WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6 :

C07D 498/18

A1

(11) International Publication Number:

WO 99/01458

(43) International Publication Date:

14 January 1999 (14.01.99)

(21) International Application Number:

PCT/EP98/03929

(22) International Filing Date:

26 June 1998 (26.06.98)

(30) Priority Data:

9713730.1

30 June 1997 (30.06.97)

GB

(71) Applicant (for all designated States except AT US): NOVARTIS AG [CH/CH]; Schwarzwaldallee 215, CH-4058 Basel (CH).

(71) Applicant (for AT only): NOVARTIS-ERFINDUNGEN VER-WALTUNGSGESELLSCHAFT MBH [AT/AT]; Brunner Strasse 59, A-1235 Vienna (AT).

(72) Inventors; and

(75) Inventors/Applicants (for US only): DOSENBACH, Cornelia [DE/DE]; Gänsmattenweg 1/1, D-79588 Efringen-Kirchen (DE). GRASSBERGER, Maximilian [AT/AT]; Schaumburgergasse 12/7, A-1040 Wien (AT). HARTMANN, Otto [CH/CH]; Spalenring 40a, CH-4055 Basel (CH). HORVATH, Amarylla [AT/AT]; Schönbrunnerstrasse 293/2/2, A-1120 Wien (AT). MUTZ, Jean-Paul [FR/FR]; 35, rue des Cigognes, F-68730 Blotzheim (FR). PENN, Gerhard [AT/CH]; Vogesenstrasse 38, CH-4056 Basel (CH). PF-EFFER, Sabine [DE/DE]; Unteres Holz 11, D-79576 Weil

(DE). WIECKHUSEN, Dierk [DE/DE]; Mühlenstrasse 78, D-79589 Binzen (DE).

(74) Agent: BECKER, Konrad; Novartis AG, Patent- und Markenabteilung, Lichtstrasse 35, CH-4002 Basel (CH).

(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

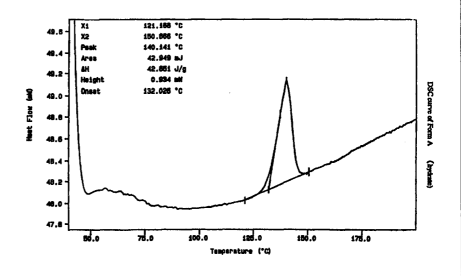
With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: CRYSTALLINE MACROLIDES AND PROCESS FOR THEIR PREPARATION

(57) Abstract

33-Epichloro-33-desoxyascomycin formula (I) and various tautomeric or solvated forms thereof, in crystalline form, such as Form A and Form B. Their preparation involves converting amorphous appropriately compound of formula (I), or compound of formula (I) in other than Form A, or compound of formula (I) in other than Form B, respectively, from a solution thereof under crystallization-inducing conditions conditions or inducing preferential crystallization of Form A or B, respectively. Such crystals are particularly indicated for use in the preparation of topical galenical forms of the compound for pharmaceutical use, e.g. creams, emulsions and ointments.



FOR THE PURPOSES OF INFORMATION ONLY

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

| AL | Albania | ES | Spain | LS | Lesotho | SI | Slovenia |
|----|--------------------------|----|---------------------|------------------------|-----------------------|----|--------------------------|
| AM | Armenia | FI | Finland | LT | Lithuania | SK | Slovakia |
| AT | Austria | FR | France | LU | Luxembourg | SN | Senegal |
| AU | Australia | GA | Gabon | LV | Latvia | SZ | Swaziland |
| ΑZ | Azerbaijan | GB | United Kingdom | MC | Monaco | TD | Chad |
| BA | Bosnia and Herzegovina | GE | Georgia | MD | Republic of Moldova | TG | Togo |
| BB | Barbados | GH | Ghana | MG | Madagascar | TJ | Tajikistan |
| BE | Belgium | GN | Guinea | MK | The former Yugoslav | TM | Turkmenistan |
| BF | Burkina Faso | GR | Greece | | Republic of Macedonia | TR | Turkey |
| BG | Bulgaria | HU | Hungary | ML | Mali | TT | Trinidad and Tobago |
| BJ | Benin | IE | Ireland | MN | Mongolia | UA | Ukraine |
| BR | Brazil | IL | Israel | MR | Mauritania | UG | Uganda |
| BY | Belarus | IS | Iceland | MW | Malawi | US | United States of America |
| CA | Canada | IT | Italy | MX | Mexico | UZ | Uzbekistan |
| CF | Central African Republic | JP | Japan | NE | Niger | VN | Viet Nam |
| CG | Congo | KE | Kenya | NL | Netherlands | YU | Yugoslavia |
| CH | Switzerland | KG | Kyrgyzstan | NO | Norway | ZW | Zimbabwe |
| CI | Côte d'Ivoire | KP | Democratic People's | NZ | New Zealand | | |
| CM | Cameroon | | Republic of Korea | PL | Poland | | |
| CN | China | KR | Republic of Korea | PT | Portugal | | |
| CU | Cuba | KZ | Kazakstan | RO | Romania | | |
| CZ | Czech Republic | LC | Saint Lucia | RU | Russian Federation | | |
| DE | Germany | LI | Liechtenstein | SD | Sudan | | |
| DK | Denmark | LK | Sri Lanka | SE | Sweden | | |
| EE | Estonia | LR | Liberia | $\mathbf{s}\mathbf{G}$ | Singapore | | |
| | | | | | | | |
| | | | | | | | |
| | | | | | | | |

WO 99/01458 PCT/EP98/03929

Crystalline macrolides and process for their preparation

The invention relates to macrolide chemistry. It concerns the compound of formula I

i.e. {[1E-(1R,3R,4S)]1R,9S,12S,13R,14S,17R,18E,21S,23S,24R,25S,27R}-12-[2-(4-chloro-3-methoxycyclohexyl)-1-methylvinyl]-17-ethyl-1,14-dihydroxy-23,25-dimethoxy-13,19,21,27-tetramethyl-11,28-dioxa-4-azatricyclo[22.3.1.0(4,9)]octacos-18-ene-2,3,10,16-tetraone,

hereinafter briefly named

"33-epichloro-33-desoxy-FR520" or "33-epichloro-33-desoxyascomycin", in crystalline form.

For simplicity, formula I as referred to herein should be understood as including the compound of formula I in the various tautomeric forms with which it is in equilibrum, particularly in solution, and solvated, e.g. hydrated forms, such as the tautomeric forms of formula

and of formula

The compound of formula I is known in amorphous form, e.g. from Sandoz EP 427 680, Example 66a in the form of a colourless foamy resin [with ¹H-NMR = 4.56 (m, H-33)], and from Merck EP 480 623, Example 53 (without any physicochemical characterization). Various names and carbon atom numberings are used in the literature.

