

# International Nonproprietary Names for Pharmaceutical Substances (INN)

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## RECOMMENDED International Nonproprietary Names: List 54

Notice is hereby given that, in accordance with paragraph 7 of the Procedure for the Selection of Recommended International Nonproprietary Names for Pharmaceutical Substances [*Off. Rec. Wild Health Org.*, 1955, **60**, 3 (Resolution EB15.R7); 1969, **173**, 10 (Resolution EB43.R9)], the following names are selected as Recommended International Nonproprietary Names. The inclusion of a name in the lists of Recommended International Nonproprietary Names does not imply any recommendation of the use of the substance in medicine or pharmacy.

Lists of Proposed (1–91) and Recommended (1–52) International Nonproprietary Names can be found in *Cumulative List No. 11, 2004* (available in CD-ROM only).

## Dénominations communes internationales des Substances pharmaceutiques (DCI)

### Dénominations communes internationales RECOMMANDÉES: Liste 54

Il est notifié que, conformément aux dispositions du paragraphe 7 de la Procédure à suivre en vue du choix de Dénominations communes internationales recommandées pour les Substances pharmaceutiques [*Actes off. Org. mond. Santé*, 1955, **60**, 3 (résolution EB15.R7); 1969, **173**, 10 (résolution EB43.R9)] les dénominations ci-dessous sont choisies par l'Organisation mondiale de la Santé en tant que dénominations communes internationales recommandées. L'inclusion d'une dénomination dans les listes de DCI recommandées n'implique aucune recommandation en vue de l'utilisation de la substance correspondante en médecine ou en pharmacie.

On trouvera d'autres listes de Dénominations communes internationales proposées (1–91) et recommandées (1–52) dans la *Liste récapitulative No. 11, 2004* (disponible sur CD-ROM seulement).

## Denominaciones Comunes Internacionales para las Sustancias Farmacéuticas (DCI)

### Denominaciones Comunes Internacionales RECOMENDADAS: Lista 54

De conformidad con lo que dispone el párrafo 7 del Procedimiento de Selección de Denominaciones Comunes Internacionales Recomendadas para las Sustancias Farmacéuticas [*Act. Of. Mund. Salud*, 1955, **60**, 3 (Resolución EB15.R7); 1969, **173**, 10 (Resolución EB43.R9)], se comunica por el presente anuncio que las denominaciones que a continuación se expresan han sido seleccionadas como Denominaciones Comunes Internacionales Recomendadas. La inclusión de una denominación en las listas de las Denominaciones Comunes Recomendadas no supone recomendación alguna en favor del empleo de la sustancia respectiva en medicina o en farmacia.

Las listas de Denominaciones Comunes Internacionales Propuestas (1–91) y Recomendadas (1–52) se encuentran reunidas en *Cumulative List No. 11, 2004* (disponible sólo en CD-ROM).

Latin, English, French, Spanish:  
Recommended INN

Chemical name or description; Molecular formula; Graphic formula

DCI Recommandée

Nom chimique ou description; Formule brute; Formule développée

DCI Recomendada

Nombre químico o descripción; Fórmula molecular; Fórmula desarrollada

**acidum salclobuzicum**

salclobuzic acid

4-(4-chloro-2-hydroxybenzamido)butanoic acid

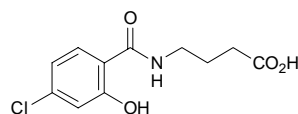
acide salclobuzique

acide 4-[(4-chloro-2-hydroxybenzoyl)amino]butanoïque

ácido salclobúxico

ácido 4-[(4-cloro-2-hidroxibenzoil)amino]butanoico

$C_{11}H_{12}ClNO_4$



**ancrivirocum**

ancriviroc

3-({4-[(Z)-(4-bromophenyl)(ethoxyimino)methyl]-4'-methyl-[1,4'-bipiperidin]-1'-yl}carbonyl)-2,4-dimethylpyridine-1-oxide

