Code 750

COMMONWEALTH OF AUSTRALIA

Patents Act 1952-1969

DECLARATION IN SUPPORT OF A CONVENTION APPLICATION FOR A PATENT OR PATENT OF ADDITION

insert (in full) Name of Company.	::CHINOIN GYOGYSZER ES VEGYESZETITERMEKEK GYARA RT
(2) Here insert title of Invention.	(harrienter referred to so the applicant) for a Potent
	(hereinafter referred to as the applicant) for a Patent for an invention entitled: (2)
	PROCESS FOR THE PREPARATION OF QUINOLINE CARBOXYLIC ACIDS
(3) Here insert full Name and Address, of Company official authorized to make declaration. (4) Here insert basic Country or Countries followed by date or dates and basic Applicant or Applicants.	We Ix" GYULA SZUK and TAMAS SZUTS, both
	of
	Or. Landa G. Budapes C. L.V., Thingar y.
	do solemnly and sincerely declare as follows:
	1 Xk am authorised by the applicant for the patent
	to make this declaration on its behalf.
	2. The basic applications as defined by Section 141 of the Act ZWXX
	WERE made in (4) Hungary
	on the 8th day of April 1987 xxkyand
	on the 26th day of February 1988, xbx both by
	CHINOIN GYOGYSZER ES VEGYESZETI TERMEKEK GYARA RT
(5) Here insert (in full) Name	3 (5) The person named on the reverse hereof
and Address of Actual Inventor or	
Inventors.	
	is anti-led to make the application are as follows:
	is entitled to make the application are as follow: the said actual inventors The applicant is the assignee of
	The applicant is the assignee of
	4. The basic application ^S referred to in paragraph 2 of this Declaration
	the first application made in a Convention country in
	respect of the invention the subject of the application.
	DECLARED at Budapest, Hungary
	this 27th day of October 1988

(12) PATENT ABRIDGMENT (11) Document No. AU-B-15721/88 (19) AUSTRALIAN PATENT OFFICE (10) Acceptance No. 612648

(54) Title PROCESS FOR THE PREPARATION OF QUINOLINE CARBOXYLIC ACIDS

International Patent Classification(s)

(51)⁴ C07D 215/56 A61K 031/47 C07F 005/02 C07F 005/04

(21) Application No.: 15721/88 (22) Application Date: 08.04.88

(87) PCT Publication Number: WO88/07993

(30) Priority Data

(31) Number (32) Date (33) Country 1505/87 08.04.87 HU HUNGARY 1505/87 26.02.88 HU HUNGARY

(43) Publication Date: 04.11.88

(44) Publication Date of Accepted Application: 18.07.91

(71) Applicant(s)
CHINOIN GYCGYSZER ES VEGYESZETI TERMEKEK GYARA RT.

(72) Inventor(s)
ISTVAN HERMECZ; GEZA KERESZTURI; LELLE VASVARI; AGNES HORVATH; MARIA BALOGH;
PETER RITLI; JUDIT SIPOS; ANIKO PAJOR; KATALIN MARMAROSI

(74) Attorney or Agent
WATERMARK PATENT & TRADEMARK ATTORNEYS, Locked Bag 5, HAWTHORN VIC 3122

(56) Prior Art Documents AU 15979/88 C07F 5/02 JP 59-122470

(57) Claim

l. Process for the preparation of compounds of the general Formula I

/wherein R stands for piperazinyl, 4-methyl-piperazinyl or 4-ethyl-piperazinyl group and pharmaceutically acceptable salts thereof which comprises reacting a compound of the general Formula II

(10) 612648

/wherein R¹ and R² stand <u>for helogen</u>, for an aliphatic acyloxy group containing 2 to 6 carbon atoms optionally substituted by halogen, or for an aromatic acyloxy group containing 7 to 11 carbon atoms/ with a piperazine derivative of the general Formula III

/wherein \mathbb{R}^3 stands for hydrogen, methyl or ethyl/ or a salt thereof and subjecting the compound of the general Formula IV

thus obtained/wherein R, R^1 and R^2 are as stated above/ to hydrolysis after or without isolation and if desired converting the compound of the general Formula I thus

(10) 612648

obtained into a salt thereof or setting free the acid from its salt.