Prior to the present invention, the compound of formula I had never been recovered in crystalline form.

It appears that the presence of a halogen atom, especially chlorine in the cyclohexyl moiety of the molecule, particularly in the 4 position thereof (also marked as position 33 in formulae I and Ic herein), has an unfavourable effect on the crystallization properties of this structural class of compounds. Thus in EP 427 680 none of the halogenated final products is obtained in crystalline form, they are colourless foams or foamy resins, and characterized by their NMR spectra.

Similarly, in EP 480 623, which covers exclusively macrolide end products halogenated in the cyclohexyl moiety, none of the specific compounds disclosed is characterized by data indicative of crystallinity, such as a melting point; most end products therein are not characterized by any physicochemical data at all, and those that are characterized, are characterized by their mass spectra, which are not indicative as regards physical state; and none of the 4-chloro end products disclosed is characterized at all.

Further analogous macrolides halogenated in the cyclohexyl moiety are also disclosed in e.g. Fisons WO 91/13889, specifically, as Examples 42a), 42b) and 49a): the compounds therein are also not obtained in crystalline form, but recovered as a foam or an oil.

Overall, the 23-membered tricyclomacrolides derived from FK 506 are obtainable in crystalline form only with difficulty, if at all, as appears also from e.g. Merck WO 97/8182, concerning a macrolide compound having a basic substituent capable of forming salts, which could be obtained in crystalline form, but as a tartrate salt. The compound of the present invention is devoid of such a basic substituent.

It is thus surprising that crystallization of the compound of formula I has now been successfully achieved.

The invention concerns the **compound of formula I in crystalline form**. The crystalline form may appear as solvated, e.g. hydrated, or anhydrous form, or be a tautomer.

While the first recovery of the compound of formula I in crystalline form occurred several years after the first synthesis of the compound, initially obtained only in amorphous form, it has turned out that subsequently to its first crystallization, the compound could be induced to crystallize from the amorphous form quite readily. The crystalline material has thus now become easily accessible, using a variety of experimental conditions extending beyond the initially used recrystallization conditions, which involved the addition of water to an ethanolic solution of the amorphous compound.

The invention also concerns a process for the preparation of the compound of formula I, or a tautomeric or solvated form thereof, in crystalline form which comprises appropriately converting amorphous compound of formula I from a solution thereof under crystallization-inducing conditions.

It also concerns the compound of formula I, or a tautomeric or solvated form thereof, in crystalline form whenever prepared by that process, and the compound of formula I in a non-crystalline, e.g. in dissolved state, or a tautomeric or solvated form thereof, whenever produced from a crystalline form.

The process of the invention is effected in conventional manner. The precise conditions under which crystals are formed may now be empirically determined and a number of methods are suitable in practice, including the initial addition of water to an ethanolic solution of the compound of formula I in amorphous form.

Crystallization-inducing conditions normally involve the use of an appropriate crystallization-inducing solvent, such as methanol, ethanol, isopropanol or water or mixtures thereof. Conveniently, the amorphous compound is dissolved in the solvent at a temperature of normally at least 10°C. The solution may be produced by dissolving in a solvent any one or more of amorphous forms of the compound, and solvates thereof, such as hydrates, methanolates, ethanolates, isopropanolates and acetonitrilates. Crystals may then be formed by conversion from solution, crystallization taking place at a temperature of between about 10°C and the boiling point of the solvent. The dissolution and crystallization may be carried out in various conventional ways. For instance, amorphous compound may be dissolved in a solvent or a mixture of solvents in which it is readily soluble at elevated temperatures but in

which it is only sparingly soluble at lower temperatures. Dissolution at elevated temperature is followed by cooling during which the desired crystals crystallize out of solution. Solvents which are suitable include esters such as methyl acetate and ethyl acetate, toluene and acetonitrile. Mixed solvents comprising a good solvent in which the compound is readily soluble, preferably, in amounts of at least 1 % by weight at 30°C, and a poor solvent in which it is more sparingly soluble, preferably in amounts of not more than about 0.01 % by weight at 30°C, may also be employed provided that crystallization from the mixture at a reduced temperature, of normally at least about 10°C, is possible using the selected solvent mixture.

Alternatively, the difference in solubility of the crystals in different solvents may be used. For example, the amorphous compound may be dissolved in a good solvent in which it is highly soluble such as one in which it is soluble in amounts of at least 1 % by weight at about 30°C, and the solution subsequently mixed with a poor solvent in which it is more sparingly soluble, such as one in which it is soluble in amounts of not more than about 0.01 % by weight at about 30°C. Thus, the solution of the compound in the good solvent may be added to the poor solvent, while maintaining normally a temperature in excess of about 10°C, or the poor solvent may be added to the solution of the compound in the good solvent, again while normally maintaining a temperature in excess of about 10°C. Examples of good solvents include lower alcohols, such as methanol, ethanol and isopropanol, as well as acetone, tetrahydrofuran and dioxane. Examples of poor solvents are water, hexane and diethyl ether. Preferably, crystallization is effected at a temperature in the range of about 10°C to about 60°C.

In an alternative embodiment of the process of the invention, solid amorphous compound is suspended at a temperature of normally at least about 10°C in a solvent in which it is incompletely soluble, preferably only sparingly soluble, at that temperature. A suspension results in which particles of solid are dispersed, and remain incompletely dissolved in the solvent. Preferably the solids are maintained in a state of suspension by agitation e.g. by shaking or stirring. The suspension is kept at a temperature of normally about 10°C or higher in order to effect a transformation of the starting solids into crystals. The amorphous solid compound suspended in a suitable solvent may be a solvate, e.g. hydrate, methanolate, ethanolate, isopropanolate or acetonitrilate. The amorphous powder may be derived by drying a solvate.

It is preferred to add "seeds" of crystalline material to the solution in order to induce crystallization.

The compound of formula I in crystalline form can readily be isolated, it can e.g. be filtered off or centrifuged from the crystallization medium, if desired after cooling, and washed and dried, and optionally further recrystallized using similar conditions.

While the initial recovery has resulted in material in a crystalline form designated as "Form A" herein, surprisingly, it has turned out upon further investigation that at least one additional crystal form of the compound may be recovered, herein designated as "Form B", which differs from Form A in various characteristics, such as its solubility. The invention thus concerns the compound of formula I or a tautomeric or solvated form thereof in **crystalline** form as such, and more particularly Form A and Form B. Form A is preferred.

Form A normally is in hydrated form at room temperature. The hydrated form can be reversibly dehydrated by heating to about 110°C. It remains in Form A thereby. The hydrated form is the more stable state of Form A at room temperature. Form B normally is not in hydrated form, even at room temperature. It is thermodynamically a more stable form than Form A.