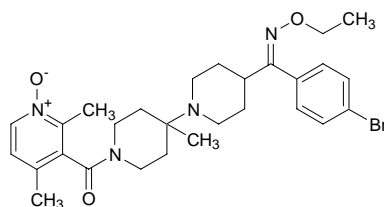
ancriviroc

4-[(Z)-(4-bromophényl)(éthoxyimino)méthyl]-1'-[(2,4-diméthyl-1-oxypyridin-3-yl)carbonyl]-4'-méthyl-1,4'-bipéridinyle

ancriviroc

4-[(Z)-(4-bromofenil)(etoxiimino)metil]-1'-[(2,4-dimetil-1-oxidopiridin-3-il)carbonil]-4'-metil-1,4'-bipéridinilo

$C_{28}H_{37}BrN_4O_3$



**aplindorum**

aplindore

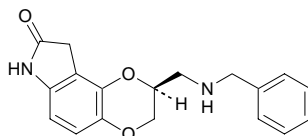
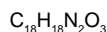
(2S)-2-[(benzylamino)methyl]-2,3,7,9-tetrahydro-8H-1,4-dioxino-[2,3-e]indol-8-one

aplindore

(2S)-2-[(benzylamino)méthyl]-2,3,7,9-tétrahydro-8H-1,4-dioxino-[2,3-e]indol-8-one

aplindor

(2S)-2-[(bencilamino)metil]-2,3,7,9-tetrahidro-8H-1,4-dioxino-[2,3-e]indol-8-ona



**atilmotinum**  
atilmotin

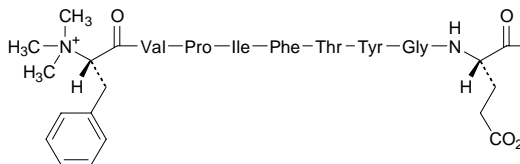
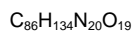
*N*-[(2*S*)-3-phenyl-2-(trimethylazaniumyl)propanoyl]-L-valyl-L-prolyl-L-isoleucyl-L-phenylalanyl-L-threonyl-L-tyrosylglycyl-L-glutamyl-L-leucyl-L-glutamyl-D-arginyl-L-leucyl-L-lysineamide

atilmotine

*N*-[(2*S*)-3-phényl-2-(triméthylammonio)propanoïl]-L-valyl-L-prolyl-L-isoleucyl-L-phénylalanil-L-thréonyl-L-tyrosylglycyl-L-glutamyl-L-leucyl-L-glutamyl-D-arginyl-L-leucyl-L-lysineamide

atilmotina

*N*-[(2*S*)-3-feníl-2-(trimetilamonio)propanoíl]-L-valil-L-prolil-L-isoleucil-L-fenilalanil-L-treonil-L-tirosilglicil-L-glutamil-L-leucil-L-glutaminiil-D-arginil-L-leucil-L-lisineamida



Leu—Gln—D-Arg—Leu—Lys—NH<sub>2</sub>

**avanafilum**  
avanafil

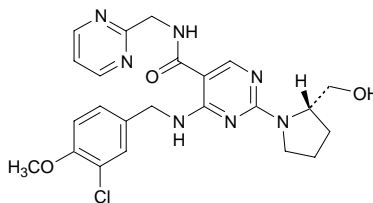
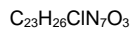
4-[[[3-chloro-4-methoxyphenyl)methyl]amino]-2-[(2*S*)-2-(hydroxymethyl)pyrrolidin-1-yl]-*N*-(pyrimidin-2-ylmethyl)pyrimidine-5-carboxamide

avanafil

4-[(3-chloro-4-méthoxybenzyl)amino]-2-[(2*S*)-2-(hydroxyméthyl)=pyrrolidin-1-yl]-*N*-(pyrimidin-2-ylméthyl)pyrimidine-5-carboxamide

avanafilo

4-[(3-cloro-4-metoxibencil)amino]-2-[(2*S*)-2-(hidroximetil)pirrolidin-1-il]-*N*-(pirimidin-2-ilmetil)pirimidina-5-carboxamida



**balicatibum**

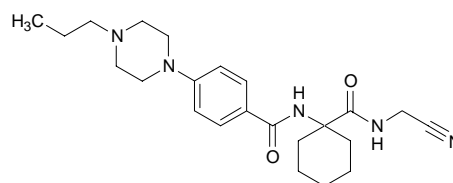
balicatib

*N*-{1-[(cyanomethyl)carbamoyl]cyclohexyl}-4-(4-propylpiperazin-1-yl)benzamide

balicatib

*N*-[1-[(cyanométhyl)carbamoyl]cyclohexyl]-4-(4-propylpipérazin-1-yl)benzamide

balicatib

*N*-[1-[(cianometil)carbamoi]ciclohexil]-4-(4-propilpiperazin-1-il)benzamidaC<sub>23</sub>H<sub>33</sub>N<sub>5</sub>O<sub>2</sub>**becatecarinum**

