International Nonproprietary Names (INN) for biological and biotechnological substances

(a review)
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0. INTRODUCTION

More than 50 years ago, WHO established the International Nonproprietary Name (INN) Expert Group / WHO Expert Committee on Specifications for Pharmaceutical Preparations, to assign nonproprietary names to medicinal substances, so that each substance would be recognized globally by a unique name. These INNs do not give proprietary rights, unlike a trade mark, and can be used freely as they are public property.

INNs have been assigned to biological products since the early days of the INN Programme. As well as many names for individual substances, animal insulin preparations were given an INN in Recommended list 3 in 1959. In the period up to 1980, names were assigned to antibiotics, synthetic peptides, hormones and other proteins. In names of compounds related by structure and/or function, specific letter groups, called stems, are included to aid recognition by health professionals. The -actide for synthetic polypeptides with a corticotrophin-like action is an example.

In 1982, the name insulin human was proposed for the recombinant protein identical to natural human insulin, and since then names have been assigned to a growing number of recombinant products. Within the INN Programme, names have not been assigned to natural human blood products or vaccines. For those groups of biological products, the WHO Expert Committee on Biological Standardization (ECBS) has been adopting the scientific names of the biological products within the definitions of respective requirements.

Since the time when insulin human became the first recommended INN (rINN) for a recombinant product, the range of biological/biotechnological products has increased in size and complexity. For example, new stems have been introduced for tissue plasminogen activators (-plase) among other groups. Analogues of recombinant glycosylated proteins produced in different cell systems have been classified using Greek letters as indicators in the sequence of product introduction: erythropoietin (epoetin alfa, beta and so on) and glycoprotein hormones (folitropin) are examples. In the 1990s, a systematic scheme for naming monoclonal antibodies was implemented, based on the stem -mab, which indicates the origin (mouse, human, etc) of the antibody and its intended use: tumour, immunomodulator and so on.

As a result of the scientific and technical developments currently taking place, new products of biotechnology and other biological products are being introduced and more products can be expected for the treatment or prevention of disease. Examples of such new products include recombinant blood products, transgenic products (human proteins expressed in animals or plants), products for gene therapy and novel vaccines.
As this area is becoming more and more complex and challenging, the INN Expert Group has requested the WHO-INN Secretariat to prepare a working document intended to summarize and review the past and present INN situation in this field.

This document, first published on the website of the INN Programme in 2006, therefore presents an inventory of the policy decisions taken by the INN Expert Group during all these years of change, and of the names assigned to biological and biotechnological substances. Considering the potential for further developments in the field of biologicals, this review is intended to be a living document which will be regularly updated to include new policies, and future INNs assigned.

Comments and suggestions from all interested parties are most welcome and will be presented to the INN Expert Group for their consideration and for possible incorporation in future updates of this review.

You are reading the current updated version, also available as pdf-copy at:

1. PHARMACOLOGICAL CLASSIFICATION OF BIOLOGICAL AND BIOTECHNOLOGICAL SUBSTANCES (1)

Alimentary tract and metabolism

insulins (see item 4.17).

Anti-infectives

antimicrobial, bactericidal permeability increasing polypeptides (see item 4.1)

human papilloma virus (see item 4.16).

Antineoplastics

peptide vaccines / recombinant vaccines (see item 4.24)

Toxins (see item 4.30).

Blood and agents acting on the haemopoietic system

antithrombins (see item 4.3)

blood coagulation cascade inhibitors (see item 4.4)

blood coagulation factors (see item 4.5)

erthropoietin type blood factors (see item 4.8)

heparin derivatives including low molecular mass heparins (see item 4.13)

hirudin derivatives (see item 4.14)

thrombomodulins (see item 4.29).
Immunomodulators and immunostimulants

colony stimulating factors (see item 4.6)
interferons (see item 4.18)
interleukin receptor antagonists (see item 4.19)
interleukin type substances (see item 4.20)
monoclonal antibodies (see item 4.21)
receptor molecules, native or modified (see item 4.27).

Hormones, hormone antagonists, hormone-release stimulating peptides or hormone-release inhibiting peptides (excluding insulins)

growth hormone (GH) derivatives (see item 4.11)
growth hormone antagonists (see item 4.12)
oxytocin derivatives (see item 4.22)
pituitary / placental glycoprotein hormones (see item 4.25)
pituitary hormone-release stimulating peptides (see item 4.26)
synthetic polypeptides with a corticotropin-like action (see item 4.28)
vasoconstrictors, vasopressin derivatives (see item 4.31).

