide, loperamide oxide, mesalazine, metoclopramide, mosapride, nizatidine, norcisapride, olsalazine, omeprazole, panto-

- 5 prazole, perprazole, prucalopride, ranitidine, rabeprazole, ridogrel, sulphasalazine); haemostatics (aminocaproic acid); lipid regulating agents (atorvastatin, lovastatin, pravastatin, probucol, simvastatin); local anaesthetics (benzocaine, lignocaine); opioid analgesics
- 10 (buprenorphine, codeine, dextromoramide, dihydrocodeine, hydrocodone, oxycodone, morphine); parasympathomimetics (eptastigmine, galanthamine, metrifonate, neostigmine, physostigmine, tacrine, donepezil, rivastigmine, milameline, sabcomeline, talsaclidine, xanomeline, meman-
- 15 tine, lazabemide); sex hormones (oestrogens: conjugated oestrogens, ethinyloestradiol, mestranol, oestradiol, oestriol, oestrone; progestogens; chlormadinone acetate, cyproterone acetate, 17-deacetyl norgestimate, desogestrel, dienogest, dydrogesterone, ethynodiol diace-
- 20 tate, gestodene, 3-keto desogestrel, levonorgestrel, lynestrenol, medroxy-progesterone acetate, megestrol, norethindrone, norethindrone acetate, norethisterone, norethisterone acetate, norethynodrel, norgestimate, norgestrel, norgestrienone, progesterone, quingestanol
- 25 acetate); stimulating agents (sildenafil); vasodilators (amlodipine, buflomedil, amyl nitrite, diltiazem, dipyridamole, glyceryl trinitrate, isosorbide dinitrate, lidoflazine, molsidomine, nicardipine, nifedipine, oxpentifylline, pentaerythritol tetranitrate).
- 30 In the preparation according to the invention, the compounding ratio between the acylated alkylated CyD and the medicament can be an any ratio so long as it meets the objects, but in view of controlling the release of the medicament from the preparation, the ratio
- 35 of the acylated alkylated CyD : the medicament can be made to be 1 : 4 to 4 : 1, preferably 1 : 2 to 2 : 1, in

terms of mole ratio.

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In the preparation of the invention, pharmaceutically acceptable other auxiliaries or additives can, if necessary, be incorporated in a range not to 5 have bad influence on the objects of the invention. As such auxiliaries or additives, there can be mentioned stabilizers, dissolution aids, suspending agents, emulsifying agents, buffering agents, preservatives, isotonizing agents, or other proper additives, which are 10 used regularly in the technical field.

The invention is specifically described below by examples, but the invention is not limited by these examples.

Example 1 Synthesis of $DMA-\beta-CyD$

- DM-β-CyD (12 g) was dissolved in 60 mL of anhydrous pyridine, and 25 mg of 4-dimethylaminopyridine was added. Then, 12 mL of acetic anhydride was gradually added dropwise, and the mixture was subjected to reaction at 80°C for 24 hours. After the completion of the reaction, the mixture was added dropwise into ice
- water to decompose the excess acid anhydride, and extracted with chloroform. Sodium carbonate was added to the organic phase to desalt it, and the mixture was subjected to separation and purification using a silica
- 25 gel column. The obtained substance was concentrated to dryness to give the desired DMA- β -CyD (D.S. 7). This CyD derivative had a melting point of 113 to 117°C, and its solubility in water at 25°C exceeded 50 mg/dl. The resulting DMA- β -CyD (D.S. 7) was recrystallized from
- 30 water to give white crystals (yield 60%) having a melting point of 126°C. Its mass spectrum and ¹H-NMR spectrum are shown in Fig. 1 and Fig. 2, respectively.

FAB MS (negative mode) *m/z* 1777 [M+*m*-nitro-

benzyl alcohol (matrix)-H); ¹H-NMR (CDCl₃) d 5.16 (t, 35 1H, CyD H-3), 5.00 (d, 1H, CyD H-1), 3.91-3.87 (m, 2H, CyD H-5 and H-6b), 3.79 (t, 1H, CyD H-4), 3.54 (d, 1H, CyD H-6a), 3.37 (s, 3H, 6-CH₃), 3.33 (s, 3H, 2-CH₃), 3.21 (dd, 1H, CyD H-2), 2.04 (s, 3H, 3-CH₃). Example 2 Synthesis of acetylated DM- β -CyD with a lower substitution at the 3-position

5 The acetylated DM- β -CyD was prepared by using a small amount of the acid anhydride (4.6 g, 45 mmol) to DM- β -CyD (10 g, 7.5 mmol). The other condition of preparation was identical to that described in Example 1, except for the recrystallization due to the face that 10 it was a mixture of components with different D.S.s. The D.S. value was determined by a peak ratio of the CyD anomeric proton (H-1) and the methyl proton of acetyl groups in ¹H-NMR spectra (see Fig. 3), and was 3.8. The mixture is hereinafter referred to as DMA4- β -CyD.

15 Example 3 Synthesis of butyrated DM-β-CyD

 $DM-\beta-CyD$ (5 g) was dissolved in 25 mL of anhydrous pyridine, and 9 mL of n-butyric anhydride was added, and the mixture was subjected to reaction at 80°C for 24 hours. After the completion of the reaction, the 20 mixture was added dropwise into ice water to decompose the excess acid anhydride, and extracted with chloroform. Sodium carbonate was added to the extract to desalt it, and the mixture was subjected to separation and purification using a silica gel column. The ob-

25 tained substance was concentrated to dryness to give the desired butyrated DM- β -CyD. This CyD derivative had a melting point of 108 to 111°C. Its ¹H-NMR spectrum is shown in Fig. 4.

Example 4 Synthesis of octanoylated DM-B-CyD

- 30 DM-β-CyD (5 g) was dissolved in 25 mL of anhydrous pyridine, and 16 mL of octanoic anhydride was added, and the mixture was subjected to reaction at 80°C for 24 hours. After the completion of the reaction, the mixture was added dropwise into ice water to decompose 35 the excess acid anhydride, and extracted with chloro-
- form. Sodium carbonate was added to the extract to

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desalt it, and the mixture was subjected to separation and purification using a silica gel column. The obtained substance was concentrated to dryness to give the desired octanoylated $DM-\beta-CyD$. This CyD derivative was 5 an oily substance at 25 °C. Its 1 H-NMR spectrum is shown in Fig. 5.

Characteristic tests

(1) Determination of stability constant of DMA- β -CyD In this test, the inclusion properties of 10DMA- β -CyD was compared with those of the parent β -CyD and DM- and TM- β -CyDs.

The stability constant between DMA- β -CyD and flurbiprofen was determined by the solubility method 15 i.e., according to the method of Higuchi, T. et al., Adv. Anal. Chem. Instr. 1965, 4, 117-212. The stability constants between β -CyD, DM- β -CyD or TM- β -CyD and flurbiprofen determined simultaneously for comparison are also shown in Table 1.

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Flurbiprofen

10010 1	
	Compound
	$\beta - CvD$

Table 1:		
	Compound	Stability constant (M^{-1})
-	β-CyD	3613
	$DM - \beta - CyD$	8055
	$TM-\beta-CyD$	1655
	$DMA - \beta - CyD$	1212

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These results suggest that DMA- β -CyD has the same inclusion ability as TM- β -CyD, although it is inferior in inclusion ability to DM- β -CyD.

(2) Test on hemolytic properties of DMA- β -CyD and

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 $DMA4 - \beta - CyD$

In this test on hemolytic properties, 6 to 7 mL of blood was taken from the auricular vein of a white rabbit, 1 mL of a preserved erythrocyte solution was added, and the mixture was gently mixed. To this was 10 added 4 to 5 mL of 10 mM isotonized phosphate buffer (pH 7.4), the mixture was gently mixed and centrifuged at 1,000 g for 5 minutes, and the supernatant was removed. This washing operation was repeated three times, 10 mM isotonized phosphate buffer (pH 7.4) was added to 1 mL 15 of the resultant solution to make the volume 20 mL, and thereby a 5 % erythrocyte suspension was prepared.

The CyD derivative of various concentrations was diluted with 10 mM isotonized phosphate buffer (pH 7.4), and the resultant each dilution was incubated at 20 37°C. Then 4 mL of this dilution was taken, 0.2 mL of the 5 % erythrocyte suspension was added, and the mixture was incubated at 37°C for 30 minutes. The mixture was centrifuged at 1,000 g for 5 minutes, 3 mL of the supernatant was measured for absorbance at 543 nm, and

- 25 thereby its hemolytic activity was determined. Further, the specimen was observed visually using a microscope. The obtained results are shown in Fig. 6. To the 5 % erythrocyte suspension was added 100 mM DMA-β-CyD (D.S. 7), DMA4-β-CyD (D.S. 3.8), β-CyD, DM-β-CyD, TM-β-CyD, 2-
- 30 HP-β-CyD (D.S. 4.8) (see Shiotani, K. et al. Pharm. Res. 1995, 12, 78-84) or subfobutyl ether β-CyD (D.S. 3.5)

(see ibid), the above treatment was carried out, and then observation by a microscope was made.

It is apparent that the hemolytic activity of DMA- β -CyDs was weaker than those of β -CyD, DM- β -CyD and 5 TM- β -CyD. For example, the hemolysis began at about 2 mM, 0.5 mM and 1 mM, and the concentrations to induce 50% hemolysis were about 4 mM, 1 mM and 2 mM for β -CyD, DM- β -CyD and TM- β -CyD, respectively. On the other hand, the hemolysis of DMA4- β -CyD with D.S. 3.8 began at about 10 12 mM, and its 50% hemolysis concentration was about 22

- mM. In the case of DMA- β -CyD with D.S. 7, no hemolyis was observed up to 100 mM. The hemolytic activity of DMA- β -CyDs was weaker than those of 2-HP (D.S. 4.8) and sulfobutyl ether of β -CyD (D.S. 3.5).
- 15 (3) Determination of the released amount of cholesterol from the intact erythrocytes of rabbits treated with $DMA-\beta-CyD$ or $DMA4-\beta-CyD$

Each DMA-β-CyD, DMA4-β-CyD, DM-β-CyD was diluted with 10 mM isotonized phosphate buffer (pH 7.4),
and the dilution was incubated at 37°C. Then, 4 mL of the dilution was taken, 0.2 mL of the 5 % erythrocyte suspension was added, and the mixture was incubated at 37°C for 30 minutes. The mixture was centrifuged at 1,000 g for 5 minutes, 5 mL of chloroform was added to 3

- 25 mL of the supernatant, and the mixture was shaken for 30 minutes to make extraction. The chloroform layer was taken, and concentrated to give a specimen. The specimen was assayed for cholesterol amount using a Cholesterol E-Test Wako (made by Wako Pure Chemical Industries,
- 30 Ltd.). The obtained results are shown in Fig. 7. One of the causes of CyD-induced hemolysis is

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known to be extractions of cholesterol and phospholipids from erythrocytes through the inclusion complex formation. Therefore, in this test, the cholesterol release behavior from rabbit erythrocytes by the addition of

- 5 DMA- β -CyD or DMA4- β -CyD was investigated and compared with the cholesterol release behaviors of a control (an isotonic buffer without CyDs) and DM- β -CyD. Figure 7 shows the released amounts of cholesterol from the intact erythrocytes of rabbits treated with β -CyDs in 10
- 10 mM phosphate buffer (pH 7.4) at 37° C. DM- β -CyD induced about 80% release of cholesterol at a concentration of 0.5 mM at which the hemolyis only slightly occurred (see Figure 6). On the other hand, DMA- β -CyDs induced only 10% release of cholesterol at the same concentration,
- 15 and this release was the same as that of the control experiment conducted in the isotonic buffer.

CLAIMS

1. An acylated alkylated cyclodextrin derivative represented by the formula (I)



wherein n is any of integers 6, 7 and 8, and the 1-position and the 4-position of the sugar residues at the both ends are mutually bound by a covalent bond,

 R^1 , R^2 and R^3 independently represent hydrogen atoms, lower alkyl groups or C_{2-20} alkanoyl groups, or in some case, represent glucosyl groups or maltosyl groups whose hydroxyl group(s) may be replaced with lower alkyloxy group(s) or C_{2-20} alkanyloxy group(s),

provided that any of R^1 , R^2 and R^3 of the number of total $3 \times n$ composed of each n are simultaneously at least one lower alkyl group and at least one C_{2-20} alkanoyl group, and the residual groups, when exist, are hydrogen atoms or the glucosyl groups or maltosyl groups of the number of up to at most 2, or a mixture of two or more of the derivatives.

2. The acylated alkylated cyclodextrin derivative

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or mixture of two or more of the derivatives according to claim 1 wherein 50 % or more of R^1 and R^3 of the number of total $2 \times n$ are lower alkyl groups, the residual R^1 and R^3 and R^2 are at least one C_{2-20} alkanoyl group, and the residual R^1 , R^2 and R^3 , when exist, are hydrogen atoms.

3. The acylated alkylated cyclodextrin derivative or mixture of two or more of the derivatives according to claim 1 wherein R^1 and R^3 are lower alkyl groups, 50 % or more of R^2 are C_{2-20} alkanoyl groups, and the residual R^2 , when exist, are hydrogen atoms.

4. The acylated alkylated cyclodextrin derivative or mixture of two or more of the derivatives according to claim 1 wherein R^1 and R^3 are lower alkyl groups, and 50 % or more of R^2 of the number of n are acyl groups. 5. The acylated alkylated cyclodextrin derivative or mixture of two or more of the derivatives according to any of claims 1 to 4 wherein the lower alkyl groups are methyl, ethyl, n-propyl, isopropyl, n-butyl or sec-butyl, and the alkanoyl groups are acetyl, n-propanoyl, n-butanoyl, n-pentanoyl, n-hexanoyl, n-heptanoyl, n-octanoyl, n-dodecanoyl, n-tetradecanoyl or n-octadecanoyl.

6. The acylated alkylated cyclodextrin derivative or mixture of two or more of the derivatives according to any of claims 1 to 5 wherein the lower alkyl groups are methyl groups, and the alkanoyl groups are acetyl groups.

7. The acylated alkylated cyclodextrin derivative or mixture of two or more of the derivatives according to any of claims 1 to 6 wherein n is 7 in the formula

(I).

8. Heptakis $(2, 6-di-0-methyl-3-0-acetyl)-\beta$ cyclodextrine; heptakis $(2, 6-di-0-methyl-3-0-butyryl)-\beta$ cyclodextrine; or heptakis (2, 6-di-0-methyl-3-0-octanoyl)- β -cyclodextrine.

9. A mixture of acetylated heptakis $(2, 6-di-0-methyl)-\beta$ -cyclodextrines having an average degree of acetyl-substitution of about 3.8 at the 3-position. 10. A process for preparing an acylated alkylated cyclodextrin derivative represented by the formula (I) according to claim 1 or a mixture of two or more of the derivatives which comprises reacting a partially alkylated cyclodextrin derivative represented by the formula (II)



wherein n is any of integers 6, 7 and 8, and the 1-position and the 4-position of the sugar residues at the both ends are mutually bound by a covalent bond,

 R^4 , R^5 and R^6 independently represent hydrogen atoms, lower alkyl groups, glucosyl groups or maltosyl groups, provided that R^4 , R^5 and R^6 of the number of total $3 \times n$ composed of each n are, simultaneously, at least one lower alkyl group and at least one hydrogen atom, and the number of the glucosyl groups

and maltosyl groups is at most 2, or a mixture of the derivatives with an activated C_{2-20} alkanoic acid in a polar solvent, if necessary in the presence of a condensing agent to acylate the compound(s) of the formula (II).

11. A pharmaceutical preparation which comprises the acylated alkylated cyclodextrin derivative or mixture of two or more of the derivatives according to any of claims 1 to 9 and a medicament in such a state that they are closely compounded.

12. The pharmaceutical preparation according to claim 11 wherein the medicament is selected from the group consisting of nonsteroidal antirheumatic agents, steroids, cardiac glycosides, benzodiazepine derivatives, benzimidazole derivatives, piperidine derivatives, piperazine derivatives, imidazole derivatives and triazole derivatives.

13. A process for preparing a pharmaceutical preparation according to claim 11 or 12 which comprises kneading closely the acylated alkylated cyclodextrin derivative or mixture of two or more of the derivatives according to any of claims 1 to 6 and a medicament in an aqueous solvent, and then, if necessary, removing the solvent.

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FIG. 1

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FIG. 3





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FIG. 5

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Concn. of β -CyDs (mM)





INTERNATIONAL SEARCH REPORT

Inter: - (unal Application No PCT/JP 99/02806

A. CLASSIFICATION OF SUBJECT MATTER IPC 6 C08B37/16 A61k A61K47/48 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) IPC 6 C08B A61K Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Category * Relevant to claim No. WO 89 09235 A (MACHEREY-NAGEL GMBH) 1 - 4, 10Х 5 October 1989 (1989-10-05) claims 1-7; example 3 χ EP 0 312 352 A (CHINOIN) 1 - 1319 April 1989 (1989-04-19) page 3, line 17,18 page 13, line 60 - line 64 page 14, line 34 - line 40 Further documents are listed in the continuation of box C. Patent family members are listed in annex. X Special categories of cited documents : "T" later document published after the international filing date or priority date and not in conflict with the application but "A" document defining the general state of the art which is not considered to be of particular relevance cited to understand the principle or theory underlying the invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention filing date cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docucitation or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or ments, such combination being obvious to a person skilled in the art. other means document published prior to the international filing date but "P" later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 9 August 1999 18/08/1999 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Lensen, H Fax: (+31-70) 340-3016

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Bibliographic data: WO9855148 (A1) --- 1998-12-10

PHARMACEUTICAL COMPOSITIONS COMPRISING CYCLODEXTRINS

Inventor(s):	VANDECRUYS ROGER PETRUS GEREBE [BE] 🟦				
Applicant(s):	JANSSEN PHARMACEUTICA NV [BE]; VANDECRUYS ROGER PETRUS GEREBE [BE] <u>*</u>				
Classification:	- international:	A61K47/12; A61K47/30; A61K47/40; A61K8/73; A61K9/14; A61K9/48; A61Q3/00; A61K47/38; (IPC1- 7): A61K47/48; A61K9/14; A61K9/48			
	- European:	A61K8/73; A61K8/73C; A61K8/73T; A61K9/00M3; A61K9/14H4; A61K9/14H6; A61K9/48H4; A61K9/48H6; A61Q3/00			
Application number:	WO1998EP03189 1	9980527			
Priority number(s):	GB19970011643 19	970605			
Also published as:	ZA9804849 (A) I NO995925 (A) K HU0004924 (A2) EP0998304 (B1) CN1258220 (A) AT247489 (T) A	JS2002150616 (A1) PT998304 (E) R20010005852 (A) JP2002511073 (A) ES2206949 (T3) EP0998304 (A1) DK998304 (T3) DE69817363 (T2) CA2292506 (A1) AU8108198 (A) R012927 (A1) less			

Abstract of WO9855148 (A1)

The invention provides a novel pharmaceutical composition comprising a no more than sparingly water-soluble drug compound, a cyclodextrin, a physiologically tolerable water-soluble acid, and a physiologically tolerable water-soluble organic polymer.

Last updated: 14.03.2012 Worldwide Database 5.7.38; 93p



WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification ⁶ :		(11) International Publication Number: WO 98/55148
A61K 47/48, 9/14, 9/48	A1	(43) International Publication Date: 10 December 1998 (10.12.98)
 (21) International Application Number: PCT/EP (22) International Filing Date: 27 May 1998 (2 (30) Priority Data: 27 May 1998 (2 (31) Priority Data: 27 May 1998 (2 (32) Priority Data: 27 May 1998 (2 (32) Priority Data: 27 May 1998 (2 (33) Priority Data: 27 May 1998 (2 (34) Priority Data: 27 May 1998 (2 (35) Inventor; and (35) Inventor/Applicant (for US only): VANDECRUYS Petrus, Gerebern [BE/BE]; Janssen Pharmaceuti Turnbuttorum 20 B 2340 Becreto (BE) 	98/031 27.05.9 C ANSSE Sweg 3 S, Rog Ca N.V	 (43) International Publication Date: 10 December 1998 (10.12.98) (81) Designated States: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, GM, GW, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM). European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG). Published With international search report. 7.,
(74) Agent: QUAGHEBEUR, Luc; Janssen Pharmaceuti Patent Dept., Turnhoutseweg 30, B-2340 Beerse (ica N.Y BE).	

(57) Abstract

The invention provides a novel pharmaceutical composition comprising a no more than sparingly water-soluble drug compound, a cyclodextrin, a physiologically tolerable water-soluble acid, and a physiologically tolerable water-soluble organic polymer.

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PHARMACEUTICAL COMPOSITIONS COMPRISING CYCLODEXTRINS

This invention relates to novel pharmaceutical compositions, in particular compositions and dosage forms providing improved drug release and uptake on administration into externally voiding body cavities (e.g. the gi tract) or on topical administration, especially

for acid solubilized drug compounds.

Many drug compounds, while possessing desired therapeutic properties, are used inefficiently due to their poor water solubilities. Thus for example where such

- 10 compounds are administered orally, only a small fraction of the drug is taken up into the blood during transit of the gi tract. As a result, to achieve adequate drug uptake it may be necessary to administer high doses of the drug compound, to prolong the period of drug administration or to make frequent administrations of the drug compound. Indeed, the poor solubility and hence poor bioavailability of a drug may cause an alternative
- 15 drug, perhaps one with undesired side effects or one which requires invasive administration (e.g. by injection or infusion), to be used in place of the poorly soluble drug.

One approach to poor solubility is to derivatise the drug molecule to introduce water solubilizing groups, e.g. ionic groups such as carboxyl groups or non-ionic groups such

- 20 as polyhydroxyalkyl groups, so as to produce a more soluble derivative. This approach however is not always successful as it may not be possible to maintain adequately high therapeutic efficacy and adequately low toxicity or other side effects. Thus one example of a poorly water soluble drug which has not been superseded by a solubilized derivative is the antifungal agent itraconazole.
- 25 Attempts have therefore been made to enhance the uptake of drugs such as itraconazole by increasing the surface area of the drug compound exposed to saliva or gastric fluid, and hence promote dissolution of the drug compound, by thinly coating the drug compound onto essentially inert carrier particles, e.g. sugar beads. This however has the drawback that the volume of solid composition required to administer a given quantity
- 30 of the drug compound is quite high since the carrier contributes significantly to the overall administration volume. Since administration of large volume capsules or tablets, or of large quantities of smaller volume capsules or tablets, provides difficulties for the patient, the drawbacks of this approach are obvious.
- 35 Yet another approach has been to administer the drug compound in the form of a solution of the drug compound and a drug complexing agent such as a cyclodextrin. This approach has limitations also in that the dosage volume is constrained by the

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solubilizing power of the complexing agent, readily unitized solid dosage forms can not be used, and there is no gradual release of the drug compound for biological uptake.

However, we have now found that by combining such drug compounds with a cyclo-

- 5 dextrin, a water-soluble acid and a water-soluble organic polymer, an administration form may be produced which surprisingly improves the biological uptake of the drug compound, in particular a form which can surprisingly improve the time profile for the drug content of the plasma of the patient (*i.e.* the pharmacokinetic profile defined by such parameters as AUC, t_{max}, C_{max}, etc.).
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Thus viewed from one aspect the invention provides a pharmaceutical composition comprising a no more than sparingly water-soluble drug compound, a cyclodextrin, a physiologically tolerable water-soluble acid, and a physiologically tolerable watersoluble organic polymer.

15 Viewed from a further aspect the invention provides the use of a no more than sparingly water-soluble drug compound, a cyclodextrin, a physiologically tolerable water-soluble acid, and a physiologically tolerable water-soluble organic polymer for the manufacture of a pharmaceutical composition according to the invention for use in a method of therapy or diagnosis of the human or non-human animal (e.g. mammalian, reptilian or avian) body.

Viewed from a still further aspect the invention provides a method of therapy or diagnosis of the human or non-human animal (e.g. mammalian, reptilian or avian) body which comprises administering to said body a therapeutically or diagnostically effective

25 dose of a pharmaceutical composition, the improvement comprising using as said composition a composition according to the present invention.

The compositions of the invention may if desired be aqueous, but in general will preferably be substantially water-free, e.g. containing up to 3% by weight, preferably

- less than 1% by weight water, and most preferably less than 0.5% water, but may be mixed with water immediately before administration or may be coated and dispersed in an aqueous medium whereby the coating is only broken down after administration.
 Such aqueous compositions are deemed to fall within the scope of the invention.
 Depending on the selection of components, the compositions of the invention may be
- liquid, solid or semi-solid e.g. gel-like. Preferably the compositions are non-freeflowing at ambient temperature (e.g. 21°C), other than as free flowing particulates.
 Thus the compositions at ambient temperature are preferably solids or semi-solids or, less preferably, highly viscous fluids.

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In the compositions of the invention the drug compound, acid, cyclodextrin and organic polymer are intimately admixed.

- 5 Thus where the composition is particulate, the acid, drug compound, cyclodextrin and organic polymer are mixed together within the particles (e.g. at the molecular level following solvent removal from a solution of these components). Granulate mixtures where individual particles do not contain all four components, or have cores of one or more components coated with other components are not preferred. This intimate admixture is important since the effects of the components are complimentary at the
- microscopic level during dissolution of the compositions of the invention.

Preferably, all components are dispersed so as to form a system that is chemically and physically uniform or homogenous throughout, or consists of one phase as defined in
thermodynamics ; such a dispersion will be called a glass thermoplastic phase or system hereinafter. The components of the glass thermoplastic system are readily bioavalaible to the organisms to which they are administered. This advantage can probably be explained by the ease with which said glass thermoplastic system can form liquid solutions when contacted with a body liquid such as gastric juice. The ease of

- 20 dissolution may be attributed at least in part to the fact that the energy required for dissolution of the components from a glass thermoplastic system is less than that required for the dissolution of components from a crystalline or microcrystalline solid phase.
- 25 As the cyclodextrin in the compositions of the invention, there may be used any of the physiologically tolerable water-soluble substituted or unsubstituted cyclodextrins or physiologically tolerable derivatives thereof, e.g. α-, β- or γ-cyclodextrins or derivatives thereof, in particular derivatives wherein one or more of the hydroxy groups are substituted, e.g. by alkyl, hydroxyalkyl, carboxyalkyl, alkylcarbonyl, carboxyalkoxyalkyl,
- 30 alkylcarbonyloxyalkyl, alkoxycarbonylalkyl or hydroxy-(mono or polyalkoxy)alkyl groups, wherein each alkyl or alkylene moiety preferably contains up to six carbons.

Substituted cyclodextrins which can be used in the invention include polyethers, e.g. as described in US Patent 3,459,731. In general, to produce these, unsubstituted

35 cyclodextrins are reacted with an alkylene oxide, preferably under superatmospheric pressure and at an elevated temperature, in the presence of an alkaline catalyst. Since a hydroxy moiety of the cyclodextrin can be substituted by an alkylene oxide which itself can react with yet another molecule of alkylene oxide, the average molar substitution

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(MS) is used as a measure of the average number of moles of the substituting agent per glucose unit. The MS can be greater than 3 and theoretically has no limit. In the cyclodextrin derivatives for use in the compositions according to the present invention the M.S. is conveniently in the range of 0.125 to 10, in particular of 0.3 to 3, or from

5 0.3 to 1.5. Preferably the M.S. ranges from about 0.3 to about 0.8, in particular from about 0.35 to about 0.5 and most particularly it is about 0.4. M.S. values determined by NMR or IR preferably range from 0.3 to 1, in particular from 0.55 to 0.75.

Further examples of substituted cyclodextrins include ethers wherein the hydrogen of
one or more cyclodextrin hydroxy groups is replaced by C₁₋₆alkyl, hydroxyC₁₋₆-alkyl,
carboxy-C₁₋₆alkyl or C₁₋₆alkyloxycarbonyl-C₁₋₆alkyl groups or mixed ethers thereof. In
particular such substituted cyclodextrins are ethers wherein the hydrogen of one or more
cyclodextrin hydroxy groups is replaced by C₁₋₃alkyl, hydroxy-C₂₋₄alkyl or carboxy-C₁₋₂alkyl or more particularly by methyl, ethyl, hydroxyethyl, hydroxypropyl,

15 hydroxybutyl, carboxymethyl or carboxyethyl.

In the foregoing definitions, the term " C_{1-6} alkyl" is meant to include straight and branched saturated hydrocarbon radicals, having from 1 to 6 carbon atoms, such as methyl, ethyl 1-methylethyl, 1,1-dimethylethyl, propyl. 2-methylpropyl, butyl, pentyl, hexyl and the like.

Such ethers can be prepared by reacting a cyclodextrin with an appropriate <u>O</u>-alkylating agent or a mixture of such agents in a concentration selected such that the desired cyclodextrin ether is obtained. The reaction is preferably conducted in a solvent in the

25 presence of a base. With such ethers, the degree of substitution (DS) is the average number of substituted hydroxy functions per glucose unit, the DS being thus 3 or less.

In the cyclodextrin derivatives for use in the compositions according to the present invention, the DS preferably is in the range of 0.125 to 3, in particular 0.3 to 2, more

30 particularly 0.3 to 1, and the MS is in the range of 0.125 to 10, in particular 0.3 to 3 and more particularly 0.3 to 1.5.

Of particular utility in the present invention are the β -cyclodextrin ethers, e.g. dimethyl- β -cyclodextrin as described in Drugs of the Future, Vol. 9, No. 8, p. 577-578 by

35 M. Nogradi (1984) and polyethers, e.g. hydroxypropyl-β-cyclodextrin and hydroxyethyl-β-cyclodextrin. Such alkyl ethers may for example be methyl ethers with a degree of substitution of about 0.125 to 3, e.g. about 0.3 to 2. Such a hydroxypropyl cyclodextrin may for example be formed from the reaction between β-cyclodextrin and propylene oxide and may have a MS value of about 0.125 to 10, e.g. about 0.3 to 3.

Especially suitable cyclodextrins are β -CD, 2,6-dimethyl- β -CD, 2-hydroxyethyl- β -CD, 2-hydroxyethyl- γ -CD, 2-hydroxypropyl- γ -CD and (2-carboxymethoxy)propyl- β -CD,

5 and in particular 2-hydroxypropyl- β -CD.

Besides simple cyclodextrins, branched cyclodextrins and cyclodextrin polymers may also be used.

Other cyclodextrins are described for example in Chemical and Pharmaceutical Bulletin 28: 1552-1558 (1980), Yakugyo Jiho No. 6452 (28 March 1983), Angew. Chem. Int. Ed. Engl. 19: 344-362 (1980), US-3,459,731, EP-A-0,149,197, EP-A-0,197,571, US-4,535,152, WO-90/12035 and GB-2,189,245. Other references describing cyclodextrins for use in the compositions according to the present invention, and which

- provide a guide for the preparation, purification and analysis of cyclodextrins include the following: "Cyclodextrin Technology" by József Szejtli, Kluwer Academic Publishers (1988) in the chapter Cyclodextrins in Pharmaceuticals; "Cyclodextrin Chemistry" by M.L. Bender et al., Springer-Verlag, Berlin (1978); "Advances in Carbohydrate Chemistry", Vol. 12, Ed. by M.L. Wolfrom, Academic Press, New York in the chapter
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More recent examples of substituted cyclodextrins include sulfobutylcyclodextrins (US-5,134,127-A). Their use is also envisaged in the present invention.

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The cyclodextrin used is preferably a β -cyclodextrin, in particular hydroxypropyl- β -cyclodextrin. The most preferred cyclodextrin derivative for use in the compositions of the present invention is hydroxypropyl- β -cyclodextrin having a M.S. in the range of from 0.35 to 0.50 and containing less than 1.5% unsubstituted β -cyclodextrin.

35 M.S. values determined by NMR or IR preferably range from 0.55 to 0.75.

Nevertheless, the choice of cyclodextrin may be directed by the ability of the selected drug compound to be complexed by a particular cyclodextrin - thus the cyclodextrins

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with greater affinity for the particular drug compound may be preferred.

In the compositions of the invention, the cyclodextrin is preferably present at 5 to 70% by weight, more preferably 8 to 55%, most preferably 10 to 45% by weight (relative to

- 5 the total weight of cyclodextrin, acid, organic polymer and drug). The quantity of cyclodextrin used however will generally be dependent on the quantity of drug and the molar ratio of cyclodextrin to drug will preferably lie in the range 100:1 to 1:5, especially 50:1 to 1:2, more especially 10:1 to 1:1.
- 10 The acid used in the compositions of the invention may be any of the water-soluble physiologically tolerable acids, in particular any of the inorganic or, more preferably, organic acids conventionally used in the preparation of acid salts of drug compounds, e.g. citric, fumaric, tartaric, maleic, malic, succinic, oxalic, malonic, benzoic, mandelic and ascorbic acids.

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Tartaric acid and more especially citric acid are preferred since the salts they form with drug compounds usually have a reduced tendency to precipitate from aqueous solutions. In general however, any acid which is not so strong as to cause degradation of the cyclodextrin and yet which is capable, on the addition of water, of generating a low pH environment, preferably lower than pH 4 and ideally about pH 2, may be used. The acid may be in liquid (e.g. solution) or solid form: however acids which are solid at ambient

In the compositions of the invention, the acid will preferably be present at 1 to 95% by weight, preferably 5 to 90% by weight, more preferably 20 to 80%, and especially preferably 35 to 60% by weight (relative to the total weight of cyclodextrin, drug compound, organic polymer and acid). The amount of acid used will be dependent upon the selected acid and drug compound, but in general an increase in the relative proportion of acid will result in an acceleration of drug dissolution on contact with

conditions in their anhydrous or hydrate forms will generally be preferred.

30 water. The amount of acid used will normally be at least the amount necessary to form a 1:1 salt with the drug compound.

In general, the acid will form a significant proportion of dosage forms that dissolve rapidly in body fluids. Typically, they will comprise from 50 to 95% by weight of acid, preferably 50 to 90% by weight, more preferably 55 to 60% by weight. Thus viewed from a further aspect the invention provides a pharmaceutical composition comprising

from a further aspect the invention provides a pharmaceutical composition comprising an organic drug compound, a water-soluble physiologically tolerable acid, a watersoluble physiologically tolerable cyclodextrin and a water-soluble physiologically -7-

tolerable organic polymer, characterised in that the weight ratios of drug compound to acid and of drug compound to cyclodextrin are no more than 2:1, preferably no more than 1.5:1, especially preferably no more than 1:1, and particularly preferably no more than 0.9:1, especially no more than 0.5:1.

5

The organic polymer used in the compositions of the invention may be any of the physiologically tolerable water soluble synthetic, semi-synthetic or non-synthetic organic polymers.

- 10 Thus for example the polymer may be a natural polymer such as a polysaccharide or polypeptide or a derivative thereof, or a synthetic polymer such as a polyalkylene oxide (e.g. PEG), polyacrylate, polyvinylpyrrolidone, etc. Mixed polymers, e.g. block copolymers and glycopeptides may of course be used.
- 15 It is believed that the effect of the organic polymer arises from an enhancement in viscosity which serves to stabilize supersaturated solutions of the drug compound on dissolution of the composition of the invention. Thus the polymer conveniently has a molecular weight in the range 500D to 2 MD, and conveniently has an apparent viscosity of 1 to 100 mPa.s when in a 2% aqueous solution at 20°C. For example, the
- 20 water-soluble polymer can be selected from the group comprising
 - alkylcelluloses such as methylcellulose,
 - hydroxyakylcelluloses such as hydroxymethylcellulose, hydroxyethylcellulose, hydroxypropylcellulose and hydroxybutylcellulose,
 - hydroxyalkyl alkylcelluloses such as hydroxyethyl methylcellulose and hydroxypropyl methylcellulose,
 - carboxyalkylcelluloses such as carboxymethylcellulose,
 - alkali metal salts of carboxyalkylcelluloses such as sodium carboxymethylcellulose,
 - carboxyalkylalkylcelluloses such as carboxymethylethylcellulose,
 - carboxyalkylcellulose esters,
- 30 starches,

- pectins such as sodium carboxymethylamylopectin,
- chitin derivates such as chitosan,
- heparin and heparinoids,
- polysaccharides such as alginic acid, alkali metal and ammonium salts thereof,
- 35 carrageenans, galactomannans, tragacanth, agar-agar, gum arabic, guargum and xanthan gum,
 - polyacrylic acids and the salts thereof,

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- polymethacrylic acids and the salts thereof, methacrylate copolymers,
- polyvinylalcohol,
- polyvinylpyrrolidone, copolymers of polyvinylpyrrolidone with vinyl acetate,
- polyalkylene oxides such as polyethylene oxide and polypropylene oxide and
- copolymers of ethylene oxide and propylene oxide, e.g. poloxamers and poloxamines.

Non-enumerated polymers which are pharmaceutically acceptable and have appropriate physico-chemical properties as defined hereinbefore are equally suited for preparing compositions according to the present invention.

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Particularly preferably the organic polymer is a cellulose ether, e.g. methyl cellulose, hydroxyethylmethylcellulose, or hydroxypropylmethylcellulose (HPMC), for example a Methocel® (available from Colorcon, England) such as Methocel A, Methocel E, Methocel F, Methocel K, Methocel J or Methocel HB or a Metolose® such as Metolose

15 SM, Metolose SH or Metolose SE. Especially preferably the organic polymer is a hydroxypropylmethylcellulose, e.g. from 5 cps Methocel E to 15000 cps Methocel K15M.

Even very small quantities of the organic polymer serve to achieve a beneficial effect in
the compositions of the invention. Thus in the compositions of the invention the organic
polymer is conveniently present at 0.05 to 35% by weight, preferably 0.1 to 20%, more
preferably 0.5 to 15%, and most preferably 2 to 11% by weight (relative to the total
weight of drug compound, acid, cyclodextrin and organic polymer). The content and
viscosity grade of the organic polymer both affect the dissolution profile for the drug

- 25 compound in the compositions of the invention, with increased organic polymer content and/or increased viscosity grade (e.g. 15000 mPa.s in place of 5 mPa.s (mPa.s values being at 2% aqueous solution at 20°)) both tending to decelerate drug compound dissolution). Accordingly the selection of the identity and quantity of the organic polymer will generally depend upon the dissolution profile that is desired. For example,
- 30 a composition that provides sustained release of the drug, will comprise a water soluble polymer having an apparent viscosity of more than 1,000 mPa.s when dissolved in a 2% aqueous solution at 20°C.

The drug compound used in the compositions of the invention may be any organic or inorganic material which is no more than sparingly soluble, *i.e.* which is sparingly soluble, slightly soluble, very slightly soluble, or practically insoluble in pure water at 21°C (ie. requiring from 30, from 100, from 1000 or from 10000 parts water to put 1 part by weight drug compound into solution). In particular, the drug is a basic compound.

Examples of such poorly water-soluble compounds that may be used in the compositions of the invention include nifedipine, itraconazole (described in EP-A-6711),

- 5 saperconazole (see US-A-4916134), (-)-[2S-[2α , 4α (S*)]]-4-[4-[4-[4-[[2-(4-chloro-phenyl]-2-[[(4-methyl-4H-1,2,4-triazol-3-yl)thio]methyl]-1,3-dioxolan-4-yl]methoxy]-phenyl]-1-piperazinyl]phenyl]-2,4-dihydro-2-(1-methylpropyl)-3H-1,2,4-triazol-3-one (Compound 40 in WO96/13499), cisapride (described in EP-A-76530), (B)-N-[4-[2-ethyl-1-(1H-1,2,4-triazol-1-yl)butyl]phenyl]-2-benzothiazolamine
- (described in WO-97/49704); methyl 6,11-dihydro-11-[1-[2-[4-(2-quinolinylmethoxy)phenyl]ethyl]-4-piperidinylidene]-5*H*-imidazo[2,1-b][3]benzazepine-3-carboxylate (described in WO-97/34897);
 4-[[4-amino-6-[(2,6-dichlorophenyl)methyl]-1,3,5-triazin-2-yl]amino]benzonitrile (described in EP-0,834,507);
- 15 (B-cis)-1-[4-[4-[4-[4-[4-[4-(2,4-difluorophenyl)-4-(1*H*-1,2,4-triazol-1-ylmethyl)-1,3-dioxolan-2-yl]methoxy]phenyl]-1-piperazinyl]phenyl]-3-(1-methylethyl)-2-imidazolidinone;

20 imidazolidinone;

3-[2-[3,4-dihydrobenzofuro[3,2-c]pyridin-2(1*H*)-yl]ethyl]-2-methyl-4*H*-pyrido-[1,2-a]pyrimidin-4-one;

N-[2-[4-(4-chlorophenyl)-1-piperazinyl]ethyl]-2-benzothiazolamine;

(B1)-*N*-[4-[2-(dimethylamino)-1-(1*H*-imidazol-1-yl)propyl]phenyl]-2-benzothiazolamine (described in WO-97/49704)

(B)-6-[amino(4-chlorophenyl)(1-methyl-1*H*-imidazol-5-yl)methyl]-4-(3-chlorophenyl)-1-methyl-2(1*H*)-quinolinone;

(B)-N-[4-[2-ethyl-1-(1H-1,2,4-triazol-1-yl)butyl]phenyl]-2-benzothiazolamine;

3-[6-(dimethylamino)-4-methyl-3-pyridinyl]-2,5-dimethyl-*N*,*N*-dipropylpyrazolo[2,3-a]pyrimidin-7-amine monohydrochloride;

(S)-[1-[2-[3-[(2,3-dihydro-1*H*-inden-2-yl)oxy]-4-methoxyphenyl]propyl]-1*H*-imidazol-2-yl]cyanamide; and

(+)-(B-trans)-4-[1-[3,5-bis(trifluoromethyl)benzoyl]-2-(phenylmethyl)-4-piperidinyl]-*N*-(2,6-dimethylphenyl)-1-piperazineacetamide (S)-hydroxybutanedioate (1:1).

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Further suitable active ingredients are those which exert a local physiological effect, as well as those which exert a systemic effect, either after penetrating the mucosa or - in

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the case of oral administration - after transport to the gastro-intestinal tract with saliva. The dosage forms prepared from the compositions according to the present invention are particularly suitable for active ingredients which exert their activity during an extended period of time, i.e. drugs having a half-life of at least several hours.

5 Examples thereof are : analgesic and anti-inflammatory drugs (celecoxib, MK966, L-745,337, NSAIDs, fentanyl, indomethacin, ketoprofen, nabumetone, oxyphenbutazone, paracetamol, phenylbutazone, piroxicam, tramadol) ; anti-arrhythmic drugs (gallopamil, procainamide, quinidine, verapamil) ; antibacterial and antiprotozoal agents

10 (amoxicillin, ampicillin, benzathine penicillin, benzylpenicillin, cefaclor, cefadroxil, cefprozil, cefuroxime axetil, cephalexin, chloramphenicol, chloroquine, ciprofloxacin, clarithromycin, clavulanic acid, clindamycin, doxyxycline, erythromycin, flucloxacillin, halofantrine, isoniazid, kanamycin, lincomycin, mefloquine, minocycline, nafcillin, neomycin, norfloxacin, ofloxacin, oxacillin, phenoxymethyl-

penicillin, pyrimethamine-sulfadoxime, quinine, streptomycin); anti-coagulants
 (warfarin); antidepressants (amitriptyline, amoxapine, butriptyline, clomipramine, desipramine, dothiepin, doxepin, fluoxetine, fluvoxamine, gepirone, imipramine, lithium carbonate, mianserin, milnacipran, nortriptyline, paroxetine, sertraline;
 3-[2-[3,4-dihydrobenzofuro[3,2-c]pyridin-2(1*H*)-yl]ethyl]-2-methyl-4*H*-

20 pyrido[1,2-a]pyrimidin-4-one) ; anti-diabetic drugs (glibenclamide, metformin) ; antiepileptic drugs (carbamazepine, clonazepam, ethosuximide, phenobarbitone, phenytoin, primidone, topiramate, valpromide) ; antifungal agents (amphotericin, clotrimazole, econazole, fluconazole, flucytosine, griseofulvin, itraconazole, ketoconazole, miconazole nitrate, nystatin, terbinafine, voriconazole) ; antigout

25 (benzbromarone, probenecid) ; antihistamines (astemizole, cinnarizine, cyproheptadine, decarboethoxyloratadine, fexofenadine, flunarizine, levocabastine, loratadine, norastemizole, oxatomide, promethazine, terfenadine) ; anti-hypertensive drugs (captopril, clonidine, cyclizine, diazoxide, dihydralazine, enalapril, fosinopril, guanethidine, ketanserin, lisinopril, minoxidil, prazosin, ramipril, rescinnamine,

- 30 reserpine, terazosin) ; anti-muscarinic agents (atropine sulphate, hyoscine) ; antivirals (acyclovir, AZT, ddC, ddI, ganciclovir, loviride, tivirapine, 3TC, delavirdine, indinavir, nelfinavir, ritonavir, saquinavir) ; antineoplastic agents and antimetabolites (adriamycine, cladribine, dactinomycin, daunorubicin, doxorubicin, etoposide, mitomycin, mitoxantrone, paclitaxel, taxol, taxotere, trimetrexate, vincristine,
- vinblastine) ; anti-migraine drugs (alniditan, naratriptan, sumatriptan) ; anti Parkinsonian drugs (bromocryptine mesylate, carbidopa, levodopa, selegiline) ;
 antipsychotic, hypnotic, anxiolytic and sedating agents (alprazolam, buspirone,

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chlordiazepoxide, chlorpromazine, chlorprothixene, clozapine, diazepam, flupenthixol, fluphenazine, flurazepam, haloperidol, 9-hydroxyrisperidone, lorazepam, mazapertine, melperone, methaqualone, olanzapine, oxazepam, pimozide, pipamperone, piracetam, promazine, risperidone, selfotel, seroquel, sertindole, sulpiride, temazepam,

- 5 thioridazine, thiothixene, triazolam, trifluoperazine, trifluperidol, triflupromazine, ziprasidone, zolpidem); anti-stroke agents (lubeluzole, lubeluzole oxide, riluzole, aptiganel, eliprodil, remacemide); antitussive (dextromethorphan, laevodropropizine, noscapine); beta-adrenoceptor blocking agents (atenolol, bupranolol, carvedilol, labetalol, metipranolol, metoprolol, nebivolol, oxprenolol, propanolol); cardiac
- inotropic agents (amrinone, digitoxin, digoxin, milrinone); corticosteroids
 (beclomethasone dipropionate, betamethasone, budesonide, cortisone, dexamethasone, fludrocortisone, hydrocortisone, methylprednisolone, paramethasone, prednisolone, prednisolone, triamcinolone); disinfectants (chlorhexidine); diuretics (acetazolamide, amiloride, benzthiazide, chlorothiazide, chlorthalidone, dichlorphenamide, ethacrynic
- 15 acid, ethoxzolamide, frusemide, hydrochlorothiazide, hydroflumethiazide, isosorbide, polythiazide, spironolactone, triamterene, trichloromethiazide) ; enzymes ; ergot alkaloids (codergocrine, ergotamine, nicergolin) ; essential oils (anethole, anise oil, caraway, cardamom, cassia oil, cineole, cinnamon oil, clove oil, coriander oil, dementholised mint oil, dill oil, eucalyptus oil, eugenol, ginger, lemon oil, mustard oil,
- 20 neroli oil, nutmeg oil, orange oil, peppermint, sage, spearmint, terpineol, thyme) ; gastro-intestinal agents (bromopride, cimetidine, cisapride, clebopride, diphenoxylate, domperidone, famotidine, lansoprazole, loperamide, loperamide oxide, mesalazine, metoclopramide, mosapride, nizatidine, norcisapride, olsalazine, omeprazole, pantoprazole, perprazole, pirenzepine, prucalopride, ranitidine, rabeprazole, ridogrel,
- 25 sulphasalazine) ; haemostatics (aminocaproic acid) ; immunosuppressants (cyclosporin A, tacrolimus) ; lipid regulating agents (atorvastatin, lovastatin, pravastatin, probucol, simvastatin) ; local anaesthetics (benzocaine, lignocaine) ; opioid analgesics (buprenorphine, codeine, dextromoramide, dextropropoxyphene, dihydrocodeine, hydrocodone, oxycodone, morphine, papaverine, pentazocine, pethidine) ;
- 30 parasympathomimetics (eptastigmine, galanthamine, metrifonate, neostigmine, physostigmine, tacrine, donepezil, rivastigmine, milameline, sabcomeline, talsaclidine, xanomeline, memantine, lazabemide) ; sex hormones (androgens : methyltestosterone, oxymetholone, stanozolol ; oestrogens : conjugated oestrogens, ethinyloestradiol, mestranol, oestradiol, oestroil, oestrone ; progestogens ; chlormadinone acetate,
- 35 cyproterone acetate, 17-deacetyl norgestimate, desogestrel, dienogest, dydrogesterone, ethynodiol diacetate, gestodene, 3-keto desogestrel, levonorgestrel, lynestrenol, medroxy-progesterone acetate, megestrol, norethindrone, norethindrone acetate,
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norethisterone, norethisterone acetate, norethynodrel, norgestimate, norgestrel, norgestrienone, progesterone, quingestanol acetate) ; stimulating agents (sildenafil) ; sympathomimetics (ephedrine, clenbuterol, fenoterol, norfenefrine, pseudoephedrine) ; vasodilators (amlodipine, amyl nitrite, buflomedil, buphenine, carbocromen, diltiazem,

5 dipyridamole, glyceryl trinitrate, isosorbide dinitrate, lidoflazine, molsidomine, nicardipine, nifedipine, nimodipine, oxpentifylline, pentaerythritol tetranitrate).

Other examples include the following:

10	alpha-Lipoic acid	lactose	methylxanthine
	8-Methoxypsoralen	lithium salts	phytomenadione
	Allopurinol	magnesium salts	propylthiouracil
	alphaTocopherol	menadione	
	iron salts	methylthiouracil	

15

Drug compounds suitable for use in the compositions of the invention include drugs of all types conventionally administered topically (e.g. in a gel patch) or into an externally voiding body duct, e.g. orally, nasally, aurally, rectally or vaginally. Such drugs include in particular antifungals, calcium channel blockers, antibacterials, antihypertensives,

antivirals, analgesics, apolipoprotein B synthesis inhibitors, and drugs which modify transit of gi tract contents (e.g. antidiarrhoea agents or motility promoters).
 Indeed, the invention is particularly applicable to poorly water-soluble imidazole, triazole, imidazo-benzazepines, nitrophenyl-pyridine, *N.N*^{*}-bisphenyl-piperazine, and *N*-phenoxyalkyl-piperidine derivatives, e.g. the compounds mentioned above and compounds as described in EP-A-6711, WO96/13499 and EP-A-76530.

•

The compositions of the invention may conveniently contain the drug compound at 0.001 to 50% by weight, preferably 0.1 to 35%, more preferably 0.5 to 30%, especially 8 to 25% and most especially 10 to 15% by weight (relative to the total weight of acid,

- 30 cyclodextrin, organic polymer and drug compound). The quantity of drug will of course depend upon the desired dissolution profile, the intrinsic solubility of the drug compound and the drug dosage required where the drug is to be delivered in dosage units (e.g. capsules, coated tablets, etc).
- 35 Thus the present invention also provides pharmaceutical dosage forms comprising a therapeutically effective amount of a composition as described hereinbefore.

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For example if the drug is to be delivered in a standard capsule (e.g. with a 900 mg capacity for a glass thermoplastic system as described in the Examples hereto, and the desired drug dose is 100 mg/capsule) then the quantities and natures of the other composition components may be selected to give the desired drug dissolution profile - in

- 5 general only a small quantity of organic polymer, e.g. 20 to 50 mg, may be necessary, and the balance may be made up from acid and cyclodextrin with the ratio of acid to cyclodextrin being set according to the required dissolution profile, e.g. with 200 to 400 mg cyclodextrin and 450 to 650 mg acid.
- 10 Besides the drug compound, the organic polymer, the acid and the cyclodextrin, the compositions of the invention may contain other conventional pharmaceutical excipients, e.g. flavours, colouring agents, antioxidants, bulking agents, fats, waxes, coating agents, dispersants, suspension fluids (e.g. where the composition coated with a gastric juice resistant coating and dispersed as particles in a suspension fluid such as water or a
- syrup), etc. Preferably such components when in intimate admixture with the drug compound will make up only a minor proportion of the composition, e.g. 0.01 to 10% by weight (relative to the total weight of acid, organic polymer, cyclodextrin and drug compound). However where the composition of the invention is encapsulated or disposed in a carrier (e.g. a fluid or a solid or semi-solid matrix), the further components not in intimate admixture with the drug compound (e.g. coating or encapsulating materials, dispersion media, etc.) may of course make up a minor or major proportion, e.g. 5 to 95% by weight, of the overall composition.
- The compositions of the invention may be prepared by making an intimate admixture of the drug compound, cyclodextrin, acid and organic polymer. This may be effected most straightforwardly by dissolving these components in a liquid solvent therefor and subsequently removing the solvent. Thus viewed from a further aspect the invention provides a process for the preparation of a pharmaceutical composition, said process comprising: dissolving a drug compound, a water-soluble cyclodextrin, a physiologically
- 30 tolerable water-soluble acid and a physiologically tolerable water-soluble organic polymer in a solvent; removing solvent from the resultant solution; optionally forming the resultant product into desired shapes; and optionally coating the resulting product with a physiologically tolerable coating material.
- The solvent used in the process of the invention is preferably a physiologically tolerable material, suitably an organic solvent such as a C_{1-6} alkanol (e.g. ethanol), acetone, DMF, a linear or cyclic ether (e.g. diethyl ether, dimethyl ether, or THF), cyclohexane, DMSO,

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etc. or a solvent mixture that also may comprise water. For an acid with a high melting point, solvents or solvent mixtures which have high boiling points may conveniently be used; generally however the boiling point of the solvent or solvent system will be no more than about 100°C. Such solvents may be used efficiently in the production of the

5 compositions of the invention and the level of residual solvent will be minimal. The solvent may conveniently be removed by evaporation, e.g. under reduced pressure, and as this may leave some solvent residue (e.g. up to 3% by weight) it is particularly desirable to use a solvent such as ethanol (or an ethanol-water mixture) which is a permitted pharmaceutical excipient.