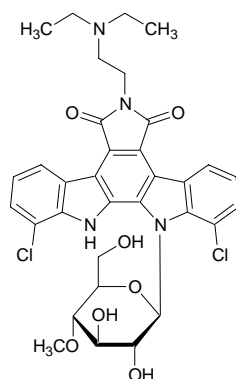
becatecarin

1,11-dichloro-6-[2-(diethylamino)ethyl]-12-(4-*O*-methyl-β-*D*-glucopyranosyl)-12,13-dihydro-5*H*-indolo[2,3-*a*]pyrrolo-[3,4-*c*]carbazole-5,7(6*H*)-dione

bécatécarine

1,11-dichloro-6-[2-(diéthylamino)éthyl]-12-(4-*O*-méthyl-β-*D*-glucopyranosyl)-12,13-dihydro-5*H*-indolo[2,3-*a*]pyrrolo-[3,4-*c*]carbazole-5,7(6*H*)-dione

becatecarina

1,11-dicloro-6-[2-(dietilamino)etil]-12-(4-*O*-metil-β-*D*-glucopiranosil)-12,13-dihidro-5*H*-indolo[2,3-*a*]pirrolo[3,4-*c*]carbazol-5,7(6*H*)-dionaC<sub>33</sub>H<sub>34</sub>Cl<sub>2</sub>N<sub>4</sub>O<sub>7</sub>

**becocalcidiolum**

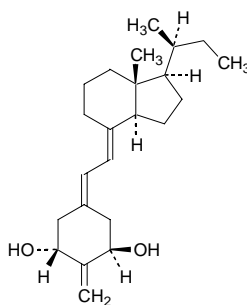
becocalcidiol

(1*R*,3*R*)-4-(2-((1*R*,3*aS*,7*aR*)-1-[(2*S*)-butan-2-yl]-7*a*-methyloctahydro-4*H*-inden-4-ylidene)ethylidene)-2-methylidencyclohexane-1,3-diol

bécocalcidiol

(1*R*,3*R*)-2-méthylidène-5-[(2*E*)-2-[(1*R*,3*aS*,7*aR*)-7*a*-méthyl-1-[(1*S*)-1-méthylpropyl]octahydro-4*H*-indén-4-ylidène]éthylidène]=cyclohexane-1,3-diol

becocalcidiol

(1*R*,3*R*)-2-metilideno-5-[(2*E*)-2-[(1*R*,3*aS*,7*aR*)-7*a*-metil-1-[(1*S*)-1-metilpropil]octahidro-4*H*-inden-4-ilideno]etilideno]=ciclohexano-1,3-diolC<sub>23</sub>H<sub>36</sub>O<sub>2</sub>**bemotrizinolum**

bemotrizinol

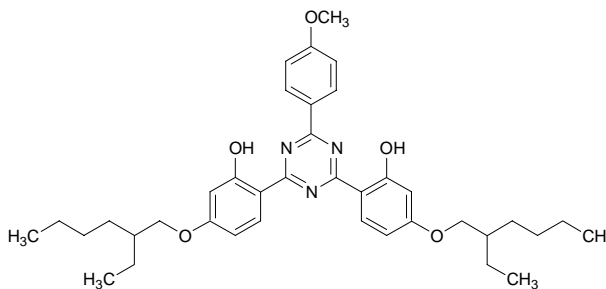
2,2'-[6-(4-methoxyphenyl)-1,3,5-triazine-2,4-diyl]bis-[5-[(2-ethylhexyl)oxy]phenol]

bémotrizinol

2,2'-[6-(4-méthoxyphényl)-1,3,5-triazine-2,4-diyl]bis-[5-[(2-éthylhexyl)oxy]phénol]

bemotrizinol

2,2'-[6-(4-metoxifenil)-1,3,5-triazina-2,4-diil]bis[5-[(2-etilhexil)=oxi]fenol]

C<sub>38</sub>H<sub>49</sub>N<sub>3</sub>O<sub>5</sub>

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