Various

antisense oligonucleotides (see item 4.2)
enzymes (see item 4.7)
gene therapy products (see item 4.9)
growth factors (see item 4.10)
peptides and glycopeptides (for special groups of peptides see -actide (see item 4.28), -pressin (see item 4.31), -relin (see item 4.26), -tocin (see item 4.22)) (see item 4.23).
2. CURRENT STATUS OF EXISTING STEMS OR SYSTEMS FOR BIOLOGICAL AND BIOTECHNOLOGICAL SUBSTANCES

2.1. Groups with respective stem

<table>
<thead>
<tr>
<th>Name of the group</th>
<th>Stem</th>
</tr>
</thead>
<tbody>
<tr>
<td>antisense oligonucleotides</td>
<td>-rsen</td>
</tr>
<tr>
<td>blood coagulation cascade inhibitors</td>
<td>-cogin</td>
</tr>
<tr>
<td>blood coagulation factors</td>
<td>-cog</td>
</tr>
<tr>
<td>colony stimulating factors</td>
<td>-stim</td>
</tr>
<tr>
<td>enzymes</td>
<td>-ase</td>
</tr>
<tr>
<td>erythropoietin type blood factors</td>
<td>-poetin</td>
</tr>
<tr>
<td>growth factors</td>
<td>-ermin</td>
</tr>
<tr>
<td>growth hormone derivatives</td>
<td>som-</td>
</tr>
<tr>
<td>heparin derivatives including low molecular mass heparins</td>
<td>-parin</td>
</tr>
<tr>
<td>hirudin derivatives</td>
<td>-irudin</td>
</tr>
<tr>
<td>pituitary hormone-release inhibiting peptides</td>
<td>-relix</td>
</tr>
<tr>
<td>interleukin receptor antagonists</td>
<td>-kinra</td>
</tr>
<tr>
<td>interleukin type substances</td>
<td>-kin</td>
</tr>
<tr>
<td>monoclonal antibodies</td>
<td>-mab</td>
</tr>
<tr>
<td>oxytocin derivatives</td>
<td>-tocin</td>
</tr>
<tr>
<td>peptides and glycopeptides (for special groups of peptides see -actide, -pressin, -relin, -tocin)</td>
<td>-tide</td>
</tr>
<tr>
<td>pituitary hormone-release stimulating peptides</td>
<td>-relin</td>
</tr>
<tr>
<td>receptor molecules, native or modified (a preceding infix should designate the target)</td>
<td>-cept</td>
</tr>
<tr>
<td>synthetic polypeptides with a corticotropin-like action</td>
<td>-actide</td>
</tr>
<tr>
<td>vasoconstrictors, vasopressin derivatives</td>
<td>-pressin</td>
</tr>
</tbody>
</table>
### 2.2. Groups with respective pre-stems

<table>
<thead>
<tr>
<th>Name of the group</th>
<th>Pre-stem</th>
</tr>
</thead>
<tbody>
<tr>
<td>antimicrobial, bactericidal permeability increasing polypeptides</td>
<td>-ganan</td>
</tr>
</tbody>
</table>

### 2.3. Groups with INN schemes

<table>
<thead>
<tr>
<th>Name of the group</th>
</tr>
</thead>
<tbody>
<tr>
<td>antithrombins</td>
</tr>
<tr>
<td>gene therapy products</td>
</tr>
<tr>
<td>insulins</td>
</tr>
<tr>
<td>interferons</td>
</tr>
<tr>
<td>pituitary / placental glycoprotein hormones</td>
</tr>
</tbody>
</table>

### 2.4. Groups without respective stems / pre-stems and without INN schemes

<table>
<thead>
<tr>
<th>Name of the group</th>
</tr>
</thead>
<tbody>
<tr>
<td>growth hormone antagonists</td>
</tr>
<tr>
<td>human papilloma virus</td>
</tr>
<tr>
<td>peptide vaccines / recombinant vaccines</td>
</tr>
<tr>
<td>thrombomodulins</td>
</tr>
<tr>
<td>toxins</td>
</tr>
</tbody>
</table>
3. GENERAL POLICIES FOR BIOLOGICAL AND BIOTECHNOLOGICAL SUBSTANCES

3.1. General policies for blood products

- INNs have not been assigned to natural human blood products.
- Many natural blood products have well-established names, so the recombinant version should have a distinctive name reflecting as much as possible the established name used in the field.
- It is essential to add "activated" to the name of the blood product when this is presented for therapeutic use in its activated form.

3.2. General policies for fusion proteins

- INNs have been assigned to some fusion proteins. If a stem exists for one or the other part of the fusion protein, this stem should be brought into the name. This allows the constant part of a fusion protein to be recognized in the name.
- At present it is considered unnecessary to indicate that the product is a fusion product within the name, but this position may need to be reviewed in the future.

3.3. General policies for gene therapy products

In 2005, the Nomenclature Scheme for Gene Therapy Products was formally adopted. The scheme is shown in Table 1.