10

If the drug compound is insoluble or poorly soluble in the solvent of choice, the process of the invention may involve dispersion of microparticles (e.g. nanoparticles having a particle size of 1 to 100 nm) of the drug compound in the solvent rather than full dissolution of the drug compound. If this is done, it is desirable that the drug compound particles be as small as possible. Nanoparticles of insoluble compounds may be prepared

15 particles be as small as possible. Nanoparticles of insoluble compounds may be prepared for example by various precipitation techniques or by milling with physiologically tolerable inorganic beads, e.g. of zirconia (EP-0,499,299).

The solvent removal may be essentially complete or it may be incomplete, in the former case to produce a solid or a gel-like solid or semi-solid, and in the latter case to produce a viscous fluid which can for example be filled into capsules.

In general, essentially complete solvent removal will be preferred as the resultant product can then readily be shaped. Shaping may be effected by spray-drying the solution (to provide the product in particulate form), by evaporation of solvent from solution disposed in molds, by molding (e.g. injection molding), by extrusion and the like. In general the product can be formed when hot and allowed to solidify on cooling. The shaped product may likewise be produced in film or sheet form by evaporation or by pouring a heated mass onto a plate and evaporating off the solvent.

30

In one preferred embodiment the product is shaped by filling into (e.g. by pouring or by extrusion) capsule shells, e.g. of gelatin.

The product may be hygroscopic, and thus may be "tacky" if touched by hand due to its absorption of moisture from the skin. Accordingly it is particularly preferred for the product to be provided with a protective coating to prevent moisture uptake during handling. Such coatings may for example take the form of capsule casings (as described above), tablet coatings, protective film or web coatings, and moisture-proof removable

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wrappings. Tablet coatings may be applied in conventional manner and may be such as to dissolve in the mouth or stomach (e.g. sugar or sugar/beeswax coatings), or alternatively may be gastric juice resistant polymers (such as the gastric juice resistant Eudragit® coatings produced by Röhm GmbH) where it is desired that drug uptake

5 should occur in the intestines. Protective films or webs may for example be used where the product is to be applied topically, e.g. for uptake across the skin or a toe or finger nail. In this event a pad of the composition will generally be disposed between an adhesive upper protective layer and a lower removable layer. An example of a topical application form for application on nails and adjoining tissue, e.g. for the treatment of fungal infection, is shown in US-A-5181914.

Where the product is produced in particulate form, e.g. by spray-drying, the particles can be loaded into water-tight administration devices (e.g. spray devices or powder dosing devices such as inhalers) for oral, nasal or topical administration of the

15 particulate. Alternatively particulates may be loaded into capsules or mixed with bulking agents such as lactose, starch, microcrystalline cellulose and mixtures thereof, and compressed to form tablets. In any event, the particles may additionally be provided with one or more coatings, e.g. to provide a delayed or prolonged release administration forms.

20

Generally however it will be preferred to shape the product into individual doses and to provide these with a protective coat, e.g. to produce a capsule, coated tablet or film covered pad single dosage unit.

- 25 While not wishing to be bound by theory it is thought that the advantageous drug compound dissolution profile for the compositions of the invention is achieved as a result of a combination of the effects of the components of the composition on exposure to water or aqueous body fluids. The water and the acid provide a highly acidic microenvironment in which the solubility of the drug compound is increased. This acidic
- 30 microenvironment contains the cyclodextrin which is capable of complexing the solubilized drug causing the production of a supersaturated solution of the drug compound and this supersaturated solution is stabilized by the viscosity enhancing effects of the organic polymer which hinders precipitation of the drug as the pH increases as the microenvironment becomes more dilute as more water enters.

35

Accordingly, in the compositions of the invention, in place of the cyclodextrin it is considered possible to use other compounds capable of complexing the drug compound, in particular host complexants capable (like cyclodextrin) of producing host:guest

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complexes with the drug compound may be used. Likewise, for base solubilized drug compounds, a physiologically tolerable water-soluble base (e.g. an inorganic or organic base such as an alkali metal carbonate (eg. sodium carbonate) ethanolamine, diethanolamine, etc.) may be used in place of the acid, and in place of the organic polymer a

5 water-soluble physiologically tolerable macromolecular (e.g. of molecular weight
 ≥ 1kD) viscosity enhancer may be used; in each case in the quantities specified above for the cyclodextrin, acid and organic polymer respectively.

While the benefits of the compositions of the invention are most pronounced where the drug compound is no more than sparingly soluble, the drug dissolution profiles achievable using the combination of drug, cyclodextrin and acid (or base) are such that particularly improved drug uptake profiles may be achieved even where the drug compound is more soluble. Thus viewed from an alternative aspect the invention provides a pharmaceutical composition comprising in intimate admixture a drug

15 compound, a cyclodextrin, a physiologically tolerable water-soluble acid, and a physiologically tolerable water-soluble organic polymer.

Moreover, where appropriate quantities of cyclodextrin, acid and drug compound are used, it is possible to achieve particularly desirable drug dissolution profiles where the

- 20 organic polymer is used in very small quantities or even omitted. Thus viewed from a further aspect the invention provides a pharmaceutical composition comprising in admixture a no more than sparingly water soluble organic drug compound, a watersoluble physiologically tolerable organic acid and a water-soluble physiologically tolerable cyclodextrin, said acid and cyclodextrin being present at 1.5 to 15 (preferably 2
- to 10, more preferably 2.5 to 6) parts by weight and 1 to 7 (preferably 1.1 to 5, more preferably 1.25 to 4) parts by weight respectively per part by weight of said drug compound.
- As has been mentioned above, the compositions according to the invention can be
 produced with particularly favourable drug dissolution profiles. Thus dissolution may be sufficiently rapid to ensure substantially complete availability of the drug compound for biological uptake (e.g. from the mouth, nose, stomach or vagina) yet sufficiently slow to provide a more prolonged plasma uptake profile (see for example Figure 1 of the accompanying drawings) e.g. by avoidance of drug reprecipitation before the
 composition reaches the stomach.

Such a dissolution profile is thus novel and advantageous in its own right and viewed from a further aspect the invention provides a pharmaceutical composition comprising

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an organic drug compound and at least one water-soluble physiologically tolerable excipient, characterised in that at 5, 15 and 45 minutes after addition of a quantity of said composition containing 100 mg of said drug compound to 600 mL of 0.1N hydrochloric acid at 37°C, from 7 to 25 (preferably 10 to 20, especially 12 to 18) %, 45

- to 70 (preferably 50 to 65, especially 54 to 63) % and at least 96 (preferably at least 97, especially at least 98) % respectively of said drug compound is in solution in said hydrochloric acid. These figures relate to *in vitro* dissolution studies conducted in accordance with the monograph USP 23, <711> Dissolution, pp. 1791-1793.
- For example, in determining the dissolution profiles set out above, the composition is placed without a coating or with a rapidly soluble coating (e.g. a gelatin capsule shell) in 0.1 N HCl (or an other appropriate medium) and the mixture is stirred using the USP-method with a paddle, apparatus 2, at a speed of 50 or 100 rpm.
- 15 The compositions according to the invention may be in any form convenient for topical administration or administration into an externally voiding body cavity (e.g. nose, lungs, mouth, ear, stomach, rectum or vagina). Typical administration forms include patches, tablets, buccal tablets, lozenges, ear-plugs, nose plugs, coated tablets, capsules, suppositories, chewing gum, gels, powders, granules, syrups and dispersions, although
- 20 patches and powders and more especially capsules and coated tablets are preferred. The drug dosage will depend upon the drug compound as well as the species and size of the subject being treated. Typically, dosages will be 0.5 to 1.2, preferably 0.8 to 1.05 times the conventional dosages for the selected drug compound administered by the same route.

25

Further, this invention comprioses a pharmaceutical composition or a pharmaceutical dosage form as described hereinbefore for use in a method of therapy or diagnosis of the human or non-human animal body.

- 30 This invention also relates to a pharmaceutical composition for use in the manufacture of a pharmaceutical dosage form for oral administration to a mammal in need of treatment, characterized in that said dosage form can be administered at any time of the day independently of the food taken in by said mammal.
- 35 Or, in other words, the present invention also concerns the se of a pharmaceutical composition as described hereinbefore for the manufacture of a pharmaceutical dosage form for oral administration to a mammal in need of treatment, characterized in that said

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dosage form can be administered at any time of the day independently of the food taken in by said mammal.

This invention also relates to a method of therapy or diagnosis of the human or non-

5 human animal body which comprises administering to said body a therapeutically or diagnostically effective dose of a pharmaceutical composition according to any one of claims 1 to 12.

This invention also relates to a pharmaceutical package suitable for commercial sale

10 comprising a container, an oral dosage form as claimed in any one of claims 12 to 17, and associated with said package written matter non-limited as to whether the dosage form can be administered with or without food.

The invention will now be described further with reference to the following non-limiting Examples and the accompanying drawings, in which:

Figures 1 and 2 are graphs showing plasma concentrations of the drug (-)-[2S- $[2\alpha,4\alpha(S^*)]]$ -4-[4-[4-[4-[4-[2-(4-chlorophenyl]-2-[[(4-methyl-4H-1,2,4-triazol-3-yl)thio]methyl]-1,3-dioxolan-4-yl]methoxy]phenyl]-1-piperazinyl]phenyl]-2,4-dihydro-2-

20 (1-methylpropyl)-3*H*-1,2,4-triazol-3-one administered in a composition according to the invention and in a conventional administration form (sugar particles coated with the drug and loaded in a gelatin capsule) [see Example 6 for further details]; and

Figure 3 is a dissolution profile for the three itraconazole compositions of Example 2.

25

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Example 1

<u>Glass thermoplastic system composition preparation</u> The following ingredients are mixed in a 250 mL glass flask:

	Drug compound (e.g. itraconazole)	20 g
30	Citric acid monohydrate	100 g

Anhydrous ethanol (100 mL) is added. The glass flask is placed on a steam bath (bain marie) and stirred at 70°C until the drug and acid are completely dissolved (about 10 minutes). Thereafter the following ingredients are added:

35Hydroxypropyl-β-cyclodextrin50 gHydroxypropylmethylcellulose (2910.5 mPa.s)10 g

The flask is placed on the steam bath and stirred at 70°C until dissolution is complete

(about 70 minutes). The solution is then poured onto cleaned stainless steel plates which are then placed in a drying oven for 2 hours at 80°C under vacuum and subsequently for 40°C under vacuum overnight. The plates are then heated to 80°C and the gel residue is scraped off and filled into 900 mg capacity gelatin capsules (size no. 0).

5

Example 2

Composition preparation

Analogously to Example 1, gelatin capsules having the following relative weights of components are prepared:

	(A)	100 mg	Itraconazole
		500 mg	citric acid monohydrate
		275 mg	hydroxypropyl- β -cyclodextrin
		25 mg	Methocel E5
15			
	(B)	100 mg	Itraconazole
		500 mg	citric acid monohydrate
		250 mg	hydroxypropyl-β-cyclodextrin
		50 mg	Methocel E5
20			
	(C)	100 mg	Itraconazole
		500 mg	citric acid monohydrate
		225 mg	hydroxypropyl-β-cyclodextrin
		75 mg	Methocel E5
25			
	(D)*	200 mg	Methyl 6,11-dihydro-11-[1-[2-[4-(2-quinolinylmethoxy)phenyl]ethyl]-
			4-piperidinylidene]-5 <i>H</i> -imidazo[2,1-b]-[3]benzazepine 3-carboxylate
		650 mg	citric acid monohydrate
		250 mg	hydroxypropyl-β-cyclodextrin
30		100	
	(E)	100 mg	$(-)-[2S-[2\alpha,4\alpha(S^*)]]-4-[4-[4-[4-[4-[2-(4-chlorophenyl]]-2-[[(4-methyl-$
			4 <i>H</i> -1,2,4-triazol-3-yl)thio]methyl]-1,3-dioxolan-4-yl]methoxy]-
			phenyl]-1-piperazinyl]phenyl]-2,4-dihydro-2-(1-methylpropyl)-3H-
			1,2,4-triazol-3-one
35		500 mg	citric acid monohydrate
		250 mg	hydroxypropyl-β-cyclodextrin
		50 mg	Methocel E5

* For example 2(D) the composition is loaded into 1100 mg gelatin capsules.

The dissolution profiles of the gels of Example 2(A), (B) and (C) are shown in Figure 3 of the accompanying drawings. These were determined by placing one capsule

- 5 containing 100 mg of itraconazole in 300 mL of stirred 0.1 N HCl at 37°C and observing the percentage of dissolved drug compound at times 0, 5, 15, 30, 45 and 60 minutes (stirring was effected using the USP-method with paddle, apparatus 2, 100 rpm). For Example 2(E), with 100 mg drug compound added to 600 mL of 0.1 N HCl at 37°C, the mean percentages of drug compound in solution at 5, 15, 30 and 45
- 10 minutes were 17.22, 61.18, 92.73 and 98.67 respectively (stirring was effected using the USP-method with paddle, apparatus 2, 100 rpm).

The dissolution profile of Example 2(E) was compared with that of a conventional capsule dosage form in which the gelatin capsule is loaded with sugar particles coated

with 100 mg of (-)-[2S-[2α,4α(S*)]]-4-[4-[4-[4-[[2-(4-chlorophenyl]-2-[[(4-methyl-4H-1,2,4-triazol-3-yl)thio]methyl]-1,3-dioxolan-4-yl]methoxy]phenyl]-1-piperazinyl]-phenyl]-2,4-dihydro-2-(1-methylpropyl)-3H-1,2,4-triazol-3-one. The capsules were placed in 10 ml of 0.1 N HCl at 37°C in glass vials and shaken in a mechanical shaker (100 strokes per minute) and the percentage of drug compound in solution after 0,

20 30 and 60 minutes was determined. The results are set out in Table 1 below.

Table 1

Percentage of drug compound in solution

	Time	Example 2(E)	Conventional Capsule
25	0	0	0
	30	91.26	15.54
	60	101.90	18.39

This clearly shows how much more readily the drug compound is made bioavailable by the compositions of the invention.

Example 3

Effect of organic polymer on supersaturation stability

Aqueous solutions of hydroxypropyl-β-cyclodextrin (HPβCD) and Methocel E5 in

35 300 ml 0.1 N HCl at 37°C were prepared having the concentrations set out in Table 2.The solutions were stirred using the USP-method with paddle, apparatus 2, 150 rpm.

	<u>Sample</u>	<u>HPβCD (mg)</u>	Methocel E5 (mg)
	1	250	250
	2	500	0
	3	250	00
5	4	500	500
	5	0	250
	6	500	250
	7	0	0
	8	250	0
10	9	0	500

To these solutions, with stirring, a concentrated solution of itraconazole in DMF (50 mg/mL) was added dropwise until precipitation was observed. Subsequently the concentration of dissolved itraconazole expressed in mg% (ie. the number of mg dissolved in 100 mL) was observed at 0, 30, 60 and 120 minutes. The results are set out

15 in Table 3 below:

Table 3

	<u>Percenta</u>	<u>ge of dr</u>	ug compo	ound in s	olution					
20	Sample Time (minutes)	1	2	3	4	5	6	7	8	9
	0	59.52	72.10	58.95	75.47	42.65	75.27	42.60	60.95	42.95
	30	62.02	74.40	62.05	78.12	44.85	80.17	44.10	62.92	45.20
25	60	62.52	70.37	62.50	79.47	45.40	80.40	44.97	64.07	46.00
	120	62.79	45.82	63.90	80.77	46.55	81.25	31.32	33.65	47.05
	HPBCD:	1:1	2:0	1:2	2:2	0:1	2:1	0;0	1:0	0:2
	Methoce	l ratio								

These results clearly demonstrate (i) the solubilizing effect of the cyclodextrin (Samples 30 2, 6 and 4 show the highest initial itraconazole concentrations, followed by Samples 8, 1 and 3, with Samples 7, 5 and 9 showing the lowest initial concentrations) and (ii) the stabilizing effect of the organic polymer (Samples 2, 8 and 7 show the greatest drop in itraconazole concentrations over 120 minutes, etc).

35

Example 4

Extended release formulation

Analogously to Example 1, gelatin capsules were prepared containing the following:

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41.55 mg	Cisapride
508.45 mg	citric acid monohydrate
250 mg	hydroxypropyl-β-cyclodextrin
100 mg	Methocel K15M

5

This formulation has a much slower dissolution rate than the compositions of Example 2. However the rate of dissolution is much more close to linear with time and shows much less dependence on the pH of the dissolution medium.

10 Example 5

Nail gel

A gel for application to nails or hooves to effect antifungal treatment is made with the following composition:

nazole	250 n	ng
acid monohydrate	2083 n	ng
xypropyl-β-cyclodextrin	333 n	ng
xypropylmethylcellulose (Methocel E5)	83 n	ng
lrous ethanol	2 n	nl
	nazole acid monohydrate xypropyl-β-cyclodextrin xypropylmethylcellulose (Methocel E5) drous ethanol	nazole250 macid monohydrate2083 moxypropyl-β-cyclodextrin333 moxypropylmethylcellulose (Methocel E5)83 mdrous ethanol2 m

20 Example 6

Body uptake

The plasma concentrations of R 103757 were determined in healthy humans at 0, $\frac{1}{2}$, 1, $\frac{1}{2}$, 2, 3, 4, 6, 8 and 12 hours after oral administration of 100 mg (-)-[2S-[2 α ,4 α (S*)]]-4-[4-[4-[4-[2-(4-chlorophenyl]-2-[[(4-methyl-4H-1,2,4-triazol-3-yl)-

- particles administered after a standard breakfast, (iv) a capsule according to Example
 2(E) administered under fasting conditions and (v) a capsule according to Example
 2(E) administered after a standard breakfast.

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The "standard breakfast" comprised four slices of bread, one slice of ham, one slice of cheese, butter, jelly and two cups of coffee or tea with milk and/or sugar if desired. The 100 mg dose of (-)-[2S-[2 α ,4 α (S*)]]-4-[4-[4-[4-[2-(4-chlorophenyl]-2-[[(4-

5 methyl-4*H*-1,2,4-triazol-3-yl)thio]methyl]-1,3-dioxolan-4-yl]methoxy]phenyl]-1piperazinyl]phenyl]-2,4-dihydro-2-(1-methylpropyl)-3*H*-1,2,4-triazol-3-one was administered just after such a breakfast.

Blood samples of 10 mL were taken to obtain 5 mL plasma. The blood samples were taken, collected in heparinized tubes, and centrifuged at 1000g for 10 minutes within 2 hours of collection. Plasma was transferred into plastic tubes, which were sealed and stored at -70°C until assayed.

The results are shown in Figures 1 and 2 which presents drug concentrations as a

15 function of time. As can be seen, the conventional capsule performs significantly worse than the solution even with fasting. However the capsule according to the invention outperforms the solution after 3 hours whether or not the recipient has fasted and, most surprisingly, completely outperforms the solution where the recipient has not fasted.

20

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Example 7

Effect of pH on dissolution rate

Following the procedure of Example 1, a placebo capsule comprising methylene blue (2,63 mg), citric acid (600 mg), hydroxypropyl- β -cyclodextrin (250 mg) and hydroxy-

25 propylmethylcellulose (Methocel E5, 50 mg) was prepared. The dissolution of these capsules was determinated at various pH values according to the USP method (600 ml medium, 37°C, Apparatus 2 with paddle, 100 rpm). The six media tested were : 0.1N HCl (pH 1.55), 0.01N HCl (pH 2.25), 0.001N HCl (pH 2.75), USP pH 4.5 (pH 4.40), USP pH 6.5 (pH 5.80) and USP pH 7.5 (pH 7.0).

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Time (min.)	0.1N HCl	0.01N HCl	0.001N HCI	USP pH 4.5	USP pH 6.5	USP pH 7.5
0	0.00	0.00	0.00	0.00	0.00	0.00
5	16.62	21.85	16.62	21.40	15.48	16.62
15	60.77	73.75	74.43	71.93	60.55	62.59
30	95.60	104.25	104.93	100.15	102.20	100.83
45	100.83	104.93	105.39	104.48	103.57	104.25
60	102.43	104.70	104.93	105.16	104.48	104.02
pH	1.55	2.25	2.75	4.40	5.80	7.00

The results are set out in table 4 below :

Example 8

5 Following the procedure of Example 1, various drug containing capsules were made having the following relative weights of components :

	А.	100 mg	itraconazole
		500 mg	citric acid
		250 mg	hydroxypropyl-\beta-cyclodextrin
10		50 mg	HPMC E5
	В.	200 mg	methyl 6,11-dihydro-11-[1-[2-[4-(2-quinolinylmethoxy)phenyl]ethyl]-4- piperidinylidene]-5 <i>H</i> -imidazo[2,1-b][3]benzazepine-3-carboxylate
		650 mg	citric acid
		250 mg	hydroxypropyl-β-cyclodextrin
15	C.	100 mg	(-)-[2S-[2α,4α(S*)]]-4-[4-[4-[4-[[2-(4-chlorophenyl)-2-[[(4-methyl- 4 <i>H</i> -1,2,4-triazol-3-yl)thio]methyl]-1,3-dioxolan-4-yl]methoxy]phenyl]-
			1-piperazinyl]phenyl]-2,4-dihydro-2-(1-methylpropyl)-3H-1,2,4-triazol-
			3-one
		500 mg	citric acid
20		250 mg	hydroxypropyl- β -cyclodextrin
		50 mg	HPMC E5
	D.	100 mg	4-[[4-amino-6-[(2,6-dichlorophenyl)methyl]-1,3,5-triazin-2-yl]amino]- benzonitrile
		500 mg	citric acid
25		250 mg	hydroxypropyl-β-cyclodextrin
		50 mg	HPMC E5

E. 5 mg (B)-N-[4-[2-ethyl-1-(1H-1,2,4-triazol-1-yl)butyl]phenyl]-2-benzo-

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		thiazolamine
	500 mg	citric acid
	395 mg	hydroxypropyl-β-cyclodextrin
F.	100 mg	(B-cis)-1-[4-[4-[4-[[4-(2,4-difluorophenyl)-4-(1 <i>H</i> -1,2,4-triazol-1-yl- methyl)-1,3-dioxolan-2-yl]methoxy]phenyl]-1-piperazinyl]phenyl]-3-(1- methylethyl)-2-imidazolidinone
	500 mg	citric acid
	250 mg	hydroxypropyl-β-cyclodextrin
	50 mg	HPMC E5

10

5

The dissolution of these compositions was determined according to the USP method (600 ml 0.1 N HCl, 37°C, Apparatus 2 with paddle, 100 ppm), except formulation (A) where only 300 ml medium was used. The results are set out in the following tables 5-10 :

15 **Table 5** : Formulation (A)

Time (min)	sample 1	sample 2	sample 3
0	0.00	0.00	0.00
5	10.75	9.99	10.69
15	56.61	57.18	59.61
30	85.89	88.98	90.24
45	95.46	99.84	96.87
60	101.94	102.06	102.87

Table 6 : Formulation (B)

Time (min)	0.1N HCl	0.01N HCl	0.001N HCl
0	0.00	0.00	0.00
5	27.00	25.17	21.39
15	92.13	86.94	84.75
30	97.11	96.63	93.09
45	98.64	99.45	94.83
60	100.29	100.08	95.28

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Table 7 : Formulation (C)

	Calculated concentration in % of the active dose						
Time (min)	sample 1	sample 2	sample 3	sample 4	sample 5	sample 6	average
0	0.00	0.00	0.00	0.00	0.00	0.00	0.00
5	13.81	17.28	17.67	19.79	18.56	16.19	17.22
15	58.44	59.10	66.60	63.42	62.46	57.06	61.18
30	92.34	92.94	93.36	92.46	92.52	92.76	92.73
45	98.28	98.94	98.82	99.30	98.52	98.16	98.67
60	100.08	99.54	99.66	100.20	100.02	99.96	99.91

Table 8 : Formulation (D)

	Calculated concentration in % of the active dose
Time (min)	Sample 1
5	0.00
5	7.41
15	49.49
30	86.92
45	99.57
60	99.84
90	101.77
120	103.52
150	103.70

Table 9 : Formulation (E)

	Calculated concentration in % of the active dose						
Time (min)	0.1N HCl	0.01N HCl	0.001 N HCl				
 0	0.00	0.00	0.00				
5	48.72	26.16	24.96				
15	100.92	96.36	94.20				
30	102.48	98.76	95.76				
 45	103.08	102.24	96.96				
60	102.00	102.00	97.80				

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	Calculate	Calculated concentration in % of the active dose					
Time (min)	sample 1	sample 2	sample 3	sample 4	sample 5	sample 6	average
0	0.00	0.00	0.00	0.00	0.00	0.00	0.00
5	12.66	14.76	12.66	12.66	15.36	17.34	14.24
15	54.36	56.82	61.98	64.80	54.78	63.78	59.42
30	94.26	93.96	97.50	98.40	95.58	97.20	96.15
45	100.98	101.28	100.50	101.16	100.68	101.34	100.99
60	101.22	101.34	101.16	101.52	100.86	101.58	101.28

Table 10 : Formulation (F)

Example 9

5 Stability testing of formulation 8 (C)

Capsules of formulation 8(C) were stored for 1 month and 3 months at 40°C, and for 1 year at room temperature. Dissolution measurements were made according to the USP method (600 ml 0.1N HCl, 37°C, paddle apparatus 2,100 rpm). The following results were obtained :

10

Table 11 : after 1 month at 40°C

	Calculated concentration in % of the active dose						
Time (min)	sample 1	sample 2	sample 3	sample 4	sample 5	sample 6	average
0	0.00	0.00	0.00	0.00	0.00	0.00	0.00
5	10.11	10.79	9.94	9.59	14.35	10.33	10.85
15	59.72	55.02	48.97	54.36	66.54	52.38	56.17
30	93.00	90.06	89.70	92.70	95.46	89.16	91.68
45	100.14	98.22	98.94	99.84	99.48	99.18	99.30
60	100.50	100.92	99.36	99.54	100.56	100.26	100.19

Table 12 : after 3 months at 40°C

	Calculated concentration in % of the active dose						
Time (min)	nin) sample 1 sample 2 sample 3 sample 4 sa				sample 5	sample 6	average
	0.00	0.00	0.00	0.00	0.00	0.00	0.00
	5.74	5.90	13.93	11.49	7.56	7.78	8.73
	43.62	45.00	56.76	48.30	43.14	47.76	47.43
	88.80	89.10	89.70	87.96	84.54	84.42	87.42
	99.36	99.96	99.54	99.78	99.18	100.08	99.65
	100.32	100.14	100.92	100.44	101.70	100.50	100.67

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	Calculated concentration in % of the active dose				
Time (min)	sample 1	sample 2	sample 3	average	
0	0.00	0.00	0.00	0.00	
5	14.94	16.14	15.48	15.52	
15	61.98	66.12	67.32	65.14	
30	92.52	91.86	96.12	93.50	
45	99.72	99.60	98.70	99.34	
60	101.10	100.80	99.36	100.42	

Table 13 : after 1 year at room temperature

5 Example 10

Variability in bioavailability of Formulation (D)

The variability in the bioavailability of Formulation (D) in beagle dogs was evaluated as follows. First, two beagle dogs received as single oral administration of a PEG-400 solution comprising 4-[[4-amino-6-[(2,6-dichlorophenyl)methyl]-1,3,5-triazin-2-

10 yl]amino]benzonitrile at a dose of 10 mg/kg. Plasm levels were measured for 32 hours. After 7 days, the same dogs were now treated with a single oral capsule comprising the formula (D) at 10 mg/kg. Plasm levels were again determined for up to 32 hours after administration. The individual results are as follows.

			Plasma lev	vels (ng/ml)
Formulation	Day	Time	Dog 1	Dog 2
PEG-400 solution	0	0 h	5.8	NQ
		0.5 h	141	63.2
		1 h	247	158
		2 h	291	141
		4 h	534	200
		6 h	368	171
	Į	8 h	246	141
	1	24 h	95.2	47.4
		32 h	36.1	20.9

			Plasma lev	vels (ng/ml)
Formulation	Day	Time	Dog 1	Dog 2
GTS capsule	7	0 h	NQ	NA
		0.5 h	24.2	68.5
		1 h	567	600
		2 h	850	859
		4 h	461	492
		6 h	288	343
		8 h	237	207
	8	24 h	74.0	32.9
		32 h	32.7	10.1

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NQ : not quantifiable by the HPLC method (< 5.0 ng/ml).

Surprisingly, the plasm levels obtained after administration of the capsules comprising formula (D) are very much more similar to one another in the two test animals than those obtained after administration of the PEG 400 solution.

Example 11

5

Permeation and accumulation of itraconazole through and in human skin A Franz cell was fitted with fresh whole human skin and its receptor filled with a 20%

- (w/v) solution of hydroxypropyl-β-cyclodextrin in water. A Finn Chambers patch was filled with Formulation 8(A) and was then placed on the skin wetted with a small amount of phosphate buffered saline. Samples of the receptor solution were withdrawn at regular intervals and the presence of itraconazole in the solution was measured using high performance liquid chromatography. At no time point could any
- trace of itraconazole be detected, indicating that this compound did not penetrate whole human skin. At the end of the experiment the skin was thoroughly washed and then extracted in order to determine the amount of itraconazole accumulated in the skin. A mean value of $12.2 \,\mu g/cm^2$ could be calculated from the results of 8 independent experiments.

<u>Claims</u>

- 1. A pharmaceutical composition comprising a no more than sparingly water-soluble drug compound, a cyclodextrin, a physiologically tolerable water-soluble acid, and a
- 5 physiologically tolerable water-soluble organic polymer.
 - 2. The composition of claim 1 characterised in that the weight ratios of drug compound to acid and of drug compound to cyclodextrin are no more than 2:1.
- 10 3. The composition of claim 1 or 2 characterized in that the physical state of said composition is a glass thermoplastic phase.
 - 4. The composition of claim 3 wherein the cyclodextrin is 2-hydroxypropyl-βcyclodextrin.

15

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- 5. The composition of claim 3 wherein the acid is selected from the group comprising citric, fumaric, tartaric, maleic, malic, succinic, oxalic, malonic, benzoic, mandelic and ascorbic acid.
- 20 6. The composition of claim 5 wherein the acid is citric acid.
 - 7. The composition of claim 3 wherein the polymer is selected from the group comprising
 - alkylcelluloses such as methylcellulose,
- hydroxyakylcelluloses such as hydroxymethylcellulose, hydroxyethylcellulose,
 hydroxypropylcellulose and hydroxybutylcellulose,
 - hydroxyalkyl alkylcelluloses such as hydroxyethyl methylcellulose and hydroxypropyl methylcellulose,
 - carboxyalkylcelluloses such as carboxymethylcellulose,
 - alkali metal salts of carboxyalkylcelluloses such as sodium
 - carboxymethylcellulose,
 - carboxyalkylalkylcelluloses such as carboxymethylethylcellulose,
 - carboxyalkylcellulose esters,
 - starches,
 - pectins such as sodium carboxymethylamylopectin,
 - chitin derivates such as chitosan,
 - heparin and heparinoids,

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- polysaccharides such as alginic acid, alkali metal and ammonium salts thereof, carrageenans, galactomannans, tragacanth, agar-agar, gum arabic, guargum and xanthan gum,
- polyacrylic acids and the salts thereof,
- polymethacrylic acids and the salts thereof, methacrylate copolymers,
 - polyvinylalcohol,
 - polyvinylpyrrolidone, copolymers of polyvinylpyrrolidone with vinyl acetate,
 - polyalkylene oxides such as polyethylene oxide and polypropylene oxide and copolymers of ethylene oxide and propylene oxide, e.g. poloxamers and
- 10 poloxamines.

5

- 8. The composition of claim 7 wherein the polymer has an apparent viscosity of 1 100 mPa.s when dissolved in a 2% aqueous solution at 20°C.
- 15 9. The composition of claim 8 wherein the polymer is hydroxypropylmethylcellulose.
 - 10. The composition of claim 3 wherein the drug is a basic compound.
- 11. A composition according to any one of the preceding claims that dissolves rapidly
 in body fluids, characterized in that it comprises from 50 to 95 % by weight of acid.
 - 12. A composition according to any one of the preceding claims that provides sustained release of the drug, characterized in that it comprises a water soluble polymer having an apparent viscosity of more than 1,000 mPa.s when dissolved in a 2% aqueous solution at 20°C.
 - 13. A pharmaceutical dosage form comprising a therapeutically effective amount of a pharmaceutical composition as defined in any one of the preceding claims.
- 30

- 14. The dosage form of claim 13 adapted for topical administration or administration into an externally voiding body cavity such as the nose, lungs, mouth, ear, stomach, rectum and vagina.
- 35 15. The dosage form of claim 13 wherein said composition is filled into a standard capsule, or alternatively is mixed with bulking agents and compressed into tablets.
 - 16. The dosage form of claim 13, characterised in that at 5, 15 and 45 minutes after

addition of said dosage form to 0.1N hydrochloric acid at 37° C in the dissolution test set forth in USP test <711> in a USP-2 dissolution apparatus equiped with a paddle, from 7 to 25%, 45 to 70% and at least 96% respectively of drug is dissolved in said 0.1 N hydrochloric acid.

5

- 17. A pharmaceutical composition according to any one of claims 1 to 12 or a pharmaceutical dosage form according to any one of claims 13 to 17 for use in a method of therapy or diagnosis of the human or non-human animal body.
- 10 18. A pharmaceutical composition according to any one of claims 1 to 12 for use in the manufacture of a pharmaceutical dosage form for oral administration to a mammal in need of treatment, characterized in that said dosage form can be administered at any time of the day independently of the food taken in by said mammal.
- 15 19. Use of a pharmaceutical composition according to any one of claims 1 to 12 for the manufacture of a pharmaceutical dosage form for oral administration to a mammal in need of treatment, characterized in that said dosage form can be administered at any time of the day independently of the food taken in by said mammal.
- 20 20. A method of therapy or diagnosis of the human or non-human animal body which comprises administering to said body a therapeutically or diagnostically effective dose of a pharmaceutical composition according to any one of claims 1 to 12.
 - 21. A pharmaceutical package suitable for commercial sale comprising a container, an oral dosage form as claimed in any one of claims 12 to 17, and associated with said package written matter non-limited as to whether the dosage form can be administered with or without food.

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0	treatment (ii)
	treatment (iii)

2/3





—	
-0	treatment (iv)
- []	treatment (v)

3/3

Figure 3



INTERNATIONAL SEARCH REPORT

Interr Dal Application No PCT/EP 98/03189

A. CLASSIFICATION OF SUBJECT MATTER IPC 6 A61K47/48 A61K A61K9/48 A61K9/14 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) IPC 6 A61K Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Belevant to claim No. Category ' Citation of document, with indication, where appropriate, of the relevant passages X,Y EP 0 689 844 A (TECNIMEDE SOCIEDADE 1.2.4-8. TECNICO ME) 3 January 1996 10,12, 13,15, 17 - 21see the whole document Χ,Υ WO 94 12217 A (INSITE VISION INC) 1,2, 9 June 1994 4-10,12, 13,15, 17-21 see the whole document WO 90 14082 A (CIRD) 29 November 1990 Х 1,2,4, 7-9, 12-14, 17-21 see the whole document -/--X Further documents are listed in the continuation of box C. X Patent family members are listed in annex. * Special categories of cited documents : "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention citation or other special reason (as specified) cannot be considered to involve an inventive step when the document is combined with one or more other such docu-"O" document referring to an oral disclosure, use, exhibition or ments, such combination being obvious to a person skilled other means in the art. "P" document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of theinternational search Date of mailing of the international search report 11 September 1998 18/09/1998 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fischer, W Fax: (+31-70) 340-3016

Form PCT/ISA/210 (second sheet) (July 1992)

INTERNATIONAL SEARCH REPORT

Inter anal Application No PCT/EP 98/03189

C.(Continu	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
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Y	WO 97 18245 A (FARMARC NEDERLAND BV ;GLINTENKAMP LUETA ANN (ZA); PENKLER LAWRENCE) 22 May 1997 see the whole document	1,2,4,5, 7,9,13, 14,17-21
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Form PCT/ISA/210 (patent family annex) (July 1992)

INTERNATIONAL SEARCH REPORT

Interer onal Application No

information on patent family members					PCT/EP 98/03189		
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Electronic Acknowledgement Receipt					
EFS ID:	13457908				
Application Number:	12986310				
International Application Number:					
Confirmation Number:	6100				
Title of Invention:	ORAL FORMULATIONS OF CLADRIBINE				
First Named Inventor/Applicant Name:	Nicholas S. Bodor				
Customer Number:	13974				
Filer:	Mary Katherine Baumeister/Lance Logan				
Filer Authorized By:	Mary Katherine Baumeister				
Attorney Docket Number:	20009904-0067				
Receipt Date:	09-AUG-2012				
Filing Date:	07-JAN-2011				
Time Stamp:	14:17:43				
Application Type:	Utility under 35 USC 111(a)				

Payment information:

Submitted wi	th Payment	no						
File Listing:								
Document Number	Document Description	File Name	File Size(Bytes)/ Message Digest	Multi Part /.zip	Pages (if appl.)			
1	Information Disclosure Statement (IDS)	IDS pdf	819507	no	4			
	Form (SB08)		91c879bacd3af8d60c8616d1b198aba73d0 1f89e					
Warnings:								
Information: 173								

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2	Foreign Reference	WO9962958A1.pdf	1414115 89f6c06179c66d2454eeffad3fe2e64373c57 e3f	no	36			
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Information:								
3	Foreign Reference	WO9855148A1.pdf	1986336	no	42			
			dcde016de9bf3453b97622aa85beec9a086 0b662					
Warnings:								
Information:								
4	Non Patent Literature	Drugs-in-Japan.pdf	409972	no	4			
			5d22071e7012d1582746081e5d59683572 3ca228					
Warnings:								
Information:					1			
5	Non Patent Literature	JPOAwTranslation.pdf	362379	no	8			
			6000e805afa0ccdaa48350addcafec83dc6e b660					
Warnings:								
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the applicati	on.							

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UNITED STATES PATENT AND TRADEMARK OFFICE United States Patent and Trademark Office Address: COMMISSIONER FOR PATENTS P.O. Box 1450 Alexandria, Virginia 22313-1450 www.uspto.gov									
APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.					
12/986,310	01/07/2011	Nicholas S. Bodor	20009904-0067	6100					
13974 SNR DENTON	7590 11/08/2012 JUSLLP		EXAMINER						
P.O. BOX 0610 Chicago II 600)80 606 1080	LAU, JONATHAN S							
Cincago, iL 00	500-1080	ART UNIT	PAPER NUMBER						
			1623						
			NOTIFICATION DATE	DELIVERY MODE					
			11/08/2012	ELECTRONIC					

Please find below and/or attached an Office communication concerning this application or proceeding.

The time period for reply, if any, is set in the attached communication.

Notice of the Office communication was sent electronically on above-indicated "Notification Date" to the following e-mail address(es):

patents@snrdenton.com martin.bruehs@snrdenton.com

	Application No.	Applicant(s)					
	12/986,310	BODOR ET AL.					
Office Action Summary	Examiner	Art Unit					
	Jonathan S. Lau	1623					
The MAILING DATE of this communication app Period for Reply	ears on the cover sheet with the	correspondence address					
 A SHORTENED STATUTORY PERIOD FOR REPLY IS SET TO EXPIRE <u>1</u> MONTH(S) OR THIRTY (30) DAYS, WHICHEVER IS LONGER, FROM THE MAILING DATE OF THIS COMMUNICATION. Extensions of time may be available under the provisions of 37 CFR 1.136(a). In no event, however, may a reply be timely filed after SIX (6) MONTHS from the mailing date of this communication. If NO period for reply is specified above, the maximum statutory period will apply and will expire SIX (6) MONTHS from the mailing date of this communication. Failure to reply within the set or extended period for reply will, by statute, cause the application to become ABANDONED (35 U.S.C. § 133). Any reply received by the Office later than three months after the mailing date of this communication, even if timely filed, may reduce any 							
Status							
 1) Responsive to communication(s) filed on <u>07 January 2011</u>. 2a) This action is FINAL. 2b) This action is non-final. 3) An election was made by the applicant in response to a restriction requirement set forth during the interview on; the restriction requirement and election have been incorporated into this action. 4) Since this application is in condition for allowance except for formal matters, prosecution as to the merits is alread in accordance with the prostate under Fx parts Over 1025 O D, 11, 450 O D, 200 							
Disposition of Claims							
5a) Of the above claim(s) is/are withdrawn from consideration. 6) □ Claim(s) is/are allowed. 7) □ Claim(s) is/are rejected. 8) □ Claim(s) is/are objected to. 9) ⊠ Claim(s) <u>1-64</u> are subject to restriction and/or election requirement.							
Application Papers							
 10) The specification is objected to by the Examiner. 11) The drawing(s) filed on is/are: a) accepted or b) objected to by the Examiner. Applicant may not request that any objection to the drawing(s) be held in abeyance. See 37 CFR 1.85(a). Replacement drawing sheet(s) including the correction is required if the drawing(s) is objected to. See 37 CFR 1.121(d). 12) The oath or declaration is objected to by the Examiner. Note the attached Office Action or form PTO-152. 							
Priority under 35 U.S.C. § 119							
 13) Acknowledgment is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f). a) All b) Some * c) None of: Certified copies of the priority documents have been received. Certified copies of the priority documents have been received in Application No 3. Copies of the certified copies of the priority documents have been received in this National Stage application from the International Bureau (PCT Rule 17.2(a)). * See the attached detailed Office action for a list of the certified copies not received. 							
Attachment(s) 1) Notice of References Cited (PTO-892) 2) Notice of Draftsperson's Patent Drawing Review (PTO-948) 3) Information Disclosure Statement(s) (PTO/SB/08) Paper No(s)/Mail Date	4) Interview Summary Paper No(s)/Mail D 5) Notice of Informal 6) Other:	y (PTO-413) bate Patent Application					

DETAILED ACTION

This Office Action details a restriction requirement and one election of species requirement.

Restriction Requirement

Restriction to one of the following inventions is required under 35 U.S.C. 121:

- Claims 1-8, 25-31, 46-61, drawn to a cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex and pharmaceutical compositions comprising thereof, classified in class 536, subclass 27.7.
- II. Claims 9-24 and 62-64, drawn to a method for enhancing the oral bioavailability of cladribine or treatment of symptoms of a cladribineresponsive condition in a subject suffering from said symptoms comprising orally administering to a subject in need thereof a pharmaceutical composition comprising said cladribine-cyclodextrin complex, classified in class 514, subclass 45.
- III. Claims 32-45, drawn to a method of making cladribine-cyclodextrin
 complex which is an intimate amorphous admixture of (a) an amorphous
 inclusion complex of cladribine with an amorphous cyclodextrin and (b)

Application/Control Number: 12/986,310 Art Unit: 1623

amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex, classified in class 536, subclass 55.3. The inventions are distinct, each from the other because of the following reasons:

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Inventions I and II are related as product and process of use. The inventions can be shown to be distinct if either or both of the following can be shown: (1) the process for using the product as claimed can be practiced with another materially different product or (2) the product as claimed can be used in a materially different process of using that product. See MPEP § 806.05(h). In the instant case the product as claimed can be used in a materially different process of using that product such as an *in vivo* efficacy study of cladribine as a function of solubility.

Inventions I and III are related as process of making and product made. The inventions are distinct if either or both of the following can be shown: (1) that the process as claimed can be used to make another and materially different product or (2) that the product as claimed can be made by another and materially different process (MPEP § 806.05(f)). In the instant case the product as claimed can be made by another and materially different process and materially different process such as by melt-extrusion to make a glassy or amorphous material.

Inventions II and III are directed to related processes. The related inventions are distinct if: (1) the inventions as claimed are either not capable of use together or can have a materially different design, mode of operation, function, or effect; (2) the inventions do not overlap in scope, i.e., are mutually exclusive; and (3) the inventions as claimed are not obvious variants. See MPEP § 806.05(j). In the instant case, the

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inventions as claimed have a materially different design, function and effect, as the process of Group II is drawn to treating a subject suffering from a disease or condition by adminstering said cladribine-cyclodextrin complex whereas the process of Group III is drawn to making said cladribine-cyclodextrin complex. Furthermore, the inventions as claimed do not encompass overlapping subject matter and there is nothing of record to show them to be obvious variants.

Restriction for examination purposes as indicated is proper because all these inventions listed in this action are independent or distinct for the reasons given above and there would be a serious search and/or examination burden if restriction were not required because at least the following reason(s) apply:

(a) the inventions have attained recognition in the art as a separate subject for inventive effort in view of their different classification;

(b) the inventions have acquired a separate status in the art due to their recognized divergent subject matter;

(c) the inventions require a different field of search (for example, employing different search queries for structural features of a cladribine-cyclodextrin complex, methods of treatment of a subject suffering from a cladribine-responsive condition, or processing conditions for making a cladribine-cyclodextrin complex).

Applicant is advised that the reply to this requirement to be complete <u>must</u> include (i) an election of a invention to be examined even though the requirement

may be traversed (37 CFR 1.143) and (ii) identification of the claims encompassing the elected invention.

The election of an invention may be made with or without traverse. To reserve a right to petition, the election must be made with traverse. If the reply does not distinctly and specifically point out supposed errors in the restriction requirement, the election shall be treated as an election without traverse. Traversal must be presented at the time of election in order to be considered timely. Failure to timely traverse the requirement will result in the loss of right to petition under 37 CFR 1.144. If claims are added after the election, applicant must indicate which of these claims are readable upon the elected invention.

Should applicant traverse on the ground that the inventions are not patentably distinct, applicant should submit evidence or identify such evidence now of record showing the inventions to be obvious variants or clearly admit on the record that this is the case. In either instance, if the examiner finds one of the inventions unpatentable over the prior art, the evidence or admission may be used in a rejection under 35 U.S.C. 103(a) of the other invention.

Election of Species Requirement

If Applicant elects the invention of Group II, Applicant is further required to elect from the following Election of Species.

Page 5
This application contains claims directed to the following patentably distinct species of cladribine-responsive condition treated. The species are independent or distinct because the different cladribine-responsive conditions treated are characterized by different symptoms and define different patient populations to be treated. In addition, these species are not obvious variants of each other based on the current record.

Examples of cladribine-responsive condition treated are:

a) multiple sclerosis disclosed in claims 19 and 20,

b) rheumatoid arthritis disclosed in claim 19, and

c) leukemia disclosed in claim 19.

Applicant is required under 35 U.S.C. 121 to elect a single disclosed species, or a single grouping of patentably indistinct species, for prosecution on the merits to which the claims shall be restricted if no generic claim is finally held to be allowable. Currently, claims 9-19, 21-24 and 62-64 are generic or subgeneric.

There is a search and/or examination burden for the patentably distinct species as set forth above because at least the following reason(s) apply:

(c) the species require a different field of search (for example, employing different search queries for different cladribine-responsive conditions treated characterized by different symptoms and different patient populations).

Applicant is advised that the reply to this requirement to be complete <u>must</u> include (i) an election of a species or a grouping of patentably indistinct species to be examined even though the requirement <u>may</u> be traversed (37 CFR 1.143) and

(ii) identification of the claims encompassing the elected species or grouping of **patentably indistinct species**, including any claims subsequently added. An argument that a claim is allowable or that all claims are generic is considered nonresponsive unless accompanied by an election.

The election may be made with or without traverse. To preserve a right to petition, the election must be made with traverse. If the reply does not distinctly and specifically point out supposed errors in the election of species requirement, the election shall be treated as an election without traverse. Traversal must be presented at the time of election in order to be considered timely. Failure to timely traverse the requirement will result in the loss of right to petition under 37 CFR 1.144. If claims are added after the election, applicant must indicate which of these claims are readable on the elected species or grouping of patentably indistinct species.

Should applicant traverse on the ground that the species, or groupings of patentably indistinct species from which election is required, are not patentably distinct, applicant should submit evidence or identify such evidence now of record showing them to be obvious variants or clearly admit on the record that this is the case. In either instance, if the examiner finds one of the species unpatentable over the prior art, the evidence or admission may be used in a rejection under 35 U.S.C. 103(a) of the other species.

Upon the allowance of a generic claim, applicant will be entitled to consideration of claims to additional species which depend from or otherwise require all the limitations of an allowable generic claim as provided by 37 CFR 1.141.

Due to the complexity of the restriction and species election requirement, no telephone communication was made. See MPEP 812.01.

Applicant is reminded that upon the cancellation of claims to a non-elected invention, the inventorship must be amended in compliance with 37 CFR 1.48(b) if one or more of the currently named inventors is no longer an inventor of at least one claim remaining in the application. Any amendment of inventorship must be accompanied by a request under 37 CFR 1.48(b) and by the fee required under 37 CFR 1.17(i).

The examiner has required restriction between product or apparatus claims and process claims. Where applicant elects claims directed to the product/apparatus, and all product/apparatus claims are subsequently found allowable, withdrawn process claims that include all the limitations of the allowable product/apparatus claims should be considered for rejoinder. All claims directed to a nonelected process invention must include all the limitations of an allowable product/apparatus claim for that process invention to be rejoined.

In the event of rejoinder, the requirement for restriction between the product/apparatus claims and the rejoined process claims will be withdrawn, and the rejoined process claims will be fully examined for patentability in accordance with 37 CFR 1.104. Thus, to be allowable, the rejoined claims must meet all criteria for patentability including the requirements of 35 U.S.C. 101, 102, 103 and 112. Until all

claims to the elected product/apparatus are found allowable, an otherwise proper restriction requirement between product/apparatus claims and process claims may be maintained. Withdrawn process claims that are not commensurate in scope with an allowable product/apparatus claim will not be rejoined. See MPEP § 821.04. Additionally, in order for rejoinder to occur, applicant is advised that the process claims should be amended during prosecution to require the limitations of the product/apparatus claims. **Failure to do so may result in no rejoinder.** Further, note that the prohibition against double patenting rejections of 35 U.S.C. 121 does not apply where the restriction requirement is withdrawn by the examiner before the patent

issues. See MPEP § 804.01.

Conclusion

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Jonathan S. Lau whose telephone number is (571)270-3531. The examiner can normally be reached on Monday - Thursday, 9 am - 4 pm EST.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Shaojia Anna Jiang can be reached on 571-272-0627. The fax phone number for the organization where this application or proceeding is assigned is 571-273-8300.

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Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free). If you would like assistance from a USPTO Customer Service Representative or access to the automated information

system, call 800-786-9199 (IN USA OR CANADA) or 571-272-1000.

/Jonathan S Lau/ Examiner, Art Unit 1623 /SHAOJIA ANNA JIANG/ Supervisory Patent Examiner Art Unit 1623

Index of Oleime					A	Application/Control No.					Applicant(s)/Patent Under Reexamination				
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PATENT Attorney Docket No.: 20009904-0067

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

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Applicant: Nicholas S. BODOR et al.

Application No.: 12/986,310

Filed: January 7, 2011

Title: ORAL FORMULATIONS OF CLADRIBINE MAIL STOP AMENDMENT

Examiner: Jonathan S. LAU

Group Art Unit: 1623

Confirmation No.: 6100

RESPONSE TO RESTRICTION REQUIREMENT

Commissioner for Patents P.O. Box 1450 Alexandria, VA 22313-1450

Sir:

In response to the Restriction Requirement dated November 8, 2012, Applicants provide the following remarks.

The Examiner has required restriction to one of the following inventions:

Group I Claims 1-8, 25-31 and 46-61, drawn to a cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex and pharmaceutical compositions comprising thereof, classified in class 536, subclass 27.7.;

Group II Claims 9-24 and 62-64, drawn to a method for enhancing the oral bioavailability of cladribine or treatment of symptoms of a cladribine-responsive condition in a subject suffering from said symptoms comprising orally administering to a subject in need thereof a pharmaceutical composition comprising said cladribine-cyclodextrin complex, classified in class 51 4, subclass 45; and

Group III Claims 32-45, drawn to a method of making a cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex, classified in class 536, subclass 55.3.

In response to the Office's Restriction Requirement, Applicants elect, with

traverse, Group I, Claims 1-8, 25-31 and 46-61, drawn to a cladribine-cyclodextrin

complex which is an intimate amorphous admixture of (a) an amorphous inclusion

complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex and pharmaceutical compositions comprising thereof, classified in class 536, subclass 27.7.

In electing the above invention, Applicants reserve the right to request rejoinder under M.P.E.P § 821.04 of process claims that depend from or otherwise include all of the features of an allowable composition claim.

For at least the reasons that follow, Applicants respectfully request reconsideration and withdrawal of the requirement.

For proper restriction between patentably distinct inventions: (1) the inventions must be independent or distinct as claimed; <u>and</u> (2) there would be a <u>serious</u> burden on the Examiner if restriction is not required. See, M.P.E.P. § 803.

Applicants further submit that it is premature to require such restrictions in the absence of having first conducted searches on the claimed inventions to determine the extent of the burden. Absent such searches, it is difficult to maintain that a review of all of the cited claims would be a <u>serious</u> burden on the Examiner. At the very least, such initial searches should be conducted for efficiency purposes so that the searches would not need to be repeated in the event the Applicant was required to file one or more divisional applications. Applicants therefore believe that such a restriction is not required.

Accordingly, for the at least the above reasons, the Requirement for Restriction is improper and should be withdrawn.

For at least these reasons and to avoid duplicate efforts at the Patent Office and undue delay and expense to Applicants, reconsideration and withdrawal of the Restriction Requirement are respectfully requested.

If there are any questions concerning this Response or the application in general,

Applicants invite the Examiner to telephone the undersigned at the below-listed number.