- 2. Process according to claim 1 which comprises reacting a compound of the general Formula II with a piperazine derivative of the general Formula III in the presence of an organic solvent.
- 3. Process according to claim 2 wherein the organic solvent is an acid amide, sulfoxide, ketone, alcohol, ether or ester.

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 4: C07D 215/56, C07F 5/02 // A61K 31/47

A1

(11) International Publication Number:

WO 88/ 07993

(43) International Publication Date: 20 October 1988 (20.10.88)

PCT/HU88/00019 (21) International Application Number:

(22) International Filing Date:

8 April 1988 (08.04.88)

(31) Priority Application Numbers:

1505/87 1505/87

(32) Priority Dates:

· 8 April 1987 (08.04.87) 26 February 1988 (26.02.88)

(33) Priority Country:

(71) Applicant (for all designated States except US): CHI-NOIN GYÓGYSZER ÉS VEGYÉSZÉTI TERMÉ-KEK GYÁRA RT.[HU/HU]; Tó utca 1-5, H-1045 Budapest IV (HU).

(72) Inventors; and

(72) Inventors; and
(75) Inventors/Applicants (for US only): HERMECZ, István
[HU/HU]; Molnár u. 53, H-1056 Budapest (HU).
KERESZTURI, Géza [HU/HU]; Széna tér 1/b, H1015 Budapest (HU). VASVÁRI, Lelle [HU/HU];
Goldmark K.u. 33, H-1122 Budapest (HU). HORVÁTH, Ágnes [HU/HU]; Budenz u. 30/a, H-1021 Budapest (HU). BALOGH, Mária [HU/HU];

Barátság u. 20, H-2120 Dunakeszi (HU). RITLI. Péter HU/HU]; Ó u. 43, H-1066 Budapest (HU). SIPOS, Judit [HU/HU]; Sáfrány u. 10, H-1116 Budapest (HU). PAJOR, Anikó [HU/HU]; Ferenc krt. 36, H-1092 Budapest (HU). MÁRMAROSI, Katalin [HU/HU]; Ybl M. st. 19, Biatorbágy, H-2051 Biatorbágy

- (74) Agent: PATENTBUREAU DANUBIA; Bajcsy-Zsilinszky út 16, H-1051 Budapest V (HU).
- (81) Designated States: AT (European patent), AU, BE (European patent), CH (European patent), DE (European patent), DK, FI, FR (European patent), GB (European patent), IT (European patent), JP, KR, LU (European patent), NL (European patent), NO, SE (European patent), SU, US.

Published

With international search report.

A. O. J. P. 8 DEC 1988

AUSTRALIAN

- 4 NOV 1988

PATENT OFFICE

(54) Title: PROCESS FOR THE PREPARATION OF QUINOLINE CARBOXYLIC ACIDS

The invention relates to a new process for the preparation of compounds of general formula (I), (wherein R stands for piperazinyl, 4-methyl-piperazinyl or 4-ethyl-piperazinyl group) and pharmaceutically acceptable salts thereof which comprises reacting a compound of general formula (II), (wherein R¹ and R² stand for halogen, for an aliphatic acyloxy group containing 2 to 6 carbon atoms and optionally substituted by halogen, or for an aromatic acyloxy group containing 7 to 11 carbon atoms) with a piperazine derivative of general formula (III), (wherein R³ stands for hydrogen, methyl or ethyl) or a salt thereof and subjecting the compound of general formula (IV) thus obtained, (wherein R, R¹ and R² are as stated above) to hydrolysis after or without isolation and if desired converting the compound of general formula (I) thus obtained into a salt thereof or setting free the same from its salt. The compounds of general formula (I) are known antibacterial agents. The advantage of the present invention is that it makes the desired compounds of general formula (I) available in a simple manner, with high yields and in a short reaction time.

PROCESS FOR THE PREPARATION OF QUINOLINE CARBOXYLIC ACIDS

This invention relates to a new process for the preparation of l-cyclopropyl-7-substituted-6-fluoro-4-oxo-1,4-dihydro-quinoline-3-carboxylic acid derivatives and pharmaceutically acceptable salts thereof.