A crystal form is defined herein as being "crystallographically pure" when it contains at most about 0.5 % (w/w), e.g. at most about 0.1 % (w/w) of other form. Thus e.g. "crystallographically pure Form A" contains about ≤ 0.5 % (w/w), .g. about ≤ 0.1 % (w/w) of Form B and/or amorphous form.

The preparation of Forms A and B may be effected using conventional means, starting either from amorphous material or from Form B or Form A, respectively, or mixtures thereof. Normally, the starting material is dissolved into an appropriate solvent and crystallized or recrystallized therefrom under conditions preferentially producing either Form A or Form B, resulting in crystallographically pure Form A or Form B.

The invention thus includes a process variant for the preparation of the compound of formula I, or a tautomeric or solvated form thereof, in crystalline Form A which comprises appropriately converting compound of formula I in other than Form A, or a tautomeric or solvated form thereof, from a solution thereof under conditions inducing

preferential crystallization of Form A. It also concerns the compound of formula I in Form A whenever prepared by that process variant.

Conversely, the invention includes a process variant for the preparation of the compound of formula I, or a tautomeric or solvated form thereof, in crystalline Form B which comprises appropriately converting compound of formula I in other than Form B from a solution thereof under conditions inducing preferential crystallization of Form B. It also concerns the compound of formula I in Form B whenever prepared by that process variant.

For the preparation of **Form A** the starting material is conveniently dissolved in an appropriate solvent, preferably ethanol/water, preferably in the proportions 9.5: 0.5. The temperature for dissolution is from about 60°C to about 75°C, preferably about 70°C. The proportion of starting material to solvent preferably is from about 1:5 to about 1:6 on a weight basis, preferably about 1:5 (w/w). The solution is filtered and then cooled to a reduced temperature, preferably of from about 70°C to about 20°C, preferably about 10°C, and a liquid in which Form A is insoluble, such as water, is carefully added. A supersaturated solution results thereby. While crystals of Form A may spontaneously be formed, preferably the supersaturated solution is seeded with a few crystals of crystallographically pure Form A. It is usually beneficial to check the purity of the seeding crystals with a melt microscope. Further addition of liquid under careful stirring leads to more crystals of Form A. Low temperature, i.e. below about 20°C, and seeding with crystallographically pure crystals of Form A appear to prevent the formation of crystals of Form B. Too lengthy stirring may be counter-productive, particularly at temperatures above 10°C, Form B being the thermodynamically more stable form.

Conveniently, as a preliminary step, the starting material is preferably thoroughly dissolved in a polar organic solvent such as an alcohol, e.g. methanol, ethanol, isopropanol, preferably ethanol, or in acetone, especially in acetone, preferably at boiling temperature, and the solvent evaporated to dryness.

For the preparation of **Form B** the starting material is again dissolved in a solvent as described above for preparing Form A, preferably ethanol/water 9.5 : 0.5 (v/v). The temperature for dissolution is again from about 60°C to about 75°C, preferably about 70°C,

and the resultant solution is filtered. The proportion of starting material to solvent is somewhat less than for preparing Form A, it is preferably about 1:7 (w/w). However, cooling is to a higher temperature than when preparing Form B, it is preferably to above 20°C, e.g. to about 25° or 30°C, and the further workup is also effected at about that temperature or a similar temperature. Seeding with crystals of Form B is optional, but greatly facilitates crystallization and allows more latitude as regards e.g. temperature. The speed of formation of the supersaturated solution appears also to exert some effect on the result obtained, speedy supersaturation resulting in increased formation of Form B.

A solvate, e.g. a hydrate, may be converted into the corresponding unsolvated form in conventional manner and vice-versa, e.g. by appropriately heating up the solvated form, or cooling down the unsolvated form of the crystal form susceptible of being solvated.

The two crystal forms identified are characterized i.a. by the following physico-chemical data:

1) **Form A**:

- appearance: white to off-white, finely crystalline powder (from ethanol/water);
- m.p. determined by DSS (10°K/min): melting onset at about 132°C;
- solubility (at 5°C): water: insoluble methanol, ethanol, ethyl acetate, diethylether, diisopropyl ether: > 100 mg/ml hexane: < 10 mg/ml;
- solubility (at 25°C): acetone, acetonitrile, ethanol, ethyl acetate, isopropanol, methanol: > 50 mg/ml; water: < 1 mg/ml;
- solubility in the oil phase of cream (oleyl alcohol/miglyol 812^R 4 : 6): 2.49 %;
- chemical purity: 98.5 %;
- thermogravimetry: loss of mass on drying up to melting: 1.46 % (Carl Fischer titration);
- morphology (SEM): sticks and agglomerates (1-100 μm);
- hygroscopicity (uptake determined by thermogravimetry):

```
1.49 % (1 day, 92 % r.h.); 1.78 % (1 week, 25°C, 75 % r.h.);
```

- DSC curve: see **Figure 1** (Perkin Elmer DSC-7 differential scanning calorimeter; measurement from 40°C to 200°C, scan heating rate 10° K/min);

- FT-IR spectrum: see Figures 3 and 5 (PE FT-IR spectrometer 1725X; KBr, paraffin oil; scan range 4000-400 cm⁻¹);
- X-ray powder diffraction pattern: see Figure 6 (Scintag XDS 2000 powder diffractometer; Scintag, Santa Clara, CA, USA); scan speed 0.5° or 1°/min (2 theta value);

2) **Form B**:

- m.p. determined by DSC (10°K/min): melting onset at about 159°C;

- solubility (at 5°C): water: 0.3 mg/ml methanol: 46.3 mg/ml ethanol: 18.1 mg/ml ethyl acetate: > 50 mg/ml diethylether: 9.3 mg/ml; diisopropylether: 1.9 mg/ml

hexane: 0.8 mg/ml;

- solubility (at 25°C): water: 0.4 mg/ml methanol: > 50 mg/ml ethanol: 34.4 mg/ml ethyl acetate: > 50 mg/ml diethyl ether: 16.3 mg/ml diisopropyl ether: 3.1 mg/ml

hexane: 1.5 mg/ml;

- solubility in the oil phase of TMF cream (oleyl alcohol/myglyol $812^R\ 4:6$): 0.37 %;
- chemical purity: 99.9 %;
- thermogravimetry: loss of mass on drying up to melting: < 0.05 %;
- morphology (SEM): needles;
- hygroscopicity (uptake determined by thermogravimetry): 1 day, 92 % r.h. and 1 week, 25°C, 75 % r.h.: none;
- DSC curve: see Figure 2 (Perkin-Elmer DSC-7; 40°C to 200°C; scan heating rate 10° K/min);
- FT-IR spectrum: see Figures 4 and 5 (PE FT-IR spectrometer 1725X; KBr; paraffin oil; scan range 4000-400 cm⁻¹);
- X-ray powder diffraction pattern: see Figure 6 (Scintag XDS 2000 powder diffractometer); scan speed 0.5° or 1°/min (2 theta value).