Respectfully submitted,

SNR Denton US LLP

Date January 8, 2013

Mary Kathouse & By:

Mary Katherine Baumeister Registration No. 26,254

Customer No. 13974 202 408 6400

Electronic Patent Application Fee Transmittal									
Application Number:	12986310								
Filing Date:	07-Jan-2011								
Title of Invention:	ORAL FORMULATIONS OF CLADRIBINE								
First Named Inventor/Applicant Name:	Nicholas S. Bodor								
Filer:	Mary Katherine Baumeister/Rebecca Brimmer								
Attorney Docket Number:	20009904-0067								
Filed as Large Entity									
Utility under 35 USC 111(a) Filing Fees									
Description		Fee Code	Quantity	Amount	Sub-Total in USD(\$)				
Basic Filing:									
Pages:									
Claims:									
Miscellaneous-Filing:									
Petition:									
Patent-Appeals-and-Interference:									
Post-Allowance-and-Post-Issuance:									
Extension-of-Time:									
Extension - 1 month with \$0 paid	-	192 1251	1	150	150				

Description	Fee Code	Quantity	Amount	Sub-Total in USD(\$)
Miscellaneous:				
	Tot	al in USD) (\$)	150

Electronic Acl	knowledgement Receipt
EFS ID:	14646869
Application Number:	12986310
International Application Number:	
Confirmation Number:	6100
Title of Invention:	ORAL FORMULATIONS OF CLADRIBINE
First Named Inventor/Applicant Name:	Nicholas S. Bodor
Customer Number:	13974
Filer:	Mary Katherine Baumeister/Rebecca Brimmer
Filer Authorized By:	Mary Katherine Baumeister
Attorney Docket Number:	20009904-0067
Receipt Date:	08-JAN-2013
Filing Date:	07-JAN-2011
Time Stamp:	18:22:19
Application Type:	Utility under 35 USC 111(a)

Payment information:

Submitted with Payment	yes					
Payment Type	Credit Card					
Payment was successfully received in RAM	\$150					
RAM confirmation Number	5468					
Deposit Account	193140					
Authorized User	BAUMEISTER, MARY KATHERINE					
The Director of the USPTO is hereby authorized to charge	e indicated fees and credit any overpayment as follows:					
Charge any Additional Fees required under 37 C.F.R. Section 1.16 (National application filing, search, and examination fees)						
Charge any Additional Fees required under 37 C.F.R. Section 194 (Patent application and reexamination processing fees)						

Charge any Additional Fees required under 37 C.F.R. Section 1.19 (Document supply fees)

Charge any Additional Fees required under 37 C.F.R. Section 1.20 (Post Issuance fees)

Charge any Additional Fees required under 37 C.F.R. Section 1.21 (Miscellaneous fees and charges)

File Listin	g:									
Document Number	Document Description	File Name	File Size(Bytes)/ Message Digest	Multi Part /.zip	Pages (if appl.)					
1	Extension of Time	EOT.pdf	57377	no	1					
			f70356080d711c1899930f69b731d4df8e3f 0874							
Warnings:										
Information:				i						
2	Response to Election / Restriction Filed	Response.pdf	112119	no	4					
			12d73a32acd51762c9bb88eb6c1a59de623 fd15f							
Warnings:										
Information:										
3	Fee Worksheet (SB06)	fee-info.pdf	30461	no	2					
			9763fd0735dc01df125006f28c0af42b22da 9504							
Warnings:										
Information:										
		Total Files Size (in bytes)	: 19	99957						
This Acknow characterized Post Card, as <u>New Applica</u> If a new appl 1.53(b)-(d) an Acknowledg <u>National Star</u> If a timely su U.S.C. 371 an national stag <u>New Internat</u> If a new inter an international of the In	This Acknowledgement Receipt evidences receipt on the noted date by the USPTO of the indicated documents, characterized by the applicant, and including page counts, where applicable. It serves as evidence of receipt similar to a Post Card, as described in MPEP 503. New Applications Under 35 U.S.C. 111 If a new application is being filed and the application includes the necessary components for a filing date (see 37 CFR 1.53(b)-(d) and MPEP 506), a Filing Receipt (37 CFR 1.54) will be issued in due course and the date shown on this Acknowledgement Receipt will establish the filing date of the application. National Stage of an International Application under 35 U.S.C. 371 If a timely submission to enter the national stage of an international application is compliant with the conditions of 35 U.S.C. 371 and other applicable requirements a Form PCT/DO/EO/903 indicating acceptance of the application as a national stage submission under 35 U.S.C. 371 will be issued in addition to the Filing Receipt, in due course. New International Application Filed with the USPTO as a Receiving Office If a new international application is being filed and the international application includes the necessary components for an international Application Number New International application is being filed and the international application includes the necessary components for an international Application Number If a new international Application Number									
national secu the applicati	urity, and the date shown on this Ack on.	nowledgement Receipt will	establish the internat	ional filing	date of					

PTO/SB/22 (09-11) Approved for use through 07/31/2012. OMB 0651-0031 U.S. Patent and Trademark Office; U.S. DEPARTMENT OF COMMERCE

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PETITION FOR EXTENSION OF TIME UN	DER 37 CFI	R 1.136(a)	Docket Number (Optional) 20009904-0067
Application Number 12/986,310	·····		Filed January 7, 2011
For ORAL FORMULATIONS OF CLADRIBINE			
Art Unit 1623 Examiner Jonathan S.	LAU		Confirmation Number 6100
This is a request under the provisions of 37 CFR application.	1.136(a) to ex	ktend the perio	od for filing a reply in the above identified
he requested extension and fee are as follows (check time pe	eriod desired a	and enter the appropriate fee below):
	Fee	Small Er	ntity Fee
One month (37 CFR 1.17(a)(1))	\$150	\$75	\$ <u>150</u>
Two month (37 CFR 1.17(a)(2))	\$570	\$285	\$
Three month (37 CFR 1.17(a)(3))	\$1290	\$645	\$
Four month (37 CFR 1.17(a)(4))	\$2010	\$1005	\$
Five month (37 CFR 1.17(a)(5))	\$2730	\$1365	\$
Applicant claims small entity status. See 37 C	FR 1.27.		
A check in the amount of the fee is enclosed.			
Payment by credit card. Fee being paid conc	urrent with th	e filing of this	EOT.
The Director has already been authorized to	nharaa faaa ir	this conligati	on to a Dononit Assount
The Director has already been additionzed to the Director is hereby authorized to charge a Deposit Account Number <u>19-3140</u> .	iny fees which	n may be requ	lired, or credit any overpayment, to
WARNING: Information on this form may on this form. Provide credit card information and autho	become pub rization on P	lic. Credit ca PTO-2038.	rd information should not be included
I am the applicant/inventor.			
assignee of record of the entire inf Statement under 37 CFR 3.73(b)	terest. See 37 is enclosed (F	CFR 3.71. Form PTO/SB/	/96).
attorney or agent of record. Regis	tration numbe	er <u>26,254</u> .	
attorney or agent acting under 37	CFR 1.34.		
Registration number if acting under	er 37 CFR 1.3	4	
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Mary Katherine Baumeister Type or printed name			202-408-6400 Telephone Number
OTE: Signatures of all the inventors or assigned	s of record of	the entire inte	erest or their representative(s) are
equired. Submit multiple forms if more than one s	signature is re	equired, see b	elow.
I Otal OI TORMS are SUDMITTED.	ne information is a	required to obtain	or retain a benefit by the public which is to file
and by the USPTO to process) an application. Confidentiality is take 6 minutes to complete, including gathering, preparing, a	governed by 35	U.S.C. 122 and 3 completed applic	7 CFR 1.11 and 1.14. This collection is estimated

Box 1450, Alexandria, VA 22313-1450. If you need assistance in completing the form, call 1-800-PTO-9199 and select option 2.

PTO/SB/06 (07-06)

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P/	Under the Par ATENT APPL	perwork Reduction ICATION FE Substitute fo	95, no persons are ERMINATION TO-875	nd to A	d to a collection of information unle Application or Docket Number 12/986,310			plays a valid ing Date 07/2011	OMB control number.		
	AF	PPLICATION /	AS FILE (Column 1	D – PART I) (Column 2)		SMALL		OR	OTH SMA	HER THAN ALL ENTITY
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	SEARCH FEE (37 CFR 1.16(k), (i), (or (m))	N/A		N/A		N/A			N/A	
	EXAMINATION FE (37 CFR 1.16(o), (p),	E or (q))	N/A		N/A		N/A			N/A	
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* lf t ** lf *** l	* If the entry in column 1 is less than the entry in column 2, write "0" in column 3. ** If the "Highest Number Previously Paid For" IN THIS SPACE is less than 20, enter "20". *** If the "Highest Number Previously Paid For" IN THIS SPACE is less than 3, enter "3".										
The	"Highest Number P	reviously Paid For	" (Total or	Independent) is th	e highest number f	oun	d in the appro	priate box in colu	mn 1.		

This collection of information is required by 37 CFR 1.16. The information is required to obtain or retain a benefit by the public which is to file (and by the USPTO to process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.14. This collection is estimated to take 12 minutes to complete, including gathering, preparing, and submitting the completed application form to the USPTO. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and Trademark Office, U.S. Department of Commerce, P.O. Box 1450, Alexandria, VA 22313-1450, DO NOT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS. **SEND TO: Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450, Alexandria, VA 22313-1450**.

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	ed States Patent	TAND TRADEMARK OFFICE	UNITED STATES DEPAR United States Patent and Address: COMMISSIONER F P.O. Box 1450 Alexandria, Virginia 22 www.uspto.gov	TMENT OF COMMERCE Trademark Office FOR PATENTS 313-1450		
APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.		
12/986,310	01/07/2011	Nicholas S. Bodor	20009904-0067	6100		
13974 SNR DENTON	7590 02/25/2013 LUSLLP		EXAMINER			
P.O. BOX 0610 Chicago II 600)80 606 1080		LAU, JONATHAN S			
Cincago, IL 60	500-1080		ART UNIT	PAPER NUMBER		
			1623			
			NOTIFICATION DATE	DELIVERY MODE		
			02/25/2013	ELECTRONIC		

Please find below and/or attached an Office communication concerning this application or proceeding.

The time period for reply, if any, is set in the attached communication.

Notice of the Office communication was sent electronically on above-indicated "Notification Date" to the following e-mail address(es):

patents@snrdenton.com martin.bruehs@snrdenton.com

	Application No.	Applicant(s)								
	12/986,310	BODOR ET AL.								
Office Action Summary	Examiner	Art Unit								
	Jonathan S. Lau	1623								
The MAILING DATE of this communication app	bears on the cover sheet with the c	correspondence address								
Period for Reply										
 A SHORTENED STATUTORY PERIOD FOR REPL' WHICHEVER IS LONGER, FROM THE MAILING D. Extensions of time may be available under the provisions of 37 CFR 1.1 after SIX (6) MONTHS from the mailing date of this communication. If NO period for reply is specified above, the maximum statutory period v. Failure to reply within the set or extended period for reply will, by statute Any reply received by the Office later than three months after the mailing earned patent term adjustment. See 37 CFR 1.704(b). 	Y IS SET TO EXPIRE <u>3</u> MONTH (ATE OF THIS COMMUNICATION 36(a). In no event, however, may a reply be tir will apply and will expire SIX (6) MONTHS from , cause the application to become ABANDONE g date of this communication, even if timely filed	(S) OR THIRTY (30) DAYS, N. nely filed the mailing date of this communication. D (35 U.S.C. § 133). d, may reduce any								
Status										
1) Responsive to communication(s) filed on $08 J_{e}$	anuary 2013.									
2a) This action is FINAL . 2b) This	action is non-final.									
3) An election was made by the applicant in resp	onse to a restriction requirement	set forth during the interview on								
; the restriction requirement and election	have been incorporated into this	s action.								
4) Since this application is in condition for allowa	nce except for formal matters, pro	osecution as to the merits is								
closed in accordance with the practice under E	Ex parte Quayle, 1935 C.D. 11, 4	53 O.G. 213.								
Disposition of Claims										
5) Claim(s) <u>1-64</u> is/are pending in the application										
5a) Of the above claim(s) <u>9-24,32-45 and 62-6</u>	<u>4</u> is/are withdrawn from considera	ation.								
6) Claim(s) is/are allowed.										
7) Claim(s) <u>1-8,25-31 and 46-61</u> is/are rejected.										
8) Claim(s) is/are objected to.										
9) Claim(s) are subject to restriction and/o	r election requirement.									
* If any claims have been determined <u>allowable</u> , you may program at a participating intellectual property office for the http://www.uspto.gov/patents/init_events/pph/index.jsp.or	y be eligible to benefit from the P the corresponding application. Fo r send an inquiry to <u>PPHfeedbac</u>	atent Prosecution Highway or more information, please see k@uspto.gov.								
Application Papers										
10) The specification is objected to by the Examine	er.									
11) The drawing(s) filed on 07 January 2011 is/are	: a) accepted or b) Objected	to by the Examiner.								
Applicant may not request that any objection to the	drawing(s) be held in abeyance. Se	e 37 CFR 1.85(a).								
Replacement drawing sheet(s) including the correct	ion is required if the drawing(s) is ob	jected to. See 37 CFR 1.121(d).								
Priority under 35 U.S.C. § 119										
12) Acknowledgment is made of a claim for foreign a) All b) Some * c) None of:	priority under 35 U.S.C. § 119(a)-(d) or (f).								
1. Certified copies of the priority document	s have been received.									
2. Certified copies of the priority document	s have been received in Applicat	ion No								
3. Copies of the certified copies of the prio	rity documents have been receive	ed in this National Stage								
application from the International Bureau	application from the International Bureau (PCT Rule 17.2(a)).									
* See the attached detailed Office action for a list of the certified copies not received.										
Attachment(s)	_									
1) 🔀 Notice of References Cited (PTO-892)	3) 🛄 Interview Summary Paper No(s)/Mail D	(PTO-413) ate.								
2) Information Disclosure Statement(s) (PTO/SB/08) Paper No(s)/Mail Date 7 Jan 2011, 9 Aug 2012	4) Other:									

DETAILED ACTION

This application is a domestic application, filed 7 Jan 2011; and claims benefit as a CON of 10/551,205, issued as Patent 7,888,328, which is a 371 of PCT/US2004/009387, filed 26 Mar 2004, which claims benefit of provisional application 60/458,922, filed 28 Mar 2003, and claims benefit of provisional application 60/484,756, filed 2 Jul 2003, and claims benefit of provisional application 60/541,247, filed 4 Feb 2004.

Claims 1-64 are pending in the current application. Claims 9-24, 32-45 and 62-64, drawn to non-elected inventions, are withdrawn. Claims 1-8, 25-31 and 46-61 are examined on the merits herein.

However, the parent provisional applications 60/458,922 and 60/484,756 upon which priority is claimed fail to provide adequate support under 35 U.S.C. 112 for the instant claims 1-8, 25-31 and 46-61 of this application since all parent applications are not seen to disclose the complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex in the independent claims 1 and 25. Written description for a cladribine freezedried cyclodextrin complex may be found in 60/458,922 at page 16-18, and a drugcyclodextrin complex in a "highest thermodynamic activity state" may be found in 60/484,756 at page 2, paragraph 2 and cladribine-cyclodextrin complex at page 6,

however no support is found for the complex which is said intimate amorphous admixture. Thus, the filing date of the instant claims is deemed to be provisional application 60/541,247, filed 4 Feb 2004. If applicant disagrees, applicant should present a detailed analysis as to why the claimed subject matter has clear support in the earlier priority applications. Applicant is reminded that such priority for the instant limitations requires written description and enablement under 35 U.S.C. § 112, first paragraph.

Election/Restrictions

Applicant's election with traverse of Group I, claims 1-8, 25-31 and 46-61 in the reply filed on 8 Jan 2013 is acknowledged. The traversal is on the ground(s) that there is no serious burden to examine the different groups of inventions. This is not found persuasive because while the groups of inventions are related to the complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex, the search to examine the different groups of inventions would involve different fields such as pharmacokinetics in the treatment of a subject or specific processing conditions to make said complex. Further, while claims 46-61 recite a product-by-process, determination of patentability is based on the product itself and the patentability of a product does not depend on its method of production. Even without having first conducted searches on the claimed invention it is

Page 3

clear that searching different fields of art, such as the field of pharmacokinetics or

processing conditions to make a complex, would constitute a serious burden.

The requirement is still deemed proper and is therefore made FINAL.

Claims 9-24, 32-45 and 62-64 are withdrawn from further consideration pursuant

Page 4

to 37 CFR 1.142(b), as being drawn to a nonelected inventions, there being no

allowable generic or linking claim. Applicant timely traversed the restriction (election)

requirement in the reply filed on 8 Jan 2013.

Claim Rejections - 35 USC § 103

The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all

obviousness rejections set forth in this Office action:

(a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.

This application currently names joint inventors. In considering patentability of the claims under 35 U.S.C. 103(a), the examiner presumes that the subject matter of the various claims was commonly owned at the time any inventions covered therein were made absent any evidence to the contrary. Applicant is advised of the obligation under 37 CFR 1.56 to point out the inventor and invention dates of each claim that was not commonly owned at the time a later invention was made in order for the examiner to consider the applicability of 35 U.S.C. 103(c) and potential 35 U.S.C. 102(e), (f) or (g) prior art under 35 U.S.C. 103(a).

Claims 1-8, 25-31 and 46-61 are rejected under 35 U.S.C. 103(a) as being unpatentable over Schultz et al. (US Patent 6,194,395, published 27 Feb 2001, provided by Applicant in IDS mailed 7 Jan 2011) in view of Baert et al. (WIPO Publication WO97/18839, published 29 May 1997, cited in PTO-892).

Schultz et al. discloses a solid pharmaceutical oral dosage form of cladribine comprising cladribine and cyclodextrin (column 2, lines 31-39). The disclosed product is substantially identical to the product-by-process. Schultz et al. discloses the use of β and γ -cyclodextrins (column 2, lines 56-58) and derivatives wherein one or more cyclodextrin hydroxy groups are replaced with groups such as methyl, hydroxypropyl, carboxymethyl (column 3, lines 26-27) or sulfobutylcyclodextrins (column 4, lines 22-24). The phrase "one or more cyclodextrin hydroxy groups" combined with the absence of specific structural details of which hydroxyl group is substituted with a methyl group meets the limitation of "randomly methylated β -cyclodextrins". Schultz et al. discloses the solid oral dosage form in the form of a tablet (column 5, lines 37-38) including the excipients sorbitol and magnesium stearate (column 6, lines 2-7), disclosing a product that is substantially identical to a product-by-process meeting the limitations of the instant claims invention. Schultz et al. discloses an oral dosage form comprising up to 15 mg cladribine and cyclodextrin from 100 to 500 mg, which would lead one of skill in the art to instantly envision a cladribine to cyclodextrin ratio ranging from 15 mg:100 mg to 15mg:500 mg, or 1:6.67 to 1:33.3 by weight (column 6, lines 23-31). The instant specification suggests that maximum amount of cladribine which can be complexed gives a weight ratio of 1:10 for the cladribine:cyclodextrin complex. Therefore a

composition comprising the cladribine:cyclodextrin complex that contains a cladribine to cyclodextrin ratio of 1:6.67 describes a composition that comprises a "saturated" complex and meets the limitations of instant claims 2 and 57. Schultz et al. incorporates-by-reference the method of making said solid oral dosage form (Schultz et al.column 5, lines 50-52) disclosed in WIPO Publication WO97/18839, Baert et al., which provides evidence in the embodiment wherein the melt-extruded forms consist essentially of amorphous material (Baert et al. page 8, lines 14-15). Therefore Baert et al. provides evidence that it was recognized in the prior art that the product disclosed by Schultz et al. inherently includes amorphous cladribine-cyclodextrin complex in a solid oral dosage form. Schultz et al. implicitly discloses an oral dosage form comprising up to 15 mg cladribine and cyclodextrin from 100 to 500 mg, or a cladribine to cyclodextrin ratio ranging from 1:6.67 to 1:33.3 by weight (column 6, lines 23-31).

To address the scientific issue regarding the equilibrium presence of both the inclusion and non-inclusion complex, while the equation for the equilibrium of the cladribine-cyclodextrin inclusion complex and the cladribine-cyclodextrin non-inclusion complex would be different for cladribine and cyclodextrin in a solvent versus cladribine and cyclodextrin in a molten state due to the lack of a solvent, the equilibrium and thus equilibrium products, the cladribine-cyclodextrin inclusion complex, would still be inherent in the product disclosed by Schultz et al.

Schultz et al. does not specifically disclose the complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an

Page 6

amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex and the composition comprising no significant amount of free crystalline cladribine therein (instant claim 1). Schultz et al. does not specifically disclose the composition corresponding to a point located on the curve of a phase solubility diagram for saturated complex cladribine-cyclodextrin complexes, said curve defining complex saturated complexes of cladribine in varying concentrations of the cyclodextrin (instant claim 7). Schultz et al. does not specifically disclose the complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex (instant claim 25). Schultz et al. does not specifically disclose the composition comprising a cladribine to cyclodextrin ratio from about 1:10 to about 1:16 (instant claims 5 and 29). Schultz et al. does not specifically disclose the complex wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b) (instant claims 8 and 31). Schultz et al. does not specifically disclose the product-by-process wherein 12.00 parts by weight of cladribine and 172.50 parts by weight of hydroxypropyl- β cyclodextrin are introduced in step (i) of the process, implying a cladribine to cyclodextrin ratio of 1:14.38, or the product-by-process wherein 16.35 parts by weight of cladribine and 172.50 parts by weight of the hydroxypropyl- β -cyclodextrin are introduced in step (i) of the process (instant claim 55), implying a cladribine to cyclodextrin ratio of 1:10.55.

Baert et al. discloses a solid mixture comprising one or more cyclodextrins and an insoluble active ingredient embedded into the cyclodextrin carrier (abstract), and teaches ratios of active ingredient to cyclodextrin of from about 1:100 to 100:1, from about 1:5 and 5:1 and from about 1:3 to 3:1 (page 11, lines 1-5). These ratios are interpreted as mole ratios because Baert et al. teaches the use of different active ingredients with different molecular weights. A mole ratio of active ingredient to cyclodextrin of about 1:3 for cladribine (MW 285.7 g/mol) and β -cyclodextrin (MW 1135 g/mol) gives a ratio by weight of approximately 1:11.9. The ratio of 1:11.9 meets the limitation of both a ratio of about 1:11 and a ratio of about 1:14 according to the nonlimiting definition of "about" as a variance of 20% provided in the instant specification page 9, lines 6-11. Such a saturated complex would consist of only (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex, and being a saturated complex corresponds to a point located on the curve of a phase solubility diagram for saturated complex cladribine-cyclodextrin complexes, said curve defining complex saturated complexes of cladribine in varying concentrations of the cyclodextrin.

It would have been obvious to one of ordinary skill in the art at the time of the invention to practice the solid pharmaceutical oral dosage form of cladribine comprising cladribine and cyclodextrin disclosed by Schultz et al. in the ratios of cladribine and cyclodextrin taught by Baert et al. One of ordinary skill in the art would be motivated to combine the Schultz et al. and Baert et al. because Schultz et al. incorporates-by-

reference Baert et al. and because Baert et al. suggests that improving a similar product according to the teachings of Baert et al. has beneficial properties such as high bioavailability and dissolution rate (Baert et al. page 7, lines 25-27). One of ordinary skill in the art would have an expectation of success because the ratios taught by Baert et al. fall within the range of ratios that is implicitly disclosed by Schultz et al. Schultz et al. in view of Baert et al. does not teach the specific cladribine to cyclodextrin ratios of 1:14.38 or 1:10.55, however these ratios are encompassed by the prior art and Baert et al. suggests optimization of the ratio (Baert et al. page 11, lines 1-5). See also MPEP 2144.05 II.A, "Generally, differences in concentration or temperature will not support the patentability of subject matter encompassed by the prior art unless there is evidence indicating such concentration or temperature is critical." One of ordinary skill in the art would be motivated to optimize the cladribine to cyclodextrin ratio to give the composition comprising no significant amount of free crystalline cladribine therein because Schultz et al. teaches undesireable recrystallization of cladribine in tissue may occur and damage the surround tissue and that complexes with cyclodextrin are known to solubilize the compound (Schultz et al. column 2, lines 1-15). Schultz et al. in view of Baert et al. does not specifically disclose the complex wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b). However, it is well known in the art that the formation of an inclusion complex from a non-inclusion complex is an equilibrium process, and the position of this equilibrium is dependent on the concentrations of the cladribine and cyclodextrin. This molecular

inclusion equilibrium is a process inherent in the formation of the inclusion complex in both aqueous solutions and hot melt liquid mixtures, and Baert et al. teaches variation of the ratio of cladribine to cyclodextrin and hence their relative concentration.

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It is noted that In re Best (195 USPQ 430) and In re Fitzgerald (205 USPQ 594) discuss the support of rejections wherein the prior art discloses subject matter which there is reason to believe inherently includes functions that are newly cited or is identical to a product instantly claimed, namely the position of the equilibrium process governing formation of an inclusion complex and a non-inclusion complex. In such a situation the burden is shifted to the applicants to "prove that subject matter shown to be in the prior art does not possess characteristic relied on" (205 USPQ 594, second column, first full paragraph).

Claims 46-61 are drawn to a product-by-process. The disclosed product is substantially identical to the instantly claimed product-by-process, an amorphous solid pharmaceutical oral dosage form comprising cladribine and cyclodextrin. "[E]ven though product-by-process claims are limited by and defined by the process, determination of patentability is based on the product itself. The patentability of a product does not depend on its method of production. If the product in the product-by-process claim is the same as or obvious from a product of the prior art, the claim is unpatentable even though the prior product was made by a different process." In re Thorpe, 777 F.2d 695, 698, 227 USPQ 964, 966 (Fed. Cir. 1985) (citations omitted) (Claim was directed to a novolac color developer. The process of making the developer was allowed. The difference between the inventive process and the prior art was the

addition of metal oxide and carboxylic acid as separate ingredients instead of adding the more expensive pre-reacted metal carboxylate. The product-by-process claim was rejected because the end product, in both the prior art and the allowed process, ends up containing metal carboxylate. The fact that the metal carboxylate is not directly added, but is instead produced in-situ does not change the end product.). See MPEP 2113.

Double Patenting

The nonstatutory double patenting rejection is based on a judicially created doctrine grounded in public policy (a policy reflected in the statute) so as to prevent the unjustified or improper timewise extension of the "right to exclude" granted by a patent and to prevent possible harassment by multiple assignees. A nonstatutory obviousness-type double patenting rejection is appropriate where the conflicting claims are not identical, but at least one examined application claim is not patentably distinct from the reference claim(s) because the examined application claim is either anticipated by, or would have been obvious over, the reference claim(s). See, e.g., *In re Berg*, 140 F.3d 1428, 46 USPQ2d 1226 (Fed. Cir. 1998); *In re Goodman*, 11 F.3d 1046, 29 USPQ2d 2010 (Fed. Cir. 1993); *In re Longi*, 759 F.2d 887, 225 USPQ 645 (Fed. Cir. 1985); *In re Van Ornum*, 686 F.2d 937, 214 USPQ 761 (CCPA 1982); *In re Vogel*, 422 F.2d 438, 164 USPQ 619 (CCPA 1970); and *In re Thorington*, 418 F.2d 528, 163 USPQ 644 (CCPA 1969).

A timely filed terminal disclaimer in compliance with 37 CFR 1.321(c) or 1.321(d) may be used to overcome an actual or provisional rejection based on a nonstatutory

double patenting ground provided the conflicting application or patent either is shown to be commonly owned with this application, or claims an invention made as a result of activities undertaken within the scope of a joint research agreement.

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Effective January 1, 1994, a registered attorney or agent of record may sign a terminal disclaimer. A terminal disclaimer signed by the assignee must fully comply with 37 CFR 3.73(b).

Claims 1-8, 25-31 and 46-61 are rejected on the ground of nonstatutory double patenting over claims 1-28 of U. S. Patent No. 7,888,328 since the claims, if allowed, would improperly extend the "right to exclude" already granted in the patent.

The subject matter claimed in the instant application is fully disclosed in the patent and is covered by the patent since the patent and the application are claiming common subject matter, as follows: claims 1-28 of U. S. Patent No. 7,888,328 recite a narrower genus of cladribine-cyclodextrin complex and composition comprising thereof wherein said cyclodextrin is hydroxypropyl- β -cyclodextrin and the weight ratio of from about 1:10 to about 1:16. Claims 12-28 of U. S. Patent No. 7,888,328 recite a product-by-process encompassed within the product-by-process of instant claims 46-61. Therefore claims 1-28 of U. S. Patent No. 7,888,328 supports a anticipation-type nonstatutory double patenting rejection over instant claims 1-8, 25-31 and 46-61.

Furthermore, there is no apparent reason why applicant was prevented from presenting claims corresponding to those of the instant application during prosecution of the application which matured into a patent. See *In re Schneller*, 397 F.2d 350, 158 USPQ 210 (CCPA 1968). See also MPEP § 804.

Conclusion

No claim is found to be allowable.

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Jonathan S. Lau whose telephone number is (571)270-3531. The examiner can normally be reached on Monday - Thursday, 9 am - 4 pm EST.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Shaojia Anna Jiang can be reached on 571-272-0627. The fax phone number for the organization where this application or proceeding is assigned is 571-273-8300.

Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free). If you would like assistance from a USPTO Customer Service Representative or access to the automated information system, call 800-786-9199 (IN USA OR CANADA) or 571-272-1000.

/SHAOJIA ANNA JIANG/ Supervisory Patent Examiner, Art Unit 1623 /Jonathan S Lau/ Examiner, Art Unit 1623

Notice of References Cited	Application/Control No. 12/986,310	Applicant(s)/Pater Reexamination BODOR ET AL.	nt Under
Notice of Melerences Cited	Examiner	Art Unit	
	Jonathan S. Lau	1623	Page 1 of 1

U.S. PATENT DOCUMENTS

*		Document Number Country Code-Number-Kind Code	Date MM-YYYY	Name	Classification
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	В	US-			
	С	US-			
	D	US-			
	Е	US-			
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FOREIGN PATENT DOCUMENTS

*		Document Number Country Code-Number-Kind Code	Date MM-YYYY	Country	Name	Classification
	Ν	WO 9718839 A1	05-1997	World Intellect	BAERT et al.	A61K 47/48
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NON-PATENT DOCUMENTS

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INFORMATION DISCLOSURE STATEMENT BY APPLICANT (Not for submission under 37 CFR 1.99)

Application Number		12986310				
Filing Date		2011-01-07				
First Named Inventor	Nicho	las S. BODOR				
Art Unit		1623				
Examiner Name Jonat		han S. LAU				
Attorney Docket Numb	er	20009904-0067				

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INFORMATION DISCLOSURE Application Number 12986310 Filing Date 2011-01-07 First Named Inventor Nicholas S. BODOR Art Unit 1623 Examiner Name Jonathan S. LAU Attorney Docket Number 20009904-0067

Examiner Initials*	Cite No	Inclue (bool publis	nclude name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of the item book, magazine, journal, serial, symposium, catalog, etc), date, pages(s), volume-issue number(s), T ⁵ publisher, city and/or country where published.									
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	Art Unit		1623									
	Examiner Name	Jonat	han S. LAU									
	Attorney Docket Number		20009904-0067									

CERTIFICATION STATEMENT

Please see 37 CFR 1.97 and 1.98 to make the appropriate selection(s):

That each item of information contained in the information disclosure statement was first cited in any communication from a foreign patent office in a counterpart foreign application not more than three months prior to the filing of the information disclosure statement. See 37 CFR 1.97(e)(1).

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See attached certification statement.

The fee set forth in 37 CFR 1.17 (p) has been submitted herewith.

X A certification statement is not submitted herewith.

SIGNATURE

A signature of the applicant or representative is required in accordance with CFR 1.33, 10.18. Please see CFR 1.4(d) for the form of the signature.

Signature	/Mary Katherine Baumeister/	Date (YYYY-MM-DD)	2012-08-09
Name/Print	Mary Katherine Baumeister	Registration Number	26,254

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Ref #	Hits	Search Query	DBs	Default Operator	Plurals	Time Stamp
S1	1	"7888328".pn.	US-PGPUB; USPAT; USOCR	ADJ	ON	2013/02/15 09:12
S2	127	((NICHOLAS) near2 (BODOR)).INV.	US-PGPUB; USPAT; USOCR	ADJ	ON	2013/02/15 09:12
S3	3	((YOGESH) near2 (DANDIKER)).INV.	US-PGPUB; USPAT; USOCR	ADJ	ON	2013/02/15 09:12
S4	127	52 or 53	US-PGPUB; USPAT; USOCR	ADJ	ON	2013/02/15 09:12
S5	5	S4 and cladribine	US-PGPUB; USPAT; USOCR	ADJ	ON	2013/02/15 09:13
S6	430	amorphous near9 cyclodextrin	US-PGPUB; USPAT; USOCR	ADJ	ON	2013/02/15 09:41
S7	168	S6 and @ad<="20040204"	US-PGPUB; USPAT; USOCR	ADJ	ON	2013/02/15 09:45
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INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification ⁶ : A61K 47/48	A1	 (11) International Publication Number: WO 97/18839 (43) International Publication Date: 29 May 1997 (29.05.97)
 (21) International Application Number: PCT/EP96 (22) International Filing Date: 20 November 1996 (20 (30) Priority Data: 95203219.1 23 November 1995 (23.11.95) (34) Countries for which the regional or international application was filed: A 	6/0511 0.11.90) E	 (81) Designated States: AL, AM, AU, BB, BG, BR, CA, CN, CZ, EE, GE, HU, IL, IS, JP, KG, KR, LK, LR, LT, LV, MD, MG, MK, MN, MX, NO, NZ, PL, RO, SG, SI, SK, TR, TT, UA, US, UZ, VN, ARIPO patent (KE, LS, MW, SD, SZ, UG), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG).
 (71) Applicant (for all designated States except US): JA: PHARMACEUTICA N.V. [BE/BE]; Turnhoutseweg 2340 Beerse (BE). (72) Inventors; and (75) Inventors/Applicants (for US only): BAERT, Lieven, Colette [BE/BE]; Gouden Boomstraat 52/3, B-8000 (BE). PEETERS, Jozef [BE/BE]; Sint Corneliusstraa 2430 Beerse (BE). VERRECK, Geert [BE/BE]; Sat 5, B-2980 Zoersel (BE). (74) Agent: DE CORTE, Filip; Janssen Pharmaceutica N.V. Dept., Turnhoutseweg 30, B-2340 Beerse (BE). 	NSSE g 30, F Brugg It 64, F Ilvialaa	Publishcd With international search report. Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments. e e in n

(57) Abstract

Process for preparing a solid mixture comprising one or more cyclodextrins and an insoluble active ingredient characterized in that said process comprises a melt-extrusion step, wherein the active ingredient is embedded into the cyclodextrin carrier.

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SOLID MIXTURES OF CYCLODEXTRINS PREPARED VIA MELT-EXTRUSION

- 5 The present invention involves a process for preparing solid mixtures by melt-extrusion comprising one or more active ingredients, preferably one or more practically insoluble active ingredients and one or more cyclodextrins. The invention further concerns pharmaceutical compositions comprising the above mixture.
- 10 WO 94/11031, published on May 5, 1994, discloses a method of manufacturing a highquality enclosure compound using extrusion techniques. In this document the extrusion of cyclodextrins together with an active ingredient is mentioned. However, the document discloses the use of a wet mixture (i.e. including water or another solvent) to feed into the extruder.

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French patent application 2,705,677 published on December 2, 1994 describes microgranules obtained by extrusion-spheronisation containing a cyclodextrin. The extrusion-spheronisation technique is the combination of an agglomeration technique, i.e. extrusion, and a shaping technique, i.e. the spheronisation. Said patent application actually teaches the formation of microgranulates containing β -cyclodextrin (Kleptose[®]) and microcrystalline cellulose (Avicel[®]) and as active ingredients ketoprofen and paracetamol. The extrusion technique used in the above-mentioned patent application consists in preforming a humid mass by forcing said human mass through a nozzle thus forming long strands of extruded material. The document does

25 not mention melt-extrusion at all.

EP 0,665,009, published as international application on April 24, 1994, discloses a method of dislocating the crystalline condition of crystalline medicine by extruding said crystalline material as such, i.e. without any excipient such as cyclodextrins.

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In J. Pharm. Pharmacolog., vol 44, No 2, pages 73-8, Uekama *et al* show how amorphous nifedipine powders were prepared by spray-drying with hydroxypropyl- β -cyclodextrins. The document does not mention melt-extrusion.

35 In Pharm. Weekbl. Sci. Ed., 1988, vol 10, No 2, page(s) 80-85, Van Doorne et al, the complex formation between β-cyclodextrins and six antimicotic imidazole derivatives was studied. In said study gels and creams comprising antimicotics were prepared

whereby a 1.8 % solution of β -cyclodextrin was added instead of purified water. There is no mentioning of extrusion at all.

In J. Antimicrob. Chemother., 1993, vol 32, No 3, pages 459-463, Hostetler *et al*describe the effect of hydroxypropyl-β-cyclodextrin on the efficacy of oral itraconazole in disseminated murine cryptococcosis. In said document the authors describe how itraconazole is solubilized in hydroxypropyl-β-cyclodextrin resulting in a 100 ml solution. There is no mentioning at all of an extrusion process.

- In Jpn. J. Med. Mycol., 1994, vol 35, No 3, page 263-267, Mikami *et al* describe the effect of carrier solvents on the efficacy of oral itraconazole therapy in aspergilossis in mice. Again this document discloses itraconazole being solubilized in hydroxypropylβ-cyclodextrin. There is no mentioning of extrusion techniques.
- 15 In "Effect of 2-Hydroxypropyl-β-cyclodextrin on Crystallization and Polymorphic Transition of Nifedipine in Solid State", Pharmaceutical research, vol 11, No 12, 1994, Uekama et al. describe a glassy mixture of 2-hydroxypropyl-β-cyclodextrin obtained by heating said mixture and immediately cooling said mixture to 0 degrees Celsius. There is no teaching that this mixture can be extruded.

20

US 5,009,900 describe glassy matrices that are useful for introducing and/or retaining and/or stabilizing the volatile and/or labile components in cooked and uncooked food products. These glassy matrices comprise chemically modified starch having a dextrose equivalent not greater than about 2; maltodextrin, corn syrup solids or a polydextrose, and a mono- or disaccharide. The document does disclose extrusion to form glassy matrices. However, there is no specific mentioning of cyclodextrins and of

None of the above mentioned documents disclose the present invention.

therapeutically or pharmaceutically active ingredients.

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Although WO 94/11031 and French patent application 2,705,677 disclose extrusion of mixtures of cyclodextrins and actives ingredients, said documents do not mention the use of meltextrusion. The technique described in WO 94/11031 and French patent application 2,705,677 has a main disadvantage, that a humid mass needs to be prepared

35 which requires adding to the cyclodextrin and the active ingredient a certain amount of water and in most cases others solvents such as ethanol or methanol. Removing the water and/or other solvents is often a troublesome production step, which often leads to

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irreproducibility because not all of the solvent can be removed. Moreover, with practically insoluble active ingredients the amounts of water and/or adjuvant solvents needed make the above technique unpractical on a production scale. Another disadvantage of the technique described in the prior art is that the drying step can

5 induce unwanted crystallization of the active ingredient.

These problems are solved in the present invention by the use of a melt-extrusion process to form solid mixtures comprising one or more cyclodextrins and insoluble active ingredients.

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The present process is advantageously applicable when said active ingredient is sensitive to a solvent such as water or an organic solvent, because it does not require any solvent. The term "sensitive" used herein means that the active ingredient is readily (e.g. within about one hour) influenced by a solvent to such an extent that its

15 physical, chemical and/or biological properties are substantially modified or changed.

The present process is further advantageous because it does not require a drying step, during which insoluble active ingredients often tend to crystallize.

20 The term "insoluble" hereinabove and hereinunder refers to three categories of compounds, i.e. the "very slightly soluble", "practically insoluble" and "insoluble".

The terms "very slightly soluble", "practically insoluble" or "insoluble" are to be understood as defined in the United States Pharmacopeia 23, NF 18 (1995) page 7, i.e.

- 25 a "very slightly soluble" compound requires from 1000 to 10,000 parts of solvent for 1 part of solute; a "practically insoluble" or "insoluble" compound requires more than 10,000 parts of solvent for 1 part of solute. The solute referred to in these cases are water or aqueous solutions.
- Three examples of this type of insoluble compounds are : itraconazole, loviride and (±)-ethyl (R*,R*)-4-[5-[1-[1-[(4-chlorophenyl)hydroxymethyl]propyl]-1,5-dihydro-5-oxo-4<u>H</u>-1,2,4-triazol-4-yl]-2-pyridinyl]-1-piperazinecarboxylate (hereinafter referred to as compound 1).
- 35 Itraconazole is an art-known antifungal. Loviride is an art-known anti-retrovirally active compound, particularly useful in treating HIV-infected patients.

10

(±)-Ethyl (R*,R*)-4-[5-[1-[1-[(4-chlorophenyl)hydroxymethyl]propyl]-1,5-dihydro-5oxo-4<u>H</u>-1,2,4-triazol-4-yl]-2-pyridinyl]-1-piperazinecarboxylate is described as compound No. 3, in WO 95/27704 published on October 19, 1995.

5 The compounds that are suitable to be used in this technique are compounds that show no appreciable decomposition at the temperatures needed to melt and extrude the mixture of said one or more active ingredients with the cyclodextrin or cyclodextrins.

The term "active ingredient" further refers to compounds or mixtures of compounds which are pharmaceutically or therapeutically or cosmetically active for treating humans or animals.

The present invention provides a process for preparing a solid mixture comprising one or more cyclodextrins and an (insoluble) active ingredient, comprising a melt-extrusion

15 step wherein one or more cyclodextrins are combined with the one or more active ingredients.

Melt-extrusion is a polymer extrusion technique which involves embedding an active ingredient in one or more carriers. In this technique the active ingredient and

20 excipients are molten in the extruder and hence embedded in thermoplastic and thermomelting polymers. The resulting molten mass is then forced through one or more nozzles resulting in a thermoplastic strand or strands.

An extruder comprises an inlet structure, a cylindrical structure called "barrel", a die and a screw or screws. A schematic overview is shown in Figure 1.

The inlet structure mostly is funnel shaped.

The barrel may comprise one or more barrel units and the screw or screws extend 30 through them.

Extruders are available in two general types, namely a single-screw extruder comprising one screw and a multi-screw extruder comprising two or more screws. While this invention can be carried out using either type of extruder, the use of a multi-

35 screw extruder, particularly a twin-screw extruder is preferred. A twin-screw extruder (and a multi-screw extruder) is more efficient in that the plural screws interferring with 5

each other precludes follow-up movement of the active ingredient and, moreover, the intermeshing of the screws provides a high energy output physically.

An interesting mode of operating the screws is to operate them in a corotating mode.

The screw or screws may have different shapes such as, for example, a trapezoidal screw, a trapezoidal cut screw, trapezoidal reverse cut screw, ball screw, kneading paddle and these may be used in the desired combination.

- 10 The load fed into the extruder via the inlet structure is forced by the screw or screws to advance, shorn and blended by the screw within the barrel and extruded from the orifice or orifices of the die. The temperature of the barrel or of the barrel units can be controlled via a heating element or even if necessary by a cooling element.
- 15 The rotational speed of the screw can be set within the allowable range of the extruder used.

A person skilled in the art is able to select the screw geometry and combination of unit screws. The principal function of the screw is to transport, crush and knead the material that is being extruded.

The orifice configuration may be circular, elliptical, rectangular or hexagonal.

Hence, said melt-extrusion step comprises the following substeps :

- a) mixing one or more cyclodextrins with the active ingredient or active ingredients,
 - b) optionally mixing additives,
 - c) heating the thus obtained mixture until melting of one of the components,
 - d) forcing the thus obtained mixture through one or more nozzles;
 - e) cooling the mixture till it solidifies.

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If desired, as mentioned above, the thermomelting mixture comprising one or more cyclodextrins and active ingredient(s) may comprise any suitable additive. When, for instance, the cyclodextrin(s) or the active ingredient(s) or one of the other possible additives is apt to be oxidized, an anti-oxidizing agent may be incorporated, preferably

in small amounts, such as, for instance 100 to 5000 ppm when compared to the total weight of the mixture. Furthermore, conventional auxiliary additives such as pigments, flavors, stabilizers, preservatives and buffers may be added.

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If necessary conventional pharmacologically acceptable plasticizers, such as long chain alcohols, ethylene glycol, propylene glycol, thriethylene glycol, butanediols, pentanols, hexanols, polyethylene glycols, aromatic carboxylates (e.g. dialkyl phthalates,

5 trimellitates, benzoates or terephthalates), aliphatic dicarboxylates or fatty acid esters can also be added. Preferably however, a plasticizer is not needed.

The term "melting" should be broadly interpreted. "Melting" can also refer to the fact that some transition is made to a glassy state, in which it is possible for one component of the mixture to get embedded into the other. In particular cases, one component will melt and the other component(s) will dissolve in the melt thus forming solid solutions, which show advantageous dissolution properties.

The possible formation of these solid solutions is one of the further advantages of the present invention. It will be appreciated by a person skilled in the art that mixing two or more solids, i.e. one or more cyclodextrins and the active ingredient or ingredients, and subsequently melting these solids together will give rise to different products than when the said solids are first brought into contact with water or another solvent and then extruded.

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A characteristic of the melt extruded mixtures of the present invention is the fact that they contain substantially less water or any other solvent than mixtures being extruded in an other way.

- 25 Preferably the present melt extruded mixtures contain no water or solvent, apart from the water or solvent that eventually is contained in the crystal structure of the active ingredient.
- It will be appreciated that the temperature inside the extruder is an important parameter. 30 When different barrel units are present, different temperatures can be applied. A person skilled in the art is able to establish the required temperatures by taking the desired type of cyclodextrin or cyclodextrins or even the complete mixture that is going to be extruded and observing the behaviour as a function of temperature with the aid of a melting point measuring instrument, such as a Kofler hot bench, a microscope hot
- 35 stage type or a differential scanning calorimeter, e.g. type DSC 7 Series Perkin Elmer.

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The cooling can be performed without using any auxiliary means, i.e. it most often suffices to let the thermoplastic strand coming out of the extruder cool down to the ambient temperature of the production site. Of course, cooling aids may be used.

- 5 Once these thermoplastic strands are cooled down these strands can be milled to obtain a powdered form of the mixture of cyclodextrin or cyclodextrins and the active ingredient.
- A person skilled in the art will appreciate that the milling can influence the physical characteristics of the extrudate. During milling the temperature of the material can rise because of the friction and also high shear forces are exerted on the material that is to be milled. Both temperature and mechanical or shear forces can result in a transition of the physical state of the material that is to be milled. A person skilled in the art has sufficient means at his disposal to control temperature and shear forces and thus to
- 15 control the milling process.

The two processes, i.e. melt extrusion and milling can be combined into one configuration as is shown in Figure 1. The mixture of one or more cyclodextrins and one or more active ingredients in combination with possible other additives is feed via

- 20 a funnel like inlet. The mixture is then melt-extruded and the mixture is forced through a nozzle onto a conveyor belt. While being transported on the conveyor belt the extrudate cools down. The cooled melt extrudate is fed into a chopper which forms pellets. These pellets may be further milled if required.
- 25 This powdered material still has the beneficial properties (high bioavailability, dissolution rate, etc.) and it can be used in the conventional way to prepare pharmaceutical, therapeutical or cosmetical solid dosage forms.
- An additional advantage of the present invention is that the active ingredient as well as 30 the cyclodextrins may be transformed in a amorphous form or even that a solid solution is formed. A person skilled in the art will appreciate that this modification of physical state from crystalline to amorphous or to solid solutions is highly advantageous for the dissolution.
- 35 The fact whether the melt extruded mixture contains amorphous material or contains a solid solution or consists essentially of amorphous material or a solid solution can be measured or checked using differential scanning calorimetry. When there is crystalline

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material present in the melt extruded mixture a differential scanning calorimeter will show an endothermic melting peak. When amorphous material or a solid solution is mainly present in the melt extruded mixture a differential scanning calorimeter will not show an endothermic melting peak. Visual inspection of the melt extrudate allows for

5 a distinction between amorphous material and a solid solution. In case the melt extrudate is opaque then both the cyclodextrin(s) and the active ingredient are in a amorphous form. In case melt extrudate is clear then a solid solution was formed.

Curves of differential scanning calorimetry are shown in Figures 2 to 7.

10

Interesting embodiments of the present invention are these melt extruded mixtures which consist mainly of amorphous material.

More interesting embodiments of the present invention are these melt extruded mixtures which consist essentially of amorphous material.

Even more interesting embodiments of the present invention are those melt extruded mixtures which consist mainly of a solid solution of the active ingredient or active ingredients in the cyclodextrin or the cyclodextrins.

20

Preferred embodiments of the present invention are those melt extruded mixtures which consist essentially of a solid solution of the active ingredient or active ingredients in the cyclodextrin or the cyclodextrins.

- 25 Another advantage of the present invention is that the granulation step in forming pharmaceutical, therapeutical or cosmetical compositions can be omitted, because the powdered material can simply be mixed with other excipients and compressed into, for instance, tablets or another solid pharmaceutical, therapeutical or cosmetical form.
- 30 Depending upon the characteristics of the melt extruded mixture, the size of the pellets of said melt extruded mixture or the mesh of the powder of said melt extruded mixture and, of course, dependent upon the other auxiliaries that are added to the unit dosage forms the unit dosage form may give immediate release or sustained release.
- 35 If desired, said solid pharmaceutical form may also be provided with a conventional coating to improve the appearance and/or the flavor (coated tablets) or additionally to target the release of the active ingredient.

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Suitable tablets may have the following compositions and may be prepared in a conventional way. The amounts given are of course dependent upon the dose required for the pharmaceutical, therapeutic or cosmetic activity.

5

Composition A

	milled melt extrudate	100 - 500 mg
	microcrystalline cellulose	100 - 300 mg
10	crospovidone	10 - 200 mg
	colloidal silicon dioxide	1 - 5 mg
	sterotex	2 - 10 mg
15	Composition B	
10	milled melt extrudate	100 - 500 mg
	Microcelac (TM) (1)	200 - 300 mg
	crospovidone	70 - 200 mg
	talc	20 - 50 mg
20	sterotex	7 - 10 mg
	colloidal silicon dioxide	1 - 5 mg
	magnesium stearate	2 - 10 mg

The cyclodextrin to be used in the aforementioned compositions include the

25 pharmaceutically acceptable unsubstituted and substituted cyclodextrins known in the art, more particularly α , β or γ cyclodextrins or the pharmaceutically acceptable derivatives thereof.

Substituted cyclodextrins which can be used in the invention include polyethers

- described in U.S. Patent 3,459,731. In general, unsubstituted cyclodextrins are reacted with an alkylene oxide, preferably under superatmospheric pressure and at an elevated temperature, in the presence of an alkaline catalyst.
 Since a hydroxy moiety of the cyclodextrin can be substituted by an alkylene oxide which itself can react with yet another molecule of alkylene oxide, the average molar
- 35 substitution (MS) is used as a measure of the average number of moles of the substituting agent per glucose unit. The MS can be greater than 3 and theoretically has no limit.

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Further substituted cyclodextrins are ethers wherein the hydrogen of one or more cyclodextrin hydroxy groups is replaced by C₁₋₆alkyl, hydroxyC₁₋₆alkyl, carboxy-C₁₋₆alkyl or C₁₋₆alkyloxycarbonylC₁₋₆alkyl or mixed ethers thereof. In particular such substituted cyclodextrins are ethers wherein the hydrogen of one or more

5 cyclodextrin hydroxy groups is replaced by C₁₋₃alkyl, hydroxyC₂₋₄alkyl or
 carboxyC₁₋₂alkyl or more in particular by methyl, ethyl, hydroxyethyl, hydroxypropyl,
 hydroxybutyl, carboxy-methyl or carboxyethyl.

In the foregoing definitions the term "C1-6alkyl" is meant to include straight and

- 10 branched saturated hydrocarbon radicals, having from 1 to 6 carbon atoms, such as, methyl, ethyl, 1-methylethyl, 1,1-dimethylethyl, propyl, 2-methylpropyl, butyl, pentyl, hexyl and the like.
- Such ethers can be prepared by reacting the starting cyclodextrin with an appropriate
 <u>O</u>-alkylating agent or a mixture of such agents in a concentration being selected so that the desired cyclodextrin ether is obtained. The said reaction is preferably conducted in a suitable solvent in the presence of an appropriate base. With such ethers, the degree of substitution (DS) is the average number of substituted hydroxy functions per glucose unit, the DS being thus 3 or less.

20

In the cyclodextrin derivatives for use in the compositions according to the present invention, the DS preferably is in the range of 0.125 to 3, in particular 0.3 to 2, more in particular 0.3 to 1 and the MS is in the range of 0.125 to 10, in particular of 0.3 to 3 and more in particular 0.3 to 1.5.

25

Of particular utility in the invention are the β -cyclodextrin ethers, e.g. dimethyl- β cyclodextrin as described in Drugs of the Future, Vol. 9, No. 8, p. 577-578 by M. Nogradi (1984) and polyethers, e.g. hydroxypropyl β -cyclodextrin and hydroxyethyl β cyclodextrin, being examples. Such an alkyl ether may be a methyl ether with a degree

30 of substitution of about 0.125 to 3, e.g. about 0.3 to 2. Such a hydroxypropyl cyclodextrin may for example be formed from the reaction between β-cyclodextrin an propylene oxide and may have a MS value of about 0.125 to 10, e.g. about 0.3 to 3.

A more novel type of substituted cyclodextrins is sulfobutylcyclodextrines. These type are also envisaged in the present invention. The ratio of active ingredient over cyclodextrin may vary widely. For example ratios of 1/100 to 100/1 may be applied. Interesting ratios of active ingredient over cyclodextrin range from about 1/10 to 10/1. More interesting ratios of active ingredient over cyclodextrin range from about 1/5 to 5/1. Most interesting ratios range from about 1/2 to 3/1. Preferred ratio is about 1/1

5 1/3 to 3/1. Preferred ratio is aabout 1/1.

The use of a mixture of cyclodextrins, either different types (α , β , γ) or different substitution (2-hydropropyl or methyl) or different substitution grades in sometimes recommendable to decrease the melting point.

10

Description of the drawings

Figure 1 is a schematic representation of a configuration for carrying out the present invention.