It is known that the l-cyclopropyl-7-substituted-6-fluoro-4-oxo-1,4-dihydro-quinoline-3-carboxylic acid derivatives of the general Formula I

10

5

- /wherein R stands for piperazinyl, 4-methyl-piperazinyl or 4-ethyl-piperazinyl group/ possess high antibacterial activity /Eur. J. Clin. Microbiol. 1983, 2, page 111;
 J. Clin. Pharmacol. 1985, 25, page 62; Drugs Exptl. Clin. Res. 1985, 5, page 317./
- The quinoline carboxylic acids of the general

 Formula I can be prepared by reacting 1-cyclopropyl-6fluoro-4-oxo-1,4-dihydro-quinoline-3-carboxylic acid and
 a cyclic amine in the presence of a solvent at a temperature of 135-140 °C for 2 hours /German Off. 3.033.157;

 German Off. 3.142.854/.

According to the present invention there is provided a new process for the preparation of 1-cyclopropyl7-substituted-6-fluoro-4-oxo-1,4-dihydro-quinoline-3carboxylic acid derivatives of the general Formula I
/wherein R has the same meaning as stated above/ which
comprises reacting a compound of the general Formula II

10

5

/wherein R¹ and R² are the same or different and stand for halogen, for an aliphatic acyloxy group containing 2 to 6 carbon atoms optionally substituted by halogen, or for an aromatic acyloxy group containing 7 to 11 carbon atoms/ with a cyclic amine of the general Formula

20

25

/wherein \mathbb{R}^3 stands for hydrogen, methyl or ethyl/ or a salt thereof and subjecting the compound of the general Formula IV



15

20

25

/wherein R, R^1 and R^2 are the same as stated above/ thus obtained to hydrolysis.

The advantage of the process of the present invention is that it makes the desired compound of the general
Formula I available in a simple manner with high yields
and in a short reaction time.

According to a preferred form of embodiment of the process of the present invention the borate derivative of the general Formula IV /wherein R, R¹ and R² are as stated above/ is converted into the desired quinoline-3-carboxylic acid of the general Formula I without isolation.

The borate derivatives of the general Formula IV are new compounds.

The borate derivatives of the general Formula II and the cyclic amine of the general Formula III can be reacted optionally in the presence of an inert organic solvent and an acid binding agent.

As inert organic solvents preferably acid amides

/e.g. dimethyl formamide, dimethyl acetamide/, ketones

/e.g. acetone, methyl ethyl ketone/, ethers /e.g. dioxane,

tetrahydrofuran, diethyl ether/, esters /e.g. ethyl acetate,

10

15

20.

25

methyl acetate, ethyl propionate/, sulfoxides /e.g. dimethyl sulfoxide/, alcohols /e.g. methanol, ethanol, ldecanol, butanol/ may be used.

As acid binding agent an organic or inorganic base may be used. From the group of organic bases trialkyl amines /e.g. triethyl amine, tributyl amine/, cyclic amines /e.g. pyridine, 1,5-diazabicyclo/5.4.0/undec-5-ene, 1,5-diazabicyclo/4.3.0/non-5-ene, 1,4-diazabicyclo-/2.2.2/octane/ can be mentioned, while as inorganic base preferably hydroxides or carbonates of alkali or alkaline earth metals can be applied. Thus as acid binding agent preferably potassium carbonate, potassium hydrogen carbonate, sodium hydroxide, calcium hydroxide, etc. or an excess of the amine of the general Formula III can be used.

The borone derivative of the general Formula II and the cyclic amine of the general Formula III can be reacted at a temperature ranging from 0 to 200 °C, depending on the solvent used. The reaction time may vary between half an hour and 10 hours depending on the reaction temperature. If the reaction is carried out at an elevated temperature, the reaction time can be shortened. The above reaction conditions are but preferable values and other conditions may be used as well.

The borates of the general Formula IV /wherein R, R¹ and R² are as stated above/ can be hydrolysed to the desired quinoline-3-carboxylic acids of the general Formula I, after or without isolation, under acidic

10

15

20

25

ĺ

or basic conditions. The compound of the general Formula IV /wherein R is as stated above/ precipitates from the reaction mixture e.g. on cooling and can be separated e.g. by filtration or centrifuging, if desired.

Basic hydrolysis may preferably be carried out by heating an aqueous solution of hydroxydes or carbonates of alkali metals or hydroxides of alkaline earth metals, One may preferably use an aqueous solution of sodium hydroxide, potassium hydroxide, sodium carbonate, potassium carbonate, calcium hydroxide, potassium hydrogen carbonate. Organic amines /e.g. triethyl amine/ may also be applied in the hydrolysis step.

