

P1 1170170

THE UNITED STAYES OF AVER CA

TO ANK TO WHOM THESE PRESENTS SHAME COME:

UNITED STATES DEPARTMENT OF COMMERCE United States Patent and Trademark Office

May 17, 2004

THIS IS TO CERTIFY THAT ANNEXED HERETO IS A TRUE COPY FROM THE RECORDS OF THE UNITED STATES PATENT AND TRADEMARK OFFICE OF THOSE PAPERS OF THE BELOW IDENTIFIED PATENT APPLICATION THAT MET THE REQUIREMENTS TO BE GRANTED A FILING DATE.

APPLICATION NUMBER: 60/458,922

FILING DATE: March 28, 2003

RELATED PCT APPLICATION NUMBER: PCT/US04/09387

REC'D **2 1 MAY 2004**WIPO FOT

By Authority of the COMMISSIONER OF PATENTS AND TRADEMARKS



N. WOODSON
Certifying Officer

PRIORITY DOCUMENT

SUBMITTED OR TRANSMITTED IN COMPLIANCE WITH RULE 17.1(a) OR (b)

Merck 2044 TWi v Merck IPR 2023-00050

=	E
	<u>0</u>
∼■	Please type a plus sign (+) inside this b
\approx	C.
3	PROVISIONAL APPL



Docket Number:

LICATION FOR PATENT COVER SHEET (Large Entity)

This is a request for filing a PROVISIONAL APPLICATION FOR PATENT under 37 CFR 1.53 (c).

			INVENTO	R(S)/APPLIC	CANT(S)				
Given Name (first and mi	ddle [if any])	Family	Name or Su	rname	Resi	dence (City and	either State or	r Foreign Country)	
Nicholas		Bodor			Miami, FL		·		,
Additional inve	Additional inventors are being named on page 2 attached hereto								
				ENTION (28		s max)			
DRAL AND TRANSMU	ICOSAL DE	LIVERY OF C	CACODEX	CTRIN BASE	FORMULA	ATIONS			
Disease all agreemendate	ano to:		CORRESI	PONDENCE	ADDRESS				,
Direct all corresponder Customer Number		236	30				ce Customer ar Code Lab		
Firm or Individual Name					<u> </u>				
Address		······································		146					
City				State			ZIP		
Country				Telephone			Fax		
	' 	ENCLOSE	D APPLIC	ATION PART	S (check a	ll that apply)			
Specification	Numi	per of Pages	23						
Drawing(s)		er of Sheets			Other (s	Po	stcard		
METHOD O	F PAYMENT	OF FILING	FEES FOR	THIS PROV	SIONAL AF	PLICATION	FOR PATEN	IT (check one)	
A check or me	oney order is	enclosed to	cover the fi	ling fees			•	FILING AMOUN	IT (\$)
The Commiss credit any over	erpayment to	Deposit Acco	ount Numbe	er:		50-1133		\$160	.00
The invention was made t	y an agency o	f the United Sta	tes Governm	ent or under a	contract with a	an agency of the	United States	Government.	
No.									
Yes, the name of	the U.S. Gove	rnment agency	and the Gove	ernment contrac	t number are:				
Respectfully submit	tea, Mas	news P) - M	W	~	DATE	Marc	ch 28, 2003	
TYPED or PRINTE	D NAME	Jeffrey J. M	iller			REGISTRA'		39,77	73
TELEPHONE	_	617-535-44	21			/ alala ala			

USE ONLY FOR FILING A PROVISIONAL APPLICATION FOR PATENT

SEND TO: Box Provisional Application, Assistant Commissioner for Patents, Washington, DC 20231

PATENT

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application:

Nicholas Bodor

Serial Number: Filing Date:

Not Yet Assigned March 28, 2003

Title:

ORAL AND TRANSMUCOSAL DELIVERY OF

CYCLODEXTRIN BASED FORMULATIONS

Docket Number:

IVAX0012-P-USA

CERTIFICATE OF MAILING UNDER 37 C.F.R.§ 1.10

"Express Mail" Mailing Label Number

EL945335370US

Date of Deposit March 28, 2003

I hereby certify that this paper or fee is being deposited with the United States Postal Service "Express Mail Post Office to Addressee" service under 37 CFR 1.10 on the date indicated above and is addressed to Box Provisional Patent Application, Assistant Commissioner for Patents, Washington, DC 20231.