15

Figure 2 is a differential scanning calorimetry curve (DSC curve) of non-milled Batch No 1 material. (see Example 1)

Figure 3 is a differential scanning calorimetry curve of milled Batch No 1 material (see Example 1)

Figure 4 is a differential scanning calorimetry curve of Batch No 2 material (see Example 1)

25 Figure 5 is a differential scanning calorimetry curve of of Batch No 3 material (see Example 1)

Figure 6 is a differential scanning calorimetry curve of of Batch No 4 material (see Example 1)

30

Figure 7 is a differential scanning calorimetry curve of of Batch No 5 material (see Example 1)

35 Example 1

Extruded samples of active ingredient with hydroxypropyl- β -cyclodextrin (HP- β -CD) were obtained using a Twin Screw Extruder type MP19 APV Baker (commercially available from the company APV Baker) with a die having a orifice of 3 mm. The process parameters for each individual experiment are shown in the table 1. This type

of extruder has a L/D ratio of 15 and a screw pattern : 4D FS - 4x30 FP - 4x60 FP - 4x90 P - 4x60 RP - 2.5D FS - 2x30 FP - 2x60 FP - 2x90 P - 3x60 RP - 3 DFS. (4D refers to a transportelement having a length of 4 times the screw diameter of the feed screw type; 4x30 FP refers to 4 forward paddles positioned with mutual angle of 30

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degrees, 4x60 RP refers to a working zone having reverse paddles positioned with a mutual angle of 60 degrees)

In this type of extruder the mixture is fed by a feeding screw turning a constant feeding

- 5 speed (v1) (A feeding speed of 10 revolutions per minute amounts to a feeding speed of 1,5 kg per hour) onto the twin transporter screws having a diameter of 18 mm turning at a transporter speed (v2). These speeds are rotational speeds (revolutions per minute).
- 10 The mixture is then transported into a first heating zone (t1). Here the rate of transport diminished by a difference of the configuration of the twin transporter screws i.e. the rotational transporter speed v2 remains the same but the material does not progress as quickly.
- 15 Subsequently, the molten mass is transported by again normal configuration twin transporter screws to a second heating zone (t2) where the rate of transport is again diminished by a difference of configuration of the twin transporter screws.

After this second heating the thermomelting mixture is transported to the nozzle of the apparatus.

mixture	Batch. No	t1 (°C)	t2 (°C)	tp (°C)	v1 (rpm)*	v2 (rpm)*
$\frac{\text{compound } 1}{\text{HP-}\beta\text{-}\text{CD}} : \frac{1}{3}$	1	256	283	280	10	100
$\frac{\text{itraconazole}}{\text{HP-}\beta\text{-CD}} \div \frac{1}{1}$	2	263	265	279	10	20
$\frac{\text{itraconazole}}{\text{HP-}\beta\text{-CD}} : \frac{1}{3}$	3	264	265	280	10	20
$\frac{\text{loviride}}{\text{HP-}\beta\text{-CD}} \div \frac{1}{1}$	4	274	285	292	10	80
$\frac{\text{loviride}}{\text{HP-}\beta\text{-CD}} : \frac{1}{3}$	5	258	265	274	10	20

Table 1

20

25 * rpm = revolutions per minute

- t1 : temperature of the first heating zone
- t2 : temperature of the second heating zone

- tp : temperature inside the barrel

- v1 : rate of feeding screw

- v2 : twin transporter screws speed(rotational).
- 5 In every case the mixture of active ingredient and 2-hydroxypropyl- β -CD gave a solid solution.

Example 2

Extruded samples of active ingredient with dimethyl- β -cyclodextrin (DM- β -CD) were

10 obtained using extruder type MP19 - APV Baker with the process parameters as shown in the table 2.

<u>Table 2</u>

15

mixture	Batch. No.	t1 (°C)	t2 (°C)	tp (°C)	v1 (1) (rpm)*	v2 (rpm)*
$\frac{\text{compound 1}}{\text{DM-}\beta\text{-}\text{CD}} : \frac{1}{1}$	6	241	245	254	0	20
$\frac{\text{itraconazole}}{\text{DM-}\beta\text{-}\text{CD}} : \frac{1}{1}$	7	239	240	253	0	20
$\frac{\text{loviride}}{\text{DM}-\beta-\text{CD}}$: $\frac{1}{1}$	8	248	250	263	0	20

* rpm = revolutions per minute

(1) The apparatus was fed manually, without using the feeding screw. In every case the mixture of active ingredient and DM- β -CD.

20 - t1 : temperature of the first heating zone

- t2 : temperature of the second heating zone

- t_p : temperature inside the barrel
- v1 : feeding screw speed (rotational)
- v₂ : twin transporter screw speed (rotational).

25

Example 3

The dissolution of the melt extrudate of Batch No 1 was compared with the dissolution of the "physical mixture" (i.e. the mixture of the two component in the ratio as shown for Batch No. 1, but not melt extruded).

An amount of 100 mg of milled melt extrudate of Batch No 1 was added to a volume of 900 ml of artificial gastric juice at a temperature of 37 degrees Celsius. The stirring method used was the peddle method with a peddle moving at 100 rotations per minute. Using UV spectrometry the relative amount of dissolved extrudate was measured

5 during 1 hour.

The same procedure for the "physical mixture" was followed.

The results of this dissolution process are shown in Table 3

10

Table 3

time mixture	milled extrudate Batch No 1	corresponding physical
(minutes)	(% of total amount dissolved)	(% of total amount dissolved)
0	0.00	0.00
5	62.10	1.71
15	70.20	14.67
30	72.63	21.06
45	74.07	26.10
60	74.25	28.35

Example 4

15 The melting behaviour was measured by using differential scanning calorimetry. The calorimeter used is the Perkin-Elmer 7 Series Thermal Analysis System. In all cases the rate of heating was set at 20 degrees Celsius per minute.

Figure 2 shows the DSC curve of melt extrudate of Batch No 1 before milling. The curve shows no endothermic or exothermic peaks and it was established by visual inspection that the molten material was a clear solution, thus indicating that the nonmilled melt extrudate of Batch No 1 is a solid solution.

Figure 3 shows the DSC curve of melt extrudate of Batch No 1 after milling. The

25 curve shows no endothermic or exothermic peaks and it was established by visual inspection that the molten material was a clear solution, thus indicating that the milled melt extrudate of Batch No 1 is a solid solution.

Figure 4 shows the DSC curve of meltextrudate of Batch No 2 before milling. The curve shows no endothermic or exothermic peaks and it was established by visual inspection that the molten material was <u>not</u> a clear solution, thus indicating that the non-milled melt extrudate of Batch No 2 is a mixture of amorphous material.

5

Figure 5 shows the DSC curve of meltextrudate of Batch No 3 before milling. The curve shows a small endothermic peak. The data on said small peak are as follows : X1 = 117.600 degrees Celsius, X2 = 143.200 degrees Celsius, Peak at 132.695 degrees Celsius, Area is 38.126 mJ, Δ H is 3.768 J/g, Height is 1.520 mW and the onset is at

- 10 125.816 degrees Celsius. Said small peak is very probably due to an impurity in the cyclodextrins. It was established that the non-milled melt extrudate of Batch No 3 is a mixture of amorphous material.
- Figure 6 shows the DSC curve of meltextrudate of Batch No 4 before milling. The
 curve shows a few small endothermic peaks. Hence, it was established that the non milled melt extrudate of Batch No 4 is a mixture of amorphous material containing
 small amounts of crystalline material

Figure 7 shows the DSC curve of meltextrudate of Batch No 5 before milling. The
curve shows no endothermic or exothermic peaks and it was established by visual
inspection that the molten material was not a clear solution, thus indicating that the
non-milled melt extrudate of Batch No 5 is a mixture of amorphous material.

Example 5

25 The melt extrudate of Batch No. 1 was milled and sieved. By mixing the appropriate amounts a tablet having the following composition was prepared in an art-known way :

	milled extrudate batch No 1	480 mg
	microcrystalline cellulose	218 mg
30	Aerosil	3 mg
	magnesium stearate	5 mg
	crospovidone	144 mg

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<u>Claims</u>

1. Process for preparing a solid mixture comprising one or more cyclodextrins and

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- one or more active ingredients characterized in that said process encompasses a melt-extrusion step, wherein the active ingredient is embedded into the cyclodextrin carrier.
- 2. A process as claimed in claim 1, wherein the melt-extrusion process comprises the following substeps :
 - a) mixing one or more cyclodextrins with one or more active ingredients, and
 - b) optionally mixing additives;
 - c) heating the thus obtained mixture until melting of one of the components;
 - d) forcing the thus obtained mixture through one or more nozzles;
- 15 e) cooling the mixture till it solidifies.
 - 3. A solid mixture obtainable by the process as described in any of claims 1 or 2, with the proviso that nifedipine in combination with 2-hydroxypropyl- β -cyclodextrin is excluded.
- 20
- 4. A solid mixture as claimed in claim 3 characterized in that the active ingredient or active ingredients are insoluble according to the definition of US Pharmacopeia.
- 5. A solid mixture as claimed in claims 3 or 4, wherein substantially only one type of 25 cyclodextrin is present.
 - 6. A solid mixture as claimed in any of claims 3 to 5 wherein a cyclodextrin is hydroxypropyl- β -cyclodextrin.
- A solid mixture as claimed in any of claim 3 to 5 wherein a cyclodextrin is dimethyl-β-cyclodextrin.
 - 8. A solid mixture as claimed in any of claims 3 to 7, wherein the active ingredient is itraconazole.

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9. A solid mixture as claimed in any of claims 3 to 7 wherein the active ingredient is loviride.

- 10. A solid mixture as claimed in any of claims 3 to 7 wherein the active ingredient is (±)-ethyl (R*,R*)-4-[5-[1-[1-[(4-chlorophenyl)hydroxymethyl]propyl]-1,5-dihydro-5-oxo-4H-1,2,4-triazol-4-yl]-2-pyridinyl]-1-piperazinecarboxylate.
- 5 11. A pharmaceutical composition comprising milled melt extrudate and other excipients.
 - 12. A process for preparing a pharmaceutical composition as claimed in claim 11 characterized by milling appropriately the solid mixture as claimed in any of

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claims 4 to 10, intimately mixing the thus obtained powdered material with other pharmaceutically acceptable excipients and further processing into pharmaceutical dosage forms.



WO 97/18839























PCT/EP96/05118



Int onal Application No PCT/EP 96/05118

A. CLASSIFICATION OF SUBJECT MATTER IPC 6 A61K47/48 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) IPC 6 A61K Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Category ' Relevant to claim No. WO 94 11031 A (NIPPON SHINYAKU CO LTD) 26 Х 1 - 12May 1994 cited in the application see abstract FR 2 705 677 A (ROQUETTE FRERES) 2 Y 1-12 December 1994 cited in the application see abstract see examples see claims Х EP 0 665 009 A (NIPPON SHINYAKU COMPANY, 1 - 12LIMITED.) 2 August 1995 cited in the application see abstract see examples see claims ----/--Further documents are listed in the continuation of box C. Patent family members are listed in annex. X X Special categories of cited documents : "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention filing date cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "Y" document of particular relevance; the claimed invention counter to considered to involve an inventive step when the document is combined with one or more other such docu-ments, such combination being obvious to a person skilled "O" document referring to an oral disclosure, use, exhibition or other means document published prior to the international filing date but later than the priority date claimed in the art '&' document member of the same patent family Date of mailing of the international search report Date of the actual completion of the international search 1 7. 03. 97 24 February 1997 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Dullaart, A Fax: (+31-70) 340-3016

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poly(vinylpyrrolidone) K-30" cited in the application * paragraph Materials and methods * see figures	
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Form PCT/ISA/218 (continuation of second sheet) (July 1992)

In uonal Application No PCT/EP 96/05118

C.(Continuz	ution) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
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ernational application No.

INTERNATIONAL	SEARCH	REPORT

PCT/EP 96/05118

Box 1 Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. X Claims Nos.: 1-7,11-12 because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically: In view of the large number of compounds, which are defined by the general definition of the active ingredient used in the claims, the search had to
be restricted for economic reasons. The search was limited to the compounds for which pharmacological data was given and/or the compounds mentioned
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
1. As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2. As all searchable claims could be searches without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4. No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

International Application No. PCT/EP 96/ 05118

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FURTHER INFORMATION CONTINUED FROM PCT/ISA/210
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in the claims, and to the general idea underlying the application (see Guidelines, chapter III, paragraph 2.3).

Claims searched incompletely: 1-7,11-12

Information on patent family members

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12/986,310	01/07/2011		514		1623		20009904-0067	
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Sheet 1 of 4

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Application Number

First Named Inventor

Attorney Docket No.

Examiner Name

Filing Date

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January 5, 2011

Nicholas S. Bodor

0056192-000067

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Sheet 3 of 4

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Examiner Signature	/Jonathan Lau/	Date Considered	02/15/2013
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*EXAMINER: Initial if reference considered, whether or not citation is in conformance with M.P.E.P. § 609. Draw line through citation if not in conformance and not considered. Include copy of this form with next communication to Applicant.

Substitute for form 1449/PTO & 1449B/PTO	Complete if Known		
FIRST	Application Number		
INFORMATION DISCLOSURE	Filing Date	January 5, 2011	
STATEMENT BY APPLICANT	First Named Inventor	Nicholas S. Bodor	
(use as many sheets as necessary)	Examiner Name		
· · · · · ·	Attorney Docket No.	0056192-000067	

Sheet 4 of 4

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Examiner Signature	/Jonathan Lau/	Date Considered	02/15/2013		
*EXAMINER: Initial if reference considered, whether or not citation is in conformance with M.P.F.P. § 609. Draw line through citation if not in					

	Application/Control No.	Applicant(s)/Patent Under Reexamination
Search Notes	12986310	BODOR ET AL.
	Examiner	Art Unit
	JONATHAN S LAU	1623

CPC- SEARCHED		
Symbol	Date	Examiner

CPC COMBINATION SETS - SEARCHED			
Symbol	Date	Examiner	

US CLASSIFICATION SEARCHED				
Class	Subclass	Date	Examiner	

SEARCH NOTES		
Search Notes	Date	Examiner
EAST - inventor name search (Nicholas Bodor, Yogesh Dandiker)	2/15/2013	JSL
EAST - see attached notes	2/15/2013	JSL
Google Scholar - see attached notes	2/15/2013	JSL

INTERFERENCE SEARCH				
US Class/ CPC Symbol	US Subclass / CPC Group	Date	Examiner	

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IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicant: Nicholas S. BODOR et al.) MAIL STOP AMENDMENT
Application No.: 12/986,310) Examiner: Jonathan S. LAU
Filed: January 7, 2011) Group Art Unit: 1623
Title: ORAL FORMULATIONS OF CLADRIBINE) Confirmation No.: 6100

REPLY AND AMENDMENT

In response to the Office Action dated February 25, 2013, please first amend the above-identified patent application as follows:

AMENDMENTS TO THE SPECIFICATION:

Please replace the first paragraph and its heading on page 1 with the following:

CROSS-REFERENCE TO EARLIER APPLICATIONS

This application is a continuation of prior copending US Application No.

10/551,205 filed November 14, 2006, now allowedU.S. Patent No. 7,888,328, which is the US national stage of International Application No. PCT/US2004/009387, filed March 26, 2004, which claims benefit under 35 U.S.C. § 119(e) of United States Provisional Application No. 60/458,922, filed March 28, 2003; of United States Provisional Application No. 60/484,756, filed July 2, 2003; and of United States Provisional Application No. 60/541,247, filed February 4, 2004, all of said applications being hereby incorporated by reference herein in their entireties and relied upon.

Electronic Patent Application Fee Transmittal					
Application Number:	129	12986310			
Filing Date:	07-	Jan-2011			
Title of Invention:	OR	ORAL FORMULATIONS OF CLADRIBINE			
First Named Inventor/Applicant Name:	Nic	holas S. Bodor			
Filer:	Ma	ry Katherine Baume	eister/Rebecca I	Brimmer	
Attorney Docket Number:	200	009904-0067			
Filed as Large Entity					
Utility under 35 USC 111(a) Filing Fees					
Description		Fee Code	Quantity	Amount	Sub-Total in USD(\$)
Basic Filing:					
Pages:					
Claims:					
Claims in Excess of 20		1202	12	80	960
Independent claims in excess of 3		1201	1	420	420
Miscellaneous-Filing:					
Petition:					
Patent-Appeals-and-Interference:					
Post-Allowance-and-Post-Issuance:		260			

Description	Fee Code	Quantity	Amount	Sub-Total in USD(\$)
Extension-of-Time:				
Miscellaneous:				
	Tot	al in USD	(\$)	1380

Electronic Acknowledgement Receipt			
EFS ID:	15887589		
Application Number:	12986310		
International Application Number:			
Confirmation Number:	6100		
Title of Invention:	ORAL FORMULATIONS OF CLADRIBINE		
First Named Inventor/Applicant Name:	Nicholas S. Bodor		
Customer Number:	13974		
Filer:	Mary Katherine Baumeister/Rebecca Brimmer		
Filer Authorized By:	Mary Katherine Baumeister		
Attorney Docket Number:	20009904-0067		
Receipt Date:	28-MAY-2013		
Filing Date:	07-JAN-2011		
Time Stamp:	18:44:25		
Application Type:	Utility under 35 USC 111(a)		

Payment information:

Submitted with Payment	yes			
Payment Type	Credit Card			
Payment was successfully received in RAM	\$1380			
RAM confirmation Number	8981			
Deposit Account 193140				
Authorized User BAUMEISTER, MARY KATHERINE				
The Director of the USPTO is hereby authorized to charge indicated fees and credit any overpayment as follows:				
Charge any Additional Fees required under 37 C.F.R. Section 1.16 (National application filing, search, and examination fees)				
Charge any Additional Fees required under 37 C.F.R. Section 262 (Patent application and reexamination processing fees)				

Charge ar	ny Additional Fees required under 37 C.F.	R. Section 1.19 (Document supply	r fees)		
Charge ar	ny Additional Fees required under 37 C.F.	R. Section 1.20 (Post issuance fees	s and charges)		
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Document Number	Document Description	File Name	File Size(Bytes)/ Message Digest	Multi Part /.zip	Pages (if appl.)
			1187619		
1		Amendment.PDF	1c400c71df815a15e645ffbd60fa36a6e5a9a	yes	58
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	Amendment/Req. Reconsiderati	on-After Non-Final Reject	1		1
	Specificati	on	2		2
	Claims		3	-	4
	Applicant Arguments/Remarks	Made in an Amendment	15	5	58
Warnings:					
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2	Fee Worksheet (SB06)	fee-info.pdf	31842	no	2
			6e5c34b2cb2eb9f9990e31b9c3495b2f282 4ea7f		
Warnings:					
Information:			1		
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This Acknowled characterized B Post Card, as d <u>New Applicatio</u> If a new applica 1.53(b)-(d) and Acknowledgen	dgement Receipt evidences receip by the applicant, and including pag escribed in MPEP 503. <u>ons Under 35 U.S.C. 111</u> ation is being filed and the applica MPEP 506), a Filing Receipt (37 CF nent Receipt will establish the filin	t on the noted date by the U ge counts, where applicable. tion includes the necessary o R 1.54) will be issued in due g date of the application.	SPTO of the indicated It serves as evidence components for a filin course and the date s	l document: of receipt s ng date (see hown on th	s, imilar to a 37 CFR is
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Under the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it displays a valid OMB control number									
PATENT APPLICATION FEE DETERMINATION RECORD Substitute for Form PTO-875							or Docket Number /986,310	Filing Date 01/07/2011	To be Mailed
(Column 1) (Column 2)									
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MULTIPLE DEPENDENT CLAIM PRESENT (37 CFR 1.16(j))									
* If the difference in column 1 is less than zero, enter "0" in column 2. TOTAL									
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	Total (37 CFR 1.16(i))	* 76	Minus	** 64	= 12		x \$80 =		960
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	Application Size Fee (37 CFR 1.16(s))								
	FIRST PRESENTATION OF MULTIPLE DEPENDENT CLAIM (37 CFR 1.16(j))								
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	Application Size Fee (37 CFR 1.16(s))								
AN	FIRST PRESENTATION OF MULTIPLE DEPENDENT CLAIM (37 CFR 1.16(j))								
							TOTAL ADD'L FE	E	
* If the entry in column 1 is less than the entry in column 2, write "0" in column 3. ** If the "Highest Number Previously Paid For" IN THIS SPACE is less than 20, enter "20". *** If the "Highest Number Previously Paid For" IN THIS SPACE is less than 3, enter "3". The "Highest Number Previously Paid For" (Total or Independent) is the highest number found in the appropriate box in column 1.									
This collection of information is required by 37 CFR 1.16. The information is required to obtain or retain a benefit by the public which is to file (and by the USPTO to									

process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.14. This collection is estimated to take 12 minutes to complete, including gathering, preparing, and submitting the completed application form to the USPTO. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and Trademark Office, U.S. Department of Commerce, P.O. Box 1450, Alexandria, VA 22313-1450. DO NOT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS. SEND TO: Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450.

If you need assistance in completing the form, call 1-800-PTO-9199 and select option 2.

AMENDMENTS TO THE CLAIMS:

This listing of claims will replace all prior versions, and listings, of claims in the application:

LISTING OF CLAIMS:

1. (Original) A pharmaceutical composition comprising a complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex, formulated into a solid oral dosage form, said composition comprising no significant amount of free crystalline cladribine therein.

2. (Original) The pharmaceutical composition according to Claim 1, wherein the complex is saturated with cladribine.

3. (Original) The composition according to Claim 1, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin, hydroxypropyl- γ -cyclodextrin, randomly methylated β -cyclodextrin, carboxymethyl- β -cyclodextrin or sulfobutyl- β -cyclodextrin.

4. (Original) The composition according to Claim 1, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin or hydroxypropyl-γ-cyclodextrin.

5. (Original) The composition according to Claim 1, wherein the weight ratio of cladribine to amorphous cyclodextrin is from about 1:10 to about 1:16.

6. (Original) The composition according to Claim 5, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin or hydroxypropyl-γ-cyclodextrin.

7. (Original) The composition according to Claim 1, wherein the approximate molar ratio of cladribine to amorphous cyclodextrin corresponds to a point

located on the curve of a phase solubility diagram for saturated complexes of cladribine in varying concentrations of the cyclodextrin.

8. (Original) The composition according to Claim 1, wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b).

9. (Withdrawn) A method for enhancing the oral bioavailability of cladribine comprising orally administering to a subject in need thereof a pharmaceutical composition comprising a complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex, formulated into a solid oral dosage form, said composition comprising no significant amount of free crystalline cladribine therein.

10. (Withdrawn) The method according to Claim 9, wherein the complex is saturated with cladribine.

11. (Withdrawn) The method according to Claim 9, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin, hydroxypropyl- γ -cyclodextrin, randomly methylated β -cyclodextrin, carboxymethyl- β -cyclodextrin or sulfobutyl- β -cyclodextrin.

12. (Withdrawn) The method according to Claim 9, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin or hydroxypropyl-γ-cyclodextrin.

13. (Withdrawn) The method according to Claim 9, wherein the weight ratio of cladribine to amorphous cyclodextrin is from about 1:10 to about 1:16.

14. (Withdrawn) The method according to Claim 13, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin or hydroxypropyl-γ-cyclodextrin.

15. (Withdrawn) The method according to Claim 9, wherein the approximate molar ratio of cladribine to amorphous cyclodextrin corresponds to a point located on the curve of a phase solubility diagram for saturated complexes of cladribine in varying concentrations of the cyclodextrin.

16. (Withdrawn) The method according to Claim 9, wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b).

17. (Withdrawn) A method for the treatment of symptoms of a cladribine-responsive condition in a subject suffering from said symptoms comprising orally administering to said subject a pharmaceutical composition comprising a complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex, formulated into a solid oral dosage form, said composition comprising no significant amount of free crystalline cladribine therein.

18. (Withdrawn) The method according to Claim 17, wherein the complex is saturated with cladribine.

19. (Withdrawn) The method according to Claim 17, wherein the cladribine-responsive condition is selected from the group consisting of multiple sclerosis, rheumatoid arthritis and leukemia.

20. (Withdrawn) The method according to Claim 19, wherein the cladribine-responsive condition is multiple sclerosis.

21. (Withdrawn) The method according to Claim 17, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin, hydroxypropyl-γ-cyclodextrin, randomly methylated β-cyclodextrin, carboxymethyl-β-cyclodextrin or sulfobutyl-β-cyclodextrin.

22. (Withdrawn) The method according to Claim 17, wherein the weight ratio of cladribine to amorphous cyclodextrin is from about 1:10 to about 1:16.

23. (Withdrawn) The method according to Claim 17, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin or hydroxypropyl-γ-cyclodextrin.

24. (Withdrawn) The method according to Claim 17, wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b).

25. (Original) A complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex.

26. (Original) The complex according to Claim 25, saturated with cladribine.

27. (Original) The complex according to Claim 25, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin, hydroxypropyl- γ -cyclodextrin, randomly methylated β -cyclodextrin, carboxymethyl- β -cyclodextrin or sulfobutyl- β -cyclodextrin.

28. (Original) The complex according to Claim 25, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin or hydroxypropyl- γ -cyclodextrin.

29. (Original) The complex according to Claim 25, wherein the weight ratio of cladribine to amorphous cyclodextrin is from about 1:10 to about 1:16.

30. (Original) The complex according to Claim 29, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin or hydroxypropyl-γ-cyclodextrin.

31. (Original) The complex according to Claim 25, wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b).

32. (Withdrawn) A process for the preparation of a complex cladribinecyclodextrin complex as claimed in Claim 25, which comprises the steps of:

 (i) combining cladribine and an amorphous cyclodextrin in water at a temperature of from about 40 to about 80°C and maintaining said temperature for a period of from about 6 to about 24 hours;

(ii) cooling the resultant aqueous solution to room temperature; and

(iii) lyophilizing the cooled solution to afford an amorphous product.

33. (Withdrawn) The process according to Claim 32, further comprising a filtration step following step (ii).

34. (Withdrawn) The process according to Claim 32, wherein step (i) is performed at a temperature of from about 45 to about 60°C.

35. (Withdrawn) The process according to Claim 32, wherein step (i) is performed at a temperature of from about 45 to about 50°C.

36. (Withdrawn) The process according to Claim 34, wherein step (i) is performed with stirring.

37. (Withdrawn) The process according to Claim 36, wherein step (i) is performed for a period of from about 6 to about 9 hours.

38. (Withdrawn) The process according to Claim 32, wherein step (ii) is performed for a period of from about 6 to about 9 hours.

39. (Withdrawn) The process according to Claim 32, wherein step (iii) comprises an initial freezing stage in which the solution is cooled to from about -40 to about -80° C, and held at said temperature for a period of from about 2 to about 4 hours.

40. (Withdrawn) The process according to Claim 39, wherein, in the initial freezing stage of step (iii), the solution is cooled to about -45°C.

41. (Withdrawn) The process according to Claim 34, wherein 12.00 parts by weight of cladribine and 172.50 parts by weight of hydroxypropyl- β -cyclodextrin are introduced in step (i), or wherein 16.35 parts by weight of cladribine and 172.50 parts by weight of hydroxypropyl- β -cyclodextrin are introduced in step (i).

42. (Withdrawn) The process according to Claim 41, wherein 825 parts by volume of water are introduced in step (i).

43. (Withdrawn) The process according to Claim 32, wherein the lyophilization step (iii) comprises:

(a) an initial freezing stage in which the complexation solution is brought to from about -40°C to about -80°C for approximately 2 to 4 hours;

(b) a primary drying stage at about -25°C for approximately 80 to 90 hours; and

(c) a secondary drying stage at about 30°C for approximately 15 to 20 hours.

44. (Withdrawn) The process according to Claim 43, wherein stage (a) of the lyophilization is conducted at about -45°C for approximately 3 to 4 hours.

45. (Withdrawn) The process according to Claim 43, wherein stage (b) of the lyophilization is conducted under a pressure of about 100 mTorr.

46. (Original) A pharmaceutical composition according to Claim 1, obtainable by a process comprising the steps of:

 (i) combining cladribine and an amorphous cyclodextrin in water at a temperature of from about 40 to about 80°C and maintaining said temperature for a period of from about 6 to about 24 hours;

(ii) cooling the resultant aqueous solution to room temperature;

(iii) lyophilizing the cooled solution to afford an amorphous product; and

(iv) formulating the amorphous product into a solid oral dosage form.

47. (Original) The pharmaceutical composition according to Claim 46, wherein the process further comprises a filtration step following step (i) or (ii).

48. (Original) The pharmaceutical composition according to Claim 46, wherein step (i) of the process is performed at a temperature of from about 45 to about 60°C.

49. (Original) The pharmaceutical composition according to Claim 46, wherein step (i) of the process is performed at a temperature of from about 45 to about 50°C.

50. (Original) The pharmaceutical composition according to Claim 48, wherein step (i) of the process is performed with stirring.

51. (Original) The pharmaceutical composition according to Claim 50, wherein step (i) of the process is performed for a period of from about 6 to about 9 hours.

52. (Original) The pharmaceutical composition according to Claim 46, wherein step (ii) of the process is performed for a period of from about 6 to about 9 hours.

53. (Original) The pharmaceutical composition according Claim 46, wherein step (iii) comprises an initial freezing stage in which the solution is cooled to

from about -40 to about -80°C, and held at said temperature for a period of from about 2 to about 4 hours.

54. (Original) The pharmaceutical composition according to Claim 53, wherein, in the initial freezing stage of step (iii), the solution is cooled to about -45°C.

55. (Original) The pharmaceutical composition according to Claim 48, wherein 12.00 parts by weight of cladribine and 172.50 parts by weight of the hydroxypropyl- β -cyclodextrin are introduced in step (i) of the process, or wherein 16.35 parts by weight of cladribine and 172.50 parts by weight of the hydroxypropyl- β -cyclodextrin are introduced in step (i) of the process.

56. (Original) The pharmaceutical composition according to Claim 55, wherein 825 parts by volume of water are introduced in step (i) of the process.

57. (Original) The pharmaceutical composition according to Claim 46, wherein the lyophilization step (iii) of the process comprises:

(a) an initial freezing stage in which the complexation solution is brought to from about -40°C to about -80°C for approximately 2 to 4 hours;

(b) a primary drying stage at about -25°C for approximately 80 to 90 hours; and

(c) a secondary drying stage at about 30°C for approximately 15 to 20 hours.

58. (Original) The pharmaceutical composition according to Claim 57, wherein stage (a) of the lyophilization is conducted at about -45°C for approximately 3 to 4 hours.

59. (Original) The pharmaceutical composition according to Claim 57, wherein stage (b) of the lyophilization is conducted under a pressure of about 100 mTorr.

60. (Original) The pharmaceutical composition according to Claim 46, wherein the formulation step (iv) of the process comprises blending the complex with magnesium stearate and compressing into tablets.

61. (Original) The pharmaceutical composition according to Claim 60, wherein magnesium stearate is pre-mixed with sorbitol powder before blending with the complex.

62-64. (Cancelled)

65. (New) A method for the treatment of symptoms of a cladribineresponsive condition in a subject suffering from said symptoms comprising orally administering to said subject a pharmaceutical composition comprising a complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex, formulated into a solid oral dosage form, said composition comprising no significant amount of free crystalline cladribine therein, wherein administering cladribine is accompanied by administering one or more additional active ingredients for treating the cladribine-responsive condition.

66. (New) The method according to Claim 65, wherein the complex is saturated with cladribine.

67. (New) The method according to Claim 65, wherein the cladribineresponsive condition is selected from the group consisting of multiple sclerosis, rheumatoid arthritis and leukemia.

68. (New) The method according to Claim 66, wherein the cladribineresponsive condition is selected from the group consisting of multiple sclerosis, rheumatoid arthritis and leukemia.

69. (New) The method according to Claim 65, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin, hydroxypropyl-γ-cyclodextrin, randomly methylated β-cyclodextrin, carboxymethyl-β-cyclodextrin or sulfobutyl-β-cyclodextrin.

70. (New) The method according to Claim 66, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin, hydroxypropyl- γ -cyclodextrin, randomly methylated β -cyclodextrin, carboxymethyl- β -cyclodextrin or sulfobutyl- β -cyclodextrin.

71. (New) The method according to Claim 67, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin, hydroxypropyl- γ -cyclodextrin, randomly methylated β -cyclodextrin, carboxymethyl- β -cyclodextrin or sulfobutyl- β -cyclodextrin.

72. (New) The method according to Claim 68, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin, hydroxypropyl- γ -cyclodextrin, randomly methylated β -cyclodextrin, carboxymethyl- β -cyclodextrin or sulfobutyl- β -cyclodextrin.

73. (New) The method according to Claim 65, wherein the weight ratio of cladribine to amorphous cyclodextrin is from about 1:10 to about 1:16.

74. (New) The method according to Claim 69, wherein the weight ratio of cladribine to amorphous cyclodextrin is from about 1:10 to about 1:16.

75. (New) The method according to Claim 65, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin.

76. (New) The method according to Claim 67, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin.

77. (New) The method according to Claim 73, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin.

78. (New) The method according to Claim 73, wherein the amorphous cyclodextrin is hydroxypropyl-γ-cyclodextrin.

79. (New) The method according to Claim 75, wherein the weight ratio of cladribine to hydroxypropyl- β -cyclodextrin is about 1:14.

80. (New) The method according to Claim 75, wherein the weight ratio of cladribine to hydroxypropyl- β -cyclodextrin is about 1:11.

81. (New) The method according to Claim 65, wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b).

82. (New) The method according to Claim 75, wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b).

83. (New) The method according to Claim 65, wherein the cladribineresponsive condition is multiple sclerosis.

84. (New) The method according to Claim 75, wherein the cladribineresponsive condition is multiple sclerosis.

85. (New) The method according to Claim 77, wherein the cladribineresponsive condition is multiple sclerosis.

86. (New) The method according to Claim 83, wherein one or more additional active ingredients for treating multiple sclerosis is/are selected from the group consisting of interferon beta, glatiramer acetate, natalizumab, alemtuzumab, 4-aminopyridine and amantadine.

87. (New) The method according to Claim 84, wherein one or more additional active ingredients for treating multiple sclerosis is/are selected from the group

consisting of interferon beta, glatiramer acetate, natalizumab, alemtuzumab, 4aminopyridine and amantadine.

88. (New) The method according to Claim 85, wherein one or more additional active ingredients for treating multiple sclerosis is/are selected from the group consisting of interferon beta, glatiramer acetate, natalizumab, alemtuzumab, 4- aminopyridine and amantadine.

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REMARKS

Entry of the foregoing and further examination and consideration of the subject application, as amended, pursuant to and consistent with 37 C.F.R. § 1.112, are respectfully requested in light of the remarks that follow.

FOREIGN PRIORITY

The Examiner's acknowledgment of the claim for foreign priority and receipt of all certified copies of the priority documents is noted, with appreciation. However, the priority claims herein are domestic priority claims, this application being a continuation of a prior application which was a national stage of a PCT application, itself claiming benefit of several U.S. provisional applications.

DRAWINGS

Applicants thank the Examiner for accepting the drawings filed herein.

OBJECTION TO THE SPECIFICATION

The disclosure has been objected to because the reference to the copending application has not been updated. The specification has been amended to include the patent number of the patent granted on the parent application.

Withdrawal of the objection is believed to be in order and is respectfully solicited.

INFORMATION DISCLOSURE STATEMENT

The acknowledgment of the Information Disclosure Statements filed January 7, 2011 and August 9, 2012 and consideration of the documents cited therein are noted, with appreciation.

STATUS OF CLAIMS

Claims 1-88 are now in this application. Claims 9-24, 32-45 and 62-64 have been withdrawn. Claims 1-8, 25-31 and 46-61 are under examination. Claims 62-64 have been cancelled in favor of new Claims 65-88, which read on the same invention as cancelled Claims 62-64.

DISCUSSION OF CLAIM AMENDMENTS

New Claims 65-88 have been added to replace cancelled Claims 62-64. They are drawn to the same subject matter, which has been withdrawn from consideration. These new claims are fully based on pages 24-25 of the as-filed specification.

It is apparent from the foregoing that no new matter has been introduced by these claim amendments.

DISCUSSION OF RESTRICTION REQUIREMENT

The restriction requirement has been made final. Nevertheless, the withdrawn claims has been maintained. It is Applicants' intention to keep the withdrawn claims

commensurate in scope with those being examined, so that the withdrawn claims can be rejoined when patentable subject matter has been found.

Although Claims 65-88 are drawn to the same invention as Claims 62-64, Applicants point out that these methods of treatment claims differ from Claims 17-24 in requiring an additional active ingredient. If as a result, the Examiner believes that Claims 65-88 could not be rejoined for examination herein with the elected claims, he is asked to so advise the Applicants so that a separate divisional application can be promptly filed.

CLAIM REJECTIONS- 35 U.S.C. § 103

Claims 1-8, 25-31 and 46-61 have been rejected under 35 U.S.C. § 103(a) as being unpatentable over Schultz (U.S. Patent No. 6,194,395) in view of Baert (International Publication No. WO97/18839). It is submitted that this rejection cannot be maintained against any of the claims now in this application.

Schultz describes two kinds of cyclodextrin formulations of cladribine, i.e., "soluble aqueous formulations of cladribine with cyclodextrin solubilizers which are injectable in humans, as well as oral solid dosage forms containing a <u>mixture</u> of cladribine and cyclodextrins." (Emphasis added). (See col., 1, lines 6-10, of the Schultz patent.) Applicants do not dispute the fact that Schultz discloses hydroxypropyl-β-cyclodextrin (HPβCD), which is amorphous, but Schultz discloses crystalline cyclodextrins as well. Indeed, Schultz teaches aqueous formulations containing a cladribine/HPβCD <u>complex</u> in solution for injectable use. On the other hand, Schultz clearly teaches that their <u>solid</u> oral dosage forms are <u>mixtures</u> of

cladribine and cyclodextrin as set forth not only in col. 1, lines 8-10 ("oral solid dosage forms containing a mixture of cladribine and cyclodextrin") but also in col. 5, lines 50-64. Nowhere does Schultz teach a sold oral dosage form containing a complex. In col. 5, lines 50-64, it is indicated that the solid oral dosage forms may be prepared as disclosed in Baert; this is in fact the only method disclosed by Schultz for preparing the solid oral dosage forms. In col. 5, beginning at line 52, it is stated that "solid mixtures of the cvclodextrins with the active ingredient are prepared via melt-extrusion....the cladribine active ingredient and the cyclodextrins are mixed with other optional ingredients and then heated until melting occurs. The mixture is then extruded through an extruder having one or more nozzles." As set forth in col. 6 of Schultz, a typical oral dosage form has a formulation containing, as a milled extrudate, 1 mg to 15 mg of cladribine and 100 to 500 mg of cyclodextrin, and as excipients, 100 to 300 mg of microcrystalline cellulose, 10 to 200 mg of crospovidone, 1 to 5 mg of colloidal silicon dioxide and 2 to 10 mg of sterotex. Schultz does not disclose or suggest to the ordinary skilled pharmaceutical scientist solid oral formulation of cladribine/cyclodextrin complexes, as claimed herein.

As explained in detail below, Applicants respectfully direct the Examiner to a reference by Van Axel Castelli (hereinafter referred to as "Van Axel Castelli"). Van Axel Castelli explains that the characteristics of a complex and a physical mixture are distinctly different. Schultz's solid <u>mixtures</u> of cladribine and cyclodextrin cannot be assumed to be the same as Applicants' product, which is a complex cladribine-cyclodextrin complex. Even if the broad ratios of Schultz's mixtures encompass the

ratios in Applicants' complexes, having the same ratio does not give a physical mixture the same properties as a complex, a fact which is clearly shown by Van Axel Castelli.

Also, Baert, which is incorporated by reference in Schultz, for its melt-extrusion method of making solid oral dosage forms, describes amorphous materials and solid solutions but does not teach or suggest that its solid dosage form products include amorphous cyclodextrin-drug complexes. On the contrary Baert teaches:

(a) Prior art problems are solved by Baert by the use of a melt-

extrusion process to form solid <u>mixtures</u> comprising one or more cyclodextrins and insoluble active ingredients (Emphasis added; see page 3, lines 7-9).

(b) The compounds suitable for use in Baert's process "are compounds

that show no appreciable decomposition at the temperatures needed to melt and

extrude the mixture of said one or more active ingredients with the cyclodextrin or

cyclodextrins" (page 4, lines 5-7).

(c) Baert teaches that the characteristics of their products are different from those of a product obtained in water, since it is stated on page 6, lines 15-19:

It will be appreciated by a person skilled in the art that mixing two or more solids, i.e., one or more cyclodextrins and the active ingredient or ingredients, and subsequently melting these solids together will give rise to different products than when the said solids are first brought into contact with water or another solvent and then extruded.

(d) Baert's process uses very high temperatures; note Table 1 on page

12, where several different drugs are mixed with HPBCD and melt-extruded, with

temperatures of around 280°C being used and the products in every case, as noted on page 13, lines 5-6, being solid solutions.

(e) Baert does not mention cladribine; indeed, Applicants have previously shown that cladribine is known to decompose at the high temperatures used by Baert and thus cladribine falls in the group of compounds Baert teaches are unsuitable for use in their process [point (b) above].

(f) Baert in no way teaches or suggests that its products contain complexes.

The Examiner correctly states that the Schultz patent incorporates by reference the method of making their solid oral dosage form by utilizing the melt-extrusion process of Baert. The Baert process, as noted above, is carried out by mixing the cyclodextrin and the active ingredient, heating until melting one of the components, forcing the mixture through one or more nozzles, and cooling until the mixture solidifies (page 5, line 24-29). Milling may follow. The term "melting" is used broadly by Baert and includes transition to a glass; in particular cases, one component melts and the other dissolves therein forming solid solutions (page 5, lines 8-12). The extruded material may contain amorphous material or a solid solution (page 7, line 35 to page 8, line 7). While amorphous products are of interest, those which are mainly a solid solution are preferred (page 8, lines 11-23). Whether or not the products are amorphous, however, they are not taught to be complexes. In Table 1, on page 30, several different mixtures of hydroxypropyl- β -cyclodextrin and selected drugs were subjected to the Baert process. As noted on page 13, lines 5-6, in every case, the mixture using this

cyclodextrin gave a solid solution. The Examiner will note from Table 1 that the temperatures used, regardless of the identity of the drug, went as high as 292°C., with the temperatures for the itraconazole/HPβCD mixtures reaching 279°C-280°C. According to *The Merck Index* (copy of excerpt enclosed), itraconazole melts at 166.2°C while HPβCD melts at 278°C according to *LookChem* (copy of excerpt also attached).

The only solid dosage form envisioned by Schultz is a melt-extrusion product of cladribine and cyclodextrin prepared according to Baert. There is no evidence that such a product was ever prepared. Indeed, cladribine melts at 220-235°C with decomposition; see the enclosed excerpt from *Linscott's Directory* (copy attached) as well as that from *The Merck Index* (also enclosed). Thus, cladribine decomposes well below the 278°C melting point of HP β CD and well below the temperature used by Baert for their melt extrusion; cladribine is therefore not suitable for the Baert process, according to Baert's teaching that suitable compounds show <u>no appreciable</u> decomposition at the temperature they use (point b above).

Furthermore, Baert's teaching on page 6 that their melt-extrusion process affords <u>different products</u> than when their solids are first brought into contact with water (point c above) militates against the Examiner's finding that a cladribine/cyclodextrin product prepared by Baert's process is inherently the same as Applicants' product, which is, in fact, prepared by first contacting cyclodextrin with water. Indeed, it is the use of water that enables the formation of cyclodextrin-drug complexes; it is by complex formation that the water solubility of many drugs has been previously improved. There is no teaching by Baert that would lead one of ordinary skill to conclude that Baert et al made

solid complexes; indeed, Baert specifically teaches on page 6 that their products are different from products obtained by first dissolving cyclodextrin and drug in water. Thus, a melt-extrusion product of cladribine and cyclodextrin cannot inherently contain Applicants' product which comprises a complex cladribine-cvclodextrin complex which is an intimate amorphous admixture consisting of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex, formulated into a solid oral dosage form, said composition comprising no significant amount of free crystalline cladribine therein. Applicants' product is prepared by preparing the complex in water; Baert teaches obtaining a different product than one that can be obtained from water; moreover, Applicants' process uses temperatures up to only about 80°C, far below the decomposition temperature for cladribine and far below the temperatures used by Baert. See Claims 32-45 herein. It is only by ignoring the teachings of Baert and by ignoring the Van Axel Castelli data that the Examiner can conclude that the Schultz product of the Baert process would inherently be Applicants' product. Indeed, Applicants have provided data in the form of the Van Axel Castelli article which clearly proves the difference between a mixture and a complex having the same ratio of ingredients.

The unsuitability of Baert's temperatures and hence for the Baert process, as applied to cladribine with cyclodextrin, specifically with HPβCD, is furthermore proved by data in Van Axel Castelli, as discussed in more detail below.

Also, to provide background with respect to drug/cyclodextrin complexation, Applicants bring to the Examiners' attention, the Loftsson and Brewster cyclodextrin review article previously made of record; see Loftsson and Brewster, "Pharmaceutical Application of Cyclodextrins. 1. Drug Solubilization and Stabilization," *Journal of Pharmaceutical Sciences*, Vol. 85, No. 10, pp. 1017-1025, 1996, American Pharmaceutical Association and the American Chemical Society, US. The Loftsson and Brewster review article shows that it was known in the art that:

(a) Cyclodextrin/drug complexation typically is carried out in aqueous media, not by mixing in the absence of water (page 1020, left column, first full paragraph).

(b) This complexation involves many different forces (van der Waals, hydrogen bonding, etc.) and the use of water is essential to the formation of complexes (page 1018, right column, second full paragraph to page 1020, left column, noting in particular the mention of release of water molecules from the cyclodextrin cavity as a driving force for drug-cyclodextrin complex formation).

(c) The complexes have different properties from mere physical mixtures of drugs and cyclodextrins, for example in terms of drug solubilization and drug stabilization (pages 1020-1024), a fact shown for HPβCD and cladribine in Van Axel Castelli discussed below.

(d) Amorphous cyclodextrins such as HPβCD have almost countless isomeric and variably substituted forms and upon complexation result in amorphous

products which are mixtures of countless complexes (page 1018, left column, line 12 from the bottom, to page 1018, right column line 2).

Applicants also take the position that the use of cladribine in Baert's melt extrusion product is inappropriate because of the fact that the decomposition temperature of cladribine is lower than the Baert process's temperature.

Furthermore, the experimental data in Van Axel Castelli and discussed below prove that cladribine and an amorphous cyclodextrin such as HP β CD do not form an eutectic mixture. Rather, Van Axel Castelli as discussed below, shows that cladribine, whether in a complex or in a mixture with HP β CD, decomposes at temperatures far below those used for this cyclodextrin in the Baert process.

Turning to Van Axel Castelli, a further copy of which is provided herewith for the Examiner's convenience, the following additional remarks are offered:

Van Axel Castelli shows that an inclusion complex of cladribine and 2hydroxypropyl-β-cyclodextrin has properties that are different from those of the cyclodextrin, those of cladribine and those of physical mixtures of the cyclodextrin with cladribine. These can be summarized as follows:

(a) Thermo gravimetric analysis (TGA) was conducted over the temperature range from 25°C to 360°C for (a) cladribine, (b) HPβCD, (c) their inclusion complex and (d) their physical mixture and the results are shown in Fig. 2. Fig. 2a, the TGA curve for cladribine itself, shows decomposition of cladribine starting at about 200°C. Fig. 2b, the TGA curve for HPβCD, shows a mass loss of 6% at 30°C to about 140°C, due to dehydration, and decomposition at about 300°C. Fig. 2c, the TGA curve

for the cladribine/HPβCD complex, shows water loss between 20°C and 100°C and a decomposition process starting at about 250°C. Fig. 2d, the TGA curve for the cladribine plus HPβCD physical mixture, shows a multi-stage decomposition pathway; the first decomposition stage between room temperature and 100°C is due to loss of water from the cyclodextrin, whereas the second decomposition stage, observed at temperatures above 200°C, is due to the decomposition of cladribine. Fig. 2d also shows that heating the mixture to high temperatures does <u>not</u> lead to complexation but rather to <u>decomposition of cladribine</u>. Comparing the TGA for the complex with that of the physical mixture shows a slower degradation for the complex than for the mixture. Nevertheless, cladribine, whether in a complex or in a mixture with HPβCD, decomposes at temperatures far below those used for HPβCD-containing products in Baert's processes.

(b) Differential scanning calorimetry (DSC) analysis was conducted and the results shown in Fig. 3 for (a) cladribine, (b) HPβCD, (c) cladribine/HPβCD complex, (d) cladribine plus HPβCD physical mixtures, and (e) cladribine plus HPβCD kneading product. The authors note that the DSC trace of cladribine shows two endothermic events (Fig. 3a), the first at 206.3°C being close to the cladribine decomposition onset temperature and corresponding to the melting transition, and the second at 211.9°C, which is during the decomposition process, and probably is due to a decomposition product of cladribine. The DSC profile for the cyclodextrin (Fig. 3b) confirms an endothermic event corresponding to water loss from about 40°C to about 100°C. The authors further note that the DSC curves of the inclusion complex (Fig. 3c), physical

mixture (Fig. 3d) and kneading product (Fig. 3e) all show a broad thermal event from about 40 to 140°C, due to water loss in the cyclodextrin. In addition, the physical mixture and the kneading product are observed to feature two endothermic events around 200°C; these can be attributed to free cladribine in the mixture and kneading product. In contrast, Fig. 3c, the DSC trace for the complex, shows only one endothermic event, which occurs in the high temperature region, at 234.5°C; this is considerably higher than the degradation onset of pure cladribine around 200°C (Fig.2a), while the latter also characterizes the mixtures. As noted by the authors, the absence of thermal events typical of pure cladribine shows a loss of cladribine crystalline character for the complex. This also confirms DSC data for the products of instant Examples 1 and 2 reported on page 31 of the instant specification and correlates well with Applicants' x-ray diffraction traces for the products of Examples 1 and 2 reported on page 31 of the specification, where no peaks for crystalline cladribine were found in the complexes;

(c) FT-IR and FT-Raman spectroscopy were also recorded by Van Axel Castelli. In Figures 5 and 6, the FT-IR and FT-Raman spectra of cladribine alone (a), cladribine/cyclodextrin physical mixture (b), cladribine/cyclodextrin complex (c) and cyclodextrin alone (d) were compared. The authors indicate that the IR spectrum of the physical mixture (Fig. 5b) can be interpreted as the sum of the spectra for pure crystalline cladribine (Fig. 5a) and pure HP β CD (Fig. 5d), also supported by Fig. 6. The authors continue:
In contrast, both IR and Raman spectra of the inclusion complex show clear differences with respect to those of the physical mixture. In particular, markers of the crystalline phase of cladribine (arrows on Figs. 4 and 5) cannot be found in the spectra of the inclusion complex.

The authors further find that their data suggest that, when part of an inclusion complex, cladribine is present in a different (non-crystalline) phase relative to that of pure crystalline cladribine and that direct interaction between cladribine molecules is prevented. Further, they note that the amorphous phase has to be attributed to the formation of molecular complexes where the interaction between cladribine and HPβCD shields cladribine molecules, thus preventing crystallization.

(d) Van Axel Castelli also used nuclear magnetic resonance

spectroscopy to better understand the molecular interactions in the cladribine/cyclodextrin complex. To obtain direct proof of complex formation, the authors conducted a 2D ROESY experiment, the results of which showed a typical inner portion of HPβCD, confirming that a host-guest inclusion complex had formed between cladribine and HPβCD. Further, the authors conducted ¹³C CP-MAS NMR experiments and reported the spectra in Fig. 10 for cladribine (a), cladribine + HPβCD physical mixture (b) and cladribine/ HPβCD complex (c). In the cladribine spectrum (Fig. 10a), the authors observed sharp peaks due to cladribine's high degree of crystallinity. The spectrum for the physical mixture (Fig. 10b) corresponds to the sum of the spectra of the two components, with no resonance peaks or line broadening, showing no intermolecular interaction in the mixture, the solid being composed of distinct ordered domains of each component. In the spectrum for the complex, no shift in the HPβCD

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signals are detected, whereas the cladribine resonances are broadened and only slightly detectable. The authors note that this indicates that no crystalline domains of cladribine are present.

(e) Van Axel Castelli also conducted DSC and TGA thermal profiles for tablets of the cladribine/ HP β CD complex and found them comparable to those for the cladribine/ HP β CD complex itself. Moreover, even stressed tablets showed no notable differences in the DSC thermal profile, which demonstrated that they were storage stable even under less than ideal conditions.

(f) Van Axel Castelli concludes that thermal analyses, vibrational analyses, and solid-state NMR all indicate that cladribine behaves differently when in the complex compared with the physical mixture or kneading product, while ROESY provides evidence for the existence of an internal complex between cladribine and HPβCD. They further conclude that tablets of the complex are not affected by their manufacturing from the complex itself and are storage stable. These tablets have been used in a successful clinical trial for oral treatment of patients with MS (CLARITY trial).

In summary, Applicants submit that the obviousness rejection based on Schultz in view of Baert is untenable and should be withdrawn. The data provided by Van Axel Castelli conclusively show that cladribine, whether alone, in a physical mixture with HP β CD, or even in a cladribine/HP β CD complex, decomposes at temperatures far lower than those used by Baert for melt extruding drugs with the same cyclodextrin. The data further show that heating a physical mixture of cladribine with HP β CD to high temperatures does not result in a complex of cladribine and in fact the cladribine in the

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mixture decomposes long before the melting point for HPβCD is reached. Thus, the Baert process is not suitable for making a melt extrudate of cladribine with hydroxypropyl-β-cyclodextrin and moreover such a product prepared according to Baert would not contain a complex of cladribine with the cyclodextrin as claimed in this application. The Schultz oral dosage form prepared by the Baert process simply cannot contain a cladribine/cyclodextrin complex.