Acidic hydrolysis may preferably be accomplished by using an aqueous mineral acid. One may preferably proceed by hydrolysing a borate of the general Formula IV by heating same with an aqueous solution of hydrochloric acid, hydrogen bromide, sulfuric acid or phosphoric acid. Hydrolysis may also be accomplished by using organic acids /e.g. acetic acid, propionic acid, etc/.

Hydrolysis of the compounds of the general Formula IV may also be carried out in aqueous medium in the presence of a water-miscible organic solvent. For this purpose e.g. alcohols /e.g. methanol, ethanol, ketones /e.g. acetone/, ethers /e.g. dioxane/, acid amides /e.g. formamide, dimethyl formamide/, sulfoxides /e.g. di-methyl sulfoxide/,or pyridine may be used.

10

15

20

25

The quinoline-3-carboxylic acid of the general Formula I thus obtained may be isolated e.g. by adjusting the pH value of the aqueous solution to a suitable value and separating the precipitated crystals e.g. by filtration or centrifuging or by liophylization of the aqueous reaction mixture.

The compounds of the general Formula I can be converted into pharmaceutically acceptable salts thereof by methods known per se. Thus preferably acid addition salts formed with hydrogen halides, sulfonic acids, sulfuric acid or organic acids. One may preferably form chlorides, bromides, 4-methyl-phenyl-sulfonates, methane sulfonates, maleates, fumarates, benzoates, etc. The compounds of the general Formula I form salts with alkali or alkaline earth metals or other metal ions as well. Accordingly sodium, potassium, magnesium, silver, copper salts, etc. may be prepared.

The compounds of the general Formula I and pharmaceutically acceptable salts thereof can be converted into hydrates /e.g. hemihydrates, trihydrates, etc./ by methods known per se.

According to a further aspect of the present invention there are provided new compounds of the general Formula IV /wherein R, R¹ and R² are as stated above/.

The starting materials of the general Formula II can be prepared e.g. by reacting 1-cyclopropyl-6-fluoro-7-chloro-4-oxo-1,4-dihydro-quinoline-3-carboxylic acil /German Off. 3.141.854/ with a borone derivative /such

as a compound of the general Formula V

$$\begin{array}{c}
R^{1} \\
B-R^{2} \\
R^{5}
\end{array}$$

/wherein R¹, R² and R⁵ stand for halogen, for an aliphatic acyloxy group containing 2 to 6 carbon atoms optionally substituted by halogen, or for an aromatic acyloxy group containing 7 to 11 carbon atoms/ or with fluoroborate in an aqueous or an organic medium.

Further details or the present invention are to be found in the following Examples without limiting the scope of protection to the said Examples.

Example 1

15

20

25

4.1 g of /l-cyclopropyl-6-fluoro-7-chloro-1,4-di-hydro-4-oxo-quinoline-3-carboxylate-03,04/-bis/aceto-0/-borone and 2.8 g of piperazine anhyiride are heated in 16 fil of dimethyl sulfoxide to 110 °C under stirring.

40 ml of a 3 W/v % aqueous sodium hydroxide solution are added to the brownish-red solution and the reaction mixture is boiled under reflux for an hour. The hot pale-yellow solution is filtered and the pH value is adjusted to 7 by adding 1.8 ml of 96 % acetic acid. The reaction mixture is cooled to room temperature, the precipitated white crystals are filtered, washed with water and methanol and dried. The crude product is purified by boiling in 10 ml water. Thus 2.99 g of 1-cyclopropyl-6-fluoro-7-/1-piperazinyl/-1,4-dihydro-4-oxo-quinoline-3-carboxylic

acid are obtained. The product decomposes at 255 °C.

Analysis for the Formula ${\rm C_{17}^{H}_{18}^{FN}_{3}^{0}_{3}}$:

Calculated: C=61.62 % H=5.48 % N=12.68 %

Found: C=61.58 % H=5.50 % N=12.61 %.

5

10

Example 2

By reacting /l-cyclopropyl-6-fluoro-4-chloro-1,4-dihydro-4-oxo-quinoline-3-carboxylate-0³,0⁴/-bis/acetato-0/-borone and N-methyl-piperazine according to Example 1.
l-cyclopropyl-6-fluoro-7-/4-methyl-piperazino/-1,4-di-hydro-4-oxo-quinoline-3-carboxylic acid is prepared. The product decomposes at 248-250 °C.