3) For <u>reference</u>, the corresponding FT-IR spectrum of the amorphous form is indicated in Figure 7.

Characterization data for all forms of the compound is further as follows:

- optical rotation: $\left[\alpha\right]_{D}^{20} = -48.0^{\circ} (\pm 0.2^{\circ}) (CDCl_3);$
- TLC: $R_f = 0.18$ (silicagel; hexane/ethyl acetate 2:1) $R_f = 0.62$ (silicagel; hexane/ethyl acetate 1:1);
- 1 H-NMR (CDCl₃): Two conformers (Z:E = 1:2). Characteristic signals of the major conformer d [ppm]: 5.35 (d,J = 1.7 Hz, H-26, 5.12 (d,J = 9.0 Hz, H-29), 5.05 (d,J = 9.4 Hz, H-20), 4.60 (d,J = 5.0 Hz, H-2), 4.56 (m,w_{1/2} = 10 Hz, H-33), 4.43 (d, J = 13.8 Hz, H-6e), 3.66 (dd, J = 9.6 Hz, J = 1.0 Hz, H-14), 3.92 (m,H-24), 2.80 (dd, J = 15.9 Hz, J = 2.7 Hz, H-23a);
- 13 C-NMR (CDCl₃): Two conformers (Z:E = 1:2). Signals of the major conformer d [ppm]: 213.7 (C-22), 196.3 (C-9), 169.1 (C-1), 164.8 (C-8), 138.8 (C-19), 132.5 (C-28), 129.2 (C-29), 122.0 (C-20), 97.0 (C-10), 79.2 (C-32), 76.7 (C-26), 75.2 (C-15), 73.7 (C-14), 72.9 (C-13), 70.2 (C-24), 59.3 (C-33), 56.7 (C-2), 54.7 (C-21), 48.6 (C-18), 39.2 (C-6), 42.7 (C-23), 39.4 (C-25), 34.7 (C-30), 34.6 (C-11), 32.8 (C-12), 32.1 (C-35), 31.7 (C-34), 27.7 (C-3), 26.4 (C-17), 25.5 (C-31), 24.5 (C-5), 24.2 (C-36), 21.1 (C-4), 20.6 (17-Me), 16.2 (11-Me), 15.9 (19-Me), 14.2 (28-Me), 11.7 (C-37), 9.3 (25-Me).

In the above NMR spectra the carbon atom numbering is as appears in formula Ic hereafter:

Abbreviations:

DMSO:

dimethylsulfoxide

DSC:

differential scanning calorimetry

FT-IR:

Fourier transformed infrared

m.p.: r.h.: melting point relative humidity

SEM:

scanning electron microscopy

T:

transmission

TG:

thermogravimetry

THF:

tetrahydrofuran

TLC:

thin layer chromatography

Explanation of the Figures:

Figure 1: DSC curve of Form A (hydrate)

Figure 2: DSC curve of Form B (anhydrous)

Figure 3: FT-IR spectrum of Form A (hydrate)

(% T = percentage transmission)

Figure 4: FT-IR spectrum of Form B

(% T = percentage transmission)

Figure 5: Comparison between FT-IR spectra of forms A (hydrate) and B

(% T = percentage transmission)

Form A (hydrate) = graph starting at 90 % T

Form B = graph starting at 82 % T

Figure 6: X-ray powder diffraction pattern of forms A (hydrate) and B

Mod. A = Form A (hydrate)

Mod. B = Form B

left ordinate: intensity (cps = counts per second) right ordinate: relative intensity (% = percentage)

top abscissa: resolution

bottom abscissa: 2 theta angle (degree)

Figure 7: FT-IR spectrum of amorphous form (% = percentage transmission)

The compound of formula I in amorphous form is known as a pharmaceutical, in particular for use as an anti-inflammatory, immunosuppressant and antiproliferative agent, both systemically or topically. The preparation of suitable galenical forms for pharmaceutical use, such as creams, emulsions and ointments is, however, difficult. Thus the amorphous form of the compound suffers from problems of instability, such as in bulk, and is generally not well-suitable for galenical processing compared to an exactly defined, crystalline form, e.g. as regards degradation of the bulk material, hygroscopicity, dissolution properties, and overall purity of the material.

The availability of well-defined crystalline forms of the compound of formula I is therefore particularly indicated for use in the preparation of galenical forms of the compound where it is indicated to overcome the above disadvantages, such as in the preparation of topical forms, e.g. creams, emulsions and ointments where it is desired to include the compound normally in dissolved state but under carefully controlled conditions. Thus in the preparation of a cream, it has been observed that Form A dissolves in the oil phase in about 10 minutes, whereas for Form B it takes about 6 hours. It is therefore very advantageous to use crystalline product of well-defined characteristics, whereby Form A or Form B is preferably employed, depending on the particular application, e.g. Form A where a lower melting point or more pronounced solubility is desired, or Form B where a higher melting point, or thermodynamically more stable product at room temperature is appropriate.

Beneficial effects gained with the crystalline form are e.g.:

- less solvent residue in the ultimate drug substance in whatever form, such as dissolved state;
- additional purification effect obtained by crystallization;
- higher stability of the drug substance; and
- easier handling in the production plant.

The compound of formula I in crystalline form may be formulated for administration in any convenient way. It preferably is in dissolved state in the ultimate galenical form.

The invention thus also includes **pharmaceutical compositions** comprising, or whenever prepared from, the compound of formula I or a tautomeric or solvated form thereof, in crystalline form, such as Form A and Form B. It also includes the compound of formula I or a tautomeric or solvated form thereof, in crystalline form **for use as a pharmaceutical**, or **for use in the preparation of a medicament** with anti-inflammatory, imunosuppressant and antiproliferative activity.

The following Examples illustrate the invention but are not limitative. All temperatures are in degrees Centigrade unless indicated otherwise.

Example 1: Crystalline 33-epichloro-33-desoxyascomycin (Form A)

(from a solution of amorphous product in ethanol)

To a solution of 27 g amorphous 33-epichloro-33-desoxyascomycin (colourless foamy resin) in 180 ml of ethanol at room temperature is carefully added water until a transient cloudiness appears (approximately 65 ml of water). The solution is left undisturbed for 16 hours at 4°. Colourless crystals are formed. 10 ml of water are added and the mixture is left undisturbed for another 4 hours at 4°. The crystalline material is separated by succion, washed with an ice-cold mixture of ethanol and water 1:1 (v/v), and dried under reduced pressure (12 mm Hg) for 20 hours at room temperature. The **title compound** is obtained (yield 18g; m.p. 135-136°; chemical purity \geq 98 %, i.e. impurity level at or below analytically detectable limits upon HPLC; hydrate).