Jeffrey J. Miller

(Person Mailing)

(Signature)

BOX PROVISIONAL PATENT APPLICATION

Assistant Commissioner for Patents Washington, DC 20231

Dear Sir:

TRANSMITTAL LETTER

Transmitted herewith for filing are the following documents:

- [] Provisional Application for Patent Cover Sheet (Large Entity);
- Provisional Patent Application (5 Pages Specification, 18 Pages Appendix);
- Authorization to charge Deposit Amount for filing fee of \$160.00; and
- [] Return Postcard

If the enclosed papers are considered incomplete, the Mail Room and/or the Application Branch is respectfully requested to contact the undersigned collect at (617) 535-4421, Boston, MA.

The Commissioner is hereby authorized to charge any additional fees which may be required or credit any overpayment to Deposit Acct. No. 50-1133.

Dated: 3-28-03

Respectfully submitted,

Jeffrey L Miller, Reg. No.39,773 McDERMOTT, WILL & EMERY

28 State Street

Boston, MA 02109-1775 617-535-4421 (Telephone) 617-535-3800 (Facsimile)

Docket Number:

IVAX0012-P-USA

PROVISIONAL APPLICATION FOR PATENT COVER SHEET (Large Entity)

	INVENTOR(S)/APPLICANT(S)						
Given Name (first and middle [if any])	Family Name or Surname	Residence (city and either State or Foreign Country)					
,							
	•						
	·						
		•					
		·					
		·					
1							
	I						

Certificate of Mailing by Express Mail

I certify that this application and enclosed fee is being deposited on March 28, 2003 with the U.S. Postal Service "Express Mail Post Office to Addressee" service under 37 C.F.R. 1.10 and is addressed to the Assistant Commissioner for Patents, Washington, D.C., 20231.

Signature of Person Mailing Correspondence

Jeffrey J. Miller

Typed or Printed Name of Person Mailing Correspondence

EL 945335370 US

"Express Mail" Mailing Label Number

USE ONLY FOR FILING A PROVISIONAL APPLICATION FOR PATENT

SEND TO: Box Provisional Application, Assistant Commissioner for Patents, Washington, DC 20231

APPENDIX

CONTENTS

I. AFI DISSUUUUU AHAIVS	1.	API	Dissolution	Analysis
-------------------------	----	-----	-------------	----------

- 1.1 Dissolution in 0.1N HCl (UV and HPLC)
- 1.2 Dissolution in Phosphate buffer pH 6.8 (UV and HPLC)
- 1.3 Dissolution in DI Water (UV)
- 1.4 Dissolution in Buffer pH 4.5

2. Intrinsic Dissolutions

- 3. API Related Substances (Comparison to Cilag)
- 4. Finished Product Related Substances
- 5. Freeze-dried complexes FB Related Substances

6. Formulations based on Fludaribine

- 4.1 Fludaribine Formulation: RDT0385
- 4.2 Enteric-coated tablets: (Fludaribine Formulation). RDT0385b
- 4.3 20% Carbomer Formulation: RDT0398a
- 5.4 Cyclodextrin Formulation: RDT0398b

7. Buccal and granule Formulations using Diclofenac as API

- 6.1 Buccal / Sublingual
- 6.2 Mucoadhesive granule for HGC fill
- 6.3 Mucoadhesive Direct compression tablet
- 6.4 Tablet within a tablet formulation:

8 Phase solubility testing

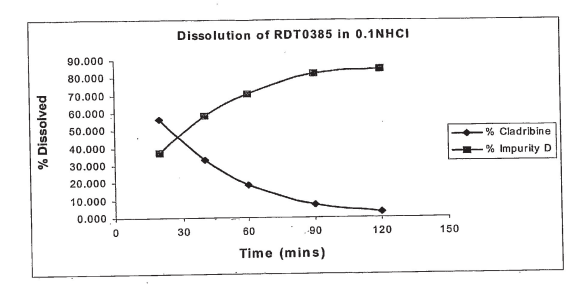
9 Cladribine freeze-dried Cyclodextrin complexes

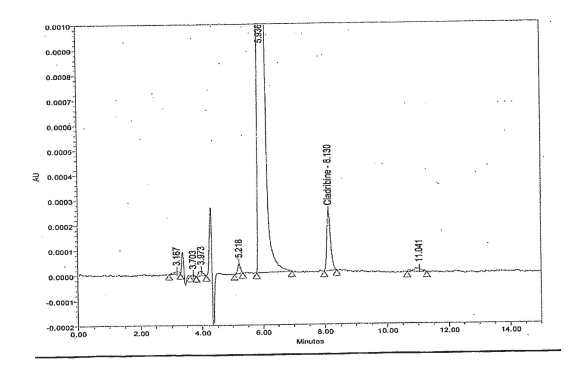
- 9.1 Cyclodextrin Complex Formulations for Buccal/Sublingual Dosage forms
- 9.2 Manufacturing Process
- 9.3 Physical Parameters
- 9.4 Dissolution profiles of Cladribine freeze-dried buccal tablets in water and salivary buffer
- 9.5 Results
- 9.6 Dissolution and Degradation profiles of Cladribine freeze-dried Cyclodextrin buccal tablets in 0.1N HCl

1. API ANALYSIS

1.1 Dissolution in 0.1N HCl

Initial analysis of active by UV showed 10% degradation of API over 2 hours. Following from this HPLC analysis of active in 0.1N HCL showed degradation of Cladribine and growth of Impurity D (RRT 0.701). Approx 3% Cladribine remaining after 120 minutes dissolution.