Applicants emphasize that Baert melt-extrusion product is not one obtained by complexation in water; Baert teach their melt-extrusion product is different from a product whose preparation begins by dissolving the drug and cyclodextrin in water. Therefore, any product that Schultz might produce from cladribine and cyclodextrin subjected to Baert's melt extrusion product cannot be the same as Applicants' complex cladribine-cyclodextrin complex which must be obtained from an aqueous solution which is treated in a specific manner. Baert never suggests that they obtain a complex by their melt-extrusion process, much less one meeting the requirements of Applicants' claims. Indeed, Baert emphasize that their process, which is different, affords a different product than that obtained by first dissolving the drug and cyclodextrin. Likewise, Applicants emphasize that Applicants' process is strikingly different from Baert's process and thus logically would not afford the product which Schultz would have expected to obtain by subjecting cladribine and cyclodextrin to Baert's process. Moreover, Applicants have formed a very special complex which contains a large amount of cladribine as an amorphous inclusion complex and as amorphous free cladribine associated with the cyclodextrin as a non-inclusion complex. Note too that

the <u>free</u> cladribine associated with the non-inclusion product is amorphous, in contrast to the cladribine starting material, which is crystalline. Note also that Applicants produce their product by first complexing in water at temperatures of from about 45°C to about 80°C, far below the temperatures used by Baert. Cladribine actually decomposes at temperatures below that used by Baert.

It is clear from the foregoing that a molecular inclusion complexation process, let alone the particular inclusion process utilized by Applicants to form their unique complex cladribine-cyclodextrin complex, is <u>not inherent</u> in Baert's melt extrusion process and that Baert's process gives a different product. To hold otherwise would be to ignore Baert's own teachings. Claims 46-61 are drawn to a product-by-process. These claims depend, directly or indirectly, from Claim 1 and thus include all of the features in the combination of features defined in Claim 1. Applicants have shown that the composition of Claim 1 is free of the art, therefore, the product-by-process claims are also patentable over the art for at least the reasons that Claim 1 is patentable.

For at least the reasons set forth above, withdrawal of the § 103 rejection is in order and is earnestly solicited.

DOUBLE PATENTING REJECTION

Claims 1-8, 25-31 and 46-61 have been rejected on the ground on non-statutory double patenting as being unpatentable over Claims 1-28 of U.S. Patent No. 7,888,328. The Examiner has erred in issuing what appears to be a "same invention" double patenting rejection, which would only be proper if the claims were of <u>identical</u> scope to

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those granted in the '328 patent. The instant claims are broader than those granted in the parent, therefore the double patenting "same invention" rejection is improper. Applicants are willing to file a terminal disclaimer with respect to the '328 patent to obviate an obviousness-type double patenting rejection, but such a rejection does not appear to have been made. There is no such thing as an "anticipation-type nonstatutory double patenting rejection." There are only two common types of double patenting rejections: 1) a "same invention" type double patenting rejection based on 35 U.S.C. § 101, which is statutory; and 2) a nonstatutory-type double patenting rejection prohibiting claims in a second patent not patentably distinguishing from claims in a first. This case fits in the second category, and the rejection clearly can be obviated by the filing of a terminal disclaimer of the commonly owned patent and application. An uncommon type of double patenting situation arose in In re Schneller but the first situation there is completely different than here. Moreover, in that case, the applicant had not filed a terminal disclaimer, whereas here an electronic terminal disclaimer has been filed today and accepted by the USPTO. It is also pointed out that Applicants did present claims corresponding to those herein in the parent application, so the Examiner's further statement relying on In re Schneller is simply incorrect. Moreover, as noted, in In re Schneller, no terminal discaimer had been filed.

CONCLUSION

The foregoing amendments and remarks are being made to place the application in condition for allowance. Applicants respectfully request entry of the amendments,

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reconsideration, and the timely allowance of the pending claims. A favorable action is awaited. Should an interview be helpful to further prosecution of this application, the Examiner is invited to telephone the undersigned.

If there are any additional fees due in connection with the filing of this response, please charge the fees to our Deposit Account No. 19-3140. If a fee is required for an extension of time under 37 C.F.R. §1.136 not accounted for above, such an extension is requested and the fee should also be charged to our Deposit Account.

Respectfully submitted,

DENTONS US LLP

Date: May 28, 2013

Mary Katherine Baumeister Registration No. 26254

Customer No. 13974 SNR Denton US LLP 1301 K Street NW, Suite 600, East Tower Washington, D.C. 20005 Phone: 202-408-9186 Fax: 202-408-6399

Attachments:

"Cladribine", The Merck Index, (2001), pp. 407-408, Thirteenth Edition, Merck & Co., Inc., Whitehouse Station, NJ

"Itraconazole", The Merck Index, (2001), p. 938, Thirteen Edition, Merck & Co., Inc. Whitehouse Station, NJ

"Hydroxypropyl-beta-cyclodextrin, CAS No. 94035-02-6" *LookChem*, http://www.lookchem.com/cas-940/94035-02-6.html, September 23, 2008

"Non-antibody Products (Kits, Proteins, Microbial Antigens, Tissues, Services, etc.) Linscott's Directory of Immunological & Biological Reagents, http://www.linscottsdirectory.com/browse/products/page:36, Records 1,751-1,800 of 130,353, September 11, 2008

Characterisation of an Inclusion Complex Between Cladribine and 2-Hydroxypropyl-β-Cyclodextrin

VALERIA VAN AXEL CASTELLI,¹ GIOVANNI TRIVIERI,¹ ILARIA ZUCCHELLI,¹ LUIGI BRAMBILLA,² TONY BARBUZZI,¹ CHIARA CASTIGLIONI,² MAURIZIO PACI,³ GIUSEPPE ZERBI,² MARGHERITA ZANOL¹

¹Merck Serono SpA, Tiburtina Site, via L. Einaudi 11, 00012 Guidonia Montecelio, Roma, Italy

²Dipartimento di Chimica, Materiali e Ingegneria Chimica "G. Natta", Politecnico di Milano, P. za Leonardo da Vinci 32, 20133 Milano, Italy

³Dipartimento di Scienze e Tecnologie Chimiche, Università di Roma "Tor Vergata", via della Ricerca Scientifica, 00133 Roma, Italy

Received 13 June 2007; revised 12 November 2007; accepted 15 November 2007

Published online in Wiley InterScience (www.interscience.wiley.com). DOI 10.1002/jps.21283

ABSTRACT: Parenterally administered cladribine (2-chloro-2'-deoxyadenosine) has demonstrated promising efficacy and safety in clinical trials in patients with multiple sclerosis (MS). An oral formulation of this small molecule would be an attractive option for patients. Here, we describe the chemical characterisation of the inclusion complex between cladribine and the drug carrier molecule 2-hydroxypropyl-ß-cyclodextrin (2-HP- β -CD). Several techniques were used to analyse the complex both in solution and in the solid state. These analyses provided evidence that the inclusion complex cannot be simply reduced to the sum of the two species, as it shows behaviour different from that of the physical mixture of the two components. Furthermore, solution nuclear magnetic resonance spectroscopy demonstrated the existence of an inclusion complex between cladribine and 2-HP-β-CD. Importantly, analysis of a tablet formulation demonstrated that the chemical characteristics of the inclusion complex are not affected by the manufacturing process, and that the complex is stable during storage. This tablet formulation is currently under investigation for the treatment of patients with MS. © 2008 Wiley-Liss, Inc. and the American Pharmacists Association J Pharm Sci 97:3897–3906, 2008 Keywords: cladribine; cyclodextrins; inclusion compounds; FT-IR; NMR spectroscopy; thermal analysis; oral drug delivery

INTRODUCTION

This article contains supplementary material, available at www.interscience.wiley.com/jpages/0022-3549/suppmat.

Correspondence to: Valeria Van Axel Castelli (Telephone: +39-774-350480; Fax: +39-774-350386;

E-mail: valeria.vanaxel@merckserono.net)

Journal of Pharmaceutical Sciences, Vol. 97, 3897-3906 (2008) © 2008 Wiley-Liss, Inc. and the American Pharmacists Association



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sclerosis (MS) are administered parenterally, which is inconvenient and can be associated with localised pain.¹ An oral agent would be an attractive option for patients, and a number of oral therapies are currently in development for the treatment of MS. Cladribine (2-chloro-2'-deoxyadenosine), an analogue of adenosine (Fig. 1), is a preferential lymphocyte-depleting therapy that targets both resting and proliferating lymphocytes.^{2,3} Cladribine is licensed for the treatment

All currently available treatments for multiple

The work described in this article was carried out at Merck Serono SpA, Tiburtina Site, via L. Einaudi 11, 00012 Guidonia Montecello, Roma, Italy. Some additional measurements were carried out at Dipartimento di Chimica, Materiali e Ingegneria Chimica "G. Natta", Politecnico di Milano, P. za Leonardo da Vinci 32, 20133 Milano, Italy.



2-HP-B-CD





of hairy cell leukaemia and B-cell chronic lymphocytic leukaemia, and has shown promising efficacy and a good safety profile in patients with MS (particularly patients with the relapsing-remitting form of the disease) when administered by intravenous infusion or subcutaneous injection.4-8 As cladribine is a small molecule (molecular weight = 285.7 Da), it is an ideal candidate for development as an oral tablet formulation.

Cyclodextrins (CDs) are rationally designed drug carriers capable of modifying the physical, chemical and biological properties of drug molecules through the formation of inclusion complexes.⁹⁻²⁵ In general, hydrophilic CDs are employed to enhance the solubility and dissolution rate of poorly water-soluble drugs, whereas hydrophobic CDs are used to slow the dissolution rate of water-soluble drugs.^{9,21,22} Advanced controlled release can be achieved by a combination of hydrophilic and hydrophobic CDs or pharmaceutical additives.²⁵ 2-Hydroxypropyl-β-cyclodextrin (2-HP- β -CD; Fig. 1) is a highly water-soluble CD, which has been used as an encapsulating agent for a variety of drugs, such as ibuprofen,¹² meloxicam,¹⁶ tanshinone compounds¹⁹ and ketoprofen,^{12,20} among others. The formation of host-guest inclusion complexes between pharmacologically active small organic molecules and

2-HP-B-CD has been shown to improve the solubility, stability and bioavailability of these molecules. It has also been proposed that formation of a 2-HP- β -CD complex of cladribine to increase its water solubility may produce an orally bioavailable formulation of this drug.²⁶

In this article the characterisation of the inclusion complex of cladribine with 2-HP-β-CD, both in solution and in the solid state, is reported. Analytical methods used include thermal analysis, mass spectrometry, Fourier-transform infrared (FT-IR) spectroscopy, FT-Raman and nuclear magnetic resonance (NMR) spectroscopy.

MATERIALS AND METHODS

Materials

Cladribine, 2-HP-\beta-CD and the cladribine/2-HP- β -CD inclusion complex²⁸ were obtained from Ivax Pharmaceuticals (Waterford, Ireland). The physical mixture of cladribine and 2-HP-B-CD was prepared by mixing the two components with a 1:1 molar ratio and subsequent homogenisation. Kneading samples were prepared²⁷ by adding a solution of methanol/water 50:50 (v/v) to the physical mixture to obtain a homogeneous paste. The cladribine/2-HP-β-CD tablet formulation was obtained from Ivax Pharmaceuticals and analysed by thermo gravimetric analyses (TGA) and differential scanning calorimetry (DSC). 'Stressed' tablets were produced by heating for 2 weeks at 40°C and 75% relative humidity.

Thermo Gravimetric Analysis

TGA was conducted using a TA Q-500 apparatus (TA Instruments, New Castle, DE). Samples were heated at a rate of 10°C/min, from 30 to 350°C in a dynamic nitrogen atmosphere.

Differential Scanning Calorimetry

DSC measurements were made on a TA Q-1000 differential scanning calorimeter (TA Instruments) using an aluminium nonhermetic pan. The heating rate was 10°C/min in the range of 20-300°C.

FT-IR and FT-Raman Spectroscopy

FT-IR and FT-Raman spectra of pure samples of cladribine, and the cladribine/2-HP-6-CD physical mixture and inclusion complex, were analysed.

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DOI 10.1002/jps

Samples were analysed in the solid state (sample as prepared, physical mixture, or after kneading) or in dimethyl sulfoxide (DMSO) solution. FT-IR spectra of solution samples were recorded on a Nicolet NEXUS (ThermoElectron, Waltham, MA) spectrometer in ATR mode ("SMART performer" module), in the spectral region from 4000 to 650 cm^{-1} . Each experiment was performed with 512 scans at a resolution of 2.0 cm^{-1} . Solid samples were analysed as pellets, prepared by grinding a few milligrams of sample with 40 mg of KBr and then compressing the powder at a pressure of 10 ton/cm². FT-IR infrared spectra of solid samples were recorded on a Nicolet NEXUS FTIR spectrometer (ThermoElectron). For each spectrum, 256 scans were collected with a resolution of 1 cm^{-1} in the spectral range 400-4000 cm⁻¹. Raman experiments were carried out using a Nicolet NXR 9650 instrument (Thermo-Electron).

Electrospray Mass Spectrometry

Mass spectra were acquired on a triple quadrupole spectrometer equipped with a turbo-ion spray source (Q-Trap from Applied Biosystems, Foster City, CA). Samples were introduced into the source at a flow rate of 10 μ L/min using a syringe pump. All the spectra were acquired in positive ion mode in a spectral range from 100 to 1700 m/z. The ionisation voltage was set at 4500 V and the declustering potential was set to 30 V.

Nuclear Magnetic Resonance

NMR experiments were carried out at 298 K on a Bruker Avance500 spectrometer (Bruker BioSpin, Rheinstetten, Germany), operating at 500.13 MHz for ¹H. Spectra in solution were performed using a 5 mm PABBI probe with a z-gradient unit. Samples for NMR experiments in solution were prepared by dissolving ~ 15 mg of cladribine, 2-HP- β -CD, or the inclusion complex in 700 μ L of D₂O or d_6 -DMSO. ¹H NMR spectra in d_6 -DMSO were performed with a pulse width (pw, 90°) of 6.8 µs. ¹H NMR spectra in D₂O were performed applying a sequence with water suppression using excitation sculpting with gradients;²⁸ pw (90°), 7.2 μ s; sculpting pw, 2 ms. ¹H-¹H 2D experiments (COSY [COrrelation SpectroscopY], TOCSY [TOtal Correlation SpectroscopY], NOESY [Nuclear Overhauser Enhancement SpectroscopY] and **ROESY** [Rotating frame Overhauser Enhance-

DOI 10.1002/jps

ment SpectroscopY]) were carried out in States-TPPI phase sensitive mode, using excitation sculpting with gradients for water suppression²⁸ in conditions analogous to those applied for 1D spectra. Time domain (TD){F2}, 2K; TD{F1}, 512; sweep width (SW), 10 ppm. ¹H-¹H TOCSY experiments were carried out with a mixing time of 40 ms, using an MLEV-17 sequence. In the case of ¹H-¹H ROESY experiments, a mixing time of 200 ms was set. Solid-state NMR spectra were performed with a MAS BB 4 mm probe. The sample-spinning rate was set at 5.0 kHz.

 13 C cross-polarisation magic angle spinning (CP-MAS) NMR spectra were acquired with 3.20 µs proton 90° pw, 1.0 ms contact time, and 60 s recycle time. Chemical shifts are expressed in ppm downfield from tetramethylsilane. No internal reference was used in order to avoid binding interference with the CD.

RESULTS AND DISCUSSION

Thermo Gravimetric Analysis

The TGA profiles of cladribine, 2-HP- β -CD, their inclusion complex, and physical mixture from 25 to 360°C are presented in Figure 2. The TGA curve of cladribine indicates a decomposition process starting at about 200°C (Fig. 2a). In the case of 2-HP- β -CD, a mass loss of 6% was observed from 30°C to about 140°C, due to dehydration (Fig. 2b). The TGA of the cladribine/2-HP- β -CD inclusion complex showed water loss between 20 and 100°C



Figure 2. Thermo gravimetric analysis profiles from 25° C to 360° C for (a) cladribine; (b) 2-hydroxypropyl- β -cyclodextrin (2-HP- β -CD); (c) cladribine/2-HP- β -CD inclusion complex; and (d) cladribine plus 2-HP- β -CD physical mixture.

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and a decomposition process starting at about 250°C (Fig. 2c). The decomposition temperature for the inclusion complex was significantly higher than the decomposition temperature observed for pure cladribine (250°C vs. 200°C, respectively). The TGA curve of the cladribine plus 2-HP-β-CD physical mixture shows a multi-stage decomposition pathway (Fig. 2d). The first stage between room temperature and 100°C is due to the water content loss of CD, whereas the second stage, observed at temperatures above 200°C, is due to the decomposition of cladribine. By comparing the decomposition stage of TGA for the complex and the physical mixture, it appears that the complex degrades slower than the mixture. The higher thermal stability of the complex is an indication of the onset of host-guest interactions.²⁹

Differential Scanning Calorimetry

In order to better understand the thermal behaviour of the samples, DSC analysis was conducted (Fig. 3). The DSC trace of cladribine shows two endothermic events (Fig. 3a). The first endothermic event at 206.3°C is close to the cladribine decomposition onset temperature and corresponds to the melting transition. The second endothermic event, occurring at 211.9°C, that is during the decomposition process, is probably due to a decomposition product of cladribine. The DSC profile of 2-HP- β -CD confirms an endothermic event corresponding to the water loss (Fig. 3b). The DSC curves of the inclusion complex, physical mixture and kneading product all show a broad



Figure 3. Differential scanning calorimetry profiles of (a) cladribine; (b) 2-HP- β -CD; (c) cladribine/2-HP- β -CD inclusion complex; (d) cladribine plus 2-HP- β -CD physical mixture; and (e) cladribine plus 2-HP- β -CD kneading product.

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thermal event (from 40 to 140°C), due to water loss in the CD (Fig. 3c-e). In addition, the physical mixture and the kneading product feature two endothermic events around 200°C, which can be attributed to the free cladribine.

Remarkably, the DSC trace of the inclusion complex (Fig. 3c) shows only one endothermic event in the high temperature region, at 234.5°C, which is a temperature considerably higher than the degradation onset of pure cladribine (~200°C; Fig. 2a). The absence of thermal events typical of pure cladribine indicates a loss of cladribine crystalline character when it is involved in the complex. This represents indirect evidence for the formation of the cladribine/2-HP- β -CD inclusion complex.

FT-IR and FT-Raman Spectroscopy

The FT-IR and FT-Raman spectra of several samples were recorded to detect spectral signals (band frequencies and band intensities) that may reveal the existence of intermolecular interactions between cladribine and 2-HP- β -CD in the inclusion complex.

In Figure 4, the FT-Raman spectra of different samples of cladribine (as prepared, after kneading, and in solution) in the region $1700-200 \text{ cm}^{-1}$ are reported. The FT-Raman spectra of the two solid-state samples (as prepared and after kneading) are practically coincident, indicating that the procedure adopted to prepare the complex does not affect the physical state of the molecule. In DMSO solution, cladribine molecules cannot



Figure 4. Fourier-transform Raman spectra of (a) cladribine as prepared; (b) cladribine after kneading; (c) cladribine in DMSO (after subtraction of DMSO spectrum). Arrows indicate peaks of interest referred to in the text.

DOI 10.1002/jps

interact with each other because of the surrounding solvent, and the Raman spectrum undergoes drastic changes. For example, the Raman lines at 1513 and 413 cm⁻¹, characteristic of pure cladribine, strongly decrease in intensity in the spectrum of solution (see arrows on Fig. 4c).

The FT-IR spectrum of cladribine (as prepared) shows many sharp bands. In the high frequency region, it is possible to identify the O-H stretching band of isolated (free) OH bonds at 3542 cm^{-1} and the four well-defined N-H stretching bands in the region $3500-3150 \text{ cm}^{-1}$ (see arrows on Fig. 5a).

Both Raman and IR spectra show very sharp vibrational bands for pure cladribine, indicating the molecules are in a structurally defined and regular environment (i.e. in a crystalline field). This observation supports the hypothesis (confirmed by NMR experiments, see below in the text) that pure cladribine forms a crystalline phase. Moreover, comparisons between solid samples and samples in DMSO solution indicate that the Raman spectroscopic signals at 1513 and 413 cm⁻¹ can be used as markers of the crystalline phase of cladribine. The bands at 3542 cm^{-1} and 3500-3150 cm⁻¹ in the IR spectrum also indicate the presence of a well-defined crystalline structure. In particular, the band observed at 3542 cm^{-1} , assigned to the stretching of free OH bonds, is in agreement with the crystal structure of cladribine as reported in the literature.¹² Mura et al.¹² noted the presence of OH bonds (one for each cladribine molecule), which do not form intermolecular hydrogen bonds.

In Figures 5 and 6, the FT-IR and FT-Raman spectra of cladribine, 2-HP- β -CD, their inclusion complex and physical mixture are compared. The



Figure 5. Fourier-transform infrared spectra of (a) cladribine as prepared; (b) cladribine plus 2-HP- β -CD physical mixture; (c) cladribine/2-HP- β -CD inclusion complex; and (d) 2-HP- β -CD. Arrows indicate peaks of interest referred to in the text.





Figure 6. Fourier transform Raman spectra of (a) cladribine as prepared; (b) cladribine plus 2-HP- β -CD physical mixture; (c) cladribine/2-HP- β -CD inclusion complex; and (d) 2-HP- β -CD. Arrows indicate peaks of interest referred to in the text.

IR spectrum of the physical mixture in Figure 5 can be interpreted as the 'sum' of the spectra for pure cladribine and pure 2-HP- β -CD. This is a clear indication of the presence of two separate phases: crystalline cladribine and 2-HP- β -CD. The same conclusion is reached from the analysis of the FT-Raman spectra in Figure 6.

In contrast, both IR and Raman spectra of the inclusion complex show clear differences with respect to those of the physical mixture. In particular, markers of the crystalline phase of cladribine (arrows on Figs. 4 and 5) cannot be found in the spectra of the inclusion complex. Moreover, the FT-Raman spectral features indicated by arrows on Figure 6 (ascribed to cladribine in the inclusion complex) show similarities with those of cladribine in DMSO solution (see Fig. 4c). This suggests that, when part of the inclusion complex, cladribine is present in a different phase relative to that of pure (crystalline) cladribine, and that direct interaction between cladribine molecules is prevented. Therefore, it seems that 2-HP-B-CD acts in a similar way to a solvent, shielding each cladribine molecule from other molecules of the same species. Cladribine samples obtained by kneading show the same spectral markers of crystallinity observed in the sample as prepared (Fig. 4), indicating that the origin of this 'amorphous' phase cannot be explained by differences in methods of preparation for the inclusion complex and crystalline cladribine. The origin of the amorphous phase, therefore, has to be attributed to the formation of molecular complexes where the interaction between cladribine and 2-HP-β-CD shields cladribine molecules, thus preventing crystallisation.

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Electrospray Ionisation Mass Spectrometry (ESI-MS) Analysis

ESI-MS is a powerful tool for the study of noncovalent complexes of CDs.^{30–33} The mild ionisation procedure allows the solution state structure of the complexes to be retained in the gaseous phase.³⁴ 2-HP- β -CD and the cladribine/2-HP- β -CD inclusion complex were analysed using ESI-MS.

The spectrum of 2-HP-\beta-CD (see Supporting Information) consists of a main distribution of single-charged ions, starting from m/z 1330 atomic mass units up to 1679 atomic mass units in which each peak is separated by 58 atomic mass units, corresponding to the 2-hydroxypropyl moiety. The spectrum reveals that 2-HP-β-CD is characterised by a degree of substitution, ranging from one to seven substitutions, in which the numerically most abundant species is CD substituted with four 2-hydroxypropyl groups. The ESI mass spectrum of the cladribine/2-HP-β-CD inclusion complex shows two main distributions (see Supporting Information). The first distribution, centred at m/z 1505 atomic mass units represents 2-HP- β -CD/Na⁺ adducts. The second distribution, centred at m/z 790 atomic mass units, consists of double-charged ions assignable to 2-HP-3-CD. The ESI-MS spectrum of the cladribine/2-HP-β-CD complex shows some differences with respect to the spectrum of free CD. However, clear evidence of the presence of the inclusion complex is not observed. Probably the ionisation process, in this case, does not allow survival of the complex.

Nuclear Magnetic Resonance Spectroscopy

Several high-resolution 1D and 2D NMR experiments were carried out in order to better understand the molecular interactions in the cladribine/ 2-HP- β -CD inclusion complex. ¹H-NMR spectra of cladribine and 2-HP- β -CD in d_{6} -DMSO are reported in Figure 7a and b, respectively. The assignment of ¹H resonances of cladribine in d_{6} -DMSO is reported in Table 1. The assignment of ¹H resonances of 2-HP- β -CD in d_{6} -DMSO is rather complex, and ¹H resonances of 2-HP- β -CD in D_{2} O are presented in Table 1. The spectrum of 2-HP- β -CD in D₂O is simplified because of the chemical exchange of OH with deuterium atoms (Fig. 7c).

Assignments shown in Table 1 for proton resonances of 2-HP- β -CD in D₂O are based on



Figure 7. 500.13 MHz ¹H NMR spectra in solution: (a) CdA in d_6 -DMSO; (b) 2-HP- β -CD in d_6 -DMSO; (c) 2-HP- β -CD in D₂O with water suppression; (d) CdA/ 2-HP- β -CD complex in D₂O with water suppression.

the literature^{35,36} and the analysis of several 2D NMR experiments,³⁷ namely, COSY, TOCSY and NOESY (see Fig. 8 and Supporting Information). The COSY pulse sequence gives rise to a 2D map in which cross-peaks appear between *J*-coupled protons. In the TOCSY experiment, cross-peaks are shown between all protons connected in a chain. The cross-peak intensity decreases with increasing distance between protons along the chain. Finally, the NOESY experiment gives correlation signals due to dipolar cross-relaxation between nuclei in a close spatial relationship. Cross-peak intensity is proportional to the distance between protons.

Two types of anomeric protons can be distinguished (due to regioisomers),³⁶ indicated as H1 and H1' in Figure 8, resonating at 5.17 and 4.99 ppm, respectively. From the TOCSY experiment (Fig. 8) it is also possible to differentiate protons H2 and H3, from protons H2' and H3'. It should be noted that for 2-HP- β -CD, inner protons (H3, 3' and H5, 5') show a downfield shift with respect to protons that are outside the cavity (H1, 1', H2, 2', H4, 4' and H6, 6'). This observation is consistent with the literature.³⁵

The ¹H NMR spectrum of the cladribine/2-HP- β -CD inclusion complex in D₂O is reported in

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DOI 10.1002/los

NH2 N b	$ \begin{array}{c} $
	3 H OR H OR R=H or −CH ₂ -CHCH ₃ 7 8 9
Cladribine	2-HP-β-CD

Table I. Assignment of "H NMR resonances of cladribine a
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oradi totino		2-111-p-60
Τγρε δ (ppm)		Multiplicity, J (Hz)
Cladribine in d_6 -DMS	0	
a	7.83	bs
b	8.35	8
C	6.26	dd $(J = 7.3, 6.2)$
d	2.64	ddd $(J = 13.3, 7.3, 5.8)$
ď	2.27	ddd $(J = 13.3, 6.2, 3.3)$
0	4.38	dddd $(J = 5.8, 4.2, 3.3, 2.8)$
ſ	5.31	d(J = 4.2)
g	3.85	dt $(J = 4.6, 2.8)$
ĥ	3.59	ddd $(J = 11.8, 5.4, 4.6)$
h'	3.51	ddd $(J = 11.8, 6.0, 4.6)$
i	4.96	dd $(J = 6.0, 5.4)$
2-HP-B-CD in D ₂ O		
1	5.17	bs
1'	4.99	bs
2	3.40	m
2'	3.5	m
3	3.96	m
3'	3.86	m
4.4'	3.55	Overlapped to other signal
5.5'	3.80-3.72	Overlapped to other signals
6.6'	3.78	Overlapped to other signals
7	3.71.3.64. 3.53.3.46	• m
8	8.94	m
9	1.08	bs

Figure 7d, and corresponding COSY and TOCSY maps are reported in the Supporting Information.

To obtain direct proof of complex formation, a 2D ROESY experiment was conducted. As with the NOESY experiments, protons that are spatially adjacent give rise to cross-peaks in ROESY spectra. An expansion of the ROESY map of the cladribine/2-HP- β -CD inclusion complex in D₂O is reported in Figure 9. In addition to the expected intramolecular cross-peaks, some intermolecular cross-peaks were observed in the spectra. In particular, cross-peaks between H3, H3', H5 and H5' of 2-HP- β -CD and protons d, d' of cladribine indicate that cladribine and CD are

in close proximity. This interaction involves a typical inner proton of 2-HP- β -CD, indicating that a host-guest inclusion complex has formed between cladribine and 2-HP- β -CD. The NOE experiment was used solely as a diagnostic tool and, therefore, no quantitative information about interaction strength was determined.

Solid-state behaviour was investigated by means of ¹³C CP-MAS NMR experiments.³⁷ The ¹³C CP-MAS spectra of cladribine, the cladribine plus 2-HP- β -CD physical mixture, and cladribine/ 2-HP- β -CD inclusion complex are reported in Figure 10. In the cladribine spectrum, sharp peaks due to its high degree of crystallinity are

DOI 10.1002/jps

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Figure 8. Enlarged region of the ¹H-¹H TOCSY NMR map of 2-HP- β -CD in D₂O. The assignment of resonances is reported in Table 1.

observed (Fig. 10a). The ¹⁸C CP-MAS spectrum of the physical mixture corresponds to the sum of the spectra of the two components, cladribine and 2-HP- β -CD (Fig. 10b). No resonance shifts or line broadening are detected in this spectrum, clearly



Figure 9. Enlarged region of the ${}^{1}H{}^{-1}H$ ROESY NMR map of CdA/2-HP- β -CD complex in D₂O. The correlation between protons d, d' of CdA and inner protons H3, H3', H5, H5' of 2-HP- β -CD (see Table 1) is indicated.

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Figure 10. ¹³C CP-MAS spectra of: (a) CdA; (b) CdA + 2-HP- β -CD physical mixture; (c) CdA/2-HP- β -CD complex.

indicating that no intermolecular interaction exists between cladribine and 2-HP- β -CD, and that the solid is composed of distinct ordered molecular domains of each component. Thus, solid-state properties of the mixture are to be considered simply as cladribine domains interspersed among domains of 2-HP- β -CD.

The ¹⁸C CP-MAS spectrum of the cladribine/ 2-HP- β -CD inclusion complex reveals some interesting features (Fig. 10c). No chemical shift perturbation of 2-HP- β -CD signals is detected, whereas the cladribine resonances are broadened and only slightly detectable (see in particular the spectral region δ 20–50 ppm). This indicates that no crystalline domains of cladribine are present. In fact, these results show that the sample consists of cladribine dispersed at a molecular level in the 2-HP- β -CD solid.

Analysis of Cladribine/2-HP- β -CD Tablet Formulation

DSC and TGA thermal profiles for the cladribine/ 2-HP- β -CD tablet formulation were comparable with those for the cladribine/2-HP- β -CD inclusion complex sample (data not shown). This indicates that the cladribine/2-HP- β -CD inclusion complex is not affected by the manufacturing procedures used to prepare the tablet. Thermal analysis of the stressed tablet formulation showed no notable differences in DSC thermal profile but higher water content in the TGA profile, compared with the cladribine/2-HP- β -CD inclusion complex. This demonstrates that the complex is stable on storage.

DOI 10.1002/jps

CONCLUSIONS

This publication reports the first comprehensive chemical analysis of the cladribine/2-HP-β-CD inclusion complex, which provides indirect evidence for the presence of intermolecular interactions between cladribine and 2-HP-B-CD in this complex. Thermal analyses (both TGA and DSC), vibrational analyses (both FT-IR and FT-Raman), and solid-state NMR all indicate that cladribine behaves differently when in the inclusion complex compared with the physical mixture or the kneading product. Finally, detection of NOE cross-peaks due to the 2-HP-B-CD inner hydrogens (H3, 3', and H5, 5') characteristically located inside the CD cavity (ROESY) provides evidence for existence of an internal complex between cladribine and 2-HP-β-CD.

Importantly, analysis of the cladribine/2-HP- β -CD tablet formulation demonstrated that the chemical characteristics of the cladribine/2-HP- β -CD inclusion complex are not affected by the manufacturing process and that the complex is stable during storage. This tablet formulation is under investigation in an ongoing clinical programme as a once-daily, intermittent oral treatment for patients with MS (CLARITY [cladribine tablets treating multiple sclerosis orally] and ONWARD [oral cladribine added on to Rebif new formulation in subjects with active multiple sclerosis] studies).

ACKNOWLEDGMENTS

The authors would like to thank Imogen Horsey (supported by Merck Serono International S.A., Geneva, Switzerland, an affiliate of Merck KGaA, Darmstadt, Germany) for her assistance in the preparation of this manuscript.

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Best Available Copy

Cladribine

2360

Secols:

B

RNS DOA

'Yellowed Rice is, A. Ajl, Eds. 357-367.

. [α]¹⁸ - 37.4 0, 418). Strong dioxane, dilute i. from lemon onous! LDm in ds). izene, dec 139°. 260, 334 mm ety sol in chlo-

benzene, der uv gnax: 260, ne, chloroform,

ydroxy-2-meth -10-carboxylic . C 57.94% H ced by Penici, Bainier and by Trans. Roy. So Nature 167, 995 oc. 1951, 2013 Money, Nanue i et al., J. Chem

u 155°, dec 290 cely sol in ethat sparingly sol table to acid md

sethyl-6-octent 6%, O 10.37% I in many out dissa: Tiem 54, 236 (1958))51, 505: Est

d 0.848-0.8%

 α -citronellal. [141-26-4] 3,7-Dimethyl-7-octenal; rhodinal. Liquid bp₁, 51°. n_D^{20} 1.4410. (α)²⁰ +9.75°. USE: In soap perfumes; insect repellent.

2354. B-CitronelloL [106-22-9] 3,7-Dimethyl-6-octen-I-ol; 2,6-dimethyl-2-octen-8-ol; citronellol; cephrol. C10H20O; mol wt 156.26. C 76.86%, H 12.90%, O 10.24%. /-Form is a constituent of rose and geranium oils. d-Form occurs in Ceylon and Java citronella oils. History: J. L. Simonsen, L. N. Owen, The Terpenes vol. I (University Press, Cambridge, 2nd ed, 1947). Prepn of (±)-form: Adams, Garvey, J. Am. Chem. Soc. 48, 477 (1926); Ofner et al., Helv. Chim. Acta 42, 2577 (1959). Prepn of (+)-form: Rienäcker, Ohloff, Angew. Chem. 73, 240 (1961); Naves, Tullen, Helv. Chim. Acta 44, 1867 (1961); Eschi-(1901); (1905; (1905); (1905); (1901); Rienticker, Chimia 27, 97 (1973); C. G. Overberger, J. L. Weise, J. Am. Chem. Soc. 90, 1525 (1968); T. Sato et al., Tetrahedron Letters 1980, 3377. Prepn of (-)-form: Ohloff, loc. clt.; Rienäcker, loc. cit.; Shono et al., Tetrahedron Letters 1974, 1295; K. Mori, T. Sugai, Synthesis 1982, 752. Synthesis of (+) or (-)-form from isoprene: Hidai et al., Chem. Commun. 1975, 170. Stereospecific prepn via microbiological (Saccharomyces cerevisice) reduction: P. Gramalica et al., Experientia 38, 775 (1982). Manuf: Woroch et al.; Bain; Webb, US 2990422; US 3005845; US 3028431 (1961, 1961, 1962, all to Glidden); Eschinasi, US 3052730 (1962 to Givaudan). Abs config of the (+)-form: Freudenberg, Hohmann, Ann. 584, 54 (1953); Freudenberg, Lwowski, ibid. 587, 213 (1954). NMR, HPLC determin of R/S enantiomer ratios: D. Valentine et al., J. Org. Chem. 41, 62 (1976). See also Rhodingl.



R-(+)-\$-Citronetto

, (+)-Form. Oily liquid, bp 224.5°, bp₁₀ 108.4°, d_4^{20} 0.8550. n_0^{20} 1.4559. (a) $_0^{20}$ +5.22°. Very slightly sol in water, miscible with with alcohol, ether.

(-)-Form. β-Rhodinol; Levocitrol. bp₁₀ 108-109°. d¹⁸₄ 1.4576. [α]²⁰₂ - 4.76°. (±)-Form. Dihydrogeraniol. di 0.851. no 1.454.

USE: In perfumery,

2355. Citrulline. [372-75-8] N⁵-(Aminocarbonyl)-L-0 biblas: Curtaine, [372-73-8] N°-(Aminocaroonyi)-L-or-biblas: Sureidonorvaline; a-anino-S-ureidovaleric acid; N⁶-carbanylomithine. C₆H₃N₃O₃; mol wt 175.19. C 41.13%, H ⁷⁴⁸%, N 23.99%, O 27.40%. H₂NCONH(CH₂)₃CH(NH₂)-COOH. An amino acid, first isolated from the juice of water-mehan Circulture with the set of the set melon, Citrullus vulgaris Schrad., Cucurbitaceae: Wada Blochem Z. 224, 420 (1930); isoln from casein: Wada, ibld. 257, 1 (1933). Synthesis from ornithine through copper complexes: Kurtz, J. Biol. Chem. 122, 477 (1938); by alkaline hydrolysis of arginine: Fox, *ibid.* **123**, 687 (1936); from cyclo-pentanone oxime: Fox *et al.*, *J. Org. Chem.* **6**, 410 (1941). Crystallization: Matsuda *et al.*, JP 71 174 (1971 to Ajinomoto). Crystalization: Matsuda et al., JP 71 174 (1971 to Ajmoniolo, CA. 74, 126056u (1971). Crystal and molecular structure: Na-ganathan, Venkatesan, Acta Crystallogr. 27B, 1079 (1971); Ashida et al., ibid. 28B, 1367 (1972). Use in asthenia and he-patic insufficiency: FR 2198739 (1974 to Hublot & Vallet). CA. 82, 144952c (1975). Clinical trial in treatment of lysinuric protein intolecones: I. Painnik et al. J. Pediatr. 97, 927 (1980); Protein integrates J. Clinical trial in treatment of systematic Protein integrates J. Rajantie *et al.*, J. Pediatr. **97**, 927 (1980); T.O. Carpenter *et al.*, N. Engl. J. Med. 312, 290 (1985). Prisms from methanol + water, mp 222°. [α]₂²⁰ + 3.7° (c = 2). pK₁ 2.43; pK₂ 9.41. Sol in water. Insol in methanol, etha-nol.

Hydrochloride. [34312-10-2] $C_{g}H_{13}N_{3}O_{3}$.HCl. Crystals, dec 185°. [α] $\frac{1}{D}$ + 17.9° (c = 2).

Malate (salt). [54940-97-5] Stimol. C6H11N3O3.C4H6O3; mol wt 309.27. THERAP CAT: Treatment of asthenia 2356. Citrullol. [1390-93-8] C12H28O4; mol wt 366.53.

C 72.09%, H 10.45%, O 17.46%. From fruit pulp of Citrullus colocynthis Schrad., Cucurbitaceae: Power, Moore, J. Chem. Soc. 97, 99 (1910); Power, Salway, Ibid. 103, 399, 1022 (1913); Khadem, Rahman, Tetrahedron Letters 1962, 1137. Crystals, mp 282-283°. uv max: 242, 272, 282 nm (log e

2.85, 2.68, 2.68). Sol in pyridine; practically insol in usual organic solvents.

Discetate. C26H42O6. Crystals, mp 162°.

2357. Clirus Red 2. [6358-53-8] 1-[(2,5-Dimethoxyphenyl)azo]-2-naphthalenol; C.I. Solvent Red 80; C.I. 12156. C18⁻ H₁₆N₂O₃; mol wt 308.33. C 70.12%, H 5.23%, N 9.09%, O 15.57%. Prepn: H. W. Elley, H. W. Daudt, US 2224904 (1940 to Du Pont). Metabolism: J. L. Radomski, J. Pharmacol. Exp. Ther. 134, 100 (1961); 136, 378 (1962). Toxicology: M. Sharratt et al., Food Cosmet. Taxicol. 4, 493 (1966). Review of carcinogenicity studies: IARC Monographs 8, 101-106. See also Colour Index vol. 4 (3rd ed., 1971) p 4033.



Crystals, mp 155-157°. Slightly sol in water; partially sol in ethanol and vegetable oils.

USE: To color orange skins.

2358. Civet. Zibeth. Unctuous secretion from receptactes between the anus and genitalia of both male and female civet cat. Constit. Civetone and similar compds.

Semi-solid, yellowish to brown unctuous substance; unpleasant, subacrid, bitter taste; fusible and burns without leaving much residue. Insol in water; partly sol in hot sloohol or in ether.

USE: As a fixative in perfumery.

2359. Civetone. [542-46-1] (2)-9-Cycloheptadecen-1one. C17H30O; mol wt 250.42. C 81.54%, H 12.07%, O 6.39%. 17-Membered macrocyclic musk, constituent of civet: Ruzicka, Helv. Chim. Acta 9, 230 (1926); Ruzicka et al., ibid. 10, 695 (1927). Occurs in nature as cis-form. Synthesis of cls-civetone: Stoll et al., ibid. 31, 543 (1948); J. Tsuji, T. Mondai, Tetraha-dron Letters 1977, 3285; E. Secane et al., Chem. & Ind. (Lon-don) 1978, 165. Synthesis of trans-form: H. Hunsdiecker, Ber. 77, 185 (1944); H. H. Mathur, S. C. Bhattacharyya, J. Chem. Soc. 1968, 114. Crystal and molecular structure of cls-civetone: G. Bernardinelli, R. Gerdil, Helv. Chim. Acta 65, 558 (1982).



Crystals, mp 31-32°. Musky odor becoming pleasant in extreme dilns. d_{3}^{13} 0.917. bp_{142} 342°; bp_2 59°. n_D^{23} 1.4830. USE: In perfumery.

2360. Cladribine. [4291-63-8] 2-Chioro-2'-deoxyadenosine; 2-chloro-6-amino-9-(2-deoxy-B-D-erythro-pentofuran-

Consult the Name Index before using this section.

"最高级最高级,我们就是你们的这些,我们还是我的爱望我的不能能,她都是你的人,你是

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2361

Clanobutin

osyl)purine; 2-chlorodeoxyadenosine; 2-CdA; CldAdo; NSC-105014-F; Leustatin. C₁₀H₁₂ClN₂O₃; mol wt 285.69. C 42.04%, H 4.23%, Cl 12.41%, N 24.51%, O 16.80%. Substituted purine nucleoside with antileukemic activity. Prepn as intermediate in synthesis of 2-deoxynucleosides: H. Venner, Ber. 93, 140 (1960); M. Ikchara, H. Tada, J. Am. Chem. Soc. 85, 2344 (1963); eldem, Ibld. 87, 606 (1965). Synthesis and blological activity: L. F. Christensen et al., J. Med. Chem. 15, 735 (1972). Stereospecific synthesis: Z. Kazimierczuk et al., J. Am. Chem. Soc. 106, 6379 (1984); R. K. Robins, G. R. Revankar, EP 173059; eldem, US 4760137 (1986, 1988 both to Brigham Young Univ.). Specific toxicity to lymphocytes: D. A. Carson et al., Proc. Nat. Acad. Sci. USA 77, 6865 (1980); eidem, Blood 62, 737 (1983). Mechanism of action: S. Seto et al., J. Clin. Invest. 75, 377 (1985). Clinical evaluation in chronic lymphocytic leukemia: L. D. Piro et al., Blood 72, 1069 (1988); in hairy cell leukemia: eidem, N. Engl. J. Med. 322, 1117 (1990).

Crystals from water, softens at 210-215°, solidifies and turns brown (Christensen). Also reported as crystals from ethanol, mp 220° (softens), resolidifies, turns brown and does not melt below 300° (Kazimierczuk). $[\alpha]_{12}^{12}$ -18.8° (c = 1 in DMF). uv max in 0.1N NaOH: 265 nm; in 0.1N HCI: 265 nm. THERAP CAT: Antineoplastic.

2361. Clanobutin. (30544-61-7) 4-[(4-Chlorobenzoyl)-(4-methoxyphenyl)mnino]butanoic acid; 4- $[\rho$ -chlorobenzoyl)- γ -(ρ anisidino)butyric acid; Bykahepar. C₁₁M₁₁ClNO₄; mol wt 347.80. C 62.16%, H 5.22%, Cl 10.19%, N 4.03%. O 18.40%. Prepn: K. Klemm *et al.*, DE 1917036 corresp to US 3780095 (1971, 1973 both to Byk-Gulden). Series of articles on synthesis, physical and pharmacological properties: Arzneimittel-Forsch. 29, 1-15 (1979). In vitro biochemical study: H. Wolf *et al.*, Biochem. Pharmacol. 29, 1649 (1980). Effect on bile excretion in mts, dogs: P. Berchtold *et al.*, Arzneimittel-Forsch. 30, 1878 (1980).



Cryst from ethyl acetate, mp 115-116°. pKa 5.04. Soly in water at 37°: 4.02×10^{-2} mol/l at pH 7. LD₅₀ in rats (mg/kg): >2000 orally; 570 i.v. (Klemm).

THERAP CAT: Cholcretic.

THERAP CAT (VET): Choleretic; in treatment of piroplasmosis and anaplasmosis.

2362. Clarithromycin. [81103-11-9] 6-O-Methylerythromycin: A-56268; TE-031; Biaxin; Clathromycin; Cyllind; Klacid; Klaricid; Macladin; Naxy; Veclam; Zeclar. $C_{12}H_{ep}$ -NO₁₃; mol wt 747.95. C 61.02%, H 9.30%, N 1.87%, O 27.81%. Semisynthetic macrolide antibiotic; derivative of erythromycin, q.v. Prepn: Y. Watanabe et al., EP 41355; eidem, US 4331803 (1981, 1982 both to Taisho); and in vitro antibacterial activity: S. Morimoto et al., J. Antibiot. 37, 187 (1984). In vitro and in vivo antibacterial activity: P. B. Fernan-

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Consult the Name Index before using this section.





Colorless needles from chloroform + diisopropyl ether (1.2) mp 217-220° (dec). Also reported as crystals from ethanol, mp 222-225° (Morimoto). uv max (CHCl₃): 288 nm (e 27.9), m max (CHCl₃): 240, 288 nm; (methanol): 211, 288 nm (e_{10}^{10} -90.4° (c = 1 in CHCl₃). Stable at acidic pH. LD₅₀ in mat, female mice, male, female rats (mg/kg): 2740, 2700, 3470, 7700 orally, 1030, 850, 669, 753 i.p., >5000 all s.c. (Abe). THERAP CAT: Antibacterial.

2363. Clathrates. Compounds that are capable of trapping other substances within their own crystal lattices. The carities of the host molecules are classified as cages, tunnels, at layered types, depending on the way they include guest molecules. The geometry of the cavities limits the guest molecules by size and shape, rather than by chemical similarity with the host molecules. Among common clathrates are molecular sieves, cyclotriphosphazenes, and Dlanin's compound, as well as hydroquinone, cyclodextrins, o-thymotide, and deoxychdir acid, q.e.v. Cavitands are organic hosts with enforced (rigid) cavities: D. J. Cram, Science 219, 1177 (1983); R. C. Helgeson et al., Chem. Commun. 1983, 101. Comprehensive boot Clathrate Compounds, V. M. Bhatnagar, Ed. (Chemical Put Co., New York, 1970) 244 pp. Reviews: D. D. MacNicol etal. Chem. Soc. Rev. 7, 65-87 (1978); E. C. Makin, "Clathratian" in Kirk-Othmer Encyclopedia of Chemical Technology Vel 6 (Wiley-Interscience, New York, 3rd ed., 1979) pp 178-189. USE: As complexing agent; stabilizing agent. In analytici separations.

2364. Clavulanic Acid. [58001-44-8] [2R-(2α, 3Z,5α]]
3-(2-Hydroxyethylidene)-7-0x0-4-0x8-1-azabicycle-[3.2.0]heptane-2-carboxylic acid; MM 14151. C₂H₃NO; md wt 199-16. C 48.25%, H 4.55%, N 7.03%, O 40.17%, Bizturnase inhibitor. Antibiotic produced by Streptomyces clavaligerus; first reported naturally occurring fused β-lactam containing oxygen. Isoln: M. Cole et al., DE 2517316 (1975 to Beecham), C.A. 84, 72635t (1976); A. G. Brown et al., J. Astibiot. 29, 668 (1976). Structure, x-ray crystallography: T. T. Howarth et al., Chem. Commun. 1976, 266. Total synthesis of antibacterial spectrum: C. Reading, M. Cole, Antimicrob. As Chemother. 11, 852 (1977). Antibacterial activity, pharmacology and clinical efficacy of combination with anoxicillin: A.P. Bal et al., Lancet 1, 620 (1980); R. N. Brogden et al., Dray 2 37-362 (1981). In vitro and In vivo synergism with ticarcilling R. Sutherland et al., Am. J. Med. 79, Suppl. 5B, 13 (1985).

Combine bodrate. antox: Clay Combin: 116876-3: Methyl (p-Nitrot 117.5-118" THERAP I ubscterial. THERAP antibacteri 2365. ayl)-4-(4,: acetonitril: 2(3H)-yl); H_@Cl₂N₄(N 15.01% eidem, US in pigeons

> mp 190 THERAT 2366. 0XY-N-{} (1-benzyl peridyi)-1 mol wt 3 8.56%. mide, a. A.V.Nc J. Prieto macolog (1980). biatica Pharma 214 (19) tinal eff (1981). Yakuri 208081 radioim 1491 (1 Duarte

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Israpafant

angina and coronary artery disease: C. E. Handler, E. Sowton, *ibid.* 27, 415 (1984); in hypertension: E. B. Nelson *et al.*, *Clin. Pharmacol. Ther.* 40, 694 (1986). Comparison of hemodynamic effects of enantiomers: R. P. Hof *et al.*, *J. Cardiovasc. Pharmacol.* 8, 221 (1986). Series of articles on pharmacology and clinical use: Am. J. Med. 86, 1-146 (1989).



mp 168-170*.

S(+)-Form. PN-205-033. Crystals from ether + hexane, mp 142° (α)^D + 6.7° (α = 1.5 in ethanol).

R(-)-Form. PN-205-034. Crystals from ether + hexane, mp 140°. (a) $\frac{1}{10^{9}}$ -6.7° (c = 1.67 in ethanol). THERAP CAT. Antihypertensive; antianginal.

5263. Israpafant. [117279-73-9] 4-(2-Chlorophenyl)-6,-9-dimethyl-2-[2-[4-(2-methylpropyl)phenyl]ethyl]-6*H*-thieno-[3.2-*f*][1,2,4]uiazolo[4.3-*a*][1,4]diazepine; (\pm)-4-(*o*-chlorophenyl)-2-(*p*-isobutylphenethyl)-6,9-dimethyl-6*H*-thieno[3,2-*f*]-*s*triazolo[4,3-*a*][1,4]diazepine; Y-24180; Pafnol. C₁₈H₂₉ClN₄S; mol wt 489.09. C 68.76%, H 5.98%, Cl 7.25%, N 11.46%, S 6.56%. Platelet activating factor (PAF) antagonist. Prepn: T. Tahara et al., EP 268242; etdem, US 4820703 (1988, 1989 both to Yoshitomi). Pharmacology: M. Terasawa et al., Prostaglandins 40, 553 (1990). Receptor binding study: S. Takchara et al., libid. 571. Clinical evaluation in asthma: S. Hozawa et al., *m. J. Respir. Crit. Care Med.* 152, 1198 (1995).



Colorless crystals from isopropyl ether, mp 129.5-131.5°. Sol in propylene glycol.

THERAP CAT: Antiasthmatic.

5264. Itaconic Acid. [97-65-4] Methylenesuccinic acid; propylenedicarboxylic acid. $C_3H_6O_4$; mol wt 130.10. C 46.16%, H 4.65%, O 49.19%. Obtained by dry distillation of citric acid and subsequent treatment of the anhydride with water. Produced on a large scale by submerged aerobic fermentation using Aspergillus terreus and low cost carbohydrates from beet or cane: Kane et al., US 2385283 (1945 to Pfizer). Synthesis from propargyl chloride, carbon monoxide, nickel carbonyl and water: Chlusoli, US 3025320 (1962 to Montecatini).



Hygroscopic crystals; characteristic odor. d 1.63. mp 162-164 with decompn. Also reported as mp 172° [Kinoshita, Acta Phytochem. (Japan) 5, 273 (1931)]. One gram dissolves in 12 ml water, 5 ml alcohol; very slightly sol in benzene, chloroform, ether, carbon disulfide, petr ether. Keep well closed.

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Consult the Name Index before using this section.

5265. Itasetron. [123258-84-4] 2,3-Dihydro-N-[(3. endo)-8-methyl-8-azabicyclo[3.2.1]oct-3-yl]-2-0xo-1H. benzimidazole-1-carboxamide; 2-0xo-N-1aH,5aH-uropan-3ayl-1-benzimidazoline-1-carboxamide. C₁₆H₂₀N,O₂: mol w 300.35. C 63.98%, H 6.71%, N 18.65%, O 10.65%. Serotonin (5-HT₃) receptor antagonist. Prepn: M. Turconi et al., EP 309423 (1989 to Istinuto De Angeli); eidem. US 5223511 (1993) to Boehringer, Ing.); M. Turconi et al., J. Med. Chem. 33, 2101 (1990). Pharmacology: idem et al., Eur. J. Pharmacol. 203, 203 (1991). Mode of action: M. B. Passani et al., Brit. J. Pharmacol. 112, 695 (1994). Clinical efficacy and tolerability: H. Goldschmidt et al., Anti-Cancer Drugs 8, 436 (1997). Review of therapeutic potential: M. B. Passani, R. Corradetti, CNS Drug Reviews 2, 195-213 (1996).



Crystals from acetonitrile, mp 205-207°. LD₅₀ in mice, rats (mg/kg): 56, 62 i.v. (Passani). Hydrochloride. [127618-28-4] DAU 6215. C₁₆H₂₀N₄.

Hydrochloride. [127618-28-4] DAU 6215. $C_{16}H_{20}N_4$. O_{2.}HCl; moi wt 336.82. Coloriess crystals, mp 270°.

THERAP CAT: Antiemetic.