Example 2: Solubilization of Forms A and B in 1 % creams

a) The solubility of Form A in a cream formulation is estimated to be about 1 % at room temperature. In the manufacturing of cream, the drug substance is completely dissolved in the oily phase at 60-75°. To assess whether and after which storage time part of the drug substance dissolved will crystallize from the 1 % cream, a series of batches is examined for crystals. The investigation of 10 batches shows that no crystals are observed after manufacturing as well as after storage at 5°, 25°, and 40°. Even in the samples which are subjected to a temperature cycling test for about 3 months no crystals are detected. Only in one preformulation (1 %) drug substance crystals very sparsely occurred after one-year-storage at 25°, which indicates that the 1 % cream is at the borderline with respect to saturation with drug substance.

The drug substance used in the batches and preformulations mentioned above contained 100 % Form A.

b) To assess whether drug substance containing Form B can be completely dissolved in the manufacturing of cream and to evaluate the long-term crystallization behaviour, several preformulations (cream, 1 %) containing 0 %, 1 %, 5 % and 10 % Form B relative to the total content of compound of formula I are manufactured and put on stability testing at different temperatures.

During the preparation of these preformulations, it is observed that the solving speed of crystal Form B in the oily phase of the cream is much slower than that of crystal Form A. Immediately after manufacturing, the forms are investigated and no remaining crystals are observed.

To get information about the long-term crystallization behaviour of these forms, the stability samples are further investigated after 6 weeks, 3, 6 and 9 months storage. The results show that no crystals form even upon prolonged storage.

Example 3: Crystalline form A of 33-epichloro-33-desoxyascomycin

(from a solution in ethanol/isopropanol/water)

10 g of 33-epichloro-33-desoxyascomycin Form B (or, alternatively, crude Form A, or amorphous material) are dissolved 1:5 (w/w) in 10 g of a mixture of ethanol/isopropanol/water 9:0.5:0.5 (v/v) at 70° and the mixture is submitted to clearfiltration on a 0.5 μm filter. The resultant solution free of crystallization germs is allowed to cool to 10°, water is added (25 % w/w based on amount of product) to supersaturate the solution. Seeding is effected with 0.06 g of crystals of Form A (following checking with a thermal microscope) and water is carefully added (7.5 times excess based on the amount of product) over a period of 4 hours at 10°, and thereafter the solution is stirred for 2 hours at that temperature. The **title compound** is obtained (hydrate; 9.3 g).

It is desirable to seed with crystals of Form A as crystallographically pure as possible. Thus seeding with crystals of Form A containing 2 % crystals of Form B as impurity can lead to recovery of crystals of Form A containing 20 % crystals of Form B as impurity. However, at low temperature the formation of the thermodynamically more stable Form B is clearly inhibited: thus seeding at 0° with a 1:1 mixture of both Forms A and B results in a product containing 75 % of Form A.

Example 4: Crystalline form B of 33-epichloro-33-desoxyascomycin

(from a solution in ethanol/isopropanol/water)

25 g of 33-epichloro-33-desoxyascomycin Form A (hydrate) (or, alternatively, crude Form B, or amorphous material) are dissolved 1:7 (w/w) in 75 g of a mixture of ethanol/isopropanol/water 9:0.5:0.5 (v/v) at 70° and the mixture is submitted to clearfiltration. The resultant solution is allowed to cool to 30°, and optionally seeded with 0.1 g of crystals of Form B. Water (3.5 times excess based on the amount of product) is carefully added over a period of 4 hours; the **title compound** is obtained (24.6 g).

Example 5: Crystalline form A of 33-epichloro-33-desoxyascomycin

(from a solution of crude product in acetone)

A solution of 94.5 g of **crude 33-epichloro-33-desoxyascomycin** (e.g. as obtained following chromatography) (containing various amounts of Form A and/or Form B and/or amorphous product) in 1500 ml of acetone is evaporated at 48-52° (400 - 50 mbar). The resultant amorphous foam is dissolved in 500 ml of ethanol/isopropanol 9.5 : 0.5 (v/v). The solution is evaporated at 48-52° (400 - 50 mbar). The resultant foam is dissolved in 400 ml of ethanol/isopropanol 9.5 : 0.5 (v/v) and the hot solution (70-75°) is filtered on a 0.45 μm filter. The solution is cooled to 20-25° over 30-40 minutes, 200 ml of water is added and the mixture is seeded with Form A crystals. Stirring is continued for 1 hour at 20-25°. A suspension is formed which is cooled to 0-5° and further stirred for 4 hours. The crystals are filtered and washed with 300 ml of precooled ethanol/water 1:3 (v/v). Drying at 45-50° (10-20 mbar, 16 hours) gives the **title compound** (hydrate; colourless crystals).

Example 6: Crystalline form A of 33-epichloro-33-desoxyascomycin

(from a solution of Form B in acetone)

94.5 g of 33-epichloro-33-desoxyascomycin Form B are dissolved in 1500 ml of boiling acetone. The acetone solution (now free of any seeds of Form B) is then treated as described in Example 5 above. The title compound is obtained (hydrate; colourless crystals).

Claims:

1. The compound of formula I

-16-

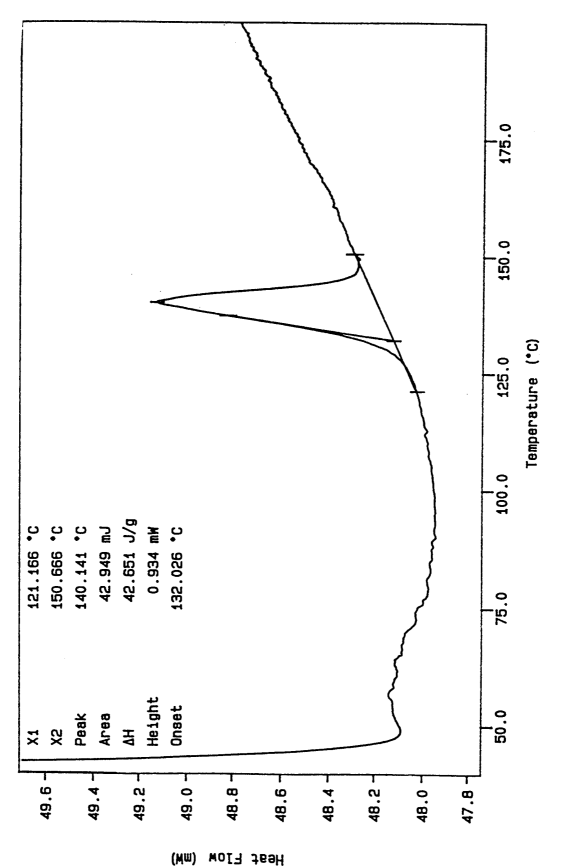
or a tautomeric or solvated form thereof, in crystalline form.