Example 3

4.1 g of /1-cyclopropyl-6-fluoro-7-chloro-1,4-di-15 hydro-4-oxo-quinoline-5-carboxylate-0³,0⁴/-bis/acetato-0/borone and 3.7 g of N-ethyl-piperazine are heated in 16 ml of dimethyl sulfoxide to 30 °C under stirring. After 10 minutes 40 ml of a 3 $\frac{W}{W}$ 3 aqueous sodium hydroxide solution are added and the reaction mixture is boiled 20 for an hour under reflux. The hot solution is filtered and the pH value is adjusted to 7 with 96 % acetic acid. The reaction mixture is cooled, the precipitated crystals are filtered and washed with water. Thus 3.3 g of 1-cyclopropyl-7-/4-ethyl-piperazinyl/-6-fluorc-1,4-dihydro-4-25 oxo-quinoline-3-carboxylic acid are obtained. M.D.: 183-185 °C.

Analysis for the Formula $C_{19}H_{22}FN_{3}O_{3}$:

Calculated: C=63.35 H=6.17 N=11.69

Found: C=63.31 H=6.21 N=11.70

5 Example 4

3.3 g of /l-cyclopropyl-6-fluoro-7-chloro-1,4-di-hydro-4-oxo-quinoline-3-carboxylate-03,04/-difluoro-borone are reacted with 3.7 g of N-ethyl-piperazine according to Example 3. Thus 3.4 g of l-cyclopropyl-7-/4-ethyl-l-piperazinyl/-6-fluoro-1,4-dihydro-4-oxo-quinoline-3-carboxylic acid are obtained which in admixture at any ratio with the product of Example 3 no depression of the melting point occurs.

WHAT WE CLAIM IS:

l. Process for the preparation of compounds of the general Formula I

5

10

/wherein R stands for piperazinyl, 4-methyl-piperazinyl or 4-ethyl-piperazinyl group/and pharmaceutically acceptable salts thereof which comprises reacting a compound of the general Formula II

15

20

25

/wherein R¹ and R² stand for halogen; for an aliphatic acyloxy group containing 2 to 6 carbon atoms optionally substituted by halogen, or for an aromatic acyloxy group containing 7 to 11 carbon atoms/ with a piperazine derivative of the general Formula III



/wherein R^3 stands for hydrogen, methyl or ethyl/ or a salt thereof and subjecting the compound of the general Formula IV

thus obtained/ wherein R, R¹ and R² are as stated above/ to hydrolysis after or without isolation and if desired converting the compound of the general Formula I thus obtained into a salt thereof or setting free the acid from its salt.

- 2. Process according to claim 1 which comprises reacting a compound of the general Formula II with a piperazine derivative of the general Formula III in the presence of an organic solvent.
- 3. Process according to claim 2 wherein the organic solvent is an acid amide, sulfoxide, ketone, alcohol, ether or ester.



- 4. Process according to claim 2, which comprises using dimethyl sulfoxide as organic solvent.
- 5. Process according to claim 1, which comprises carrying out the reaction of the compounds of the general Formulae II and III in the presence of an acid binding agent.
- 6. Process according to claim 4, which comprises using an amine or an excess of the compound of the general Formula III as acid binding agent.
- 7. Process according to claim 1 ,which comprises carrying out the hydrolysis in acidic medium.
- 8. Process according to claim 6, which comprises carrying out the reaction by using as acid, an organic or inorganic acid.
- 9. Process according to claim 8, wherein the acid is selected from hydrochloric acid, sulfuric acid or acetic acid.
- 10. Process according to claim 1, which comprises carrying out the hydrolysis in a basic medium.
- 11. Process according to claim—8, which comprises using as a base an alkali metal hydroxide, an alkaline earth metal hydroxide or an organic base.
- 12. Process according to claim 11, wherein the organic base is an aqueous triethylamine solution.

Disk 0099/1.57



13. Compounds of the general Formula IV

/wherein R stands for piperazinyl, 4-methyl-piperazinyl or 4-ethyl-piperazinyl group, R¹ and R² stand for an aliphatic acyloxy group containing 2 to 6 carbon atoms optionally substituted by halogen, or for an aromatic acyloxy group containing 7 to 11 carbon atoms/.