5266. Itraconazole. [84625-61-6] 4-[4-[4-[4-[2-(2,4-Dichlorophenyl)-2-(1H-1,2,4-triazol-1-ylmethyl)-1,3-dioxolan.4yl]methoxy]phenyl]-1-piperazinyl]phenyl]-2,4-dibydro-2-[1methylproyl]-3H-1,2,4-triazol-3-one; (\pm) -1-sec-butyl-4[p-[4-[p-[[(2R*,4S*)-2-(2,4-dichlorophenyl)-2-(1H-1,2,4-triazol-1-ylmethyl)-1,3-dioxolan.4-yl]methoxy]phenyl]-1-piperazinyl]phenyl]- Δ^2 -1,2,4-triazolin-5-one; oriconazole: R-51211; ltrizole; Sporanox; Triasporin. C₁₃H₁₄Cl₂N₆O₄; mel wt 705.65. C 59.57%, H 5.43%, Cl 10.05%, N 15.88%, O 9.07%. Orally active antimycotic structurally related to ketoconazole; q.v Prepri: J. Heeres, L. J. Backx, EP 6711; eidem, US 4267179 (1980, 1981 both to Janssen); J. Heeres et al., J. Med Chem. 27, 894 (1984). In vliro activity: A. Espinel-Ingroff et al., Antimicrob. Ag. Chemother. 26, 5 (1984). HFLC determo in biological sumples: R. Woestenborghs et al., J. Chromatog. 413, 332 (1987). Symposium on pharmacology and clinical efficacy: Rev. Infect. Dis. 9, Suppl 1, S1-S152 (1987). Toxicity data: H. Van Cautern et al., ibid. S43. Review of clinical pharmacokinetics: J. Heykants et al., Mycoses 32, Suppl 1, 67-87 (1989); of clinical efficacy in dermatophytosis: P. De Doncker, G. Cauwenbergh, Brit. J. Clin. Pract. Suppl. 71, 118-122 (1990). Review: A. M. Sugar, Curr. Clin. Topics Inf. Diu-13, 74-98 (1993).



Crystals from toluene, mp 166.2°, pKa 3.7. Lipophilic: $p_{\rm H}^{\rm ar}$ titlon coefficient (*n*-octanol/aq buffer of pH 8.1): 5.66. Practically insol in water and dil acidic solns. LD₃₀ (14 day) in mice rats, dog's (mg/kg): >320, >320, >200 orally (Van Cautern). THERAP CAT: Antifungal.

5267. Itramin Tosylate. [13445-63-1] 2-Aminocihano nitrate mono(4-methylbenzenesulfonate); 2-aminocihano m trate mono-p-toluenesulfonate; 2-nitratoethylaminotohueno sulfonate; Cardisan; Tostram; Nilatii. $C_9H_{14}N_2O_5$: mol 4° 278.28. C 38.85%, H 5.07%, N 10.07%, O 34.50%, S 11.52% Prepn: SE 168308 (1959 to Aktiebolaget Pharmacia), CA.54, 24405d (1960). Too

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Look for Chemical	S	Search Help
urrent position: <u>Home</u> > Prod	u <u>cts</u> > Hydroxypropyl-beta-cyclodextrin	nna an a'
	Hydroxypropyl-beta CAS No:9403	a-cyclodextrin 35-02-6
Name:	Hydroxypropyl-beta-cyclodextrin	
Synonyms:	beta-Hydroxypropylcyclodextrin	
	beta-Cyclodextrin, 2-hydroxypropyl ether	
	HPB 2-Hydroxypropyd-beta-cyclodeytrin	
	128446-35-5	
CAS Number:	94035-02-6	
Molecular Formula:	C ₄₂ (H) _{70-n} O ₃₅ (C ₃ H ₇) _n	
Melting Point:	278 °C	
Safety Description:	S24/25 <u>Details</u>	
], inquire now List of	Suppliers for Hydroxypropyl-beta-cyclodextrin	Countr
Onbio Inc. Introduction:HYDROXYPF	OPYL-BETA-CYCLODEXTRIN	氏 册 United S
Yiming Fine Chemicals Introduction:mp : 267 °C (<u>.Co., Ltd.</u> ec.)	China (Ma
storage temp.: 2-8°C		
solubility : H2O, 45 % (w/v		
form · solution (clear, color	less)	
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Name	Description	Size	Catalog #	Supplier	
СКВВ	Recombinant Human Creatine Kinase BB Isoenzyme	10µg, 50µg, 1mg	СКІ- 268	PROSPEC-TANY TECHNOGENE LTD.	More In
<u>Ckdk6</u>	The RP-39008 Ckdk6 protein is a partial length (aa 1-327) bacterially expressed recombinant protein.RP-39008 is suitable for use as a control in ELISA and Western blot applications.The RP- 39008 protein is GST- tagged.	10 ug	RP-39008	ABR - AFFINITY BIOREAGENTS INC.	More In
СКММ	Human Creatine Kinase MM	200µg, 1mg, 10mg	СКІ- 273	PROSPEC-TANY TECHNOGENE LTD.	More In
<u>CKS-17</u>	Sequence: Leu-Gin-Asn- Arg-Arg-Gly-L eu-Asp- Leu-Leu-Phe-Leu-Ly s- Glu-Gly-Gly-LeuStorage and Stability: Lyophilized powder may be stored at 4?C for short-term only. Reconstitute to nominal volume by adding sterile 40-50% glycerol and store at -20?C. R	1mg	C5818-05	UNITED STATES BIOLOGICAL	More In
CV(5-17 /7 17)	Sequence: Leu-Asp-Leu- Leu-Phe-LeuSt orage and Stability: Lyophilized powder may be stored at 4?C for short-term only. Reconstitute to nominal volume by adding sterile 40-S0% glycerol and store at -20?C. Reconstituted product is stable for 12 months	25mg			More Te

<u>CKS-17</u>	This Peptide CKS-17 is considered as the major immunosuppressive site of retroviral TM protein. It can suppress T effector cell function in vitro. It has been shown that CKS-17 causes an imbalance of human types 1 and 2 cytokines and Inhibition of	0.5 mg	06-271- 83162	GENWAY BIOTECH INC.	More In
<u>CKS-17</u>	This Peptide CKS-17 is considered as the major immunosuppressive site of retroviral TM protein. It can suppress T effector cell function in vitro. It has been shown that CKS-17 causes an imbalance of human types 1 and 2 cytokines and inhibition of	1 mg	06-271- 83162	GENWAY BIOTECH INC.	More In
CARCINOEMBRYONIC ANTIGEN (CL)	testing/assay service	n/a	n/a	RDL REFERENCE LABORATORY INC.	More In
CASPASE-3/7 (CL)	n/a	n/a	n/a	PROMEGA CORPORATION	More In
CASPASE-8 (CL)	n/a	n/a	n/a	PROMEGA CORPORATION	More In
CASPASE-9 (CL)	n/a	n/a	n/a	PROMEGA CORPORATION	More In
CASPASE (CL)	n/a	n/a	n/a	PROMEGA CORPORATION	More In
<u>CL 218872</u>	Benzodiazepine agonist displaying selectivity for a1 subunit-containing GABAA receptors (Ki values are 130, 1820, 1530, > 10000, 490 and > 10000 nM for a1, a2, a3, a4, a5 and a6- subunit containing re	10mg, 50mg	1709	TOCRIS BIOSCIENCE	More In
<u>CL-387,785</u>	Irreversibly inhibits EGF- receptor (EGFR) kinase activity in vivo (IC50 = 250-490 pM) as well as EGF-stimulated autophosphorylation of tyrosine residues in the EGFR in vivo (IC50 = 5 nM). Blocks EGF- mediated growth in A431 cells. Inhibits prolifer	n/a	233100	CALBIOCHEM/EMD BIOSCIENCES	More In
	A selective inhibitor of MMP-13 (IC50 = 10 μ M).				

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<u>CL-82198</u>	Binds to the S1' pocket of MMP-13 with its morpholine ring adjacent to the catalytic zinc atom. Does not inhibit MMP-1, MMP-9, and TACE.	n/a	233105	CALBIOCHEM/EMD BIOSCIENCES	More In
Calphostin C, Cladosporium Cladosporioides	A cell permeable, highly specific inhibitor of protein kinase C (IC50 = 50 nM) that interacts with the protein's regulatory domain by competing at the binding site of diacylglycerol and phorbol esters. Does not compete with Ca2+ or phospholi	n/a	208725	CALBIOCHEM/EMD BIOSCIENCES	More In
<u>Cladribine</u>	It is a substituted purine nucleoside with antileukemic activity.Melting Point: 220-2357C dec.Solubility: Methanol, Water	50mg	C5819-75	UNITED STATES BIOLOGICAL	<u>More In</u>
Clarithromycin	A semi-synthetic macrolide antibiotic. A derivative of erythromycin.Melting Point: 217-220?C dec.Solubility: Chloroform, Ethanol	50mg	C5829	UNITED STATES BIOLOGICAL	More In
Clavulanic Acid	A B-Lactamase inhibitor.	10mg	C5836	UNITED STATES BIOLOGICAL	More In
CLIC3	The RP-39009 CLIC3 protein is a full length bacterially expressed recombinant protein.RP- 39009 is suitable for use as a control in ELISA and Western blot applications.The RP- 39009 amino acid sequence corresponds to the NCBI accession number NP_004	10 ug	RP-39009	ABR - AFFINITY BIOREAGENTS INC.	More In
<u> Clidinium Bromide</u>	An anticholinergic. Used as an antispasmodic.Melting Point: 240-241?C	59	C5840-75	UNITED STATES BIOLOGICAL	More In
	A metal ion chelator that crosses the blood brain barrier and acts as a neurotoxic antibiotic. Reported to dissolve				

		<u> </u>			
Clofulbicyne	n/a	1 mg.	TXL9001-1	ACCURATE CHEMICAL & SCIENTIFIC CO.	More In
Clofibrate	An anti- hyperlipoproteinemic agent believed to act by inhibiting cholesterol biosynthesis. Activates PPARa and induces cytochrome P450 4A1 and 4A3. Imparts protection against acetaminophen toxicity and increases hepatic glutathione levels.	n/a	231405	CALBIOCHEM/EMD BIOSCIENCES	More In
<u>Clofibric acid</u>	PPAR agonist. Antihyperlipoproteinemic.	1g	0825	TOCRIS BIOSCIENCE	More_In
Clofibrate	PPAR agonist (EC50 values are 50, 500 and > 100 μM at PPAR α, PPAR γ and PPAR δ respectively). Antihyperlipoproteinemic.	1g	0824	TOCRIS BIOSCIENCE	More In
<u>CLOFIBRATE</u>	n/a	n/a	n/a	CAYMAN CHEMICAL CO.	More In
Clofarabine	Deoxycytidine klnase (dCK) substrate. Phosphorylated to form clofarabine triphosphate, which competes with dATP for DNA polymerase- α and - ε and potently inhibits ribonucleotide reductase (IC50 = 65 nM). Induces apoptosis by directl	10mg, 50mg	2600	TOCRIS BIOSCIENCE	More In
<u>Clofarabine</u>	ISecond generation purine nucleoside analog; antimetabolite that inhibits DNA synthesis and resists deamination by adenosine deaminase. Antineoplastic.Melting Point: 225-227?C	10mg	C5843-55	UNITED STATES BIOLOGICAL	More_In
CLK2, active	n/a	10 ug	14-774	MILLIPORE	Моге Іп
	senile plaques and reduce amyloid's ability to clump together, apparently by trapping the Cu2+ and Zn2+ that stud these depos	n/a	233165	CALBIOCHEM/EMD BIOSCIENCES	More In
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-		5x1		ACCURATE CHEMICAL &	
Clofulbicyne	n/a	mg.	TXL9001-5	SCIENTIFIC CO.	More In
<u>Clomifene citrate</u>	International Chemical Reference Substances are established upon the advice of the WHO Expert Committee on Specifications for Pharmaceutical Preparations. They are supplied primarily for use In physical and chemical tests and assays described In t	100 mg	9930259	W.H.O. COLLABORATING CENTRE	More In
<u>Clomiphene, Citrate</u>	An unducer of ovulation. A gonad-stimulating principle.Melting Point: 116.5-118?CSolubility: Methanol	10g	C5843-65	UNITED STATES BIOLOGICAL	More In
Cloning	>1500 bp into 3 different expression vectors	n/a	PE05-0003	HYPEROMICS FARMA INC.	More In
Cloning	<1500 bp into 3 different expression vectors	n/a	PE05-0002	HYPEROMICS FARMA INC.	More In
<u>Clopidogrel</u> Carboxylic Acid	A metabolite of the drug Clopidogrel.Solubility: Methanol, Water	5mg	C5849-01	UNITED STATES BIOLOGICAL	More In
<u>CLOSTRIPAIN</u> <u>Clostridium</u>	n/a	n/a	n/a	PROMEGA CORPORATION	More In
<u>Clotrimazole</u>	An antifungal agent that acts as a potent and specific inhibitor of the Ca2+-activated K+ channel (Gardos channel; IC50 = 650 nM). Prevents K+ loss and dehydration of sickled erythrocytes.	n/a	233230	CALBIOCHEM/EMD BIOSCIENCES	More In
<u>Clozapine</u>	An antipsychotic.Melting Point: 183-184? CSolubility: Acetone, Ether	250mg	C5866	UNITED STATES BIOLOGICAL	More In
Clozapine	Atypical antipsychotic drug, with a much lower tendency to cause extrapyramidal side effects than conventional neuroleptics. Displays a broad range of pharmacological actions; the antipsychotic effects are thought to be mediated principally by 5-H	50mg, 500mg	0444	TOCRIS BIOSCIENCE	More In

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CLTB	The RP-39010 CLTB protein is a full length bacterially expressed recombinant protein.RP- 39010 is suitable for use as a control in ELISA and Western blot applications.The RP- 39010 amino acid sequence corresponds to the NCBI accession number NP_0018	10 ug	RP-39010	ABR - AFFINITY BIOREAGENTS INC.	<u>More In</u>
<u>Aldosterone-3 CMO (BSA)</u>	The major mineralcorticoid, which is secreted almost independently of ACTH from the pitutitary, is aldosterone. Aldosterone secretion is controlled mostly by the levels of potassium and sodium in serum and a blood pressure control system called th	5mg	A1350-04	UNITED STATES BIOLOGICAL	More In
Androstenedione-3 (CMO)	Androstenedione was discovered in 1935. It is naturally produced in men and women. It is a direct precursor to the hormone testosterone. The liver converts androstenedione to testosterone.Precursor:4- Androsten-3,17-dione- 3Sto rage and Stability:Lyo	10mg	A2292-02	UNITED STATES BIOLOGICAL	More In
CMPD-1	Non-ATP-competitive, selective inhibitor of p38 a-mediated MK2a (mitogen-activated protein kinase-activated protein kinase 2a) phosphorylation (apparent Ki = 330 nM). Does not inhibit p38 a- mediated phosphorylation of the two other kno	10mg, 50mg	2186	TOCRIS BIOSCIENCE	More In
CMV	Glycine Extract	mL	0810003GE	ZEPTOMETRIX CORP.	More In
CMV	Cytomegalovirus (AD169) Infected Cell Extract. Used for IgG assays - Control is NHDF AV043	n/a	CV001	EASTCOAST BIO INC.	More In
	Cytomegalovirus				

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CMV	Gradient Purified. Used for IgM assays.	n/a	CV046	EASTCOAST BIO INC.	More In
CMV	Cytomegalovirus Ag slides for FA. Made to Order	n/a	CG015	EASTCOAST BIO INC.	More In
<u>CMV</u>	Part Pure	n/a	J43010	BIOSPACIFIC INC.	More In

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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
12/986,310	01/07/2011	Nicholas S. Bodor	20009904-0067	6100
13974 7590 09/11/2013 DENTONS US LLP			EXAMINER	
P.O. BOX 0610)80		LAU, JONATHAN S ART UNIT PAPER NUMBER	
Chicago, IL 60	506-1080			
			1623	
			NOTIFICATION DATE	DELIVERY MODE
			09/11/2013	ELECTRONIC

Please find below and/or attached an Office communication concerning this application or proceeding.

The time period for reply, if any, is set in the attached communication.

Notice of the Office communication was sent electronically on above-indicated "Notification Date" to the following e-mail address(es):

martin.bruehs@dentons.com patents.us@dentons.com

	Application No. 12/986,310	Applicant(s) BODOR ET AL.			
Office Action Summary	Examiner Jonathan S. Lau	Art Unit 1623	AIA (First Inventor to File) Status		
The MAILING DATE of this communication ap	bears on the cover sheet with t	he corresponde	nce address		
 Period for Reply A SHORTENED STATUTORY PERIOD FOR REPL WHICHEVER IS LONGER, FROM THE MAILING D Extensions of time may be available under the provisions of 37 CFR 1.1 after SIX (6) MONTHS from the mailing date of this communication. If NO period for reply is specified above, the maximum statutory period Failure to reply within the set or extended period for reply will, by statute Any reply received by the Office later than three months after the mailin earned patent term adjustment. See 37 CFR 1.704(b). 	Y IS SET TO EXPIRE <u>3 MON</u> ATE OF THIS COMMUNICAT 36(a). In no event, however, may a reply will apply and will expire SIX (6) MONTHS a, cause the application to become ABANE g date of this communication, even if timel	TH(S) OR THIF FION. be timely filed From the mailing date OONED (35 U.S.C. § - y filed, may reduce an	RTY (30) DAYS, of this communication. 133). y		
Status					
 1) Responsive to communication(s) filed on <u>28 M</u> A declaration(s)/affidavit(s) under 37 CFR 1. 	<u>1ay 2013</u> . 1 30(b) was/were filed on	<u>.</u>			
2a) This action is FINAL . 2b) This	s action is non-final.				
3) An election was made by the applicant in resp	onse to a restriction requirem	ent set forth dui	ring the interview on		
 ; the restriction requirement and election have been incorporated into this action. 4) ☐ Since this application is in condition for allowance except for formal matters, prosecution as to the merits is closed in accordance with the practice under <i>Ex parte Quayle</i>, 1935 C.D. 11, 453 O.G. 213. 					
Disposition of Claims					
 5) ∑ Claim(s) <u>1-61 and 65-88</u> is/are pending in the application. 5a) Of the above claim(s) <u>9-24,32-45 and 65-88</u> is/are withdrawn from consideration. 6) □ Claim(s) is/are allowed. 7) ∑ Claim(s) <u>1-8,25-31 and 46-61</u> is/are rejected. 8) □ Claim(s) is/are objected to. 9) □ Claim(s) are subject to restriction and/or election requirement. * If any claims have been determined <u>allowable</u>, you may be eligible to benefit from the Patent Prosecution Highway program at a participating intellectual property office for the corresponding application. For more information, please see http://www.uspto.gov/patents/init_events/pph/index.jsp or send an inquiry to <u>PPHfeedback@uspto.gov</u>. Application Papers 10) □ The specification is objected to by the Examiner. 11) □ The drawing(s) filed on is/are: a) □ accepted or b) □ objected to by the Examiner. 					
Applicant may not request that any objection to the drawing(s) be held in abeyance. See 37 CFR 1.85(a). Replacement drawing sheet(s) including the correction is required if the drawing(s) is objected to. See 37 CFR 1.121(d).					
Priority under 35 U.S.C. § 119 12) ☐ Acknowledgment is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f). Certified copies: a) ☐ All b) ☐ Some * c) ☐ None of the: 1. ☐ Certified copies of the priority documents have been received. 2. ☐ Certified copies of the priority documents have been received in Application No 3. ☐ Copies of the certified copies of the priority documents have been received in this National Stage application from the International Bureau (PCT Rule 17.2(a)). * See the attached detailed Office action for a list of the certified copies not received.					
Attachment(s) 1)	3) Interview Sum Paper No(s)/M 4) Other:	mary (PTO-413) ail Date			

DETAILED ACTION

The present application is being examined under the pre-AIA first to invent provisions.

This Office Action is responsive to Applicant's Amendment and Remarks, filed 28 May 2013, in which claims 62-64 are canceled and new claims 65-88 are added.

This application is a domestic application, filed 7 Jan 2011; and claims benefit as a CON of 10/551,205, issued as Patent 7,888,328, which is a 371 of PCT/US2004/009387, filed 26 Mar 2004, which claims benefit of provisional application 60/458,922, filed 28 Mar 2003, and claims benefit of provisional application 60/484,756, filed 2 Jul 2003, and claims benefit of provisional application 60/541,247, filed 4 Feb 2004. The filing date of the instant claims is deemed to be provisional application 60/541,247, filed 4 Feb 2004.

Claims 1-61 and 65-88 are pending in the current application. Claims 9-24, 32-45 and new claims 65-88, drawn to non-elected inventions, are withdrawn. Claims 1-8, 25-31 and 46-61 are examined on the merits herein.

Election/Restrictions

Newly submitted claims 65-88 are directed to an invention that is independent or distinct from the invention originally claimed for the following reasons: New claims 65-88

Application/Control Number: 12/986,310 Art Unit: 1623

are encompassed within the invention of Group II detailed in the Restriction Requirement mailed 8 Nov 2012, drawn to a method of treatment of symptoms of a cladribine-responsive condition in a subject suffering from said symptoms comprising orally administering to a subject in need thereof a pharmaceutical composition comprising said cladribine-cyclodextrin complex.

New Claims 65-88 are withdrawn from further consideration pursuant to 37 CFR 1.142(b), as being drawn to a nonelected invention, there being no allowable generic or linking claim. Applicant timely traversed the restriction requirement in the reply filed on 8 Jan 2013.

Regarding Applicant's remarks filed 28 May 2013, if the method of use of claims 65-88 require all limitations of an allowable product then they will be rejoined for examination on the merits. New claims 65-88 are encompassed within the invention of Group II detailed in the Restriction Requirement mailed 8 Nov 2012, and will be considered for rejoinder upon finding allowable claims.

Rejections Withdrawn

Applicant's Amendment, filed 28 May 2013, with respect that claims 1-8, 25-31 and 46-61 are rejected under 35 U.S.C. 103(a) as being unpatentable over Schultz et al. (US Patent 6,194,395, published 27 Feb 2001, provided by Applicant in IDS mailed 7 Jan 2011) in view of Baert et al. (WIPO Publication WO97/18839, published 29 May 1997, of record) has been fully considered and is persuasive, as Applicant provides

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evidence in Van Axel Castelli comparing the physical mixture of cladribine-CD prepared by physical mixture and kneading (Van Axel Castelli, page 3898, left column, paragraph 3) with the complex of the instant invention. In view of newly discovered reference to Redenti et al. (Int. J. Pharm., 1996, 129, p289-294, cited in PTO-892), Applicant's remarks are persuasive that the mixture taught by Schultz in view of Baert can reasonably be concluded to be the physical mixture, and the comparative evidence provided by Van Axel Castelli is applicable and persuasive.

This rejection has been **withdrawn**. However, upon further consideration, a new ground(s) of rejection is made in view of Vandercruys (US Patent Application Publication 2002/0150616, published 17 Oct 2002, filed 27 May 1998, cited in PTO-892).

Claim Rejections - 35 USC § 103

The following is a quotation of pre-AIA 35 U.S.C. 103(a) which forms the basis for all obviousness rejections set forth in this Office action:

(a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.

This application currently names joint inventors. In considering patentability of the claims under pre-AIA 35 U.S.C. 103(a), the examiner presumes that the subject matter of the various claims was commonly owned at the time any inventions covered therein were made absent any evidence to the contrary. Applicant is advised of the obligation

under 37 CFR 1.56 to point out the inventor and invention dates of each claim that was not commonly owned at the time a later invention was made in order for the examiner to consider the applicability of pre-AIA 35 U.S.C. 103(c) and potential pre-AIA 35 U.S.C. 102(e), (f) or (g) prior art under pre-AIA 35 U.S.C. 103(a).

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Claims 1-8, 25-31 and 46-61 are rejected under pre-AIA 35 U.S.C. 103(a) as being unpatentable over Vandercruys (US Patent Application Publication 2002/0150616, published 17 Oct 2002, filed 27 May 1998, cited in PTO-892) in view of Schultz et al. (US Patent 6,194,395, published 27 Feb 2001, provided by Applicant in IDS mailed 7 Jan 2011), Baert et al. (WIPO Publication WO97/18839, published 29 May 1997, of record), Redenti et al. (Int. J. Pharm., 1996, 129, p289-294, cited in PTO-892), and Pitha et al. (Life Sci., 1998, 43, p493-502, cited in PTO-892).

Vandercruys teaches pharmaceutical composition comprising a cyclodextrin and a sparingly water-soluble drug (abstract). Vandercruys teaches the components are preferably in the glass thermoplastic phase without a crystalline or microcrystalline phase (paragraph 13 spanning pages 1-2), or an amorphous phase. Vandercruys teaches the cyclodextrin can be 2-hydroxypropyl-gamma-CD, and in particular 2hydroxypropyl-beta-CD (page 2, paragraph 20-21). Vandercruys teaches the cyclodextrin includes sulfobutylcyclodextrins (page 2, paragraph 24). Vandercruys teaches varying the molar ratio of cyclodextrin to drug in the range of 100:1 to 5:1, especially 50:1 to 1:2, more especially 10:1 to 1:1 (page 3, paragraph 27). Vandercruys teaches an extensive list of possible drugs which includes cladribine (paragraph 70 spanning pages 4-5). Vandercruys teaches the composition to be an intimate admixture

prepared by dissolving the mixture and removing the solvent (page 6, paragraph 77). Vandercruys implies the mixture results in complexing of the drug in the CD (page 7, paragraphs 87-88). Vandercruys teaches the complex prepared by a vacuum-distillation process (page 8, paragraph 104-105). Vandercruys teaches the pharmaceutical composition in a dosage form for oral administration (page 7, paragraph 95).

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Vandercruys does not specifically disclose the complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex and the composition comprising no significant amount of free crystalline cladribine therein (instant claim 1). Vandercruys does not specifically disclose the composition corresponding to a point located on the curve of a phase solubility diagram for saturated complex cladribine-cyclodextrin complexes, said curve defining complex saturated complexes of cladribine in varying concentrations of the cyclodextrin (instant claim 7).

Vandercruys does not specifically disclose the complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex (instant claim 25). Vandercruys does not specifically disclose the composition comprising a cladribine to cyclodextrin ratio from about 1:10 to about 1:16 (instant claims 5 and 29). Vandercruys does not specifically disclose the complex wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by

weight of the cladribine is in the non-inclusion complex (b) (instant claims 8 and 31). Vandercruys does not specifically disclose the product-by-process wherein 12.00 parts by weight of cladribine and 172.50 parts by weight of hydroxypropyl- β -cyclodextrin are introduced in step (i) of the process, implying a cladribine to cyclodextrin ratio of 1:14.38, or the product-by-process wherein 16.35 parts by weight of cladribine and 172.50 parts by weight of step (i) of the hydroxypropyl- β -cyclodextrin are introduced in step (i), implying a cladribine to cyclodextrin ratio of 1:14.38, or the product-by-process wherein 16.35 parts by weight of cladribine and 172.50 parts by weight of the hydroxypropyl- β -cyclodextrin are introduced in step (i) of the process (instant claim 55), implying a cladribine to cyclodextrin ratio of 1:10.55.

Schultz et al. discloses a solid pharmaceutical oral dosage form of cladribine comprising cladribine and cyclodextrin (column 2, lines 31-39). The disclosed product is substantially identical to the product-by-process. Schultz et al. discloses the use of β and γ -cyclodextrins (column 2, lines 56-58) and derivatives wherein one or more cyclodextrin hydroxy groups are replaced with groups such as methyl, hydroxypropyl, carboxymethyl (column 3, lines 26-27) or sulfobutylcyclodextrins (column 4, lines 22-24). The phrase "one or more cyclodextrin hydroxy groups" combined with the absence of specific structural details of which hydroxyl group is substituted with a methyl group meets the limitation of "randomly methylated β -cyclodextrins". Schultz et al. discloses the solid oral dosage form in the form of a tablet (column 5, lines 37-38) including the excipients sorbitol and magnesium stearate (column 6, lines 2-7), disclosing a product that is substantially identical to a product-by-process meeting the limitations of the instant claims invention. Schultz et al. discloses an oral dosage form comprising up to 15 mg cladribine and cyclodextrin from 100 to 500 mg, which would lead one of skill in the art to instantly envision a cladribine to cyclodextrin ratio ranging from 15 mg:100 mg

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to 15mg:500 mg, or 1:6.67 to 1:33.3 by weight (column 6, lines 23-31). The instant specification suggests that maximum amount of cladribine which can be complexed gives a weight ratio of 1:10 for the cladribine:cyclodextrin complex.

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Therefore a composition comprising the cladribine:cyclodextrin complex that contains a cladribine to cyclodextrin ratio of 1:6.67 describes a composition that comprises a "saturated" complex. Schultz et al. incorporates-by-reference the method of making said solid oral dosage form (Schultz et al.column 5, lines 50-52) disclosed in WIPO Publication WO97/18839, Baert et al., which provides evidence in the embodiment wherein the melt-extruded forms consist essentially of amorphous material (Baert et al. page 8, lines 14-15). Therefore Schultz et al. suggests an amorphous cladribine-cyclodextrin complex in a solid oral dosage form. Schultz et al. implicitly discloses an oral dosage form comprising up to 15 mg cladribine and cyclodextrin from 100 to 500 mg, or a cladribine to cyclodextrin ratio ranging from 1:6.67 to 1:33.3 by weight (column 6, lines 23-31).

Baert et al. discloses a solid mixture comprising one or more cyclodextrins and an insoluble active ingredient embedded into the cyclodextrin carrier (abstract), and teaches ratios of active ingredient to cyclodextrin of from about 1:100 to 100:1, from about 1:5 and 5:1 and from about 1:3 to 3:1 (page 11, lines 1-5). These ratios are interpreted as mole ratios because Baert et al. teaches the use of different active ingredients with different molecular weights. A mole ratio of active ingredient to cyclodextrin of about 1:3 for cladribine (MW 285.7 g/mol) and β -cyclodextrin (MW 1135 g/mol) gives a ratio by weight of approximately 1:11.9. The ratio of 1:11.9 meets the

limitation of both a ratio of about 1:11 and a ratio of about 1:14 according to the nonlimiting definition of "about" as a variance of 20% provided in the instant specification page 9, lines 6-11. Such a saturated complex would consist of only (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex, and being a saturated complex corresponds to a point located on the curve of a phase solubility diagram for saturated complex cladribine-cyclodextrin complexes, said curve defining complex saturated complexes of cladribine in varying concentrations of the cyclodextrin.

Redenti et al. teaches the one of ordinary skill in the art recognizes the differences between a piroxicam:CD complex prepared by melt-spinning and freezedrying amorphous piroxicam:CD complex to show that the freeze-dried complex is a true inclusion complex and not a dispersed mixture of two amorphous components (abstract). Redenti et al. suggests the piroxicam:CD complex prepared by melt-spinning is a physical mixture of amorphous piroxicam and amorphous CD, which is subject to recrystallization upon standing (page 290, paragraph spanning left and right column). Redenti et al. teaches the freeze-dried complex does not recrystallize upon standing (page 291, left column, paragraph 2).

Pitha et al. teaches the level of skill in the art regarding the formulation of 2hydroxypropyl CD to form amorphous mixtures through formation of inclusion complexes (page 493, abstract).

invention to combine Vandercruys in view of Schultz et al., Baert et al., Redenti et al., and Pitha et al. in order to select the pharmaceutical composition comprising a complex of cyclodextrin and a sparingly water-soluble drug taught by Vandercruys to be a solid pharmaceutical oral dosage form of cladribine comprising cladribine and cyclodextrin and to optimize the ratio of cyclodextrin to the drug. One of ordinary skill in the art would have been motivated to combine Vandercruys in view of Schultz et al., Baert et al., Redenti et al., and Pitha et al. to select a complex of cyclodextrin and cladribine with a reasonable expectation of success because Vandercruys teaches the list of sparingly water-soluble drug includes cladribine and Schultz et al. and Baert et al. provide guidance for selecting the pharmaceutical composition comprising a cyclodextrin and cladribine. One of ordinary skill in the art would have been motivated to make an amorphous complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex and the composition comprising no significant amount of free crystalline cladribine therein because Vandercruys teaches the components are preferably in the glass thermoplastic phase without a crystalline or microcrystalline phase and teaches preparation by dissolving the mixture and removing the solvent by a vacuum-distillation process and implies the mixture results in complexing of the drug in the CD, Redenti et al. suggests a piroxicam:CD complex prepared by melt-spinning is a physical mixture of amorphous piroxicam and amorphous CD, which is subject to recrystallization upon standing while

a freeze-dried complex does not recrystallize upon standing, and Pitha et al. teaches it is known in the art that 2-hydroxypropyl CD forms amorphous mixtures through formation of inclusion complexes, suggesting of the dissolved and vacuum-distilled product of Vandercruys be a complex that does not recrystallize upon standing with a reasonable expectation of success.

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It would have been routine experimentation to optimize the ratio of cladribine to cyclodextrin within prior art conditions because Vandercruys teaches varying the molar ratio of cyclodextrin to drug in the range of 100:1 to 5:1, especially 50:1 to 1:2, more especially 10:1 to 1:1 and Schultz et al. discloses an oral dosage form comprising up to 15 mg cladribine and cyclodextrin from 100 to 500 mg, which would lead one of skill in the art to instantly envision a cladribine to cyclodextrin ratio ranging from 15 mg:100 mg to 15mg:500 mg, or 1:6.67 to 1:33.3 by weight. See also MPEP 2144.05 II.A, providing "[W]here the general conditions of a claim are disclosed in the prior art, it is not inventive to discover the optimum or workable ranges by routine experimentation."

Claims 46-61 are drawn to a product-by-process. The disclosed product is substantially identical to the instantly claimed product-by-process, an amorphous solid pharmaceutical oral dosage form comprising cladribine and cyclodextrin. "[E]ven though product-by-process claims are limited by and defined by the process, determination of patentability is based on the product itself. The patentability of a product does not depend on its method of production. If the product in the product-by-process claim is the same as or obvious from a product of the prior art, the claim is unpatentable even though the prior product was made by a different process." In re

Thorpe, 777 F.2d 695, 698, 227 USPQ 964, 966 (Fed. Cir. 1985) (citations omitted) (Claim was directed to a novolac color developer. The process of making the developer was allowed.

The difference between the inventive process and the prior art was the addition of metal oxide and carboxylic acid as separate ingredients instead of adding the more expensive pre-reacted metal carboxylate. The product-by-process claim was rejected because the end product, in both the prior art and the allowed process, ends up containing metal carboxylate. The fact that the metal carboxylate is not directly added, but is instead produced in-situ does not change the end product.). See MPEP 2113. In the instant case, Vandercruys teaches the components are preferably in the glass thermoplastic phase without a crystalline or microcrystalline phase (paragraph 13 spanning pages 1-2) and Vandercruys implies the mixture results in complexing of the drug in the CD (page 7, paragraphs 87-88). Therefore the product in the product-byprocess claim is obvious from the product taught by Vandercruys in view of Schultz et al., Baert et al., Redenti et al., and Pitha et al. even though the prior product was made by a different process of Vandercruys teaching the composition to be an intimate admixture prepared by dissolving the mixture and removing the solvent by a vacuumdistillation process.

Response to Applicant's Remarks:

Applicant's Remarks, filed 28 May 2013, have been fully considered and not found to be persuasive regarding the new grounds of rejection.

Applicant's remarks are persuasive regarding rejection over Schultz et al. (US Patent 6,194,395, published 27 Feb 2001, provided by Applicant in IDS mailed 7 Jan 2011) in view of Baert et al. (WIPO Publication WO97/18839, published 29 May 1997, of record). Applicant argues that Schultz et al. is non-operable because the process of Baert et al. will result in decomposition of cladribine. However, Schultz et al. in claim 13 and 14 at column 8, lines 40-60 explicitly claims the composition prepared by meltextrusion. A patent shall be presumed valid and each claim of a patent (whether in

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independent, dependent, or multiple dependent form) shall be presumed valid independently of the validity of other claims. See 35 USC 282. However Redenti et al. (Int. J. Pharm., 1996, 129, p289-294, cited in PTO-892) provides evidence supporting Applicant's remarks that the composition prepared by melt-extrusion is a physical mixture and not a complex.

Applicant's remarks are moot in view of Vandercruys teaching the composition to be an intimate admixture prepared by dissolving the mixture and removing the solvent by a vacuum-distillation process and implying the intimate admixture to be a complex. Vandercruys teaches the components are preferably in the glass thermoplastic phase without a crystalline or microcrystalline phase, or an amorphous complex, and Pitha et al. teaches it is known to one of ordinary skill in the art to use 2-hydroxypropyl CD to form amorphous mixtures through formation of inclusion complexes. Therefore the new grounds of rejection over Vandercruys in view of Schultz et al., Baert et al., Redenti et al., and Pitha et al. teaches all structural limitations of the product and product-byprocess as instantly claimed.

The following grounds of rejection are maintained.

Double Patenting

The nonstatutory double patenting rejection is based on a judicially created doctrine grounded in public policy (a policy reflected in the statute) so as to prevent the unjustified or improper timewise extension of the "right to exclude" granted by a patent and to prevent possible harassment by multiple assignees. A nonstatutory obviousness-type double patenting rejection is appropriate where the conflicting claims are not identical, but at least one examined application claim is not patentably distinct from the reference claim(s) because the examined application claim is either anticipated by, or would have been obvious over, the reference claim(s). See, e.g., *In re Berg*, 140 F.3d 1428, 46 USPQ2d 1226 (Fed. Cir. 1998); *In re Goodman*, 11 F.3d 1046, 29 USPQ2d 2010 (Fed. Cir. 1993); *In re Longi*, 759 F.2d 887, 225 USPQ 645 (Fed. Cir. 1985); *In re Van Ornum*, 686 F.2d 937, 214 USPQ 761 (CCPA 1982); *In re Vogel*, 422 F.2d 438, 164 USPQ 619 (CCPA 1970); and *In re Thorington*, 418 F.2d 528, 163 USPQ 644 (CCPA 1969).

A timely filed terminal disclaimer in compliance with 37 CFR 1.321(c) or 1.321(d) may be used to overcome an actual or provisional rejection based on a nonstatutory double patenting ground provided the conflicting application or patent either is shown to be commonly owned with this application, or claims an invention made as a result of activities undertaken within the scope of a joint research agreement.

Effective January 1, 1994, a registered attorney or agent of record may sign a terminal disclaimer. A terminal disclaimer signed by the assignee must fully comply with 37 CFR 3.73(b).

Claims 1-8, 25-31 and 46-61 are rejected on the ground of nonstatutory double patenting over claims 1-28 of U. S. Patent No. 7,888,328 since the claims, if allowed, would improperly extend the "right to exclude" already granted in the patent.

The subject matter claimed in the instant application is fully disclosed in the patent and is covered by the patent since the patent and the application are claiming common subject matter, as follows: claims 1-28 of U. S. Patent No. 7,888,328 recite a narrower genus of cladribine-cyclodextrin complex and composition comprising thereof wherein said cyclodextrin is hydroxypropyl- β -cyclodextrin and the weight ratio of from about 1:10 to about 1:16. Claims 12-28 of U. S. Patent No. 7,888,328 recite a product-by-process encompassed within the product-by-process of instant claims 46-61. Therefore claims 1-28 of U. S. Patent No. 7,888,328 supports an anticipation-type nonstatutory double patenting rejection over instant claims 1-8, 25-31 and 46-61.

Furthermore, there is no apparent reason why applicant was prevented from presenting claims corresponding to those of the instant application during prosecution of the application which matured into a patent. See *In re Schneller*, 397 F.2d 350, 158 USPQ 210 (CCPA 1968). See also MPEP § 804.

Response to Applicant's Remarks:

Applicant's Remarks, filed 28 May 2013, have been fully considered and not found to be persuasive.

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Applicant notes that the distinction between 1) "same invention" type double patenting statutorily based on 35 USC 101 and 2) nonstatutory-type double patenting rejection, commonly called an "obviousness-type double patenting rejection". To clarify the language of the above rejection, it is meant that anticipation is the epitome of obviousness in the context of patent examination, therefore the above rejection which is stated to be "on the ground of <u>nonstatutory</u> double patenting" (emphasis added) in the rejection statement is performed with an anticipation-type analysis which is meant to be the epitome of obviousness. Applicant is correct that the nonstatutory-type double patenting rejection detailed herein falls within the 2nd category and would be overcome by filing of a terminal disclaimer. Applicant's remarks state that a terminal disclaimer was filed, however a review of the record of the instant application shows <u>no terminal</u>

disclaimer has been filed. Clarification is requested regarding the terminal disclaimer.

This rejection is maintained because the record shows no terminal disclaimer has been filed.

Conclusion

This Office Action details new grounds of rejection not necessitated by Applicant's Amendment. Accordingly, this Office Action is Non-Final.

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Jonathan S. Lau whose telephone number is (571)270-

3531. The examiner can normally be reached on Monday - Thursday, 9 am - 4 pm EST.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Shaojia Anna Jiang can be reached on 571-272-0627. The fax phone number for the organization where this application or proceeding is assigned is 571-273-8300.

Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free). If you would like assistance from a USPTO Customer Service Representative or access to the automated information system, call 800-786-9199 (IN USA OR CANADA) or 571-272-1000.

/SHAOJIA ANNA JIANG/ Supervisory Patent Examiner, Art Unit 1623 /Jonathan S Lau/ Examiner, Art Unit 1623

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U.S. PATENT DOCUMENTS

*		Document Number Country Code-Number-Kind Code	Date MM-YYYY	Name	Classification
*	А	US-2002/0150616	10-2002	VANDECRUYS, ROGER PETRUS GEREBERN	424/464
	В	US-			
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NON-PATENT DOCUMENTS

* Include as applicable: Author, Title Date, Publisher, Edition or Volume, Pertinent Pages) U Redenti et al., Int. J. Pharm., 1996, 129, p289-294. Pitha et al., Life Sci., 1998, 43, p493-502. ۷ W Х

*A copy of this reference is not being furnished with this Office action. (See MPEP § 707.05(a).) Dates in MM-YYYY format are publication dates. Classifications may be US or foreign.

	Application/Control No.	Applicant(s)/Patent Under Reexamination
Search Notes	12986310	BODOR ET AL.
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	JONATHAN S LAU	1623

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Google Scholar - see attached notes	2/15/2013	JSL								
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	Articles	Drug solubilizers to aid pharmacologists: amorphous cyclodextrin derivatives		
8	Legal documents	J Pitha, T trie, PB Sktar, JS Nye - Lite sciences, 1988 - Elsevier Abstract Conversion of crystalline α-, β-, or γ- cyclodextrins into amorphous mixtures of water soluble derivatives yields non-toxic solubilizers which dissolve drugs through the formation		
	Any time	of inclusion complexes. From these types of compounds 2-hydroxypropyl ethers of Clied by 213 Related articles Ali 3 versions Cite		
	Since 2013 Since 2012 Since 2009 Custom range	A non-surfactant formulation for alfaxalone based on an amorphous cyclodextrin : Activity studies in rats and dogs KS Estes, ME Brewster, Al Webb, N Bodor - International journal of, 1990 - Elsevier Abstract The steroid anesthetic, alfaxalone, is commercially available for veterinary use in a		
	Sort by relevance Sort by date	formulation containing Cremophor-EL® and alfadalone acetate. The allergic sensitivity of some species to this formulation limits its veterinary application and also led to withdrawal Cited by 28 Related articles All 3 versions Cite	N7	
	include patents	DSC GP Betlinetti, M Sorrenti, S Rossi, F Ferrari • of pharmaceutical and, 2002 - Elsevier A microcalorimetric method based on differential scanning calorimetry (DSC) of drug-	¥.	
	🗰 Create alert	(NAP) in combinations with amorphous hydroxypropyl β- cyclodextrin (HPβCd), β Cited by 11 Related anticles All 5 versions Cite		
		Amorphous water soluble derivatives of cyclodextrins: Nontoxic dissolution enhancing excipients J Pliha, J Pliha - Journal of pharmaceutical sciences, 1985 - Wiley Online Library Abstract Dissolution properties of drugs may be improved by their conversion to an amorphous state or by complexation with cyclodextrins. The present report describes the preparation of cyclodextrin derivatives which are intrinsically amorphous , and water Cited by 184 Related articles All 4 versions Cite		
		Carbon-13 CP/MAS NMR studies of amylose inclusion complexes, cyclodextrins, and the amorphous phase of starch granules: relationships between glycosidic MJ Gidley, SM Bociek - Journal of the American Chemical Society, 1988 - ACS Publications Abstract: In order to characterize molecular conformations within starch granules and to examine the relationships between polysaccharide conformation and solid state I3C chemical shifts, a range of polymeric and oligomeric a-(1-4) glucans has been examined Cited by 251 Related articles All 4 versions Cite		
		<u>Use of 2-hydroxypropyl-B-cyclodextrin as a solubilizing and stabilizing excipient for protein drug</u> ME Brewster, MS Hora, JW Simpkins, N Bodor - Pharmaceutical research, 1991 - Springer T. Irle, K. Fukanaga, A. Yoshida, K. Uekama, H. Fales, and J. Pitha. Amorphous water-soluble cyclodextrin derivatives. Pharm. Res. 5:713-717 (198 3)	2	

water-soluble derivatives of cyclodextrins: Nontoxic dissolution enhancing excipients. ... Cited by 125 Related anticles All 6 versions. Cite

An intravenous toxicity study of 2-hydroxypropyl-B-cyclodextrin. a useful drug solubilizer. in rats [PDF] from addiandcassi.com

and monkeys

ME Brewster, KS Estes, N Bodor - International journal of pharmaceutics, 1990 - Elsevier ... Pharmaceutical preparations containing cyclodextrin derivatives. (1988) US Patent 4, 727, 064. Pitha and Pitha, 1985; J. Pitha, J. Pitha; Amorphous water-soluble derivatives of cyclodextrins: nontoxic dissolution enhancing excipients. J. Pharm. Sci., 74 (1985), pp. 987–990. ... Cited by 92 Related articles All 6 versions Cite

Characterization of physicochemical properties of naproxen systems with amorphous

B-cyclodextrin-epichlorohydrin polymers

Evaluation of the cytotoxicity of cyclodextrins and hydroxypropylated derivatives

F Leroy-Lechat, D Wouessidjewe, JP Andreux... - International journal of ..., 1994 - Elsevier ... 373–376. Pitha et al., 1988; J. Pitha, T. Irie, PB Sklar, JS Nye; Drug solubilizers to aid pharmacologists. **Amorphous cyclodextrin** derivatives. Life Sci., 43 (1988), pp. 493–502. ... Pitha, 1990; J. Pitha; Hydroxypropyl **cyclodextrins** in pharmacy and pharmacology, Progress from toy ... Cited by 57 Related articles. All 2 versions. Cite

A study on the differentiation between **amorphous** piroxicam: <u>B-cyclodextrin</u> complex and a mixture of the two **amorphous** components

E Redenti, T Peveri, M Zanol, P Ventura... - International journal of ..., 1996 - Elsevier **Amorphous** piroxicam was prepared by the melt spinning method to prove that freeze-dried **amorphous** piroxicam: β -cyclodextrin is a true inclusion compound and not a dispersed mixture of the two **amorphous** components. Differential scanning calorimetry (DSC) and ... Cited by 42 Related articles All 3 versions Cite

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Application Number	12986310				
Filing Date	07-Jan-2011				
First Named Inventor	Nicholas Bodor				
Attorney Docket Number	20009904-0067				
Title of Invention	ORAL FORMULATIONS OF CLAD	DRIBINE			
Filing of terminal disclaimer does Office Action	not obviate requirement for resp	onse unde	r 37 CFR 1.111 to outstanding		
This electronic Terminal Disclaim	er is not being used for a Joint Res	search Agro	eement.		
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- is reissued; or - is in any manner terminated prior to t	the expiration of its full statutory t	erm as pre	sently shortened by any terminal disclaimer.		
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I hereby declare that all statements r belief are believed to be true; and fu the like so made are punishable by fi that such willful false statements ma	made herein of my own knowledge are true and that all statements made on information and rther that these statements were made with the knowledge that willful false statements and ine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and y jeopardize the validity of the application or any patent issued thereon.				
THIS PORTION MUST BE COMPLETE	D BY THE SIGNATORY OR SIGNATORIES				
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 An attorney or agent registered this application 	An attorney or agent registered to practice before the Patent and Trademark Office who is of record in this application				
Registration Number26254					
A sole inventor					
A joint inventor; I certify that I am authorized to sign this submission on behalf of all of the inventors					
A joint inventor; all of whom are signing this request					
O The assignee of record of the entire interest that has properly made itself of record pursuant to 37 <u>CFR 3.7</u> 1					
Signature	/Mary Katherine Baumeister/				
Name	Mary Katherine Baumeister				

*Statement under 37 CFR 3.73(b) is required if terminal disclaimer is signed by the assignee (owner). Form PTO/SB/96 may be used for making this certification. See MPEP § 324.

Electronic Patent Application Fee Transmittal							
Application Number:		12986310					
Filing Date:	07-	Jan-2011					
Title of Invention:		ORAL FORMULATIONS OF CLADRIBINE					
First Named Inventor/Applicant Name:	Nic	holas S. Bodor					
Filer:	Ma	ry Katherine Baume	eister/Rebecca E	Brimmer			
Attorney Docket Number:	20009904-0067						
Filed as Large Entity							
Utility under 35 USC 111(a) Filing Fees							
Description		Fee Code	Quantity	Amount	Sub-Total in USD(\$)		
Basic Filing:							
Statutory or Terminal Disclaimer		1814	1	160	160		
Pages:							
Claims:							
Miscellaneous-Filing:							
Petition:							
Patent-Appeals-and-Interference:							
Post-Allowance-and-Post-Issuance:							
Extension-of-Time:		348					

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Miscellaneous:				
	Tot	al in USD	(\$)	160

Doc Code: DISQ.E.FILE Document Description: Electronic Terminal Disclaimer – Approved

Application No.: 12986310

Filing Date: 07-Jan-2011

Applicant/Patent under Reexamination: Bodor et al.

Electronic Terminal Disclaimer filed on December 10, 2013

APPROVED

This patent is subject to a terminal disclaimer

DISAPPROVED

Approved/Disapproved by: Electronic Terminal Disclaimer automatically approved by EFS-Web

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Electronic Acknowledgement Receipt					
EFS ID:	17619390				
Application Number:	12986310				
International Application Number:					
Confirmation Number:	6100				
Title of Invention:	ORAL FORMULATIONS OF CLADRIBINE				
First Named Inventor/Applicant Name:	Nicholas S. Bodor				
Customer Number:	13974				
Filer:	Mary Katherine Baumeister/Rebecca Brimmer				
Filer Authorized By:	Mary Katherine Baumeister				
Attorney Docket Number:	20009904-0067				
Receipt Date:	10-DEC-2013				
Filing Date:	07-JAN-2011				
Time Stamp:	17:12:04				
Application Type:	Utility under 35 USC 111(a)				

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National Stage of an International Application under 35 U.S.C. 371 If a timely submission to enter the national stage of an international application is compliant with the conditions of 35 U.S.C. 371 and other applicable requirements a Form PCT/DO/EO/903 indicating acceptance of the application as a national stage submission under 35 U.S.C. 371 will be issued in addition to the Filing Receipt, in due course. <u>New International Application Filed with the USPTO as a Receiving Office</u> If a new international application is being filed and the international application includes the necessary components for an international filing date (see PCT Article 11 and MPEP 1810), a Notification of the International Application Number and of the International Filing Date (Form PCT/RO/105) will be issued in due course, subject to prescriptions concerning national security, and the date shown on this Acknowledgement Receipt will establish the international filing date of the application.								

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applica	ant: Nicholas S. BODOR et al.)	MAIL STOP AMENDMENT
Applic	ation No.: 12/986,310)	Examiner: Jonathan S. LAU
Filed:	January 7, 2011)	Group Art Unit: 1623
Title:	ORAL FORMULATIONS OF CLADRIBINE)))	Confirmation No.: 6100

REPLY TO OFFICIAL ACTION

In response to the Office Action dated September 11, 2013, Applicants submit the remarks below:

Attorney Docket No. 20009904-0067 Application No. 12/986,310 Page 2

REMARKS

Further examination and consideration of the subject application, as pursuant to and consistent with 37 C.F.R. § 1.112, are respectfully requested in light of the remarks that follow.

DOMESTIC PRIORITY

The Examiner's acknowledgment of the several claims for domestic priority is noted, with appreciation.

STATUS OF CLAIMS

Claims 1-61 and 65-88 are now in this application. Claims 9-24, 32-45, and 65-88 have been withdrawn. Claims 1-8, 25-31 and 46-61 are under examination. No amendments have been made.

DISCUSSION OF RESTRICTION REQUIREMENT

Newly submitted Claims 65-88 have been withdrawn from consideration. Applicants appreciate the Examiner's statement that these claims will be considered for rejoinder upon finding allowable claims.

CLAIM REJECTIONS- 35 U.S.C. § 103

Claims 1-8, 25-31 and 46-61 have been rejected under 35 U.S.C. § 103(a) as being unpatentable over Vandercruys (U.S. Publication No. 2002/0150616) in view of Schultz et al., (US Patent No. 6,194,395), Baert (International Publication No.

WO97/18839), Redenti et al., (Int. J. Pharm., 1996, 129, p289-294) and Pitha et al., (Life Sci., 1998, 43, p493-502). It is submitted that this rejection cannot be maintained against any of the claims in this application.

The teaching of Vandercruys (US 2002/0150616) is limited to compositions, wherein the drug compound, an acid, a cyclodextrin and an organic polymer are intimately admixed [0011]. Such limitation is also stated in the abstract. As set forth in the present application as filed at page 1, line 15, cladribine is an acid-labile drug. In view of this and the clear teaching that Vandercruys deals with formulations which mandatorily contain an acid ingredient which is detrimental to the stability of cladribine, a person skilled in the art would not have considered Vandercruys if he wanted to develop a formulation for cladribine, despite its mention in a "laundry list" of active ingredients.

All components being present in the composition taught by Vandercruys, i.e. the acid, drug compound, cyclodextrin and organic polymer, are mixed together, whereby it is emphasized that all four components are present and indeed that "mixtures where individual particles do not contain all four components are not preferred". Such intimate admixture is taught as complimentary at the microscopic level [0012] and it is further taught that all components are dispersed to form the chemically and physically uniform homogeneous material throughout which is called a glass thermoplastic phase or system [0013]. Such system is exemplified in Example 1 and is made by solvation of drug compound and acid in the solvent, addition and dissolution of the cyclodextrin and polymer and subsequent evaporation of the solvent. Therefore, Vandercruys provides a clear teaching to combine the drug, an acid, a cyclodextrin and an organic polymer into

one physically and chemically homogeneous thermoplastic system composition preparation.