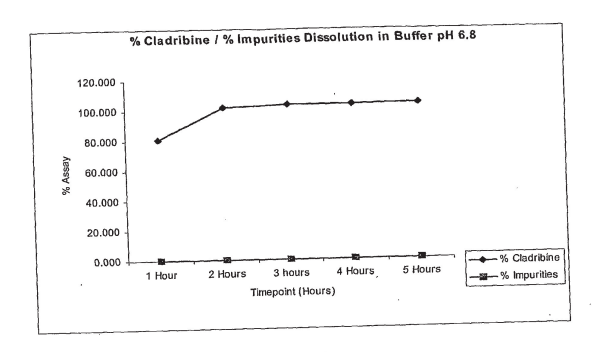




Chromatogram of Cladribine after 2 hours dissolution in 0.1N HCl. Growth of impurity D at retention time 5.936 minutes

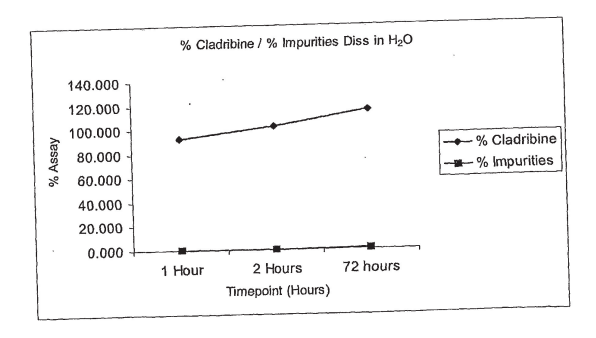
1.2 Dissolution in Phosphate buffer pH 6.8 (HPLC)

102% dissolved after 2 hours. No observed degradation after 5 hours. No increase in related substances.



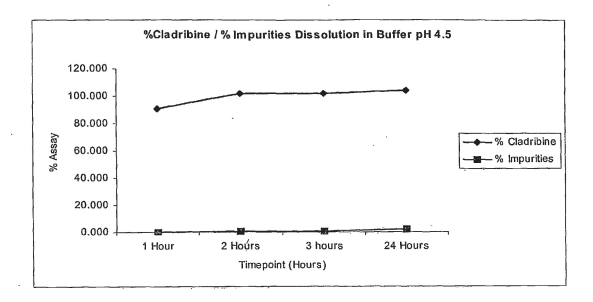
1.3 Dissolution in DI Water (HPLC)

102% dissolved after 2 hours. No observed degradation after 5 hours. Increase in assay of Cladribine after 72 hours due to evaporation of medium. No increase in related substances



1.4 Dissolution in Buffer pH 4.5 (HPLC)

102% dissolved after 2 hours. No observed degradation after 2 hours. Increase in impurities (0.1%) of Cladribine after 24 hours.



INTRINSIC DISSOLUTIONS 2

IDR of 0.1 mg/min/cm² corresponds to solubility of 1 mg/ml. Note:

Cilag estimate solubility of 5mg/ml in water

IDR of API in DI water:

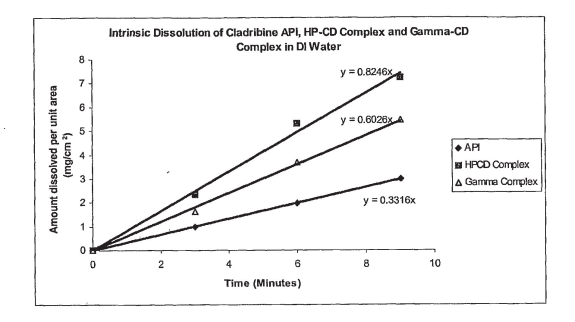
0.3316 mg/min/cm²

IDR of Gamma-CD complex in DI water:

