

## WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



## INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification <sup>7</sup> : A61K 47/10, C07D 498/18, A61K 31/715	A2	(11) International Publication Number:WO 00/33878(43) International Publication Date:15 June 2000 (15.06.00)	
<ul> <li>(21) International Application Number: PCT/EP99/09521</li> <li>(22) International Filing Date: 6 December 1999 (06.12.99)</li> <li>(30) Priority Data: 9826882.4 7 December 1998 (07.12.98) GB 9904934.8 4 March 1999 (04.03.99) GB</li> <li>(71) Applicant (for all designated States except AT US): NOVAR- TIS AG [CH/CH]; Schwarzwaldallee 215, CH-4058 Basel (CH).</li> </ul>		<ul> <li>(81) Designated States: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, TZ, 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).</li> </ul>	
(71) Applicant (for AT only): NOVARTIS-ERFINDUNGEN VER- WALTUNGSGESELLSCHAFT M.B.H. [AT/AT]; Brunner Strasse 59, A-1230 Vienna (AT).			
<ul> <li>(72) Inventors; and</li> <li>(75) Inventors/Applicants (for US only): NAVARRO, François [FR/FR]; 53, Rue Principale, F-68440 Bruebach (FR). PETIT, Samuel [FR/FR]; 11, Parc de la Risle, F-76130 Mont-Saint-Aignan (FR). STONE, Guy [US/CH]; March- bachstrasse 9, CH-4107 Ettingen (CH).</li> </ul>		(). 30	
(74) Agent: BECKER. Konrad; Novartis AG, Corporate In Property, Patent & Trademark Department, CH–4 (CH).			

#### (54) Title: MACROLIDES

(57) Abstract

Δ

The invention relates to the stabilization of poly-ene macrolides and to a particular macrolide obtained in crystalline form.

#### FOR THE PURPOSES OF INFORMATION ONLY

LS

LT

LU LV

MC MD

MG

МК

ML

MN

MR

MW

МΧ NE

NL

NO

 $\mathbf{NZ}$ PL РТ RO RU SD SE SG

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

Ibania rmenia ustria zerbaijan osnia and Herzegovina arbados elgium urkina Faso ulgaria enin razil elarus anada	ES FI GA GB GE GH GN GR HU IE IL IS
ustria ustria zerbaijan osnia and Herzegovina arbados ełgium urkina Faso ulgaria enin razil elarus	FR GA GB GH GN GR HU IE IL
ustralia zerbaijan osnia and Herzegovina arbados ełgium urkina Faso ulgaria enin razil elarus	GA GB GE GN GR HU IE IL
zerbaijan osnia and Herzegovina arbados elgium urkina Faso ulgaria enin razil elarus	GB GE GH GN GR HU IE IL
osnia and Herzegovina arbados elgium urkina Faso ulgaria enin razil elarus	GE GH GN GR HU IE IL
arbados elgium urkina Faso ulgaria enin razil elarus	GH GN GR HU IE IL
elgium eugium ulgaria enin razil elarus	GN GR HU IE IL
urkina Faso oulgaria enin razil elarus	GR HU IE IL
ulgaria enin razil elarus	HU IE IL
enin razil elarus	IE IL
razil elarus	IL
elarus	
enardo	IS
anada	
anaua	IT
entral African Republic	JP
ongo	KE
witzerland	KG
ôte d'Ivoire	KP
ameroon	
hina	KR
uba	KZ
zech Republic	LC
ermany	LI
•	LK
	LR
	ameroon hina uba zech Republic eirmany Denmark stonia

DOCKE.