Vandercruys does not teach a combination of the drug and a cyclodextrin without an acid and an organic polymer. In Example 1, the acid is even present in a severalfold amount of the drug (weight ratio drug : acid = 1 : 5) and in a double amount compared to the cyclodextrin. Example 2 discloses similar weight ratios. It is impermissible to directly deduce a teaching given for a composition containing all components to a part of it consisting of only two components (drug and cyclodextrin). This is especially the case if another ingredient (here an acid) is mandatory and is present in an amount exceeding that which is picked out of it (here cyclodextrin).

The foregoing is even true for the result achieved by a process which is described to be used for the preparation of the composition, i.e. the vacuum distillation process described in the Examples 1 and 2 on page 8.

Further, Vandercruys teaches that the cyclodextrin is preferably present at 5 to 70% by weight relative to the total weight of cyclodextrin, acid, organic polymer and drug, and that the molar ratio of cyclodextrin to drug preferably lies in the range of 100 :1 [0027], This clearly shows that the cyclodextrin forms part of a composition of multiple ingredients and that <u>Vandercruys provides no teaching with respect to complexation</u> and association of amorphous drug to cyclodextrin complex as provided by the present invention.

In summary, Vandercruys discloses compositions which in addition to the drug and the cyclodextrin mandatorily contain further ingredients (an acid and an organic polymer) which form an integral part of the glass thermoplastic system and do not

reasonably allow one to pick out parts of it and to combine it with other teachings. Consideration of Vandercruys is further prohibited in that it teaches that an acid must be present (in an amount strongly exceeding the amount of the drug) and a person skilled in the art would reasonably expect that the presence of such acid would be detrimental to the stability of cladribine in view of the fact that cladribine is an acid-labile compound.

The secondary references do not supply the deficiencies in the primary reference, Vandercruys.

Schultz (US 6194395) teaches the use of cyclodextrin as a solubilizer and stabilizer of cladribine in aqueous solution only, which is quite different for the invention. As far as solid oral dosage forms are mentioned, such disclosure is limited to solid mixtures of cladribine and cyclodextrin (see column 5, lines 50-58). The Examiner argues that the melt-extruding of the mixture refers to Baert (WO 97/118839) and that, according to Baert, such melt-extruding leads to amorphous material. However, the fact that melt-extruding of cyclodextrin and cladribine might lead to amorphous material does not imply that such material consists of an amorphous inclusion complex of cladribine in cyclodextrin and more than ever does not imply that amorphous free cladribine is associated with amorphous cyclodextrin complex. Amorphousness can also be provided by different arrangements such as by a mixture of the amorphous cladribine and cyclodextrin. Further, a mixture is explicitly taught by Schultz. Indeed. as the Examiner himself has acknowledged on page 4 of the September 11, 2013 Official Action, "Applicant's remarks are persuasive that the mixture taught by Schultz in view of Baert can reasonably be concluded to be the physical mixture, and the comparative evidence provided by Van Axel Castelli is applicable and persuasive."

The Examiner's allegation set forth on page 9, second paragraph, regarding Rendenti et al., that piroxicam/ β -cyclodextrin complex prepared by melt-spinning is a physical mixture of amorphous piroxicam and amorphous cyclodextrin supports such view. This is because melt extruding comprises the steps a) mixing of cyclodextrin and active ingredient, c) heating of the mixture until melting of one of the components and e) cooling the mixture till it solidifies (See Baert, page 5 steps a), c) and e)). As piroxicam has the melting point of 205 - 210°C (page 290, right column, line 1) and β -cyclodextrin melts with decomposition of 250°C (page 291, left column, lines 13 - 25), melt-spinning of a piroxicam β -cyclodextrin mixture must also comprise the step of heating of the mixture until melting of one of the components (here piroxicam only, melting of β cyclodextrin can be excluded as this is accompanied by its decomposition), as described in step c) of Baert for melt extruding.

Further, with respect to instant Claims 5-6 and 29-30, while Baert teaches that ratios of active ingredient to cyclodextrin in their <u>mixtures</u> may vary widely and mentions ratios of 1/100 to 100/1 (page 11, first paragraph), these claims require specific ratios of cladribine to cyclodextrin in the claimed complexes. It is clear that no teaching is given in the art to achieve the technical teaching of the invention, i.e. to provide an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex. The Examiner's calculations as set forth on page 8 are nothing else than cherry picking based on hindsight analysis, using Applicants' own invention as a guide through the maze of Vandercruys, Schultz et al., Baert et al., Rendenti et al. and Pitha et al. This is impermissable "picking and

choosing". Applicants submit that withdrawal of the rejection is in order.

With respect to product-by-process Claims 46-61, these claims afford the same product as that of Claim 1 and are patentable for the same reason that Claim 1 is patentable. Vandercruys does not provide the claimed composition comprising a complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a noninclusion complex, formulated into a solid oral dosage form, said composition comprising no significant amount of free crystalline cladribine therein. The secondary references fail to provide what Vandercruys lacks. The references in combination fail to teach or suggest Applicants' invention.

DOUBLE PATENTING

The Examiner has maintained the double patenting rejection over Claims 1-28 of U.S. Patent No. 7,888,328. Applicants' representative sincerely apologizes for the fact that the Terminal Disclaimer which had been meant to be filed on the same day as the filing of the previous response was inadvertently and unintentionally not filed at that time. Applicants' representative also apologizes for any inconvenience this has caused to the Examiner. An e-Terminal Disclaimer with respect to the '328 patent has now been filed and approved. Therefore, it is believed that the double patenting rejection has now been obviated.

As to *In re Schneller*, 158 USPQ 210 (CCPA 1968), that case was one in which a Terminal Disclaimer had not been filed. Further, Applicants here did in fact present

claims corresponding to those herein in the parent application.

CONCLUSION

The foregoing remarks are being made to place the application in condition for allowance. Applicants respectfully request reconsideration and the timely allowance of the pending claims. A favorable action is awaited. Should an interview be helpful to further prosecution of this application, the Examiner is invited to telephone the undersigned.

If there are any additional fees due in connection with the filing of this response, please charge the fees to Deposit Account No. 19-3140. If a fee is required for an extension of time under 37 C.F.R. §1.136 not accounted for above, such an extension is requested and the fee should also be charged to the Deposit Account.

Respectfully submitted,

DENTONS US LLP

Date: December 11, 2013

New Katherin Baumsister

Mary Katherine Baumeister Registration No. 26254

Customer No. 13974 SNR Denton US LLP 1301 K Street NW, Suite 600, East Tower Washington, D.C. 20005 Phone: 202-408-9186 Fax: 202-408-6399
Electronic Acl	cnowledgement Receipt	
EFS ID:	17634350	
Application Number:	12986310	
International Application Number:		
Confirmation Number:	6100	
Title of Invention:	ORAL FORMULATIONS OF CLADRIBINE	
First Named Inventor/Applicant Name:	Nicholas S. Bodor	
Customer Number:	13974	
Filer:	Mary Katherine Baumeister/Rebecca Brimmer	
Filer Authorized By:	Mary Katherine Baumeister	
Attorney Docket Number:	20009904-0067	
Receipt Date:	11-DEC-2013	
Filing Date:	07-JAN-2011	
Time Stamp:	18:55:02	
Application Type:	Utility under 35 USC 111(a)	

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New Applications Under 35 U.S.C. 111

If a new application is being filed and the application includes the necessary components for a filing date (see 37 CFR 1.53(b)-(d) and MPEP 506), a Filing Receipt (37 CFR 1.54) will be issued in due course and the date shown on this Acknowledgement Receipt will establish the filing date of the application.

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If a timely submission to enter the national stage of an international application is compliant with the conditions of 35 U.S.C. 371 and other applicable requirements a Form PCT/DO/EO/903 indicating acceptance of the application as a national stage submission under 35 U.S.C. 371 will be issued in addition to the Filing Receipt, in due course.

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If a new international application is being filed and the international application includes the necessary components for an international filing date (see PCT Article 11 and MPEP 1810), a Notification of the International Application Number and of the International Filing Date (Form PCT/RO/105) will be issued in due course, subject to prescriptions concerning national security, and the date shown on this Acknowledgement Receipt will establish the international filing date of the application.

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicant: Nicholas S. BODOR et al.) MAIL STOP AMENDMENT
Application No.: 12/986,310)) Examiner: Jonathan S. LAU
Filed: January 7, 2011)) Group Art Unit: 1623
Title: ORAL FORMULATIONS OF CLADRIBINE) Confirmation No.: 6100

SUPPLEMENTAL AMENDMENT IN RESPONSE TO TELEPHONE INTERVIEW

Commissioner for Patents P.O. Box 1450 Alexandria, VA 22313-1450

Madame:

In supplemental response to the Office Action dated September 11, 2013, please

first amend the above-identified patent application as follows:

Attorney Docket No. 20009904-0067 Application No. 12/986,310 Page 2

AMENDMENTS TO THE CLAIMS:

This listing of claims will replace all prior versions, and listings, of claims in the application:

LISTING OF CLAIMS:

1. (Original) A pharmaceutical composition comprising a complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex, formulated into a solid oral dosage form, said composition comprising no significant amount of free crystalline cladribine therein.

2. (Original) The pharmaceutical composition according to Claim 1, wherein the complex is saturated with cladribine.

3. (Original) The composition according to Claim 1, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin, hydroxypropyl- γ -cyclodextrin, randomly methylated β -cyclodextrin, carboxymethyl- β -cyclodextrin or sulfobutyl- β -cyclodextrin.

4. (Original) The composition according to Claim 1, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin or hydroxypropyl-γ-cyclodextrin.

5. (Original) The composition according to Claim 1, wherein the weight ratio of cladribine to amorphous cyclodextrin is from about 1:10 to about 1:16.

6. (Original) The composition according to Claim 5, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin or hydroxypropyl- γ -cyclodextrin.

7. (Original) The composition according to Claim 1, wherein the approximate molar ratio of cladribine to amorphous cyclodextrin corresponds to a point located on the curve of a phase solubility diagram for saturated complexes of cladribine in varying concentrations of the cyclodextrin.

8. (Original) The composition according to Claim 1, wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b).

9. (Withdrawn) A method for enhancing the oral bioavailability of cladribine comprising orally administering to a subject in need thereof a pharmaceutical composition comprising a complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous

cyclodextrin as a non-inclusion complex, formulated into a solid oral dosage form, said composition comprising no significant amount of free crystalline cladribine therein.

10. (Withdrawn) The method according to Claim 9, wherein the complex is saturated with cladribine.

11. (Withdrawn) The method according to Claim 9, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin, hydroxypropyl- γ -cyclodextrin, randomly methylated β -cyclodextrin, carboxymethyl- β -cyclodextrin or sulfobutyl- β -cyclodextrin.

12. (Withdrawn) The method according to Claim 9, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin or hydroxypropyl-γ-cyclodextrin.

13. (Withdrawn) The method according to Claim 9, wherein the weight ratio of cladribine to amorphous cyclodextrin is from about 1:10 to about 1:16.

14. (Withdrawn) The method according to Claim 13, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin or hydroxypropyl-γ-cyclodextrin.

15. (Withdrawn) The method according to Claim 9, wherein the approximate molar ratio of cladribine to amorphous cyclodextrin corresponds to a point located on

the curve of a phase solubility diagram for saturated complexes of cladribine in varying concentrations of the cyclodextrin.

16. (Withdrawn) The method according to Claim 9, wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b).

17. (Withdrawn-Currently Amended) A method for the treatment of symptoms of a cladribine-responsive condition <u>selected from the group consisting of multiple</u> <u>sclerosis, rheumatoid arthritis and leukemia</u> in a subject suffering from said symptoms comprising orally administering to said subject a pharmaceutical composition comprising a complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex, formulated into a solid oral dosage form, said composition comprising no significant amount of free crystalline cladribine therein.

18. (Withdrawn) The method according to Claim 17, wherein the complex is saturated with cladribine.

19. (Canceled)

20. (Withdrawn-Currently Amended) The method according to Claim 19<u>17</u>, wherein the cladribine-responsive condition is multiple sclerosis.

21. (Withdrawn) The method according to Claim 17, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin, hydroxypropyl- γ -cyclodextrin, randomly methylated β -cyclodextrin, carboxymethyl- β -cyclodextrin or sulfobutyl- β -cyclodextrin.

22. (Withdrawn) The method according to Claim 17, wherein the weight ratio of cladribine to amorphous cyclodextrin is from about 1:10 to about 1:16.

23. (Withdrawn) The method according to Claim 17, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin or hydroxypropyl-γ-cyclodextrin.

24. (Withdrawn) The method according to Claim 17, wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b).

25. (Original) A complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex.

26. (Original) The complex according to Claim 25, saturated with cladribine.

27. (Original) The complex according to Claim 25, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin, hydroxypropyl- γ -cyclodextrin, randomly methylated β -cyclodextrin, carboxymethyl- β -cyclodextrin or sulfobutyl- β -cyclodextrin.

28. (Original) The complex according to Claim 25, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin or hydroxypropyl- γ -cyclodextrin.

29. (Original) The complex according to Claim 25, wherein the weight ratio of cladribine to amorphous cyclodextrin is from about 1:10 to about 1:16.

30. (Original) The complex according to Claim 29, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin or hydroxypropyl- γ -cyclodextrin.

31. (Original) The complex according to Claim 25, wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b).

32. (Withdrawn) A process for the preparation of a complex cladribinecyclodextrin complex as claimed in Claim 25, which comprises the steps of:

(i) combining cladribine and an amorphous cyclodextrin in water at a temperature of from about 40 to about 80°C and maintaining said temperature for a period of from about 6 to about 24 hours;

(ii) cooling the resultant aqueous solution to room temperature; and

(iii) Iyophilizing the cooled solution to afford an amorphous product.

33. (Withdrawn) The process according to Claim 32, further comprising a filtration step following step (ii).

34. (Withdrawn) The process according to Claim 32, wherein step (i) is performed at a temperature of from about 45 to about 60°C.

35. (Withdrawn) The process according to Claim 32, wherein step (i) is performed at a temperature of from about 45 to about 50°C.

36. (Withdrawn) The process according to Claim 34, wherein step (i) is performed with stirring.

37. (Withdrawn) The process according to Claim 36, wherein step (i) is performed for a period of from about 6 to about 9 hours.

38. (Withdrawn) The process according to Claim 32, wherein step (ii) is performed for a period of from about 6 to about 9 hours.

39. (Withdrawn) The process according to Claim 32, wherein step (iii) comprises an initial freezing stage in which the solution is cooled to from about -40 to about -80° C, and held at said temperature for a period of from about 2 to about 4 hours.

40. (Withdrawn) The process according to Claim 39, wherein, in the initial freezing stage of step (iii), the solution is cooled to about -45°C.

41. (Withdrawn) The process according to Claim 34, wherein 12.00 parts by weight of cladribine and 172.50 parts by weight of hydroxypropyl- β -cyclodextrin are introduced in step (i), or wherein 16.35 parts by weight of cladribine and 172.50 parts by weight of hydroxypropyl- β -cyclodextrin are introduced in step (i).

42. (Withdrawn) The process according to Claim 41, wherein 825 parts by volume of water are introduced in step (i).

43. (Withdrawn) The process according to Claim 32, wherein the lyophilization step (iii) comprises:

(a) an initial freezing stage in which the complexation solution is brought to
 from about -40°C to about -80°C for approximately 2 to 4 hours;

(b) a primary drying stage at about -25°C for approximately 80 to 90 hours; and

(c) a secondary drying stage at about 30°C for approximately 15 to 20 hours.

44. (Withdrawn) The process according to Claim 43, wherein stage (a) of the lyophilization is conducted at about -45°C for approximately 3 to 4 hours.

45. (Withdrawn) The process according to Claim 43, wherein stage (b) of the lyophilization is conducted under a pressure of about 100 mTorr.

46. (Original) A pharmaceutical composition according to Claim 1, obtainable by a process comprising the steps of:

 (i) combining cladribine and an amorphous cyclodextrin in water at a temperature of from about 40 to about 80°C and maintaining said temperature for a period of from about 6 to about 24 hours;

(ii) cooling the resultant aqueous solution to room temperature;

- (iii) lyophilizing the cooled solution to afford an amorphous product; and
- (iv) formulating the amorphous product into a solid oral dosage form.

47. (Original) The pharmaceutical composition according to Claim 46, wherein the process further comprises a filtration step following step (i) or (ii).

48. (Original) The pharmaceutical composition according to Claim 46, wherein step (i) of the process is performed at a temperature of from about 45 to about 60°C.

49. (Original) The pharmaceutical composition according to Claim 46, wherein step (i) of the process is performed at a temperature of from about 45 to about 50°C.

50. (Original) The pharmaceutical composition according to Claim 48, wherein step (i) of the process is performed with stirring.

51. (Original) The pharmaceutical composition according to Claim 50, wherein step (i) of the process is performed for a period of from about 6 to about 9 hours.

52. (Original) The pharmaceutical composition according to Claim 46, wherein step (ii) of the process is performed for a period of from about 6 to about 9 hours.

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53. (Original) The pharmaceutical composition according Claim 46, wherein step (iii) comprises an initial freezing stage in which the solution is cooled to from about -40 to about -80°C, and held at said temperature for a period of from about 2 to about 4 hours.

54. (Original) The pharmaceutical composition according to Claim 53, wherein, in the initial freezing stage of step (iii), the solution is cooled to about -45°C.

55. (Original) The pharmaceutical composition according to Claim 48, wherein 12.00 parts by weight of cladribine and 172.50 parts by weight of the hydroxypropyl-β-cyclodextrin are introduced in step (i) of the process, or wherein 16.35 parts by weight of cladribine and 172.50 parts by weight of the hydroxypropyl-β-cyclodextrin are introduced in step (i) of the process.

56. (Original) The pharmaceutical composition according to Claim 55, wherein 825 parts by volume of water are introduced in step (i) of the process.

57. (Original) The pharmaceutical composition according to Claim 46, wherein the lyophilization step (iii) of the process comprises:

(a) an initial freezing stage in which the complexation solution is brought to
 from about -40°C to about -80°C for approximately 2 to 4 hours;

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(b) a primary drying stage at about -25°C for approximately 80 to 90 hours; and

(c) a secondary drying stage at about 30°C for approximately 15 to 20 hours.

58. (Original) The pharmaceutical composition according to Claim 57, wherein stage (a) of the lyophilization is conducted at about -45°C for approximately 3 to 4 hours.

59. (Original) The pharmaceutical composition according to Claim 57, wherein stage (b) of the lyophilization is conducted under a pressure of about 100 mTorr.

60. (Original) The pharmaceutical composition according to Claim 46, wherein the formulation step (iv) of the process comprises blending the complex with magnesium stearate and compressing into tablets.

61. (Original) The pharmaceutical composition according to Claim 60, wherein magnesium stearate is pre-mixed with sorbitol powder before blending with the complex.

62. - 64. (Canceled)

65. (Withdrawn-Currently Amended) A method for the treatment of symptoms of a cladribine-responsive condition <u>selected from the group consisting of multiple</u> <u>sclerosis, rheumatoid arthritis and leukemia</u> in a subject suffering from said symptoms comprising orally administering to said subject a pharmaceutical composition comprising a complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin as a non-inclusion complex, formulated into a solid oral dosage form, said composition comprising no significant amount of free crystalline cladribine therein, wherein administering cladribine is accompanied by administering one or more additional active ingredients for treating the cladribine-responsive condition.

66. (Withdrawn) The method according to Claim 65, wherein the complex is saturated with cladribine.

67. - 68. (Canceled)

69. (Withdrawn) The method according to Claim 65, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin, hydroxypropyl- γ -cyclodextrin, randomly methylated β -cyclodextrin, carboxymethyl- β -cyclodextrin or sulfobutyl- β -cyclodextrin.

70. (Withdrawn) The method according to Claim 66, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin, hydroxypropyl- γ -cyclodextrin, randomly methylated β -cyclodextrin, carboxymethyl- β -cyclodextrin or sulfobutyl- β -cyclodextrin.

71. - 72. (Canceled)

73. (Withdrawn) The method according to Claim 65, wherein the weight ratio of cladribine to amorphous cyclodextrin is from about 1:10 to about 1:16.

74. (Withdrawn) The method according to Claim 69, wherein the weight ratio of cladribine to amorphous cyclodextrin is from about 1:10 to about 1:16.

75. (Withdrawn) The method according to Claim 65, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin.

76. (Canceled)

77. (Withdrawn) The method according to Claim 73, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin.

78. (Withdrawn-Currently Amended) The method according to Claim 73.74, wherein the amorphous cyclodextrin is hydroxypropyl- γ -cyclodextrin.

79. (Withdrawn) The method according to Claim 75, wherein the weight ratio of cladribine to hydroxypropyl- β -cyclodextrin is about 1:14.

80. (Withdrawn) The method according to Claim 75, wherein the weight ratio of cladribine to hydroxypropyl- β -cyclodextrin is about 1:11.

81. (Withdrawn) The method according to Claim 65, wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b).

82. (Withdrawn) The method according to Claim 75, wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b).

83. (Withdrawn) The method according to Claim 65, wherein the cladribineresponsive condition is multiple sclerosis.

84. (Withdrawn) The method according to Claim 75, wherein the cladribineresponsive condition is multiple sclerosis.

85. (Withdrawn) The method according to Claim 77, wherein the cladribineresponsive condition is multiple sclerosis.

86. (Withdrawn) The method according to Claim 83, wherein one or more additional active ingredients for treating multiple sclerosis is/are selected from the group consisting of interferon beta, glatiramer acetate, natalizumab, alemtuzumab, 4-aminopyridine and amantadine.

87. (Withdrawn) The method according to Claim 84, wherein one or more additional active ingredients for treating multiple sclerosis is/are selected from the group consisting of interferon beta, glatiramer acetate, natalizumab, alemtuzumab, 4-aminopyridine and amantadine.

88. (Withdrawn) The method according to Claim 85, wherein one or more additional active ingredients for treating multiple sclerosis is/are selected from the group consisting of interferon beta, glatiramer acetate, natalizumab, alemtuzumab, 4-aminopyridine and amantadine.

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REMARKS

Further examination and consideration of the subject application, as amended, pursuant to and consistent with 37 C.F.R. § 1.112, are respectfully requested in light of the amendments to the claims and following remarks.

STATEMENT OF SUBSTANCE OF TELEPHONE INTERVIEW

The Applicants thank Examiner Lau for the courtesy of the Examiner-initialed telephone interview with the undersigned representative on February 18, 2014.

In the telephone interview, Examiner Lau indicated that the withdrawn method of treatment claims (17-24 and 65-88) could be rejoined and allowed, if the independent method of treatment claims were amended to specify that the cladribine-responsive condition is selected from the group consisting of multiple sclerosis, rheumatoid arthritis and leukemia (as originally specified in Claims 19, 67 and 68). Applicants' representative agreed to make the specified amendment to each of the independent method of treatment claims and to amend their dependent claims consistently with these amendments.

STATUS OF CLAIMS AND DISCUSSION OF AMENDMENTS

Claims 1-18, 20-61, 65-66, 69-70, 73-75 and 77-88 are now in this application. By the foregoing amendment, Claims 17, 20, 65 and 78 have been amended; and Claims 19, 67-68, 71-72 and 76 have been canceled as redundant in light of the amendment of the independent method of treatment claims. Claim 20 has been amended so that it no longer depends from a canceled claim. The dependency of Claim 78 has also been corrected; because of a typographical error, it was originally duplicative of Claim 77.

It is clear from the foregoing that no new matter has been introduced by these claim amendments and that the amendments place the method of treatment claims in rejoinable and allowable form.

CONCLUSION

The foregoing remarks and amendments are being made to place the method of treatment claims in the application in condition for rejoinder and allowance. Applicants respectfully request reconsideration and the timely allowance of the pending claims. A favorable action is awaited. Should a further interview be helpful to expedite prosecution of this application, the Examiner is invited to telephone the undersigned as soon as possible.

If there are any additional fees due in connection with the filing of this response, please charge the fees to counsel's Deposit Account No. 19-3140. If a fee is required

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for an extension of time under 37 C.F.R. §1.136 not accounted for above, such an

extension is requested and the fee should also be charged to counsel's Deposit

Account.

Respectfully submitted,

DENTONS US LLP

Date: February 20, 2014

/Mary Katherine Baumeister/ Mary Katherine Baumeister Registration No. 26254

Customer No. 13974

Dentons US LLP 1301 K Street NW, Suite 600, East Tower Washington, D.C. 20005 Phone: 202-408-9186 Fax: 202-408-6399

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Electronic Acl	cnowledgement Receipt
EFS ID:	18248155
Application Number:	12986310
International Application Number:	
Confirmation Number:	6100
Title of Invention:	ORAL FORMULATIONS OF CLADRIBINE
First Named Inventor/Applicant Name:	Nicholas S. Bodor
Customer Number:	13974
Filer:	Mary Katherine Baumeister/Louie Malloy
Filer Authorized By:	Mary Katherine Baumeister
Attorney Docket Number:	20009904-0067
Receipt Date:	20-FEB-2014
Filing Date:	07-JAN-2011
Time Stamp:	09:09:10
Application Type:	Utility under 35 USC 111(a)

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Document Number	Document Description	File Name	File Size(Bytes)/ Message Digest	Multi Part /.zip	Pages (if appl.)
1		0067SuppAmdt.pdf	180392 224c3b7e8b1fdf7d59900df2c6bdf960ee7c 2e5c	yes	20

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EXAMINER

LAU, JONATHAN S

ART UNIT PAPER NUMBER
1673

DATE MAILED: 03/13/2014

APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
12/986,310	01/07/2011	Nicholas S. Bodor	20009904-0067	6100

TITLE OF INVENTION: ORAL FORMULATIONS OF CLADRIBINE

APPLN. TYPE	ENTITY STATUS	ISSUE FEE DUE	PUBLICATION FEE DUE	PREV. PAID ISSUE FEE	TOTAL FEE(S) DUE	DATE DUE
nonprovisional	UNDISCOUNTED	\$960	\$0	\$0	\$960	06/13/2014

THE APPLICATION IDENTIFIED ABOVE HAS BEEN EXAMINED AND IS ALLOWED FOR ISSUANCE AS A PATENT. <u>PROSECUTION ON THE MERITS IS CLOSED</u>. THIS NOTICE OF ALLOWANCE IS NOT A GRANT OF PATENT RIGHTS. THIS APPLICATION IS SUBJECT TO WITHDRAWAL FROM ISSUE AT THE INITIATIVE OF THE OFFICE OR UPON PETITION BY THE APPLICANT. SEE 37 CFR 1.313 AND MPEP 1308.

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13974 7590 DENTONS US LLP P.O. BOX 061080 Chicago, IL 60606-1080 03/13/2014

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Certificate of Mailing or Transmission I hereby certify that this Fee(s) Transmittal is being deposited with the United States Postal Service with sufficient postage for first class mail in an envelope addressed to the Mail Stop ISSUE FEE address above, or being facsimile transmitted to the USPTO (571) 273-2885, on the date indicated below.

(Depositor's name
(Signature
(Date

APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
12/986.310	01/07/2011	Nicholas S. Bodor	20009904-0067	6100

TITLE OF INVENTION: ORAL FORMULATIONS OF CLADRIBINE

APPLN. TYPE ENTITY STATUS ISSUE FEE DUE PUBLICATION FEE DUE PREV. PAID ISSUE FEE TOTAL FEE(S) DUE DATE DUE nonprovisional UNDISCOUNTED \$960 \$0 \$0 \$0 \$960 06/13/2014 EXAMINER ART UNIT CLASS-STIECLASS I. Change of correspondence address or indication of "Fee Address" (37) CFR 1.360. 2. <t< th=""><th></th><th></th><th></th><th></th><th></th><th></th><th></th></t<>								
nonprovisional UNDISCOUNTED \$960 \$0 \$0 \$960 \$06132014 EXAMINER ART UNIT CLASS-SUBCLASS	APPLN. TYPE	ENTITY STATUS	ISSUE FEE DUE	PUBLICATION FEE DUE	PREV. PAID ISSUE FEE	TOTAL FEE(S) DUE	DATE DUE	
EXAMINER ART UNIT CLASS-SUBCLASS LAU, JONATHAN S 1673 514-046000 I. Change of correspondence address or indication of "Fee Address" (37 (CFR 1.50). 2. For printing on the patent front page, list (1) The names of up to 3 registered patent attorneys address form PTO/SB/122) attached. 1 I. Tee, Address" indication (or "Fee Address" indication form PTO/SB/47, Rev G3-Q2 or more recent) attached. Use of a Castomer 2 Wintber is required. 2. The reductor of a Castomer 2 3. ASSIGNEE NAME AND RESIDENCE DATA TO BE PRINTED ON THE PATENT! (print or type) PLEASE NOTE: Unless an assignee is identified below, no assignee data will appear on the patent. If an assignee is identified below, the document has been filed recordation as set forth in 37 CFR 3.11. Completion of this form is NOT a substitute for filing an assignment. (A) NAME OF ASSIGNEE Please check the appropriate assignee category or categories (will not be printed on the patent): Individual Corporation or other private group entity Governm data. The following fee(s) are submitted: I asser fee Ophication Fee (No small entity discount permitted) A check is enclosed. Optication or Micro Entity Status. See 37 CFR 1.29 I Applicant asserting small entity status. See 37 CFR 1.27 NOTE: Checking this box will be taken to micro entity status. See 37 CFR 1.27 I applicant changing to regular undiscounted fee status. NOTE: Checking this box will be taken to be a notifi	nonprovisional	UNDISCOUNTED	\$960	\$0	\$0	\$960	06/13/2014	
LAU, JONATHAN S 1673 514-046000 I Change of correspondence address or indication of "Fee Address" (37 CFR 1.53). 2. For printing on the patent front page, list (TFR 1.563). 0. The address" indication (or "Fee Address" indication form PIONSB/122) attached. 1. The Address" indication (or "Fee Address" indication form PIONSB/17, Rev 03-02 or more recent) attached. Use of a Customer 2. 3. ASSIGNEE NAME AND RESIDENCE DATA TO BE PRINTED ON THE PATENT (print or type) PLEASE NOTE: Unless an assignee is identified below, no assignee data will appear on the patent. If an assignee is identified below, the document has been filed recordation as set forth in 37 CFR 3.11. Completion of this form is NOT a substitute for filing an assignment. (A) NAME OF ASSIGNEE Please check the appropriate assignee category or categories (will not be printed on the patent): Individual Corporation or other private group entity Governn 4a. The following fee(s) are submitted: 4b. Payment of Fee(s): (Please first reapply any previously paid issue fee shown above) Acheck is enclosed. Payment by credit card. From TPO-2038 is attached. Change in Entity Status (from status indicated above) Acheck is enclosed. Payment by credit card. Form PTO-2038 is attached. Payment in the rice contification of Micro Entity Status, checking this box will be take to be a notification of loss of entitlement to micro entity status. See 37 CFR 1.27 NOTE: Absent a valid certification of Micro Entity Status, see 37 CFR 1.27 NOTE: Absent a val	EXAMINER		ART UNIT	CLASS-SUBCLASS				
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CIRN 1:303). (1) The names of up to 3 registered patent attorneys 1 CIRN 1:303). (1) The names of up to 3 registered patent attorneys 2 Green Address' indication for "Fee Address' indication form PTO/SB/122) attached. (2) The name of a single firm (having as a member a registered patent attorneys or agents and the names of up to 2 registered patent attorneys or agents. If no name is listed, no name will be printed. 2 3. ASSIGNEE NAME AND RESIDENCE DATA TO BE PRINTED ON THE PATENT (print or type) 9 2 3 PLEASE NOTE: Unless an assignce is identified below, no assignce data will appear on the patent. (A) NAME OF ASSIGNEE (B) RESIDENCE: (CITY and STATE OR COUNTRY) Please check the appropriate assignee category or categories (will not be printed on the patent): Individual Corporation or other private group entity Governm 4a. The following fee(s) are submitted: 4b. Payment of Fee(s): (Please first reapply any previously paid issue fee shown above) 4b. Payment by credit card. Form PTO-2038 is attached. 1 Check is enclosed. Publication Fee (No small entity status. See 37 CFR 1.29 NOTE: Absent a valid certification of Micro Entity Status (see forms PTO/SB/15A and 15B), iss fee papilicant casting small entity status. See 37 CFR 1.21 NOTE: Absent a valid certification of loss of entitlement to small or micro entity status. See 37 CFR 1.29 Applicant changing to regular undiscounted fee status. NOTE: Abs	1. Change of correspondence address or indication of "Fee Address" (37			2. For printing on the patent front page, list				
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Please check the appropriate assignee category or categories (will not be printed on the patent): Individual Corporation or other private group entity Governm 4a. The following fee(s) are submitted: 4b. Payment of Fee(s): (Please first reapply any previously paid issue fee shown above) A check is enclosed. Publication Fee (No small entity discount permitted) A check is enclosed. Payment by credit card. Form PTO-2038 is attached. Advance Order - # of Copies The Director is hereby authorized to charge the required fee(s), any deficiency, or credits any overpayment, to Deposit Account Number (enclose an extra copy of this forr 5. Change in Entity Status (from status indicated above) NOTE: Absent a valid certification of Micro Entity Status (see forms PTO/SB/15A and 15B), iss fee payment in the micro entity amount will not be accepted at the risk of application abandonme Applicant certifying micro entity status. See 37 CFR 1.27 NOTE: Absent a valid certification of loss of entitlement to micro entity status, checking this box will be take to be a notification of loss of entitlement to small or micro entity status. NOTE: This form must be signed in accordance with 37 CFR 1.31 and 1.33. See 37 CFR 1.4 for signature requirements and certifications. Date Authorized Signature Date Date Tyreed or printed name Resistration No.	(A) NAME OF ASS	IGNEE		(B) RESIDENCE: (CITY	and STATE OR COUNT	(RY)		
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Advance Order - # of Copies The Director is hereby authorized to charge the required fee(s), any deficiency, or credits any overpayment, to Deposit Account Number(enclose an extra copy of this form overpayment, to Deposit Account Number	Publication Fee (No small entity discount j	permitted)	Payment by credit card. Form PTO-2038 is attached.				
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	Typed or printed name				Registration No.			

Page **387** 3

PTOL-85 Part B (10-13) Approved for use through 10/31/2013.

OMB 0651-0033 U.S. Patent and Trademark Office; U.S. DEPARTMENT OF COMMERCE

UNITED STATES PATENT AND TRADEMARK OFFICE United States Patent and Trademark Office Address: COMMISSIONER FOR PATENTS P.O. Box 1450 Alexandria, Virginia 22313-1450 www.uspio.gov						
APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.		
12/986,310	01/07/2011	Nicholas S. Bodor	20009904-0067	6100		
13974 75	90 03/13/2014		EXAM	IINER		
DENTONS US LLP P.O. BOX 061080 Chicago, IL 60606-1080			LAU, JONATHAN S			
			ART UNIT	PAPER NUMBER		
			1673			
			DATE MAILED: 03/13/201	4		

Determination of Patent Term Adjustment under 35 U.S.C. 154 (b)

(application filed on or after May 29, 2000)

The Patent Term Adjustment to date is 172 day(s). If the issue fee is paid on the date that is three months after the mailing date of this notice and the patent issues on the Tuesday before the date that is 28 weeks (six and a half months) after the mailing date of this notice, the Patent Term Adjustment will be 172 day(s).

If a Continued Prosecution Application (CPA) was filed in the above-identified application, the filing date that determines Patent Term Adjustment is the filing date of the most recent CPA.

Applicant will be able to obtain more detailed information by accessing the Patent Application Information Retrieval (PAIR) WEB site (http://pair.uspto.gov).

Any questions regarding the Patent Term Extension or Adjustment determination should be directed to the Office of Patent Legal Administration at (571)-272-7702. Questions relating to issue and publication fee payments should be directed to the Customer Service Center of the Office of Patent Publication at 1-(888)-786-0101 or (571)-272-4200.

OMB Clearance and PRA Burden Statement for PTOL-85 Part B

The Paperwork Reduction Act (PRA) of 1995 requires Federal agencies to obtain Office of Management and Budget approval before requesting most types of information from the public. When OMB approves an agency request to collect information from the public, OMB (i) provides a valid OMB Control Number and expiration date for the agency to display on the instrument that will be used to collect the information and (ii) requires the agency to inform the public about the OMB Control Number's legal significance in accordance with 5 CFR 1320.5(b).

The information collected by PTOL-85 Part B is required by 37 CFR 1.311. The information is required to obtain or retain a benefit by the public which is to file (and by the USPTO to process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.14. This collection is estimated to take 12 minutes to complete, including gathering, preparing, and submitting the completed application form to the USPTO. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and Trademark Office, U.S. Department of Commerce, P.O. Box 1450, Alexandria, Virginia 22313-1450. DO NOT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS. SEND TO: Commissioner for Patents, P.O. Box 1450, Alexandria, Virginia 22313-1450. Under the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it displays a valid OMB control number.

Privacy Act Statement

The Privacy Act of 1974 (P.L. 93-579) requires that you be given certain information in connection with your submission of the attached form related to a patent application or patent. Accordingly, pursuant to the requirements of the Act, please be advised that: (1) the general authority for the collection of this information is 35 U.S.C. 2(b)(2); (2) furnishing of the information solicited is voluntary; and (3) the principal purpose for which the information is used by the U.S. Patent and Trademark Office is to process and/or examine your submission related to a patent application or patent. If you do not furnish the requested information, the U.S. Patent and Trademark Office may not be able to process and/or examine your submission, which may result in termination of proceedings or abandonment of the application or expiration of the patent.

The information provided by you in this form will be subject to the following routine uses:

- 1. The information on this form will be treated confidentially to the extent allowed under the Freedom of Information Act (5 U.S.C. 552) and the Privacy Act (5 U.S.C 552a). Records from this system of records may be disclosed to the Department of Justice to determine whether disclosure of these records is required by the Freedom of Information Act.
- 2. A record from this system of records may be disclosed, as a routine use, in the course of presenting evidence to a court, magistrate, or administrative tribunal, including disclosures to opposing counsel in the course of settlement negotiations.
- 3. A record in this system of records may be disclosed, as a routine use, to a Member of Congress submitting a request involving an individual, to whom the record pertains, when the individual has requested assistance from the Member with respect to the subject matter of the record.
- 4. A record in this system of records may be disclosed, as a routine use, to a contractor of the Agency having need for the information in order to perform a contract. Recipients of information shall be required to comply with the requirements of the Privacy Act of 1974, as amended, pursuant to 5 U.S.C. 552a(m).
- 5. A record related to an International Application filed under the Patent Cooperation Treaty in this system of records may be disclosed, as a routine use, to the International Bureau of the World Intellectual Property Organization, pursuant to the Patent Cooperation Treaty.
- 6. A record in this system of records may be disclosed, as a routine use, to another federal agency for purposes of National Security review (35 U.S.C. 181) and for review pursuant to the Atomic Energy Act (42 U.S.C. 218(c)).
- 7. A record from this system of records may be disclosed, as a routine use, to the Administrator, General Services, or his/her designee, during an inspection of records conducted by GSA as part of that agency's responsibility to recommend improvements in records management practices and programs, under authority of 44 U.S.C. 2904 and 2906. Such disclosure shall be made in accordance with the GSA regulations governing inspection of records for this purpose, and any other relevant (i.e., GSA or Commerce) directive. Such disclosure shall not be used to make determinations about individuals.
- 8. A record from this system of records may be disclosed, as a routine use, to the public after either publication of the application pursuant to 35 U.S.C. 122(b) or issuance of a patent pursuant to 35 U.S.C. 151. Further, a record may be disclosed, subject to the limitations of 37 CFR 1.14, as a routine use, to the public if the record was filed in an application which became abandoned or in which the proceedings were terminated and which application is referenced by either a published application, an application open to public inspection or an issued patent.
- 9. A record from this system of records may be disclosed, as a routine use, to a Federal, State, or local law enforcement agency, if the USPTO becomes awars a violation or potential violation of law or regulation.

	Application No.	Applicant	s)		
	12/986,310	BODOR ET	BODOR ET AL.		
Notice of Allowability	Examiner	Art Unit	AIA (First Inventor to File) Status		
	Junaman S. Lau	10/3	No		
The MAILING DATE of this communication appears on the cover sheet with the correspondence address All claims being allowable, PROSECUTION ON THE MERITS IS (OR REMAINS) CLOSED in this application. If not included herewith (or previously mailed), a Notice of Allowance (PTOL-85) or other appropriate communication will be mailed in due course. THIS NOTICE OF ALLOWABILITY IS NOT A GRANT OF PATENT RIGHTS. This application is subject to withdrawal from issue at the initiative of the Office or upon petition by the applicant. See 37 CFR 1.313 and MPEP 1308.					
1. This communication is responsive to <u>See Continuation She</u>	<u>et</u> . s/were filed on				
 2. An election was made by the applicant in response to a res requirement and election have been incorporated into this a 	triction requirement set forth o	during the interview o	n; the restriction		
3. ☑ The allowed claim(s) is/are <u>1-18,20-61,65,66,69,70,73-75 and 77-88</u> . As a result of the allowed claim(s), you may be eligible to benefit from the Patent Prosecution Highway program at a participating intellectual property office for the corresponding application. For more information, please see http://www.uspto.gov/patents/init_events/pph/index.jsp or send an inquiry to PHFeedback@uspto.gov/patents/init_events/pph/index.jsp or send an inquiry to http://www.uspto.gov/patents/init_events/pph/index.jsp or send an inquiry to PHFeedback@uspto.gov/patents/init_events/pph/index.jsp or send an inquiry to PHFeedback@uspto.gov/patents/init_events/pph/index.jsp or send an inquiry to <a a="" abandonment="" application.="" below.="" communication="" comply="" complying="" date"="" extendable.<="" failure="" file="" href="http://www.uspto.gov/patents/init_events/ppi/init_events/ppi/init_events/ppi/init_events/init_events/init_events/ppi/init_eve</td></tr><tr><td>4. Acknowledgment is made of a claim for foreign priority under</td><td>er 35 U.S.C. § 119(a)-(d) or (f</td><td><sup>;</sup>).</td><td></td></tr><tr><td colspan=5>a) All b) Some *c) None of the: 1 Destribute soft the unique to the uni</td></tr><tr><td>2. Certified copies of the priority documents have</td><td colspan=5> Certified copies of the priority documents have been received in Application No. </td></tr><tr><td>3. Copies of the certified copies of the priority do</td><td>cuments have been received</td><td>in this national stage</td><td>application from the</td></tr><tr><td>International Bureau (PCT Rule 17.2(a)).</td><td></td><td></td><td></td></tr><tr><td>* Certified copies not received:</td><td></td><td></td><td></td></tr><tr><th colspan=6>Applicant has THREE MONTHS FROM THE " in="" is="" mailing="" not="" noted="" of="" period="" reply="" requirements="" result="" th="" the="" this="" three-month="" timely="" to="" will="" with="">					
5. CORRECTED DRAWINGS (as "replacement sheets") mus	t be submitted.				
including changes required by the attached Examiner's Amendment / Comment or in the Office action of Paper No./Mail Date					
Identifying indicia such as the application number (see 37 CFR 1.84(c)) should be written on the drawings in the front (not the back) of each sheet. Replacement sheet(s) should be labeled as such in the header according to 37 CFR 1.121(d).					
6. DEPOSIT OF and/or INFORMATION about the deposit of E attached Examiner's comment regarding REQUIREMENT FC	BIOLOGICAL MATERIAL must OR THE DEPOSIT OF BIOLO	st be submitted. Note DGICAL MATERIAL.	the		
Attachment(s)					
1. Notice of References Cited (PTO-892)	5. 🔲 Examiner's	Amendment/Comme	nt		
2. Information Disclosure Statements (PTO/SB/08),	6. 🛛 Examiner's	Statement of Reason	s for Allowance		
 3. Examiner's Comment Regarding Requirement for Deposit of Biological Material 	7. 🗌 Other				
 Interview Summary (PTO-413), Paper No./Mail Date <u>1 pg</u>. 					
	/SHAOJIA ANN	IA JIANG/			
	Supervisory Pat	ent Examiner, Art l	Jnit 1673		

Continuation Sheet (PTOL-37)

Continuation of Item 1. This communication is responsive to : Applicant's Amendment and Remarks, filed 11 Dec 2013, and Applicant's Supplement Amendment and Remarks, filed 20 Feb 2014.

DETAILED ACTION

The present application is being examined under the pre-AIA first to invent provisions.

This Office Action is responsive to Applicant's Amendment and Remarks, filed 11 Dec 2013, and Applicant's Supplement Amendment and Remarks, filed 20 Feb 2014, in which claims 17, 20, 65 and 78 are amended and claims 19, 67, 68, 71, 72 and 76 are canceled.

Applicant's Supplement Amendment and Remarks, filed 20 Feb 2014, will be entered because the amendment is deemed to place the application in condition for allowance.

This application is a domestic application, filed 7 Jan 2011; and claims benefit as a CON of 10/551,205, issued as Patent 7,888,328, which is a 371 of PCT/US2004/009387, filed 26 Mar 2004, which claims benefit of provisional application 60/458,922, filed 28 Mar 2003, and claims benefit of provisional application 60/484,756, filed 2 Jul 2003, and claims benefit of provisional application 60/541,247, filed 4 Feb 2004. The filing date of the instant claims is deemed to be provisional application 60/541,247, filed 4 Feb 2004. Application/Control Number: 12/986,310 Art Unit: 1673

Claims 1-18, 20-61, 65, 66, 69, 70, 73-75 and 77-88 are pending in the current application. Claims 9-18, 20-24, 32-45, 65, 66, 69, 70, 73-75 and 77-88, drawn to non-elected inventions, are rejoined herein. Claims 1-18, 20-61, 65, 66, 69, 70, 73-75 and 77-88 are allowed herein.

Terminal Disclaimer

The terminal disclaimer filed on 10 Dec 2013 disclaiming the terminal portion of any patent granted on this application which would extend beyond the expiration date of US Patent 7,888,328 has been reviewed and is accepted. The terminal disclaimer has been recorded.

Election/Restrictions

Claims 1-8, 25-31 and 46-61 are directed to an allowable product. Pursuant to the procedures set forth in MPEP § 821.04(B), claims 9-18, 20-24, 32-45, 65, 66, 69, 70, 73-75 and 77-88, directed to the process of making or using an allowable product, previously withdrawn from consideration as a result of a restriction requirement, are hereby rejoined and fully examined for patentability under 37 CFR 1.104.

Because all claims previously withdrawn from consideration under 37 CFR 1.142 have been rejoined, **the restriction requirement as set forth in the Office action mailed on 8 Nov 2012 is hereby withdrawn**. In view of the withdrawal of the restriction requirement as to the rejoined inventions, applicant(s) are advised that if any claim presented in a continuation or divisional application is anticipated by, or includes all the Application/Control Number: 12/986,310 Art Unit: 1673

limitations of, a claim that is allowable in the present application, such claim may be subject to provisional statutory and/or nonstatutory double patenting rejections over the claims of the instant application. Once the restriction requirement is withdrawn, the provisions of 35 U.S.C. 121 are no longer applicable. See *In re Ziegler*, 443 F.2d 1211, 1215, 170 USPQ 129, 131-32 (CCPA 1971). See also MPEP § 804.01.

REASONS FOR ALLOWANCE

The following is an examiner's statement of reasons for allowance:

Rejections Withdrawn

Applicant's Remarks, filed 11 Dec 2013, with respect that claims 1-8, 25-31 and 46-61 are rejected under pre-AIA 35 U.S.C. 103(a) as being unpatentable over Vandercruys (US Patent Application Publication 2002/0150616, published 17 Oct 2002, filed 27 May 1998, cited in PTO-892) in view of Schultz et al. (US Patent 6,194,395, published 27 Feb 2001, provided by Applicant in IDS mailed 7 Jan 2011), Baert et al. (WIPO Publication WO97/18839, published 29 May 1997, of record), Redenti et al. (Int. J. Pharm., 1996, 129, p289-294, cited in PTO-892), and Pitha et al. (Life Sci., 1998, 43, p493-502, cited in PTO-892) has been fully considered and is persuasive, as Applicant's remarks are persuasive that Vandercruys requires the essential feature of a physiologically tolerable water-soluble acid which preferably gives a pH 2 (abstract and page 3, paragraph 0028-0029) and that cladribine is known to be acid-labile, for example as a adenosine analog subject to acid-catalyzed glycolysis. While Vandercruys includes cladribine in an extensive list of possible drugs at paragraph 0070 spanning

Application/Control Number: 12/986,310 Art Unit: 1673

pages 4-5, Vandercruys does not disclose a working example of the invention of Vandercruys wherein the sparingly water-soluble drug is an acid-labile drug such as an adenosine analog. Further, Applicant's remarks of record, filed 28 May 2013, suggest unpredictability with regard to specifically the amorphous cladribine-cyclodextrin complex as taught by Shultz and Baert. Therefore applicant's remarks are persuasive that there is no reasonable expectation of success to combine Vandercruys in view of Schultz et al., Baert et al., Redenti et al., and Pitha et al. in a manner to render obvious the instant invention as claimed. The instant invention as claimed is not taught or fairly suggested with a reasonable expectation of success by the closest prior art.

This rejection has been withdrawn.

Applicant's terminal disclaimer, filed 10 Dec 2013, with respect that claims 1-8, 25-31 and 46-61 are rejected on the ground of nonstatutory double patenting over claims 1-28 of U. S. Patent No. 7,888,328 has been fully considered and is persuasive, as the terminal disclaimer has been recorded.

This rejection has been withdrawn.

Any comments considered necessary by applicant must be submitted no later than the payment of the issue fee and, to avoid processing delays, should preferably accompany the issue fee. Such submissions should be clearly labeled "Comments on Statement of Reasons for Allowance."

Conclusion

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Jonathan S. Lau whose telephone number is (571)270-3531. The examiner can normally be reached on Monday - Thursday, 9 am - 4 pm EST.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Shaojia Anna Jiang can be reached on 571-272-0627. The fax phone number for the organization where this application or proceeding is assigned is 571-273-8300.

Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free). If you would like assistance from a USPTO Customer Service Representative or access to the automated information system, call 800-786-9199 (IN USA OR CANADA) or 571-272-1000.

/SHAOJIA ANNA JIANG/ Supervisory Patent Examiner, Art Unit 1673 /Jonathan S Lau/ Examiner, Art Unit 1673
[Application No.	Applicant(a)				
Examiner-Initiated Interview Summary	12/986,310 Examiner	Art Unit				
		1672				
	Jonathan S. Lau	1073				
All participants (applicant, applicant's representative, PTO	personnel):					
(1) <u>Jonathan S. Lau</u> .	(3)					
(2) <u>Mary Katherine Baumeister</u> .	(4)					
Date of Interview: <u>18 February 2014</u> .						
Type: 🛛 Telephonic 🔲 Video Conference 🗌 Personal [copy given to: 🗌 applicant [applicant's representative]					
Exhibit shown or demonstration conducted: Yes If Yes, brief description: <u>none</u> .	X No.					
Issues Discussed \Box 101 \boxtimes 112 \Box 102 \Box 103 \Box Othe (For each of the checked box(es) above, please describe below the issue and detail	PrS ed description of the discussion)					
Claim(s) discussed: <u>17 and 65</u> .						
Identification of prior art discussed: none.						
Substance of Interview (For each issue discussed, provide a detailed description and indicate if agreement reference or a portion thereof, claim interpretation, proposed amendments, argume	was reached. Some topics may include: i ents of any applied references etc)	dentification or clarification of a				
The product claims were indicated as allowable, and rejoin breadth of the claim language of "cladribine-responsive con written description or scope of enablement if rejoined. In orc specific conditions of claims 19, 67 and 68 into claims 17 ar	der of the withdrawn method o dition" in claims 17 and 65 wo der to facilitate allowance of th nd 65 was discussed.	elaims was discussed. The uld likely raise issues of e application, incorporating				
Applicant recordation instructions: It is not necessary for applicant to p	rovide a separate record of the subst	ance of interview.				
Examiner recordation instructions : Examiners must summarize the substance of any interview of record. A complete and proper recordation of the substance of an interview should include the items listed in MPEP 713.04 for complete and proper recordation including the identification of the general thrust of each argument or issue discussed, a general indication of any other pertinent matters discussed regarding patentability and the general results or outcome of the interview, to include an indication as to whether or not agreement was reached on the issues raised.						
Attachment						
U.S. Patent and Trademark Office PTOL-413B (Rev. 8/11/2010) Interview 39	Summary 7	Paper No. 20140225				

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FIRST INFORMATION DISCLOSURE STATEMENT BY APPLICANT

(use as many sheets as necessary)

Sheet 1 of 4

Examiner

Complete if KnownApplication Number12/986310Filing DateJanuary 5, 2011First Named InventorNicholas S. BodorExaminer Name0056192-000067

U.S. PATENT DOCUMENTS

Examiner Initials	Document Number- Kind Code	Publication Date MM-DD-YYYY	Name of Patentee or Applicant of Cited Document	Pages, Columns, Lines Where Relevant Passages or Figures Appear
	US-4,383,992	05-17-1983	Lipari	
	US-6,239,118 B1	05-29-2001	Shatz et al.	
	US-5,424,296	06-13-1995	Saven et al.	
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	US-6,699,849	03-02-2004	Loftsson et al.	