0.6026 mg/min/cm²

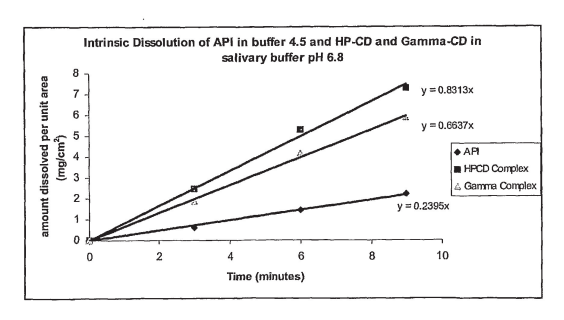
0.8246 mg/min/cm²

IDR of HP-CD complex in DI water:

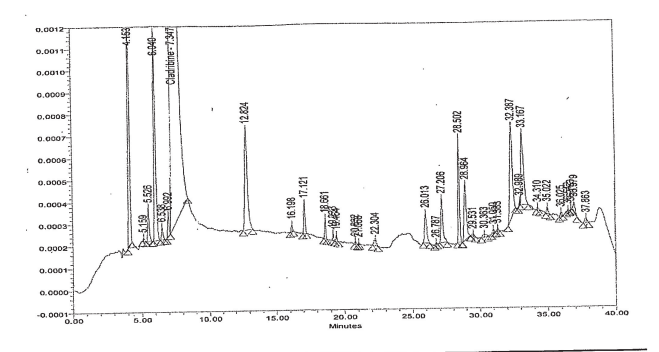


IDR of API in phosphate buffer pH 4.5: IDR of Gamma-CD complex in salivary buffer pH 7.0: IDR of HP-CD complex in salivary buffer pH 7.0:

0.2395 mg/min/cm² 0.6637 mg/min/cm² 0.8313 mg/min/cm²



3. API RELATED SUBSTANCES



Name	Specification	RRT	Cilag Assay	IVAX Assay
2-Amino-2 deoxyadenosine (Impurity B)	NMT 0.3%	0.563	0.200	0.060
2-Chloro-adenine (Impurity D)	NMT 0.3%	0.701	<0.1	0.002
2-Methoxy-2-deoxyadenosine (Impurity E)	NMT 0.2%	0.821	0.200	0.082
2-Chloro-9-(2 deoxy-α-D- ribofuranosyl)-adenine (Impurity F)	NMT 0.2%	0.951	<0.1	0.01
Cladribine	98% - 102%	1.000	99.8	98.5
Unknown 1	NMT 0.1%	1.763		0.088
Unlnown Impurity RRT (Cilag RRT) = 1.85	NMT 0.2%	1.85		ND
Impurity G	NMT 0.1%	2.123		ND
RWJ-47753-000	NMT 0.1%	3.877		0.043
RWJ-47754-000	NMT 0.1%	4.511		0.056
TOTAL IMPURITIES	NMT 1.0%		0.6%	0.3%

4. FINISHED PRODUCT RELATED SUBSTANCES

Name	RRT	Specification	RDT0385 (Fludaribine formulation)	RDT0398a (Carbomer formulation)	RDT039 (Cyclode formulat
2-Amino-2 deoxyadenosine (Impurity B)	0.563	NMT 0.3%	0.059	0.067	0.056
2-Chloro-adenine (Impurity D)	0.701	NMT 0.3%	0.002	0.002	0.002
2-Methoxy-2- deoxyadenosine (Impurity E)	0.821	NMT 0.2%	0.083	0.093	0.076
2-Chloro-9-(2 deoxy-α- D-ribofuranosyl)- adenine (Impurity F)	0.951	NMT 0.2%	0.010	0.012	0.009
Cladribine	1.000	98% - 102%	96	114	90
Unknown 1	1.763	NMT 0.1%	0.086	0.101	0.082
Unlnown Impurity RRT (Cilag RRT) = 1.85		NMT 0,2%			
Impurity G	2.123	NMT 0.1%	0.001	0.000	0.000
RWJ-47753-000	3.877	NMT 0.1%	0.042	0.050	0.039
RWJ-47754-000	4.511	NMT 0.1%	0.049	0.059	0.047
TOTAL IMPURITIES		NMT 1.0%	0.33%	0.38%	0.24%