- 2. The compound according to claim 1 in crystalline Form A.
- 3. The compound according to claim 1 in crystalline Form B.
- 4. A process for the preparation of the compound according to claim 1 which comprises appropriately converting amorphous compound of formula I from a solution thereof under crystallization-inducing conditions.
- 5. A variant of the process according to claim 4 for the preparation of the compound according to claim 2 which comprises appropriately converting compound of formula I in other than Form A, or a tautomeric or solvated form thereof, from a solution thereof under conditions inducing preferential crystallization of Form A.

- 6. A variant of the process according to claim 4 for the preparation of the compound according to claim 3 which comprises appropriately converting compound of formula I in other than Form B, or a tautomeric or solvated form thereof, from a solution thereof under conditions inducing preferential crystallization of Form B.
- 7. A pharmaceutical composition comprising, or whenever prepared from, the compound according to any one of claims 1 to 3.
- 8. The compound according to any one of claims 1 to 3 for use as a pharmaceutical, or for use in the preparation of a medicament.
- 9. The compound according to any one of claims 1 to 3 whenever prepared by a process according to any one of claims 4 to 6.
- 10. The compound of formula I depicted in claim 1, in a non-crystalline state, or a tautomeric or solvated form thereof, whenever produced from a crystalline form according to claim 1.

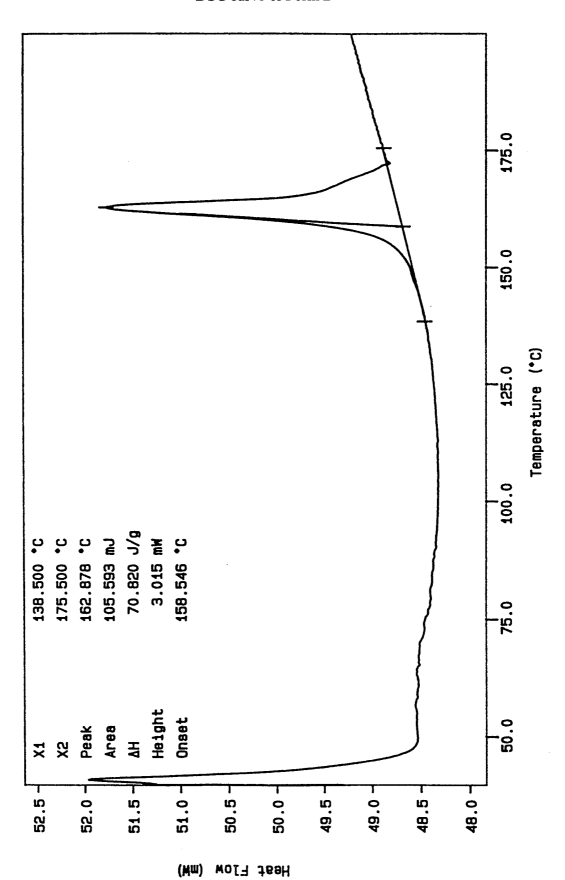
1/7
Figure 1

DSC curve of Form A (hydrate)

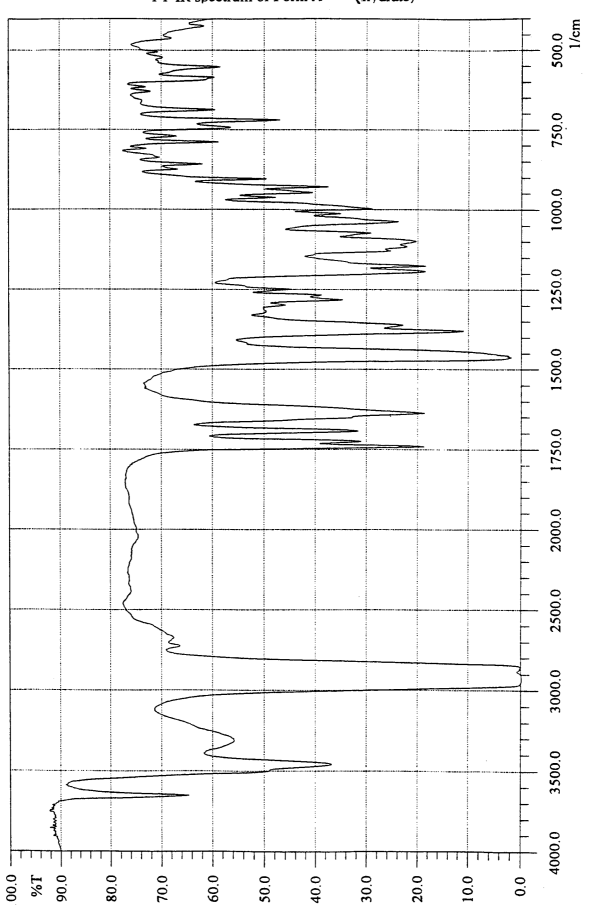


2 / 7 <u>Figure 2</u>

DSC curve of Form B

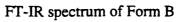


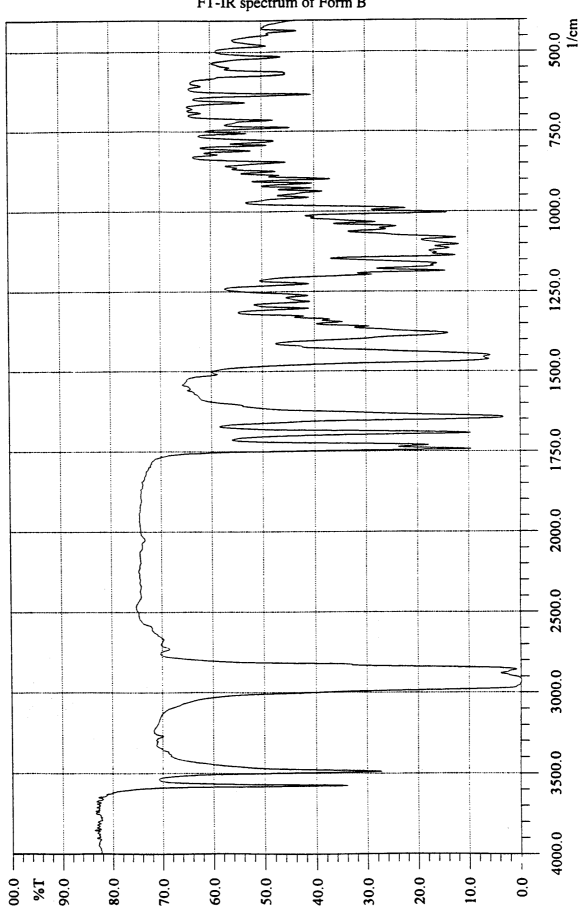
FT-IR spectrum of Form A (hydrate)