	FOREIGN PATENT DOCUMENTS										
	Foreign Patent Document			STATUS							
Examiner Initials	Country Code ¹ , Number, Kind Code	Publication Date (MM-DD-YYYY)	Name of Patentee or Applicant of Cited Document	Translation	Partial Translation	Eng. Lang. Summary	Search Report	IPER	Abstract	Cited Pç	in Spec. / . No(s).
	EP 0 197 571 A2	10-15-1986	Janssen Pharmaceutica N.V.							x	
	WO 90/12035A1	10-18-1990	Janssen Pharmaceutica N.V.							x	
	DE 31 18 218 A1	04-22-1982	Chinoin Gyogyszer Es Vegyeszet						x	x	
	DE 33 17 064 A1	11-15-1984	Consortium Elektrochem Ind						x	x	
	GB 2 189 245 A	10-21-1987	American Maize- Products Company							x	
	EP 0 149 197 B1	07-24-1985	Janssen Pharmaceutica N.V.	Х						x	
	EP 0 094 157 A1	11-16-1983	Janssen Pharmaceutica N.V.							x	
	WO 99/42111	08-26-1999	Cyclops, ehf								

 Signature
 Considered

 *EXAMINER: Initial if reference considered, whether or not citation is in conformance with M.P.E.P. § 609. Draw line through citation if not in conformance and not considered. Include copy of this form with next communication to Applicant.

Date

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Application Number	12/986310		
Filing Date	January 5, 2011		
First Named Inventor	Nicholas S. Bodor		
Examiner Name			
Attorney Docket No.	0056192-000067		

Sheet 2 of 4

FOREIGN PATENT DOCUMENTS

¹Enter Office that issued the document, by the two-letter code.

	NON-PATENT LITERATURE DOCUMENTS
Examiner Initials	Include name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of the item (book, magazine, journal, serial, symposium, catalog, etc.), date, page(s), volume-issue number(s), publisher, city and/or country where published.
	TARASIUK et al., "Stability of 2-Chloro-2'-Deoxyadenosine at Various pH and Temperature",
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	Birkhauser Publishers Ltd., Basel, Switzerland
	ROMINE et al., "A Double-Blind, Placebo-Controlled, Randomized Trial of Cladribine in Relapsing-
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Examiner		Date					
Signature		Considered					
*EXAMINER: Initial if reference considered, whether or not citation is in conformance with M.P.E.P. § 609. Draw line through citation if not in							
conformance	conformance and not considered. Include copy of this form with next communication to Applicant.						

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Complete if Known			
Application Number	12/986310		
Filing Date	January 5, 2011		
First Named Inventor	Nicholas S. Bodor		
Examiner Name			
Attorney Docket No.	0056192-000067		

Sheet 3 of 4

NON-PATENT LITERATURE DOCUMENTS

Examiner Initials	Include name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of the item (book, magazine, journal, serial, symposium, catalog, etc.), date, page(s), volume-issue number(s), publisher, city and/or country where published.
	BAKTHIAR et al., "A study of the complexation between dimethyl-β-cyclodextrin and steroid
	hormones using electrospray ionization mass spectrometry", Rapid Communications in Mass
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Examiner		Date	
Signature		Considered	
*EXAMINER	Initial if reference considered, whether or not citation is in co	nformance with M.P.E.I	P. § 609. Draw line through citation if not in

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Application Number	12/986310		
Filing Date	January 5, 2011		
First Named Inventor	Nicholas S. Bodor		
Examiner Name			
Attorney Docket No.	0056192-000067		

Sheet	4	of	4	
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SUZUKI et al., "A Study of 1:1 Plus 1:2 Complexes Between Barbiturate and ά-Cyclodextrin Using the Freezing Point Depression Method," <i>Chem. Pharm. Bull.</i> , 1993, 41(8), P1444-1447

OTHER
International Search Report dated October 12, 2004 for PCT/US2004/009387, filed March 26, 2004
PCT International Preliminary Report on Patentability and Written Opinion for International Application No. PCT/US2004/009387, International Filing date March 26, 2004.

Examiner Signature	/Jonathan Lau/	Date Considered	03/06/2014							
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	Application/Control No.	Applicant(s)/Patent Under Reexamination
Issue Classification	12986310	BODOR ET AL.
	Examiner	Art Unit
	JONATHAN S LAU	1673

CPC			
Symbol		Туре	Version

CPC Combination Sets											
Symbol	Туре	Set	Ranking	Version							

/JONATHAN S LAU/ Examiner.Art Unit 1673	02/25/2014	Total Claims Allowed:				
(Assistant Examiner)	(Date)	7	9			
/SHAOJIA ANNA JIANG/ Supervisory Patent Examiner, Art Unit 1673		O.G. Print Claim(s)	O.G. Print Figure			
(Primary Examiner)	(Date)	1	none			

U.S. Patent and Trademark Office

	Application/Control No.	Applicant(s)/Patent Under Reexamination
Issue Classification	12986310	BODOR ET AL.
	Examiner	Art Unit
	JONATHAN S LAU	1673

US ORIGINAL CLASSIFICATION					INTERNATIONAL CLASSIFICATION									
	CLASS		ę	SUBCLASS		CLAIMED NON-CLAIMED								N-CLAIMED
514			46			А	6	1	к	31 / 7076 (2006.01.01)				
CROSS REFERENCE(S)					A	6	1	к	47 / 40 (2006.01.01)					
CLASS	SUB	CLASS (ONE	SUBCLAS	S PER BLO	CK)									
514	58													

/JONATHAN S LAU/ Examiner.Art Unit 1673	02/25/2014	Total Claims Allowed:			
(Assistant Examiner)	(Date)	7	9		
/SHAOJIA ANNA JIANG/ Supervisory Patent Examiner, Art Unit 1673		O.G. Print Claim(s)	O.G. Print Figure		
(Primary Examiner)	(Date)	1	none		

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Part of Paper No. 20140225

	Application/Control No.	Applicant(s)/Patent Under Reexamination
Issue Classification	12986310	BODOR ET AL.
	Examiner	Art Unit
	JONATHAN S LAU	1673

	Claims renumbered in the same order as presented by applicant								СР	A D] Т.D.	[R.1. 4	47	
Final	Original	Final	Original	Final	Original	Final	Original	Final	Original	Final	Original	Final	Original	Final	Original
22	1	54	17	9	33	37	49	61	65	77	81				
23	2	55	18	10	34	33	50	62	66	74	82				
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25	4	56	20	12	36	38	52		68	75	84				
26	5	57	21	13	37	39	53	64	69	69	85				
27	6	58	22	14	38	40	54	63	70	79	86				
28	7	59	23	15	39	35	55		71	76	87				
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49	12	4	28	20	44	44	60		76						
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52	15	7	31	31	47		63	72	79						
53	16	8	32	32	48		64	73	80						

/JONATHAN S LAU/ Examiner.Art Unit 1673	02/25/2014	Total Claims Allowed:	
(Assistant Examiner)	(Date)	7	9
/SHAOJIA ANNA JIANG/ Supervisory Patent Examiner, Art Unit 1673		O.G. Print Claim(s)	O.G. Print Figure
(Primary Examiner)	(Date)	1	none

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Part of Paper No. 20140225

EAST Search History

EAST Search History (Prior Art)

Ref #	Hits	Search Query	DBs	Default Operator	Plurals	Time Stamp
L1	1463	514/46.ccls.	US-PGPUB; USPAT; USOCR	NEAR	ON	2014/02/25 15:47
L2	1423	514/58.ccls.	US-PGPUB; USPAT; USOCR	NEAR	ON	2014/02/25 15:48
L3	56	2 and cladribine	US-PGPUB; USPAT; USOCR	NEAR	ON	2014/02/25 15:48
L4	91	1 and cladribine	US-PGPUB; USPAT; USOCR	NEAR	ON	2014/02/25 15:48
L5	58	4 and (cyclodextrin or complex)	US-PGPUB; USPAT; USOCR	NEAR	ON	2014/02/25 15:48
L6	1692	a61k31/7076.cpc.	US-PGPUB; USPAT; USOCR	NEAR	ON	2014/02/25 15:50
L7	1416	a61k47/40.cpc. or a61k47/48969.cpc.	US-PGPUB; USPAT; USOCR	NEAR	ON	2014/02/25 15:52
L8	19	6 and 7	US-PGPUB; USPAT; USOCR	NEAR	ON	2014/02/25 15:52
L9	49	3 and amorphous	US-PGPUB; USPAT; USOCR	NEAR	ON	2014/02/25 15:52
L10	12	5 and amorphous	US-PGPUB; USPAT; USOCR	NEAR	ON	2014/02/25 15:52
L13	36	cladribine.ti,ab.	US-PGPUB; USPAT; USOCR	NEAR	ON	2014/02/25 15:57
L14	15	13 AND ((A61K31/7076 OR A61K31/724 OR A61K47/40 OR A61K9/205).CPC.)	US-PGPUB; USPAT; USOCR	NEAR	ON	2014/02/25 15:58
L15	11	A61K31/7076.cpc. and (A61K31/724 OR A61K47/40 OR A61K9/205).CPC.	US-PGPUB; USPAT; USOCR	NEAR	ON	2014/02/25 15:59

EAST Search History (Interference)

Ref #	Hits	Search Query	DBs	Default Ope	erator	Plurals	Time Stamp
L11	0	514/58.ccls.	UPAD	NEAR		ON	2014/02/25 15:56
L12	1	514/46.ccls.	UPAD	NEAR		ON	2014/02/25 15:56

2/25/2014 4:00:01 PM

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 $\label{eq:c:loss} C: \ Users \ jlau1 \ Documents \ EAST \ Workspaces \ 12986310 \ - \ cladribine \ cd \ complex.wsp$

Attorney Docket No. 20009904-0067 Application No. 12/986,310 Page 2

AMENDMENTS TO THE CLAIMS:

This listing of claims will replace all prior versions, and listings, of claims in the application:

LISTING OF CLAIMS:

1. (Original) A pharmaceutical composition comprising a complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex, formulated into a solid oral dosage form, said composition comprising no significant amount of free crystalline cladribine therein.

2. (Original) The pharmaceutical composition according to Claim 1, wherein the complex is saturated with cladribine.

3. (Original) The composition according to Claim 1, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin, hydroxypropyl- γ -cyclodextrin, randomly methylated β -cyclodextrin, carboxymethyl- β -cyclodextrin or sulfobutyl- β -cyclodextrin.

4. (Original) The composition according to Claim 1, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin or hydroxypropyl-γ-cyclodextrin.

5. (Original) The composition according to Claim 1, wherein the weight ratio of cladribine to amorphous cyclodextrin is from about 1:10 to about 1:16.

6. (Original) The composition according to Claim 5, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin or hydroxypropyl- γ -cyclodextrin.

7. (Original) The composition according to Claim 1, wherein the approximate molar ratio of cladribine to amorphous cyclodextrin corresponds to a point located on the curve of a phase solubility diagram for saturated complexes of cladribine in varying concentrations of the cyclodextrin.

8. (Original) The composition according to Claim 1, wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b).

9. (Withdrawn) A method for enhancing the oral bioavailability of cladribine comprising orally administering to a subject in need thereof a pharmaceutical composition comprising a complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous

cyclodextrin as a non-inclusion complex, formulated into a solid oral dosage form, said composition comprising no significant amount of free crystalline cladribine therein.

10. (Withdrawn) The method according to Claim 9, wherein the complex is saturated with cladribine.

11. (Withdrawn) The method according to Claim 9, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin, hydroxypropyl- γ -cyclodextrin, randomly methylated β -cyclodextrin, carboxymethyl- β -cyclodextrin or sulfobutyl- β -cyclodextrin.

12. (Withdrawn) The method according to Claim 9, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin or hydroxypropyl-γ-cyclodextrin.

13. (Withdrawn) The method according to Claim 9, wherein the weight ratio of cladribine to amorphous cyclodextrin is from about 1:10 to about 1:16.

14. (Withdrawn) The method according to Claim 13, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin or hydroxypropyl-γ-cyclodextrin.

15. (Withdrawn) The method according to Claim 9, wherein the approximate molar ratio of cladribine to amorphous cyclodextrin corresponds to a point located on

the curve of a phase solubility diagram for saturated complexes of cladribine in varying concentrations of the cyclodextrin.

16. (Withdrawn) The method according to Claim 9, wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b).

17. (Withdrawn-Currently Amended) A method for the treatment of symptoms of a cladribine-responsive condition <u>selected from the group consisting of multiple</u> <u>sclerosis, rheumatoid arthritis and leukemia</u> in a subject suffering from said symptoms comprising orally administering to said subject a pharmaceutical composition comprising a complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex, formulated into a solid oral dosage form, said composition comprising no significant amount of free crystalline cladribine therein.

18. (Withdrawn) The method according to Claim 17, wherein the complex is saturated with cladribine.

19. (Canceled)

20. (Withdrawn-Currently Amended) The method according to Claim 19<u>17</u>, wherein the cladribine-responsive condition is multiple sclerosis.

21. (Withdrawn) The method according to Claim 17, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin, hydroxypropyl- γ -cyclodextrin, randomly methylated β -cyclodextrin, carboxymethyl- β -cyclodextrin or sulfobutyl- β -cyclodextrin.

22. (Withdrawn) The method according to Claim 17, wherein the weight ratio of cladribine to amorphous cyclodextrin is from about 1:10 to about 1:16.

23. (Withdrawn) The method according to Claim 17, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin or hydroxypropyl-γ-cyclodextrin.

24. (Withdrawn) The method according to Claim 17, wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b).

25. (Original) A complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin and (b) amorphous free cladribine associated with amorphous cyclodextrin as a non-inclusion complex.

26. (Original) The complex according to Claim 25, saturated with cladribine.

27. (Original) The complex according to Claim 25, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin, hydroxypropyl- γ -cyclodextrin, randomly methylated β -cyclodextrin, carboxymethyl- β -cyclodextrin or sulfobutyl- β -cyclodextrin.

28. (Original) The complex according to Claim 25, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin or hydroxypropyl-γ-cyclodextrin.

29. (Original) The complex according to Claim 25, wherein the weight ratio of cladribine to amorphous cyclodextrin is from about 1:10 to about 1:16.

30. (Original) The complex according to Claim 29, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin or hydroxypropyl-γ-cyclodextrin.

31. (Original) The complex according to Claim 25, wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b).

32. (Withdrawn) A process for the preparation of a complex cladribinecyclodextrin complex as claimed in Claim 25, which comprises the steps of:

(i) combining cladribine and an amorphous cyclodextrin in water at a temperature of from about 40 to about 80°C and maintaining said temperature for a period of from about 6 to about 24 hours;

(ii) cooling the resultant aqueous solution to room temperature; and

(iii) lyophilizing the cooled solution to afford an amorphous product.

33. (Withdrawn) The process according to Claim 32, further comprising a filtration step following step (ii).

34. (Withdrawn) The process according to Claim 32, wherein step (i) is performed at a temperature of from about 45 to about 60°C.

35. (Withdrawn) The process according to Claim 32, wherein step (i) is performed at a temperature of from about 45 to about 50°C.

36. (Withdrawn) The process according to Claim 34, wherein step (i) is performed with stirring.

37. (Withdrawn) The process according to Claim 36, wherein step (i) is performed for a period of from about 6 to about 9 hours.

38. (Withdrawn) The process according to Claim 32, wherein step (ii) is performed for a period of from about 6 to about 9 hours.

39. (Withdrawn) The process according to Claim 32, wherein step (iii) comprises an initial freezing stage in which the solution is cooled to from about -40 to about -80° C, and held at said temperature for a period of from about 2 to about 4 hours.

40. (Withdrawn) The process according to Claim 39, wherein, in the initial freezing stage of step (iii), the solution is cooled to about -45°C.

41. (Withdrawn) The process according to Claim 34, wherein 12.00 parts by weight of cladribine and 172.50 parts by weight of hydroxypropyl- β -cyclodextrin are introduced in step (i), or wherein 16.35 parts by weight of cladribine and 172.50 parts by weight of hydroxypropyl- β -cyclodextrin are introduced in step (i).

42. (Withdrawn) The process according to Claim 41, wherein 825 parts by volume of water are introduced in step (i).

43. (Withdrawn) The process according to Claim 32, wherein the lyophilization step (iii) comprises:

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(a) an initial freezing stage in which the complexation solution is brought to
 from about -40°C to about -80°C for approximately 2 to 4 hours;

(b) a primary drying stage at about -25°C for approximately 80 to 90 hours; and

(c) a secondary drying stage at about 30°C for approximately 15 to 20 hours.

44. (Withdrawn) The process according to Claim 43, wherein stage (a) of the lyophilization is conducted at about -45°C for approximately 3 to 4 hours.

45. (Withdrawn) The process according to Claim 43, wherein stage (b) of the lyophilization is conducted under a pressure of about 100 mTorr.

46. (Original) A pharmaceutical composition according to Claim 1, obtainable by a process comprising the steps of:

 (i) combining cladribine and an amorphous cyclodextrin in water at a temperature of from about 40 to about 80°C and maintaining said temperature for a period of from about 6 to about 24 hours;

(ii) cooling the resultant aqueous solution to room temperature;

(iii) lyophilizing the cooled solution to afford an amorphous product; and

(iv) formulating the amorphous product into a solid oral dosage form.

47. (Original) The pharmaceutical composition according to Claim 46, wherein the process further comprises a filtration step following step (i) or (ii).

48. (Original) The pharmaceutical composition according to Claim 46, wherein step (i) of the process is performed at a temperature of from about 45 to about 60°C.

49. (Original) The pharmaceutical composition according to Claim 46, wherein step (i) of the process is performed at a temperature of from about 45 to about 50°C.

50. (Original) The pharmaceutical composition according to Claim 48, wherein step (i) of the process is performed with stirring.

51. (Original) The pharmaceutical composition according to Claim 50, wherein step (i) of the process is performed for a period of from about 6 to about 9 hours.

52. (Original) The pharmaceutical composition according to Claim 46, wherein step (ii) of the process is performed for a period of from about 6 to about 9 hours.

53. (Original) The pharmaceutical composition according Claim 46, wherein step (iii) comprises an initial freezing stage in which the solution is cooled to from about -40 to about -80°C, and held at said temperature for a period of from about 2 to about 4 hours.

54. (Original) The pharmaceutical composition according to Claim 53, wherein, in the initial freezing stage of step (iii), the solution is cooled to about -45°C.

55. (Original) The pharmaceutical composition according to Claim 48, wherein 12.00 parts by weight of cladribine and 172.50 parts by weight of the hydroxypropyl-β-cyclodextrin are introduced in step (i) of the process, or wherein 16.35 parts by weight of cladribine and 172.50 parts by weight of the hydroxypropyl-β-cyclodextrin are introduced in step (i) of the process.

56. (Original) The pharmaceutical composition according to Claim 55, wherein 825 parts by volume of water are introduced in step (i) of the process.

57. (Original) The pharmaceutical composition according to Claim 46, wherein the lyophilization step (iii) of the process comprises:

(a) an initial freezing stage in which the complexation solution is brought to
 from about -40°C to about -80°C for approximately 2 to 4 hours;

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(b) a primary drying stage at about -25°C for approximately 80 to 90 hours; and

(c) a secondary drying stage at about 30°C for approximately 15 to 20 hours.

58. (Original) The pharmaceutical composition according to Claim 57, wherein stage (a) of the lyophilization is conducted at about -45°C for approximately 3 to 4 hours.

59. (Original) The pharmaceutical composition according to Claim 57, wherein stage (b) of the lyophilization is conducted under a pressure of about 100 mTorr.

60. (Original) The pharmaceutical composition according to Claim 46, wherein the formulation step (iv) of the process comprises blending the complex with magnesium stearate and compressing into tablets.

61. (Original) The pharmaceutical composition according to Claim 60, wherein magnesium stearate is pre-mixed with sorbitol powder before blending with the complex.

62. - 64. (Canceled)

65. (Withdrawn-Currently Amended) A method for the treatment of symptoms of a cladribine-responsive condition <u>selected from the group consisting of multiple</u> <u>sclerosis, rheumatoid arthritis and leukemia</u> in a subject suffering from said symptoms comprising orally administering to said subject a pharmaceutical composition comprising a complex cladribine-cyclodextrin complex which is an intimate amorphous admixture of (a) an amorphous inclusion complex of cladribine with an amorphous cyclodextrin as a non-inclusion complex, formulated into a solid oral dosage form, said composition comprising no significant amount of free crystalline cladribine therein, wherein administering cladribine is accompanied by administering one or more additional active ingredients for treating the cladribine-responsive condition.

66. (Withdrawn) The method according to Claim 65, wherein the complex is saturated with cladribine.

67. - 68. (Canceled)

69. (Withdrawn) The method according to Claim 65, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin, hydroxypropyl- γ -cyclodextrin, randomly methylated β -cyclodextrin, carboxymethyl- β -cyclodextrin or sulfobutyl- β -cyclodextrin.

70. (Withdrawn) The method according to Claim 66, wherein the amorphous cyclodextrin is hydroxypropyl- β -cyclodextrin, hydroxypropyl- γ -cyclodextrin, randomly methylated β -cyclodextrin, carboxymethyl- β -cyclodextrin or sulfobutyl- β -cyclodextrin.

71. – 72. (Canceled)

73. (Withdrawn) The method according to Claim 65, wherein the weight ratio of cladribine to amorphous cyclodextrin is from about 1:10 to about 1:16.

74. (Withdrawn) The method according to Claim 69, wherein the weight ratio of cladribine to amorphous cyclodextrin is from about 1:10 to about 1:16.

75. (Withdrawn) The method according to Claim 65, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin.

76. (Canceled)

77. (Withdrawn) The method according to Claim 73, wherein the amorphous cyclodextrin is hydroxypropyl-β-cyclodextrin.

78. (Withdrawn-Currently Amended) The method according to Claim 73_{74} , wherein the amorphous cyclodextrin is hydroxypropyl- γ -cyclodextrin.

79. (Withdrawn) The method according to Claim 75, wherein the weight ratio of cladribine to hydroxypropyl- β -cyclodextrin is about 1:14.

80. (Withdrawn) The method according to Claim 75, wherein the weight ratio of cladribine to hydroxypropyl- β -cyclodextrin is about 1:11.

81. (Withdrawn) The method according to Claim 65, wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b).

82. (Withdrawn) The method according to Claim 75, wherein from about 30 to about 40 percent by weight of the cladribine is in the inclusion complex (a) and from about 70 to about 60 percent by weight of the cladribine is in the non-inclusion complex (b).

83. (Withdrawn) The method according to Claim 65, wherein the cladribineresponsive condition is multiple sclerosis.

84. (Withdrawn) The method according to Claim 75, wherein the cladribineresponsive condition is multiple sclerosis.

85. (Withdrawn) The method according to Claim 77, wherein the cladribineresponsive condition is multiple sclerosis.

86. (Withdrawn) The method according to Claim 83, wherein one or more additional active ingredients for treating multiple sclerosis is/are selected from the group consisting of interferon beta, glatiramer acetate, natalizumab, alemtuzumab, 4-aminopyridine and amantadine.

87. (Withdrawn) The method according to Claim 84, wherein one or more additional active ingredients for treating multiple sclerosis is/are selected from the group consisting of interferon beta, glatiramer acetate, natalizumab, alemtuzumab, 4- aminopyridine and amantadine.

88. (Withdrawn) The method according to Claim 85, wherein one or more additional active ingredients for treating multiple sclerosis is/are selected from the group consisting of interferon beta, glatiramer acetate, natalizumab, alemtuzumab, 4- aminopyridine and amantadine.

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Case law My library ^{New1}	ligand, anxiogenic agent 1.1 antipsychotic doparnine antagonist 10 adenosine antagonist, antiasthmatic are often bi-phasic, that is, 502 Solubilization by Amorphous Cyclodextrins Vol above mixtures contain hundreds of isomeric and enantiomeric cyclodextrin derivatives there Cited by 207 Related articles. All 4 versions. Cite. Save	
Any time Since 2014 Since 2013 Since 2010 Custom range	Pharmaceutical preparations containing cyclodextrin derivatives J Pitha - US Patent 4,727,064, 1989 - Google Patents a stablizing amorphous complex of a drug and a mixture of cyclodextrins which comprises the steps of: 1. Dissolving an intrinsically amorphous mixture of cyclodextrin deratives which are water soluble and capable of forming inclusion complexes with drugs in water; and Cited by 239 Related articles All 2 versions Cite Save	
- 2004	Pharmaceutical compositions containing drugs which are instable or sparingly soluble in water and methods for their preparation SW Müller, U Brauns - US Patent 6,407,079, 2002 - Google Patents Mar 26, 2004, Jan 12, 2011, Ares Trading SA, Oral formulations of cladribine Daniel B. Carr, Formulations of low dose non-steroidal anti-inflammatory drugs and beta-cyclodextrin.	
Sort by relevance Sort by date	WO200808681641 ", Jan 16, 2008, Jul 24, 2008, Adenosine Therapeutics, Lic, Adenosine Cited by 41 Related articles All 2 versions Cite Save	
include patents	In vivo and In vitro Correlation for Delayed release Behaviour of a Molsidomine/O carboxymethyl O ethyl <u>B</u> cyclodextrin Complex in Gastric Acidity controlled T HORIKAWA, F HIRAYAMA Journal of pharmacy and, 1995 - Wiley Online Library	
‱ Create alert	Omeprazole, an H+, K+- adenosine triphosphatase inhibi- tor, was donated by Yoshitomi Pharmaceutical Ind of I-hexylcarbamoyI-5-fluorouracil (HCFU) by 0-carboxymethyI-0-ethyI-P- cyclodextrin Miiller, B. W., Brauns, U. (1985) Solubilization of drugs by modified P- cyclodextrins Cited by 22 Related articles All 4 versions Cite Save	
	Pharmaceutical applications of cyclodextrins. 1. Drug solubilization and stabilization <u>Licitation</u> , ME Brewster - Journal of pharmaceutical sciences, 1996 - Wiley Online Library β-CD Adenosine 7 -21 -53 32 been moved away from the cavity by a butyl ether spacer group, is an excellent solubilizer.71 (Carboxymethyl)-β-cyclodextrin is another in- teresting anionic cyclodextrin derivative.72 Compared to neutral cyclodextrins, enhanced complexation Cited by 1534 Related articles All 6 versions Cite Save	[PDF] from researchgate.net
	Subject Index to Volume 18 (2001) AN Drug, A Adenosine - Pharmaceutical Research, 2001 - Springer Crystallization, macromolecular, 1483 Crystallization, supercritical fluid, 852 Cyclic adenosine monophosphate (cAMP), 1651 Cyclic voltammetry, 702 Cyclodextrins, 667, 886 1455 Hydrophobic drugs, 510 Hydrophobicity, 104 Hydropropyl cyclodext#P3667 Hydroxyethyl	

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Method and compositions for solubilization and stabilization of polypeptides, especially proteins MS Hora, J Rubinfeld, W Stern, GJ Wong - US Patent 5,730,969, 1998 - Google Patents ... 2) in increasing concentrations (percent w/v) of hydroxypropyl-β-cyclodextrin (HPBCD ... affect their biological function, aqueous solubility and ability to interact with the cyclodextrins. ... of proteins and immobilized proteases), cell surface recognition proteins (**adenosine** and analogs ... Cited by 28 Related articles All 5 versions Cite Save

Role of cyclic adenosine 3', 5'-monophosphate and serum albumin in head-to-head agglutination

of boar spermatozoa

H Harayama, M Miyake, S Kato - Reproduction, Fertility and Development, 2001 - CSIRO ... mKRB, modified Krebs-Ringer bicarbonate; mKRB-P, mKRB containing 0.1% polyvinyl alcohol; BSA, bovine serum albumin; dbcAMP, dibutyryl cyclic **adenosine** 3',5'-monophosphate; MBC, methyl-β-**cyclodextrin**. Page 10. Mechanism of sperm agglutination 315 ... Cited by 14 Related articles All 7 versions Cite Save

Improvement of gliquidone hypoglycaemic effect in rats by cyclodextrin formulations A Miro, <u>E Quaglia</u>, U Sorrentino, MI La Rotonda... - European journal of ..., 2004 - Elsevier ... These compounds display similar chemical features and seem to act by blocking the pancreatic islet β-cell **adenosine** 5'triphosphate (ATP ... Pharmaceutical applications of cyclodextrins. ... Physico-chemical and pharmacological properties of nimesulide/β-cyclodextrin formulations. ...

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Cyclodextrin complexes of benzodiazepines

<u>I Lottsson. M Masson. E Stefansson</u> - US Patent 6,699,849, 2004 - Google Patents ... isomeric products, chemical modification can transform the crystalline **cyclodextrins** into **amorphous** mixtures increasing ... In aqueous solutions, **cyclodextrins** form complexes with many drugs through a process ... Once included in the **cyclodextrin** cavity, the drug molecules may be ... Cited by 8 Related articles All 2 versions Cite Save

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Search Notes	12986310	BODOR ET AL.
	Examiner	Art Unit
	JONATHAN S LAU	1623

CPC- SEARCHED		
Symbol	Date	Examiner
A61K 31/7076	2/25/2014	JSL
A61K 47/40, 47/48969	2/25/2014	JSL

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Symbol	Date	Examiner

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Class	Subclass	Date	Examiner
514	46, 58	2/25/2014	JSL

SEARCH NOTES		
Search Notes	Date	Examiner
EAST - inventor name search (Nicholas Bodor, Yogesh Dandiker)	2/15/2013	JSL
EAST - see attached notes	2/15/2013	JSL
Google Scholar - see attached notes	2/15/2013	JSL
Google Scholar - see attached notes	9/3/2013	JSL
EAST - inventor name search (Nicholas Bodor, Yogesh Dandiker) - updated	2/25/2014	JSL
Review prosecution history of parent case 10/551205	2/25/2014	JSL
Google Scholar - see attached notes	2/25/2014	JSL

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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.	
12/986,310	01/07/2011	Nicholas S. Bodor	20009904-0067	6100	
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P.O. BOX 0610	80		LAU, JONATHAN S		
Chicago, IL 606	06-1080		ART UNIT	PAPER NUMBER	
			1673		
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			04/17/2014	ELECTRONIC	

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The time period for reply, if any, is set in the attached communication.

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martin.bruehs@dentons.com patents.us@dentons.com



Commissioner for Patents United States Patent and Trademark Office P.O. Box 1450 Alexandria, VA 22313-1450 www.uspto.gov

Application No. : 12986310 Applicant : Bodor Filing Date : 01/07/2011 Date Mailed : 04/17/2014

NOTICE TO FILE CORRECTED APPLICATION PAPERS

Notice of Allowance Mailed

This application has been accorded an Allowance Date and is being prepared for issuance. The application, however, is incomplete for the reasons below.

Applicant is given two (2) months from the mail date of this Notice within which to respond. This time period for reply is extendable under 37 CFR 1.136(a) for only TWO additional MONTHS.

The informalities requiring correction are indicated in the attachment(s). If the informality pertains to the abstract, specification (including claims) or drawings, the informality must be corrected with an amendment in compliance with 37 CFR 1.121 (or, if the application is a reissue application, 37 CFR 1.173). Such an amendment may be filed after payment of the issue fee if limited to correction of informalities noted herein. See Waiver of 37 CFR 1.312 for Documents Required by the Office of Patent Publication, 1280 Off. Gaz. Patent Office 918 (March 23, 2004). In addition, if the informality is not corrected until after payment of the issue fee, for purposes of 35 U.S.C. 154(b)(1)(iv), "all outstanding requirements" will be considered to have been satisfied when the informality has been corrected. A failure to respond within the above-identified time period will result in the application being ABANDONED.

See attachment(s).

A copy of this notice <u>MUST</u> be returned with the reply. Please address response to "Mail Stop Issue Fee, Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450".

/Lois Stone/ Publication Branch Office of Data Management (571) 272-4200

Application No. <u>12986310</u>

	IDENTIFICATION OF APPLICATION DEFICIENCIES
	Applicant must provide legible text for the following item(s).
	Specification filed , page(s) .
	Claims filed, claim(s).
	Oath/declaration filed .
	Other: .
	Applicant must provide missing information on the following page(s) of the specification by amending the specification to add the missing text. No new matter may be added.
X	The specification refers to one or more applications by attorney docket number and does not show the U.S. application number(s). Applicant must supply the U.S. application number in place of each attorney docket number. <u>page 23</u> , lines 25 and 27
	Applicant must provide an Abstract of the Disclosure.
	Applicant has submitted a DECLARATION (37 CFR 1.63) FOR A UTILITY OR DESIGN APPLICATION USING AN APPLICATION DATA SHEET (37 CFR 1.76) (e.g., form PTO/SB/01A). The Application Data Sheet, however, is not present with the filed application. Applicant must submit an Application Data Sheet or file a new oath or declaration (e.g., PTO/SB/01) executed by the inventors and containing the information required in 37 CFR 1.63.
	Applicant must provide an executed declaration.
	Applicant must provide the missing page(s) of the oath/declaration or Application Data Sheet filed
	Applicant must provide a declaration signed by inventor(s).
	The oath/declaration filed shows non-initialed and/or non-dated alterations. Applicant must file a new oath/declaration in compliance with 37 CFR 1.67(a).
	Applicant(s) in the latest-filed oath/declaration or Application Data Sheet (ADS) did not show the inventor's residence at all, or did not show both a city and state in the U.S. inventor's residence, or did not show both a city and country in the non-U.S. inventor's residence. Applicant must supply an oath/declaration or Application Data Sheet (ADS) that shows each U.S. inventor's city and state of

residence and each non-U.S. inventor's city and country of residence.

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applic	ant: Nicholas S. BODOR et al.)	MAIL STOP ISSUE FEE
Applic	ation No.: 12/986,310)	Examiner: Jonathan S. LAU
Filed:	January 7, 2011)))	Group Art Unit: 1623
Title:	ORAL FORMULATIONS OF CLADRIBINE))	Confirmation No.: 6100

REPLY AND AMENDMENT PURSUANT TO 37 C.F.R. § 1.312

In response to the Notice to File Corrected Application Papers dated April 17, 2014, Applicants hereby submit the following amendment and remarks pursuant to 37 C.F.R. §1.312.

Attorney Docket No. 20009904-0067 Application No. 12/986,310 Page 2

AMENDMENTS TO THE SPECIFICATION:

Please replace page 23 of the as-filed specification with the attached new page 23, which deleted the final sentence in the paragraph at lines 7-29.

Attached: Marked-Up and Clean Copies of Page 23

REMARKS

After allowance of this application, it was noted that page 23 of the as-filed specification contained references to applications for which no application numbers were given. These were provisional applications which were abandoned without the filing of non-provisional applications based thereon. Further, they were not for inventions of the present inventors and belonged to a former assignee. They were not made available to the public. In the parent case, now patented (Application No. 12/986, 310), the sentence in question was deleted during prosecution, by an amendment made October 3, 2008. Accordingly, page 23 of the specification has been amended to delete the final sentence on page 23, consistent with the parent.

This amendment was not proposed sooner because it was not realized that the error had not been corrected until after allowance. This amendment does not raise any new issues and thus is appropriate at this time. Entry is respectfully requested.

Respectfully submitted,

DENTONS US LLP

Date: June 12, 2014

<u>/Mary Katherine Baumeister/</u> Mary Katherine Baumeister Registration No. 26254

Customer No. 13974 Dentons US LLP 1301 K Street NW, Suite 600, East Tower Washington, D.C. 20005 Phone: 202-408-9186 Fax: 202-408-6399

-23-

Physicians, Vol. 111, No. 1, 35-44 (1999); Selby et al., The Canadian Journal of Neurological Sciences, 25, 295-299 (1998); Tortorella et al., Current Opinion in Investigational Drugs, 2 (12), 1751-1756 (2001); Rice et al., Neurology, 54, 1145-1155 (2000); and Karlsson et al., British Journal of Haematology, 116, 538-548 (2002); all of which are incorporated by reference herein in their entireties and relied upon.

Moreover, the route of administration for which the therapeutically effective dosages are taught in the literature should be taken into consideration. While the instant compositions optimize the bioavailability of cladribine following oral administration, it will be appreciated that even optimal bioavailability from oral dosage forms is not expected to approach bioavailability obtain after intravenous administration, particularly at early time points. Thus, it is often appropriate to increase a dosage suggested for intravenous administration to arrive at a suitable dosage for incorporation

- into a solid oral dosage form. At the present time, it is envisioned that, for 15 the treatment of multiple sclerosis, 10 mg of cladribine in the instant complex cladribine-cyclodextrin complex in the instant solid dosage form would be administered once per day for a period of five to seven days in the first month, repeated for another period of five to seven days in the second
- 20 month, followed by ten months of no treatment. Alternatively the patient would be treated with 10 mg of cladribine in the instant complex cladribinecyclodextrin complex in the instant dosage form once per day for a period of five to seven days per month for a total of six months, followed by eighteen months of no treatment. For further dosing information, see also U.S.
- Provisional Patent Application No._____ IIVAX0021-P-USA/Attorney-Docket No. 033935-011], and U.S. Provisional Patent Application No. -IIVAX0022-P-USA/Attomey Docket No. 033935-012], both entitled "Cladribine Regimen for Treating Multiple Sclerosis", both filed on-March 25, 2004 and incorporated by reference herein in their entireties.

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-23-

Physicians, Vol. 111, No. 1, 35-44 (1999); Selby et al., The Canadian
Journal of Neurological Sciences, 25, 295-299 (1998); Tortorella et al.,
Current Opinion in Investigational Drugs, 2 (12), 1751-1756 (2001); Rice et al., Neurology, 54, 1145-1155 (2000); and Karlsson et al., British Journal of Haematology, 116, 538-548 (2002); all of which are incorporated by
reference herein in their entireties and relied upon.

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Moreover, the route of administration for which the therapeutically effective dosages are taught in the literature should be taken into consideration. While the instant compositions optimize the bioavailability of cladribine following oral administration, it will be appreciated that even optimal bioavailability from oral dosage forms is not expected to approach bioavailability obtain after intravenous administration, particularly at early time points. Thus, it is often appropriate to increase a dosage suggested for intravenous administration to arrive at a suitable dosage for incorporation

- 15 into a solid oral dosage form. At the present time, it is envisioned that, for the treatment of multiple sclerosis, 10 mg of cladribine in the instant complex cladribine-cyclodextrin complex in the instant solid dosage form would be administered once per day for a period of five to seven days in the first month, repeated for another period of five to seven days in the second
- 20 month, followed by ten months of no treatment. Alternatively the patient would be treated with 10 mg of cladribine in the instant complex cladribinecyclodextrin complex in the instant dosage form once per day for a period of five to seven days per month for a total of six months, followed by eighteen months of no treatment.

433

Electronic Acknowledgement Receipt			
EFS ID:	19292491		
Application Number:	12986310		
International Application Number:			
Confirmation Number:	6100		
Title of Invention:	ORAL FORMULATIONS OF CLADRIBINE		
First Named Inventor/Applicant Name:	Nicholas S. Bodor		
Customer Number:	13974		
Filer:	Mary Katherine Baumeister/Rebecca Brimmer		
Filer Authorized By:	Mary Katherine Baumeister		
Attorney Docket Number:	20009904-0067		
Receipt Date:	12-JUN-2014		
Filing Date:	07-JAN-2011		
Time Stamp:	18:18:08		
Application Type:	Utility under 35 USC 111(a)		

Payment information:

Submitted wi	th Payment	no			
File Listin	g:				
Document Number	Document Description	File Name	File Size(Bytes)/ Message Digest	Multi Part /.zip	Pages (if appl.)
1		312-Amendment.pdf	189700	yes	5
			1aaf1c6d1026e3403772a0cb4aacbef87c09 39d9		

	Multipart Description/PDF files in .z	ip description	
	Document Description	Start	End
	Amendment after Notice of Allowance (Rule 312)	1	1
	Specification	2	2
	Applicant Arguments/Remarks Made in an Amendment	3	3
	Specification	4	5
Narnings:	in the DDE is too large. The pages should be 9.5 y 11 or 4.4. If this DDE is submit	tod the pages will be resi-	ad upon ontry into
mage File Wra	pper and may affect subsequent processing	ted, the pages will be resiz	eu upon entry into
Information			
	Total Files Size (in bytes):	189	700
characterize Post Card, a	d by the applicant, and including page counts, where applicable. I s described in MPEP 503.	t serves as evidence of	f receipt similar t
<u>New Applica</u> If a new app 1.53(b)-(d) a Acknowledg <u>National Sta</u>	<u>itions Under 35 U.S.C. 111</u> lication is being filed and the application includes the necessary of nd MPEP 506), a Filing Receipt (37 CFR 1.54) will be issued in due o ement Receipt will establish the filing date of the application. ge of an International Application under 35 U.S.C. 371	omponents for a filing ourse and the date sh	date (see 37 CFF own on this
New Applica If a new app 1.53(b)-(d) a Acknowled <u>c</u> National Sta If a timely su U.S.C. 371 an national sta	ntions Under 35 U.S.C. 111 lication is being filed and the application includes the necessary of nd MPEP 506), a Filing Receipt (37 CFR 1.54) will be issued in due o gement Receipt will establish the filing date of the application. <u>ge of an International Application under 35 U.S.C. 371</u> obmission to enter the national stage of an international application of other applicable requirements a Form PCT/DO/EO/903 indicatir ge submission under 35 U.S.C. 371 will be issued in addition to the	omponents for a filing ourse and the date sho on is compliant with th ng acceptance of the a Filing Receipt, in due	date (see 37 CFI own on this e conditions of 3 pplication as a course.

If a new international application is being filed and the international application includes the necessary components for an international filing date (see PCT Article 11 and MPEP 1810), a Notification of the International Application Number and of the International Filing Date (Form PCT/RO/105) will be issued in due course, subject to prescriptions concerning national security, and the date shown on this Acknowledgement Receipt will establish the international filing date of the application.

PART B - FEE(S) TRANSMITTAL

Complete and send this form, together with applicable fee(s), to: <u>Mail</u> Mail Stop ISSUE FEE **Commissioner for Patents** P.O. Box 1450 Alexandria, Virginia 22313-1450

(571)-273-2885 or <u>Fax</u>

INSTRUCTIONS: This form should be used for transmitting the ISSUE FEE and PUBLICATION FEE (if required). Blocks 1 through 5 should be completed where appropriate. All further correspondence including the Patent, advance orders and notification of maintenance fees will be mailed to the current correspondence address as indicated unless corrected below or directed otherwise in Block 1, by (a) specifying a new correspondence address; and/or (b) indicating a separate "FEE ADDRESS" for maintenance fee notifications.

CURRENT CORRESPONDENCE ADDRESS (Note: Use Block 1 for any change of address)

13974 7590 DENTONS US LLP P.O. BOX 061080 Chicago, IL 60606-1080 03/13/2014

Note: A certificate of mailing can only be used for domestic mailings of the Fee(s) Transmittal. This certificate cannot be used for any other accompanying papers. Each additional paper, such as an assignment or formal drawing, must have its own certificate of mailing or transmission.

Certificate of Mailing or Transmission I hereby certify that this Fee(s) Transmittal is being deposited with the United States Postal Service with sufficient postage for first class mail in an envelope addressed to the Mail Stop ISSUE FEE address above, or being facsimile transmitted to the USPTO (571) 273-2885, on the date indicated below.

(Depositor's nar	ne)
(Signatu	re)
(Da	ite)

APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
12/986,310	01/07/2011	Nicholas S. Bodor	20009904-0067	6100

TITLE OF INVENTION: ORAL FORMULATIONS OF CLADRIBINE

APPLN. TYPE	ENTITY STATUS	ISSUE FEE DUE	PUBLICATION FEE DUE	PREV. PAID ISSUE FEE	TOTAL FEE(S) DUE	DATE DUE			
nonprovisional	UNDISCOUNTED	\$960	\$0	\$0	\$960	06/13/2014			
EXAM	AINER	ART UNIT	CLASS-SUBCLASS						
LAU, JON	JATHAN S	1673	514-046000						
1. Change of correspond CFR 1.363). Change of corresp Address form PTO/S "Fee Address" inc PTO/SB/47; Rev 03- Number is required	lence address or indicatio condence address (or Cha B/122) attached. dication (or "Fee Address 02 or more recent) attach	n of "Fee Address" (37 nge of Correspondence " Indication form ed. Use of a Customer	 2. For printing on the patent front page, list (1) The names of up to 3 registered patent attorneys or agents OR, alternatively, (2) The name of a single firm (having as a member a registered attorney or agent) and the names of up to 2 registered patent attorneys or agents. If no name is listed, no name will be printed. 						
3. ASSIGNEE NAME A PLEASE NOTE: Un recordation as set for (A) NAME OF ASSI	AND RESIDENCE DATA less an assignee is ident th in 37 CFR 3.11. Comp GNEE	A TO BE PRINTED ON ⁷ ified below, no assignee oletion of this form is NO	THE PATENT (print or typ data will appear on the pa T a substitute for filing an (B) RESIDENCE: (CITY	be) atent. If an assignee is ic assignment. ' and STATE OR COUNT	lentified below, the doc	ument has been filed for			
ARES TRA Please check the approp	DING S.A.	categories (will not be p	AUBONNE, SWITZ rinted on the patent): \Box	ERLAND Individual 🖄 Corporati	on or other private group	pentity 🖵 Government			
4a. The following fee(s) X Issue Fee Publication Fee (1 X Advance Order -	are submitted: No small entity discount p # of Copies4	4 permitted) 4	 b. Payment of Fee(s): (Plea A check is enclosed. Payment by credit car The Director is hereby overpayment, to Depo 	tse first reapply any prev d. Form PTO-2038 is attact vauthorized to charge the r sit Account Number <u>19</u>	riously paid issue fee sh ched. required fee(s), any defic -3140 (enclose an o	own above) ciency, or credits any extra copy of this form).			
 5. Change in Entity Sta Applicant certifyi Applicant assertin Applicant changin 	ntus (from status indicate ng micro entity status. Se ng small entity status. See ng to regular undiscounte	d above) ee 37 CFR 1.29 37 CFR 1.27 d fee status.	<u>NOTE:</u> Absent a valid ce fee payment in the micro <u>NOTE:</u> If the application to be a notification of loss <u>NOTE:</u> Checking this boy entity status, as applicable	rtification of Micro Entity entity amount will not be was previously under mic s of entillement to micro e x will be taken to be a noti e.	Status (see forms PTO/S accepted at the risk of ap ro entity status, checking ntity status. fication of loss of entitle	SB/15A and 15B), issue oplication abandonment. g this box will be taken ement to small or micro			
NOTE: This form must	be signed in accordance v	vith 37 CFR 1.31 and 1.3	3. See 37 CFR 1.4 for signa	ature requirements and cer	tifications.				
Authorized Signature	/MARTIN A. BRI	JEHS/		Date JUNE	13, 2014				
Typed or printed nam	ne MARTIN A. BRI	JEHS		Registration No.	45635				

Page **439** 3

PTOL-85 Part B (10-13) Approved for use through 10/31/2013.

OMB 0651-0033 U.S. Patent and Trademark Office; U.S. DEPARTMENT OF COMMERCE

Electronic Patent Application Fee Transmittal					
Application Number:	12	986310			
Filing Date:	07.	07-Jan-2011			
Title of Invention:	ORAL FORMULATIONS OF CLADRIBINE				
First Named Inventor/Applicant Name:	Nicholas S. Bodor				
Filer:	Martin A. Bruehs/Louie Malloy				
Attorney Docket Number:	20	009904-0067			
Filed as Large Entity					
Utility under 35 USC 111(a) Filing Fees					
Description		Fee Code	Quantity	Amount	Sub-Total in USD(\$)
Basic Filing:					
Pages:					
Claims:					
Miscellaneous-Filing:					
Petition:					
Patent-Appeals-and-Interference:					
Post-Allowance-and-Post-Issuance:					
Utility Appl Issue Fee		1501	1	960	960
Extension-of-Time:		437			

Description	Fee Code	Quantity	Amount	Sub-Total in USD(\$)
Miscellaneous:				
Printed Copy of Patent - No Color	8001	4	3	12
	Tot	al in USD) (\$)	972

Electronic Acknowledgement Receipt			
EFS ID:	19296893		
Application Number:	12986310		
International Application Number:			
Confirmation Number:	6100		
Title of Invention:	ORAL FORMULATIONS OF CLADRIBINE		
First Named Inventor/Applicant Name:	Nicholas S. Bodor		
Customer Number:	13974		
Filer:	Martin A. Bruehs/Louie Malloy		
Filer Authorized By:	Martin A. Bruehs		
Attorney Docket Number:	20009904-0067		
Receipt Date:	13-JUN-2014		
Filing Date:	07-JAN-2011		
Time Stamp:	11:55:10		
Application Type:	Utility under 35 USC 111(a)		

Payment information:

Submitted wi	th Payment	yes					
Payment Type	2	Credit Card	Credit Card				
Payment was	successfully received in RAM	\$972	\$972				
RAM confirma	ation Number	9528	9528				
Deposit Acco	Deposit Account						
Authorized U	ser						
File Listin	g:						
Document Number	Document Description	4 59 e Name	File Size(Bytes)/ Message Digest	Multi Pages Part /.zip (if appl.)			

This Acknow	/ledgement Receipt evidences receip	t on the noted date by the US	SPTO of the indicated	document	S,
Information	:			70040	
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1	Issue Fee Dayment (DTO SEP)	0067lssupEpp.pdf	138250		1

New Applications Under 35 U.S.C. 111

If a new application is being filed and the application includes the necessary components for a filing date (see 37 CFR 1.53(b)-(d) and MPEP 506), a Filing Receipt (37 CFR 1.54) will be issued in due course and the date shown on this Acknowledgement Receipt will establish the filing date of the application.

National Stage of an International Application under 35 U.S.C. 371

If a timely submission to enter the national stage of an international application is compliant with the conditions of 35 U.S.C. 371 and other applicable requirements a Form PCT/DO/EO/903 indicating acceptance of the application as a national stage submission under 35 U.S.C. 371 will be issued in addition to the Filing Receipt, in due course.

New International Application Filed with the USPTO as a Receiving Office

If a new international application is being filed and the international application includes the necessary components for an international filing date (see PCT Article 11 and MPEP 1810), a Notification of the International Application Number and of the International Filing Date (Form PCT/RO/105) will be issued in due course, subject to prescriptions concerning national security, and the date shown on this Acknowledgement Receipt will establish the international filing date of the application.

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applica	ant: Nicholas S. BODOR et al.)	MAIL STOP ISSUE FEE
Annlie	ation No - 12/086 310)	Examinar Innathan S Al
ryymu)	
Filed:	January 7, 2011)	Group Art Unit: 1623
)	
Title:	ORAL FORMULATIONS OF)	Confirmation No.: 6100
	CLADRIBINE)	

REPLY AND AMENDMENT PURSUANT TO 37 C.F.R. § 1.312

In response to the Notice to File Corrected Application Papers dated April 17, 2014, Applicants hereby submit the following amendment and remarks pursuant to 37 C.F.R. §1.312.

Unit	ed States Patent	and Trademark Office	UNITED STATES DEPAR United States Patent and Address: COMMISSIONER F P.O. Box 1450 Alexandria, Virginia 22: www.uspto.gov	TMENT OF COMMERCE Trademark Office OR PATENTS 313-1450
APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
12/986,310	01/07/2011	Nicholas S. Bodor	20009904-0067	6100
13974 DENTONS US	7590 06/25/2014 LLP	EXAMINER		
P.O. BOX 0610 Chiango II 600)80 606 1080	LAU, JONATHAN S		
Cincago, 12 00000-1080			ART UNIT	PAPER NUMBER
			1673	
			NOTIFICATION DATE	DELIVERY MODE
			06/25/2014	ELECTRONIC

Please find below and/or attached an Office communication concerning this application or proceeding.

The time period for reply, if any, is set in the attached communication.

Notice of the Office communication was sent electronically on above-indicated "Notification Date" to the following e-mail address(es):

martin.bruehs@dentons.com patents.us@dentons.com

	Application No	Applicant(s)					
Response to Rule 312 Communication	12/986,310	BODOR ET AL.					
	Examiner	Art Unit					
	Jonathan S. Lau	1673					
The MAILING DATE of this communication	n appears on the cover sheet	with the correspondence address –					
1. 🛛 The amendment filed on <u>12 June 2014</u> under 37 CFF	1.312 has been considered, ar	nd has been:					
a) entered.							
b) 🛛 entered as directed to matters of form not affect	ing the scope of the invention.						
c) 🔲 disapproved because the amendment was filed	after the payment of the issue f	ee.					
Any amendment filed after the date the issue fee is paid must be accompanied by a petition under 37 CFR 1.313(c)(1)							
and the required fee to withdraw the application from issue.							
d) 🔲 disapproved. See explanation below.	d) 🔲 disapproved. See explanation below.						
e) 🔲 entered in part. See explanation below.							
airected to matters of form not affecting the scope of th	e invention.						
/SHAOJIA ANNA JIANG/	/Jonathan S Lau/	1670					
Supervisory Patent Examiner, Art Unit 1673	⊂xaminer, Art Unit	10/3					





APPLICATION NO.	ISSUE DATE	PATENT NO.	ATTORNEY DOCKET NO.	CONFIRMATION NO.
12/986,310	07/22/2014	8785415	20009904-0067	6100
13974	7590 07/02/2014	4		

DENTONS US LLP P.O. BOX 061080 Chicago, IL 60606-1080

ISSUE NOTIFICATION

The projected patent number and issue date are specified above.

Determination of Patent Term Adjustment under 35 U.S.C. 154 (b)

(application filed on or after May 29, 2000)

The Patent Term Adjustment is 357 day(s). Any patent to issue from the above-identified application will include an indication of the adjustment on the front page.

If a Continued Prosecution Application (CPA) was filed in the above-identified application, the filing date that determines Patent Term Adjustment is the filing date of the most recent CPA.

Applicant will be able to obtain more detailed information by accessing the Patent Application Information Retrieval (PAIR) WEB site (http://pair.uspto.gov).

Any questions regarding the Patent Term Extension or Adjustment determination should be directed to the Office of Patent Legal Administration at (571)-272-7702. Questions relating to issue and publication fee payments should be directed to the Application Assistance Unit (AAU) of the Office of Data Management (ODM) at (571)-272-4200.

APPLICANT(s) (Please see PAIR WEB site http://pair.uspto.gov for additional applicants):

Nicholas S. Bodor, Bal Harbour, FL; Yogesh Dandiker, Toronto, CANADA;

The United States represents the largest, most dynamic marketplace in the world and is an unparalleled location for business investment, innovation, and commercialization of new technologies. The USA offers tremendous resources and advantages for those who invest and manufacture goods here. Through SelectUSA, our nation works to encourage and facilitate business investment. To learn more about why the USA is the best country in the world to develop technology, manufacture products, and grow your business, visit <u>SelectUSA.gov</u>.