5. ASSAY AND RELATED SUBSTANCES OF FREEZED DRIED COMPLEX RAW MATERIAL AND TABLETS

Identity	Chemical Name	RRT	Gammá – CD Raw Material	HP-β-CD Raw Material	FD02 (5mg Gamma- CD Tablets)	FD03 (5mg HPCD Tablets)
Imp B	2-Amino-2'- deoxyadenosine	0.54	0.28	0.19	0.31	0.29
Imp D	2-Chloroadenine	0.73	<0.05	ND	ND	ND
Imp E	2-Methoxy-2'- deoxyadenosine	0.83	0.14	0.12	0.13	0.13
Imp F	2-Chloro-9-(2'- deoxy-α-D- ribofuranosyl)- adenine	0.93	ND	ND	ND	ND
API	Cladribine	1.00	108	100	105	102
Theoretical % Active in Complex	Cladribine		2.128	2.347		
Actual % Active in Complex	Cladribine		2.293	2.353		
Unknown	Not Known	1.89	0.06	0.09	0.07	0.07
RWJ- 49616-000	Not Known	2.60	ND	ND	ND	ND
Unknown	Not Known	3.06	<0.05	1.56*	<0.05	<0.05
Unknown	Not Known	3.43	0.05	0.07	0.08	0.06
RWJ- 47753-000	Not Known	3.90	ND	ND	ND	ND
Unknown	Not Known	4.18	ND	ND	0.26	ND
Unknown	Not Known	4.39	ND	ND	0.98	0.31
Unknown	Not Known	4.63	ND	0.33	ND	ND
RWJ- 47754-000	Not Known	4.68	0.22	0.15	0.34	0.21
TOTAL			0.75	2.51	2.17	1.01

^{*} To be investigated. Possible solvent or carryover.

SUMMARY

No differences observed in assay for related substances for API and any formulations. Recommended PDA analysis on API also.

6. FORMULATIONS BASED ON FLUDARIBINE

Three 100g batches using Cladribine API have been manufactured using the following formulations:

Batch	- RDT0385	= RDT0398a	RDT0398b
	Fludaribine Formulation	Carbomer	Cyclodextrin Formulation
Ingredient/mg/batch			
Cladribine API	10.00	10.00	10.00
Hydroxypropyl -β= Cyclodextrin			41.79
Carbomer 974P		20.00	
Avicel PH101	21.80	16.7	11.25
Lactose DC11	65.00	50.1	33.76
Crosacarmellose Sodium	2.00	2.00	2.00
Collidol Silicon Dioxide	0.20	0.20	0.20
Magnesium Sterate	1.00	1.00	1.00
Total	100.00	100.00	· 100.00

Measurement	RD(F0385	RDT0398a (20%-Carbomer)	RDT0398b (Gyclodextrin)
Average tablet weight (mg)	100.1	101.1	103.3
Average Hardness (Kp)	4.9	4.4	3.7
Friability (%)	0.18	0.03	0.18
Thickness (mm)	2.86	3.24	2.92
Disintegration (min)	0.50	> 15.00	6.60

6.1 Fludaribine Formulation: RDT0385

• Assay - 101.4%

• CU - 100.5%, RSD = 3.17%

UV Dissolution (0.1N HCl)
 Max 91% 30 minutes.

HPLC analysis carried out on dissolution in HCl showed breakdown of Cladribine into impurity
 D. Only 3% Cladribine remaining after 2 hours dissolution.

• UV Dissolution (buffer pH 6.8) - Slow release. 85% after 240 minutes

UV Dissolution (Water)
 Fast release. 101% after 2 hours.

6.2 Enteric-coated tablets: (Fludaribine Formulation). RDT0385b

- UV Dissolution in 0.1N HCl followed by buffer pH 6.8 7.0.
- 7% dissolution after 2 hours in acid, (min 5%, max 18%). On addition of pH 7.0 conditions dissolution increased to 97% after 2 hours (min 84%, Max 107%). After 4 hours in acid, dissolution was 116%.

6.3 20% Carbomer Formulation: RDT0398a

Results may be related to tablet weight i.e. heavier tablet gives higher dissolutions

Assay
 CU
 UV Dissolution (0.1N HCl)
 UV Dissolution (buffer pH 6.8)
 I13.9%
 105.7%, RSD = 6.4%. One result at 123.1%
 Max 80%, 240 minutes. Slow release profile
 Slow release. 86% after 10 hours. 0.1%

Carbomer interference. Further HPLC analysis shows possible Carbomer peak at 4-5 minutes.

0.2% - 1.0%.

UV Dissolution (Water)

Fast release. 97% after 2 hours.

6.4 Cyclodextrin Formulation: RDT0398b

Cyclodextrin formulation is sub-potent due to extra Mag Stearate added. Estimated potency at 95%.

• Assay - 89.9%

• CU - 83.2%, RSD = 3.3%

• UV Dissolution (0.1N HCl) - Max 83%, 48 minutes. Degradation occurs.

UV Dissolution (buffer pH 6.8)
 Max 76%, 1 hour. No Cyclodextrin interference

UV Dissolution (Water)
 Max 86% after 1 hour.