Δ

5	Spain
	Finland
R I	France
A	Gabon
В	United Kingdom
E	Georgia
H	Ghana
N	Guinea
R	Greece
U	Hungary
;	Ireland
,	Israel
	Iceland
•	Italy
•	Japan
Е	Kenya
G	Kyrgyzstan
P	Democratic People's
	Republic of Korea
R	Republic of Korea
Z	Kazakstan
C [	Saint Lucia
-	Liechtenstein
К	Sri Lanka
R	Liberia

Lesotho	
Lithuania	
Luxembourg	
Latvia	
Monaco	
Republic of Moldova	
Madagascar	
The former Yugoslav	
Republic of Macedoni	ia
Mali	
Mongolia	
Mauritania	
Malawi	
Mexico	
Niger	
Netherlands	
Norway	
New Zealand	
Poland	
Portugal	
Romania	
Russian Federation	
Sudan	
Sweden	
Singapore	

SI	Slovenia
~-	01010111
SK	Slovakia
SN	Senegal
SZ	Swaziland
TD	Chad
TG	Togo
TJ	Tajikistan
ТМ	Turkmenistan
TR	Turkey
тт	Trinidad and Tobago
UA	Ukraine
UG	Uganda
US	United States of America
UZ	Uzbekistan
VN	Viet Nam
YU	Yugoslavia
ZW	Zimbabwe

Find authenticated court documents without watermarks at docketalarm.com.

WO 00/33878

PCT/EP99/09521

#### MACROLIDES

DOCKE.

The present invention relates to the stabilization of a pharmaceutically active ingredient sensitive to oxidation, e.g. a poly-ene macrolide, preferably a poly-ene macrolide having immunosuppressant properties, particularly rapamycins.

- 1 -

The handling and storage particularly in the bulk form of pharmaceutically active ingredients which are sensitive to oxidation is difficult. Special handling is necessary and often the oxidation-sensitive ingredient is stored in air-tight packaging under protective gas. Substantial amounts of stabilizers are added during the formulating process of such pharmaceutically active ingredients.

Poly-ene macrolides have satisfactory stability properties. However, it has now been found that their stability to oxygen may substantially be improved by the addition of a stabilizer, e.g. an antioxidant, during their isolation step.

According to the invention, there is provided

1. A process for stabilizing a poly-ene macrolide comprising adding an antioxidant to the purified macrolide, preferably at the commencement of its isolation step.

This process is particularly useful for the production of a stabilized poly-ene macrolide in bulk. The amount of antioxidant may conveniently be up to 1%, more preferably from 0.01 to 0.5 % (based on the weight of the macrolide). Such a small amount is referred to hereinafter as a catalytic amount.

As alternatives to the above the present invention also provides:

2. A mixture, e.g. a bulk mixture, comprising a poly-ene macrolide and an anti-oxidant, preferably a catalytic amount thereof, preferably in solid form.

The mixture may be in particulate form e.g. cristallized or amorphous form. It may be in a sterile or substantially sterile condition, e.g. in a condition suitable for pharmaceutical use. 3. Use of a mixture as defined above in 2. in the manufacture of a pharmaceutical composition.

Examples of poly-enes macrolides are e.g. molecules comprising double bonds, preferably conjugated double bonds, for example such having antibiotic and/or immunosuppressant properties, e.g. macrolides comprising a lactam or lactone bond and their derivatives, e.g. compounds which have a biological activity qualitatively similar to that of the natural macrolide, e.g. chemically substituted macrolides. Suitable examples include e.g. rapamycins and ascomycins. A preferred poly-ene macrolide is a macrolide comprising at least 2 conjugated double bonds, e.g. 3 conjugated double bonds.

Rapamycin is a known lactam macrolide produceable, for example by <u>Streptomyces</u> <u>hygroscopicus.</u> The structure of rapamycin is given in Kessler, H. et al.; 1993; Helv. Chim. Acta, <u>76</u> : 117. Rapamycin has antibiotic and immunosuppressant properties. Derivatives of rapamycin are known, e.g. 16-O-substituted rapamycins, for example as disclosed in WO 94/02136 and WO 96/41807, 40-O-substituted rapamycins, for example as disclosed in WO 94/09010, WO 92/05179, WO 95/14023, 94/02136, WO 94/02385 and WO 96/13273, all of which being incorporated herein by reference. Preferred rapamycin derivatives are e.g. rapamycins wherein the hydroxy in position 40 of formula A illustrated at page 1 of WO 94/09010 is replaced by -OR wherein R is hydroxyalkyl, hydroxyalkoxyalkyl, acylaminoalkyl or aminoalkyl, e.g. 40-O-(2-hydroxy)ethyl-rapamycin, 40-O-(3-hydroxy)propyl-rapamycin, and 40-O-[2-(2-hydroxy)ethoxy]ethyl-rapamycin.