4/7

Figure 4





5 / 7 <u>Figure 5</u>

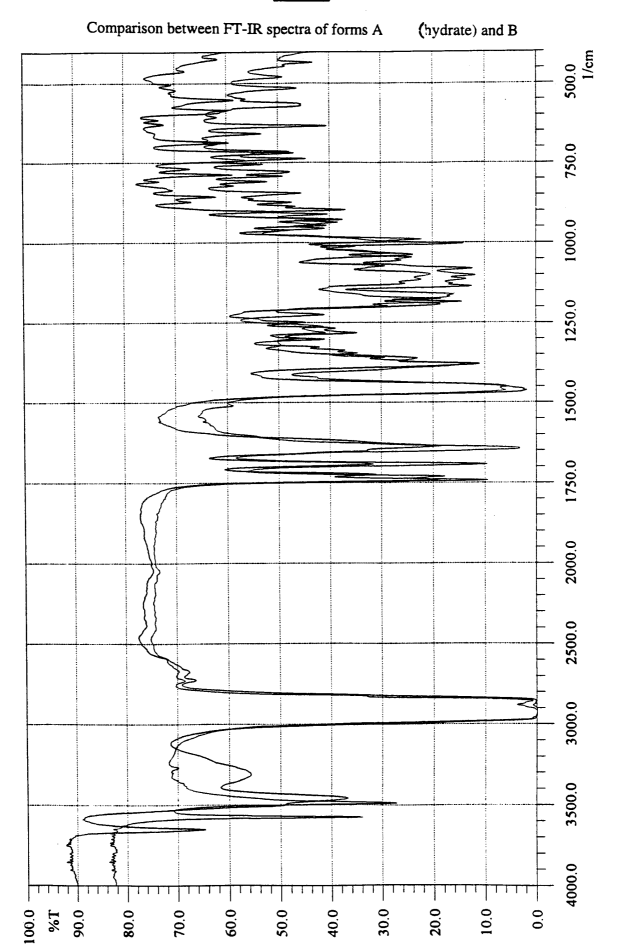
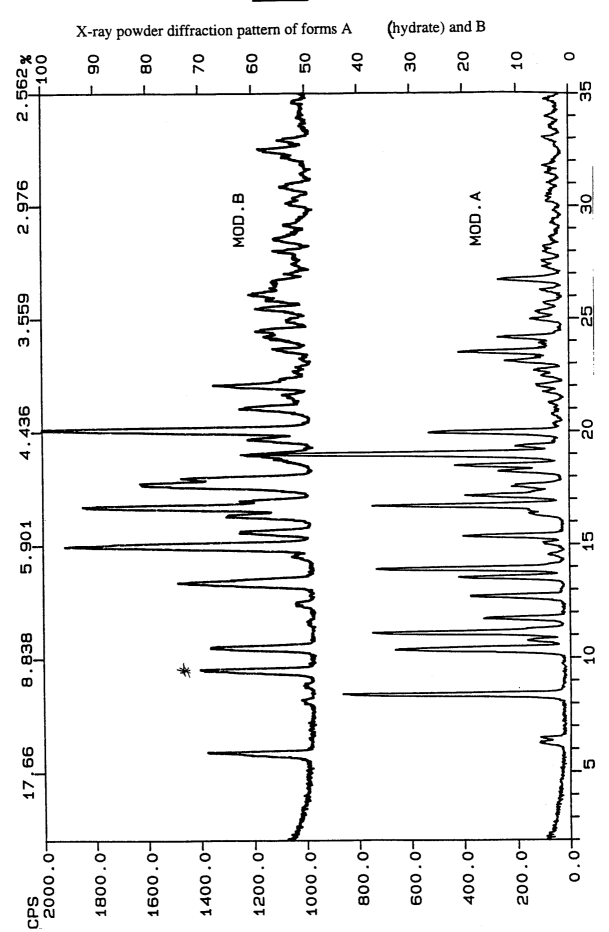