SUMMARY

- Cladribine API is acid labile. Formulation needed to avoid acidic stomach conditions.
- No degradation observed in water, buffer pH 4.5 and buffer pH 6.8
- API IDR matches Cilag estimated solubility. Best IDR in water.
- Solubility issue in buffer pH 6.8. Dissolution values are less than assay results.
- Solubility does not seem to be a problem in water. Dissolution results matching assay and CU.
- Fludaribine formulation shows fast release in water and slow release in buffer pH 6.8.
- Carbomer formulation allows for slow release. Carbomer impurity (approx 1.0%) present in chromatography.

Some spurious CU results (121%) indicating possible processing problems with Carbomer 974P or high levels of Carbomer.

• Possible potency issue with Cyclodextrin formulation. Only getting 90% assay and dissolution. Immediate release in buffer and water.

7 BUCCAL AND GRANULE FORMULATIONS WITH DICLOFENAC API

Six batches using Diclofenac Sodium in place of Cladribine API were manufactured to explore the development of buccal / sublingual and mucoadhesive tablets as patentable cladribine formulations.

Formulation:

	#RDT0399a=	RDT0399b	RDT0399d	-RDT0399e-	■RDT0399f =	RDT0399g
	Buccal	Buccal	Granule :	Granule -	DC tablet	DC tablet
	tablet	tablet	(Carbopol	(Carbopol	(Carbopol	(Carbopol
			974P)	974P)	≥ = 71G) = .	71G
Ingredient/						
mg/tablet	10.00	10.00	10.00	10.00	10.00	10.00
Diclotenac	10.00	10.00	10.00	10.00	10.00	10.00
Sodium CMC	2.50	5.00				
Sorbitol	87.00	84.50			的图制造造	
Carbopol 974P			2.50	10.00		
Carbopol-71G				能理測度	2.50	10.00
Avicel PH101			86.80	79.30		
Avicel PH102					21.75	19.88
Lactose DG11.				對為對於	65.25	59.63
Aërosil 🗐 💮	海(新疆)表	多數學學	0.20	0.20		自己
_=Mag. stearate=	0.50	0.50*	0.50	0.50	0.50	0.50

^{*}Extra 0.5mg/tablet added to minimise picking. RDT0399c was manufactured as RDT0399a placebo.

Physical parameters:

Measurement -	RDT0399a	RDT0399b/4	RDT0399f	RDT0399g
Tooling//shape =	Concave	Flat /Concave	Concave	Concave
Average tablet weight (mg)	95.8	94.5	95.1	99.7
Average Hardness (Kp)	3.76	2.10	2.94	2.46
Friability (%)	1,35	0.60	0.00	0.00
Thickness (mm)	3.07	2.90	2.95	3.10
Disintegration (min)	2min 34sec	4min 45sec	>15min*	>15min**

^{*}Tablet formed a soft globular mass with adhesive properties

NOTE: Diclofenac has solubility problems in 0.1N HCl. Diclofenac Na dissolves 16% - 20% in 0.1N HCl.

^{**} Tablet formed a globular mass with strong adhesive properties. Mass was dry in center after 15 mins.

7.1 Buccal / Sublingual:

RDT0399a + RDT0399b:

Manufactured using Sodium CMC at 2.5 - 5 % respectively.

- UV Dissolution of approx 70% after 10 hours in simulated saliva solution. 68% dissolution after 30 minutes.
- Assay of 70%.
- No obvious reason for low results.
- Poor taste from tablets. Possible Diclofenac Na taste. Recommend 2mg drug formulation per 100 mg tablet to inhibit possible taste issues.

7.2 Mucoadhesive granule for HGC fill:

NOTE:

Carbopol 71G may offer better flow properties due to its granular nature which may alleviate possible processing problems.

RDT0399d:

Manufactured using Carbopol 974P at 2.5%.

• 5% dissolution in 0.1N HCl after 2 hours. 97% dissolution after 3 hours in pH 7.0 buffer.

RDT0399e:

Manufactured using Carbopol 974P at 10%.

• 6% dissolution in 0.1N HCl after 2 hours. 91% - 99% after 3 hours in pH 7.0

7.3 Mucoadhesive Direct compression tablet:

RDT0399f:

Manufactured using Carbopol 71G at 2.5%.

• 5% dissolution in 0.1N HCl after 2 hours. 76% after 3 hours in pH 7.0

RDT0399g:

Manufactured using Carbopol 71G at 10%.

2% dissolution in 0.1N HCl after 2 hours. 90% after 3 hours in pH 7.0

All tablet formulations flowed and compressed well.

The granulated product produced a good strong granule. Milled through a 0.075 inch comil screen.