Ascomycins, of which FK-506 and ascomycin are the best known, form another class of lactam macrolides, many of which have potent immunosuppressive and anti-inflammatory activity. FK506 is a lactam macrolide produced by <u>Streptomyces tsukubaensis</u>. The structure of FK506 is given in the Appendix to the Merck Index, 11th ed. (1989) as item A5. Ascomycin is described e.g. in USP 3,244,592. Ascomycin, FK506, other naturally occurring macrolides having a similar biological activity and their derivatives, e.g. synthetic analogues and derivatives are termed collectively "Ascomycins". Examples of synthetic analogues or derivatives are e.g. halogenated ascomycins, e.g. 33-epi-chloro-33-desoxy-ascomycin such as disclosed in EP-A-427,680, tetrahydropyran derivatives, e.g. as disclosed in EP-A-626,385.

DOCKET

Particularly preferred macrolides are rapamycin and 40-O-(2-hydroxy)ethyl-rapamycin.

Preferred antioxidants are for example 2,6-di-tert.-butyl-4-methylphenol (hereinafter BHT), vitamin E or C, BHT being particularly preferred.

- 3 -

A particularly preferred mixture of the invention is a mixture of rapamycin or 40-O-(2hydroxy)ethyl-rapamycin and 0.2% (based on the weight of the macrolide) of antioxidant, preferably BHT.

The antioxidant may be added to the poly-ene macrolide at the commencement of the isolation steps, preferably the final isolation step, more preferably just prior to the final precipitation step. The macrolide is preferably in a purified state. It may be dissolved in an inert solvent and the antioxidant is added to the resulting solution, followed by a precipitation step of the stabilized macrolide, e.g. in an amorphous form or in the form of crystals. Preferably the mixture of the invention is in amorphous form.

The resulting stabilized macrolide exhibits surprisingly an improved stability to oxidation and its handling and storage, e.g. in bulk form prior to its further processing for example into a galenic composition, become much easier. It is particularly interesting for macrolides in amorphous form.

The macrolide stabilized according to the invention may be used as such for the production of the desired galenic formulation. Such formulations may be prepared according to methods known in the art, comprising the addition of one or more pharmaceutically acceptable diluent or carrier, including the addition of further stabilizer if required.

Accordingly there is further provided:

DOCKE.

4. A pharmaceutical composition comprising, as active ingredient, a stabilized mixture as disclosed above, together with one or more pharmaceutically acceptable diluent or carrier.

The composition of the invention may be adapted for oral, parenteral, topical (e.g. on the skin), occular, nasal or inhalation (e.g. pulmonary) administration. A preferred

## DOCKET A L A R M



# Explore Litigation Insights

Docket Alarm provides insights to develop a more informed litigation strategy and the peace of mind of knowing you're on top of things.

## **Real-Time Litigation Alerts**



Keep your litigation team up-to-date with **real-time alerts** and advanced team management tools built for the enterprise, all while greatly reducing PACER spend.

Our comprehensive service means we can handle Federal, State, and Administrative courts across the country.

## **Advanced Docket Research**



With over 230 million records, Docket Alarm's cloud-native docket research platform finds what other services can't. Coverage includes Federal, State, plus PTAB, TTAB, ITC and NLRB decisions, all in one place.

Identify arguments that have been successful in the past with full text, pinpoint searching. Link to case law cited within any court document via Fastcase.

## **Analytics At Your Fingertips**



Learn what happened the last time a particular judge, opposing counsel or company faced cases similar to yours.

Advanced out-of-the-box PTAB and TTAB analytics are always at your fingertips.

## API

Docket Alarm offers a powerful API (application programming interface) to developers that want to integrate case filings into their apps.

### LAW FIRMS

Build custom dashboards for your attorneys and clients with live data direct from the court.

Automate many repetitive legal tasks like conflict checks, document management, and marketing.

### FINANCIAL INSTITUTIONS

Litigation and bankruptcy checks for companies and debtors.

## E-DISCOVERY AND LEGAL VENDORS

Sync your system to PACER to automate legal marketing.