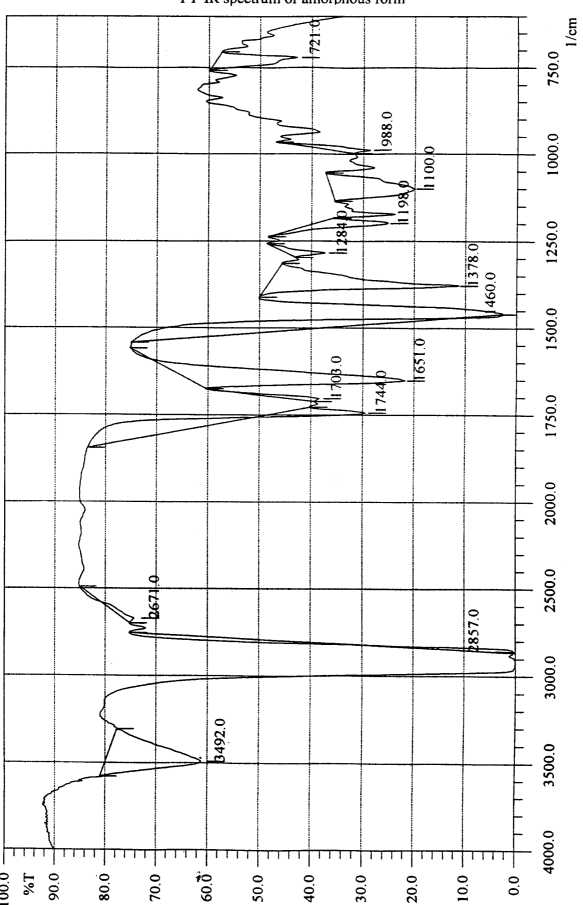
Figure 6



7/7

Figure 7

FT-IR spectrum of amorphous form



national Application No PCT/EP 98/03929

| A. CLASSIFICATION OF SUBJECT MATTER IPC 6 C07D498/18 | | | | | | | | |
|--|---|---|---|--|--|--|--|--|
| | | | | | | | | |
| According to | According to International Patent Classification (IPC) or to both national classification and IPC | | | | | | | |
| | SEARCHED | A. A. Market - 1-3 | | | | | | |
| IPC 6 | ocumentation searched (classification system followed by classificati C07D | on symbols) | | | | | | |
| Documental | tion searched other than minimumdocumentation to the extent that s | such documents are included in the fields sear | rched | | | | | |
| | | | | | | | | |
| Electronic d | ata base consulted during the international search (name of data ba | ise and, where practical, search terms used) | | | | | | |
| C. DOCUM | ENTS CONSIDERED TO BE RELEVANT | , | | | | | | |
| Category ° | Citation of document, with indication, where appropriate, of the rel | evant passages | Relevant to claim No. | | | | | |
| Х | EP 0 427 680 A (SANDOZ LTD ;SAND (DE); SANDOZ AG (AT)) 15 May 199 cited in the application | | 1-3,9,10 | | | | | |
| Υ | see the whole document | | 4-8 | | | | | |
| Х | EP 0 480 623 A (MERCK & CO INC) 15 April 1992 | | 1-3,9,10 | | | | | |
| Υ | cited in the application see the whole document | | 4-8 | | | | | |
| Υ | GIRON,D.: "Thermal analysis and calorimetric methods in the characterisation of polymorphs a solvates" THERMOCHIM.ACTA, vol. 248, 1995, pages 1-59, XPOO AMSTERDAM see the whole document | 1-10 | | | | | | |
| | | -/ | | | | | | |
| X Furt | her documents are listed in the continuation of box C. | X Patent family members are listed in | ı annex. | | | | | |
| "A" docume consid "E" earlier of filing of "L" docume which | stegories of cited documents : ant defining the general state of the art which is not dered to be of particular relevance document but published on or after the international date ant which may throw doubts on priority claim(s) or is cited to establish the publicationdate of another n or other special reason (as specified) | "T" later document published after the interror priority date and not in conflict with to cited to understand the principle or the invention "X" document of particular relevance; the cleannot be considered novel or cannot involve an inventive step when the document of particular relevance; the cleans to the comment of particular relevance. | the application but ory underlying the aimed invention be considered to current is taken alone aimed invention | | | | | |
| "O" document referring to an oral disclosure, use, exhibition or document is combined with one or more other such docu- | | | | | | | | |
| other means "P" document published prior to the international filing date but later than the priority date claimed "at document member of the same patent family | | | | | | | | |
| Date of the | actual completion of theinternational search | Date of mailing of the international sear | ch report | | | | | |
| 2 | 2 October 1998 | 10/11/1998 | | | | | | |
| Name and r | mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2046, Tx. 31 651 epo nl, | Authorized officer Stellmach. J | | | | | | |

2

national Application No
PCT/EP 98/03929

| C.(Continu | ation) DOCUMENTS CONSIDERED TO BE RELEVANT | |
|------------|---|-----------------------|
| Category ° | Citation of document, with indication, where appropriate, of the relevant passages | Relevant to claim No. |
| Y | WO 91 13889 A (FISONS PLC) 19 September 1991 cited in the application see the whole document | 1-10 |
| Y | WO 97 08182 A (MERCK & CO INC ;MATHRE DAVID J (US); SHUMAN RICHARD F (US); SOHAR) 6 March 1997 cited in the application see the whole document | 1-10 |
| Υ | EP 0 652 219 A (AMERICAN HOME PROD) 10 May 1995 see the whole document | 4-6 |
| Υ | WO 95 22390 A (UNIV MISSOURI ;ARMSTRONG DANIEL (US)) 24 August 1995 see the whole document | 4-6 |
| Υ | US 4 127 647 A (KOBAYASHI TOSHIYUKI ET AL) 28 November 1978 see the whole document | 4-6 |
| Y | CHEMICAL ABSTRACTS, vol. 114, no. 5, 4 February 1991 Columbus, Ohio, US; abstract no. 43482y, MURATA,H. ET AL.: "Purification of macrolide antibiotis by crystallization from aqueous solution" page 812; XP002081481 | 4-6 |
| Y | see abstract & JP 02 196796 A (KYOWA HAKKO CO.LTD.) 3 August 1990 | 4-6 |
| | | |

Information on patent family members

national Application No
PCT/EP 98/03929

| Patent document cited in search repo | rt | Publication date | | atent family nember(s) | Publication date |
|--------------------------------------|-------|------------------|--|--|--|
| EP 0427680 | A | 15-05-1991 | AT AU CA DE DK ES GR HU IE JP | 126803 T 640963 B 6584390 A 2029694 A 69021833 D 69021833 T 427680 T 2077663 T 3017858 T 30096 A 210900 B 69974 B 96268 A 2750302 B | 15-09-1995 09-09-1993 23-05-1991 10-05-1991 28-09-1995 21-03-1996 18-12-1995 01-12-1995 31-01-1996 23-02-1996 28-09-1995 16-10-1996 23-07-1996 13-05-1998 |
| EP 0480623 | А | | JP LV LV US CA JP | 3223291 A 11621 A 11621 B 5352671 A | 02-10-1991 20-12-1996 20-04-1997 04-10-1994 |
| WO 9113889 | А | 19-09-1991 | US EP US PT | 5143918 A | 01-09-1992 04-05-1994 22-03-1994 31-01-1992 |
| WO 9708182 | Α | 06-03-1997 | AU EP | 6900596 A 0848717 A | 19-03-1997 24-06-1998 |
| EP 0652219 | A | 10-05-1995 | AU AU BR CA CN CZ FI HU JP NZ | 6071498 A 7759594 A 9404265 A 2134844 A 1109058 A,B 9402674 A 945186 A 71122 A 7184674 A 264852 A | 04-06-1998 18-05-1995 04-07-1995 06-05-1995 27-09-1995 17-05-1995 06-05-1995 28-11-1995 25-07-1995 21-12-1995 |

Information on patent family members

national Application No
PCT/EP 98/03929

| Patent document cited in search report | | Publication date | Patent family member(s) | | Publication date | |
|--|---|------------------|--|---|--|--|
| EP 0652219 | Α | | SG SK US ZA | 48919 A 130494 A 5508398 A 9408641 A | 18-05-1998 11-07-1995 16-04-1996 02-05-1996 | |
| WO 9522390 | Α | 24-08-1995 | EP JP US | 0748247 A 10501727 T 5626757 A | 18-12-1996 17-02-1998 06-05-1997 | |
| US 4127647 | A | 28-11-1978 | JP JP CA DE FR GB MX NL | 1049411 C 51118816 A 55045042 B 1063936 A 2615403 A 2306705 A 1534070 A 3401 E 7603635 A,B, | 26-06-1981 19-10-1976 15-11-1980 09-10-1979 28-10-1976 05-11-1976 29-11-1978 30-10-1980 12-10-1976 | |