7.4 Tablet within a tablet formulation:

Outer tablet coat used to protect Cladribine from acidic stomach conditions.

Dissolution in 0.1N HCL followed by buffer pH 6.8. Tablets completely disssolved in acid (86% - 95%) after 25 minutes. No advantage.

8 PHASE SOLUBILITY TESTING

Table 1. Solubility of cyclodextrins in water (g/100 ml)

Temperatur e (°C)		BCD		HPCD
20.0	10.1	1.55	23.2	360.0
25.0	13.0	1.85	30.0	
30.0	16.0	2.25	38.5	
40.0	25.6	3.52	63.5	

PROTOCOL FOR PHASE SOLUBILITY STUDIES OF CLADRIBINE IN PRESENCE OF CYCLODEXTRIN

Reported Solubility of Cladribine in Water is 5 mg/ml

TABLE 1

SOLUTION SYSTEMS	Solution of CD, 800 mg in 4ml B.soln		DRUG ADDED
A	2ml B. Soln	(400 mg)	25 mg
В	2ml B.soln. + 2ml D.Water	(200 mg)	25 mg
С	2 ml soln. B + 2 ml D. Water	(100 mg)	25 mg
D	2 ml soln. C + 2 ml D.Water	(50 mg)	25 mg
E	2 ml soln. D + 2 ml D.Water	(25 mg)*	25 mg
	* Use only 2 ml of solution for		
F	2 ml D.Water	(0.0 mg)	25 mg

Cyclodextrin

B.soln. - Bulk Solution

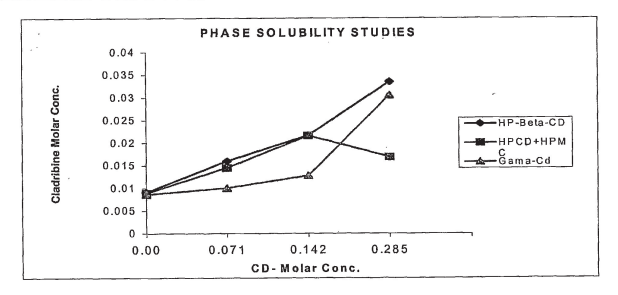
D. Water - Deionised Water

Method for preparation.

- 1. In screw capped vials take 2 ml Cyclodextrin solutions as mentioned in Table 1.
- 2. Add respective quantity of drug in each vial.
- 3. Allow the samples to sonicate for 30 minutes.
- 4. Remove the samples from sonicator and place on shaker for 8 hrs.
- 5. The sample after shaking is filtered to get clear supernant.
- 6. Analyse the sample by UV at 265 nm wavelength.

RESULTS:

CD Conc.	Cladribine –HP betaCD (Trial A)		Cladribine -HP betaCD + HPMC(0.1%) (Trial B)			Cladribine -gama- CD (Trial C)			
CD Conc.	Absorbance	mg/ml	Molar concn.	Absorbance	mg/ml	Molar concn.	Absorbance	mg/ml	Molar c
0.00	0.140	2.610	0.0091	0.137	2.550	0.0089	0.132	2.459	0.0086
0.018	0.169	3,139	0.011	0.146	2.711	0.0095	0.1352	2.519	0.0088
0.035	0.191	3.554	0.0124	0.175	3.262	0.0114	0.1531	2.852	0.0100
0.071	0.245	4.570	0.016	0.223	4.149	0.0145	0.1542	2.873	0.0101
0.142		6.211	0.0217	0.332	6.185	0.0216	0.1965	3.661	0.0128
0.285	0.514	9.581	0.0335	0.259	4.831	0.0169	0.4688	8.733	0.0306



Observations:

- The best solubility results are obtained with HP-beta CD as complexing agent.
- With HP-beta CD + HPMC (0.1%) results are similar to HP-beta CD, at higher concentration fine precipitation was observed in the vials at the end of the study.
 - Absorbance of this sample is low and indicates precipitation of solubised drug
- Absorbance with Gama-Cyclodextrin is low as compared with HP-beta CD.
- Ball park solubility of 9.581 mg/ml in comparison to 5 mg/ml solubility with API alone.

SUMMARY

- Cyclodextrin/Cladribine complex showed increased Cladribine solubility
- Complex sent for freeze-drying.
- Cladribine API ground to decrease particle size (10g)
- Process buccal and sublingual tablets using freeze-dried material
 Issues regarding taste and poor assay, dissolution on previous buccal tablets.
 Information on buccal formulation work in Miami.
- Continued investigation into oral dosage formulations:
 - 1. Tablet-within-tablet:

High viscosity HPMC in outer formulation for protection against acidic stomach conditions.