DATED this 17th day of April, 1991.

CHINOIN GYOGYSZER ES VEGYESZETA TERMEKEK GYARA RT

WATERMARK PATENT ATTORNEYS

2ND FLOOR "THE ATRIUM"

290 BURWOOD ROAD

HAWTHORN, VIC. 3122

AUSTRALIIA



INTERNATIONAL SEARCH REPORT

International Application No PCT/HU 88/00019

	IFICATION OF SUBJECT MATTER (if several classification symbols apply, ind		10 007 00019
According	te International Patant Classification (IPC) or to both National Classification and IPC		
IPC ⁴	: C 07 D 215/56; C 07 F 5/02 // A 61 K 3	31/47	
	SEARCHED		
	Minimum Documentation Searched 7		
Classificatio	on System Classification Symbols		
Int.C	1. ⁴ C 07 D 215/56; C 07 F 5/02,5/04.		
	Documentation Searched other than Minimum Documentation to the Extent that such Documents are included in the Fields		
III. DOCU	MENTS CONSIDERED TO BE RELEVANT		
Category •		esages 12	Relevant to Claim No. 13
Y	Chemical Abstracts, vol. 103, no. 15,		(1,3,8)
	1985, October 14 (Columbus, Ohio, USA)		
i	Daiichi Seiyaku "Oxazines" see page 73		
	column 1, abstract-no. 123 491p & JP,		
	60-78 986 (DAIICHI SEIYAKU) 04 May 198	35.	
أر			
Y	Chemical Abstracts, vol. 102, no. 7, i		(1,3,8)
	1985, February 18 (Columbus, Ohio, USA		
	Daiichi Seiyaku "1-Ethyl-6-fluoro-4-ox		
	(1-piperazinyl)-1,4-dihydro-quinoline-		
	carboxylic acids" see page 605, column		
	abstract-no. 62 272y & JP, A, 59-122 4	170	
	(DAIICHI SEIYAKU) 14 July 1984.		
i			
· .			
"A" doc con "E" earl filin "L" doc whi cita "O" doc oth "P" doc	cument defining the general state of the art which is not neidered to be of particular relevance. Iter document but published on or after the international ng date cument which may throw doubts on priority claim(s) or ich is cited to establish the publication date of another stion or other special reason (as specified) cument referring to an oral disclosure, use, exhibition or discounties or considered.	nd not in confi and the principal ricular relevant dered novel or ive step ricular relevant lered to involve bined with one binetion being	the international filing date let with the application but to or theory underlying the ce; the claimed invention cannot be considered to ce; the claimed invention an inventive step when the or more other such docupobvious to a person skilled patent family
IV. CERT	PIFICATION		
Date of th	ne Actual Completion of the International Search Date of Mailing of this	International S	earch Réport
27 J	une 1988 (27.06.88) 01 July 19	88 (01.	07.88)
Internation	nal Searching Authority Signature of Authorize	d Officer	
AUST	RIAN PATENT OFFICE	vi n	

Anhang zum internationalen Recherchenbericht über die internationale Patentanmeldung Nr.

In diesem Anhang sind die Mitglieder der Patentfamilien der im obengenannten internationalen Recherchenbericht angeführten Patentdokumente angegeben. Diese Angaben dienen nur zur Unterrichtung und erfolgen ohne Gewähr.

Annex to the International Search Report on International Patent Application No. PCT/HU 88/00019

This Annex lists the patent family members relating to the patent documents cited in the above-mentioned International search report. The Austrian Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

Annexe au rapport de recherche internationale relatif à la demande de brevet international n°.

La présente annexe indique les membres de la famille de brevets relatifs aux documents de brevets cités dans le rapport de recherche internationale visé ci-dessus. Les renseignements fournis sont donnés à titre indicatif et n'engagent pas la responsabilité de l'Office autrichien des brevets.

Im Recherchenbericht
angeführtes Patentdokument
Patent document cited
in search report
Document de brevet cité
dans le rapport
de recherche

Datum der
Veröffentlichung
Publication
date
Date de
publication

Mitglied(er) der Patentfamilie Patent family member(s) Membre(s) de la famille de brevets Datum der
Veröffentlichung
Publication
date
Date de
publication

JP-A2-60-078 986

JP-A2-59-122 470

04/05/1985

None

14/07/1984

None