2. Soft gel capsule:

10g API sent to Czechslovakia for trials.

3. Dry emulsion formulation:

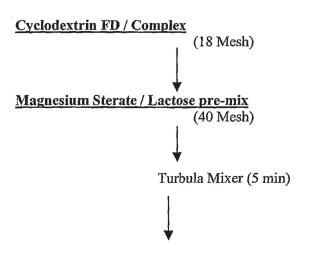
Dummy emulsion to be made with freeze-dried sample

9 CLADRIBINE FREEZE-DRIED CYCLODEXTRIN COMPLEXES

9.1 Cyclodextrin Complex Formulations for Buccal/Sublingual Dosage forms

	PRODUCT		Gamma = . CD Tablets	Gamma=CD Sorbitol Tablets	Gamma-CD: + Cladribine - Complex - Tablets	
22 - 43 F14 20 MINHAD DE CITE MAN AC	Batch No. Ingredient	Lot no.	the billion and Combined to 150 160-160-100 and the company	RDT 0418B	The first of the last in the state of the st	RDT 0418D Mg/Tablet
FD-01	Gamma -CD	N/A	213	213	-	
FD-02	Gamma-CD +Cladribine	N/A	-	-	235	-
FD-03	HPCD + Cladribine	N/A	_	-	-	218
RE0484	Sorbitol	1F290	-	5.0	-	· -
RE0541	Magnesium Stearate	1C130	2.0	2.0	2.0	2.0

9.2 Manufacturing Process



Manesty Single station F-press (220 mg, 10.0 mm round concave UP/ Flat Bevelled LP)

OBSERVATIONS

Flow and compressibility good for all fractions.

No picking noticed

9.3 Physical Parameters

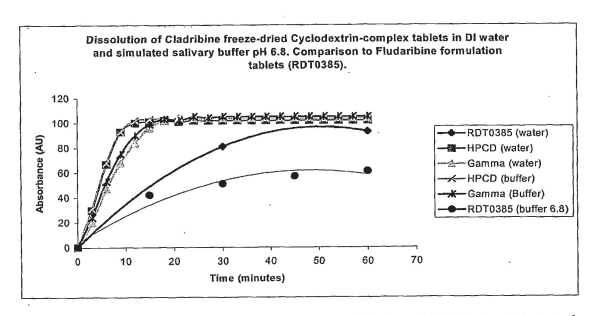
Average weight: A) 215 mg, B) 220 mg, C) 237mg, D)220mg.

Average Hardness: 3- 4 Kp

• Thickness: 3.2 mm - 3.4 mm

Disintegration Time: 6 – 7 minutes (Water/Simulated Saliva Buffer)

9.4 Dissolution profiles of freeze-dried buccal tablets in water and simulated salivary buffer solution



Simulated Saliva Solution: 2.38g Na₂HPO₄, 0.19g KH₂PO₄ and 8g NaCl in 1 litre of distilled water, pH 6.75, at 37°C

9.5 Results

Increased Dissolution time.

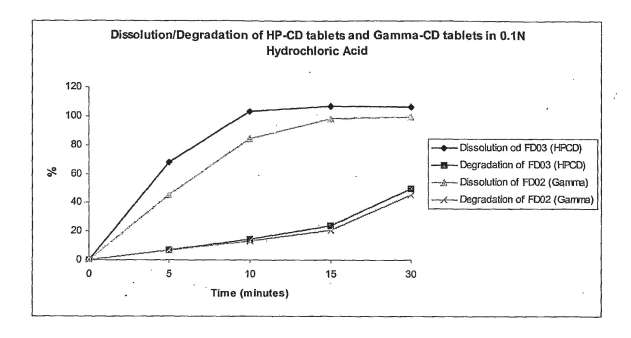
HP-CD, 100% dissolution in salivary buffer after 10 minutes. Gamma-CD, 100% dissolution in salivary buffer after 15 minutes

HP-CD. 100% dissolution in water after 10 minutes. Gamma-CD. 100% dissolution in salivary buffer after 15 - 18 minutes

Increased Solubility.

100% dissolution attained for both tablet types in both buffers. Comparison to Fludaribine formulation dissolution in water and buffer show faster dissolution and greater solubility.

9.6 Dissolution and Degradation profiles of freeze-dried Cladribine-Cyclodextrin complex buccal tablets in 0.1N HCl



- Degradation of Cladribine peak to Impurity D observed. 10 15% after 10 minutes. 100% dissolution after 10 15 minutes.
- By optimising complexation, we can further inhibit acidic degradation of the drug in the stomach whilst increasing drug availability for absorption.