---

1 The list of fusion proteins published is given in Annex 1
### Table 1 Two-word scheme for gene therapy products

<table>
<thead>
<tr>
<th>word 1</th>
<th>prefix</th>
<th>infix</th>
<th>suffix</th>
</tr>
</thead>
<tbody>
<tr>
<td>(gene component)</td>
<td>to contribute to the distinctive name</td>
<td>to identify the gene using, when available, existing infixes for biological products or using similar infix as for the protein for which the gene codes.</td>
<td>-(a vowel)gene e.g. -(o)gene</td>
</tr>
<tr>
<td></td>
<td>e.g. al-; bet-; val-</td>
<td>e.g. -ermin-: growth factor</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>-kin-: interleukin</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>-lim-: immunomodulator</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>-mul-: multiple gene</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>-tusu-: tumour suppression</td>
<td></td>
</tr>
<tr>
<td>word 2</td>
<td>to contribute to the distinctive name</td>
<td>e.g. -adeno-: adenovirus</td>
<td></td>
</tr>
<tr>
<td>(vector component)</td>
<td></td>
<td>-cana-: canarypox virus</td>
<td>-vec (non-replicating viral vector)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>-herpa-: herpes virus</td>
<td>-repvec (replicating viral vector)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>-lenti-: lentivirus</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>-morbilli-: paramyxoviridae morbillivirus</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>-parvo-: adeno-associated virus</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>(paroviridae dependovirus)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>-retro-: other retrovirus</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>-vaci-: vaccinia virus</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>-plasmid (plasmid vector)</td>
<td></td>
</tr>
</tbody>
</table>

In the case of naked DNA, there is no need for a second word in the name.

### 3.4. General policies for glycosylated compounds (8)

**For glycoproteins / glycopeptides**

- Identification of the group with a stem, e.g. for erythropoietin: -poetin, indication of differences in the amino acid chain by using a random prefix and indication of differences in the glycosylation pattern by another designator, expressed by a Greek letter spelt in full and added as second word to the name (e.g. epoetin alfa (66)). The Greek letters\(^2\) are used in the Greek alphabetical order.

---

\(^2\) The transliteration of Greek letters in English, French and Spanish is given in Annex 2
• Identification of the group with a word, e.g. interferon. Subgroups are identified by a Greek letter spelt in full and added as second word to the name; differences in the composition of the amino acid sequence are indicated by using an Arabic figure; different compounds, including different glycosylation pattern, are indicated by a small letter (e.g. interferon beta (73), peginterferon alfa-2a (84)).

3.5. **General policies for immunoglobulins** (9) (10)

Not to select an INN for each of immunoglobulins is the current policy.

The "systematic" or descriptive name is essential since the prescriber must know all the information conveyed by it and there is no benefit in assigning an INN from which it will not be readily apparent.

3.6. **General policies for monoclonal antibodies** (1) (3) (11)

• INN for monoclonal antibodies (mAbs) are composed of a prefix, a substem A, a substem B and a suffix.
• The common stem for mAbs is -mab, placed as a suffix.
• The stem -mab is to be used for all products containing an immunoglobulin variable domain which binds to a defined target.
• Substem B indicates the species on which the immunoglobulin sequence of the mAb is based (shown in Table 2).

---

3 It contains the revised naming scheme for monoclonal antibodies; the previous naming scheme for monoclonal antibodies is given in Annex 3
The distinction between chimeric and humanized antibodies is as follows:

A **chimeric** antibody is one that contains contiguous foreign-derived amino acids comprising the entire variable domain of both heavy and light chains linked to heavy and light constant regions of human origin.

A **humanized** antibody has segments of foreign-derived amino acids interspersed among variable domain segments of human-derived amino acid residues and the humanized variable heavy and variable light domains are linked to heavy and light constant regions of human origin.

The –*xizu*- infix is used for an antibody having both chimeric and humanized chains.

The –*axo*- infix is used for an antibody having both rat and mouse chains.

- Substem A indicates the target (molecule, cell, organ) class (shown in Table 3).
Table 3 Substem A for target class

<table>
<thead>
<tr>
<th>Substem A</th>
<th>Target Class</th>
</tr>
</thead>
<tbody>
<tr>
<td>-b(a)</td>
<td>bacterial</td>
</tr>
<tr>
<td>-c(i)</td>
<td>cardiovascular</td>
</tr>
<tr>
<td>-f(u)</td>
<td>fungal</td>
</tr>
<tr>
<td>-k(i)</td>
<td>interleukin</td>
</tr>
<tr>
<td>-l(i)</td>
<td>immunomodulating</td>
</tr>
<tr>
<td>-n(e)  (under discussion)</td>
<td>neural</td>
</tr>
<tr>
<td>-s(o)</td>
<td>bone</td>
</tr>
<tr>
<td>-tox(a)</td>
<td>toxin</td>
</tr>
<tr>
<td>t(u)</td>
<td>tumour</td>
</tr>
<tr>
<td>-v(i)</td>
<td>viral</td>
</tr>
</tbody>
</table>

In principle, a single letter, e.g. -b- for bacterial is used as substem A. Whenever substem B starts with a consonant (e.g. x or z), to avoid problems in pronunciation, an additional vowel indicated in the table, e.g. -ba- is inserted.

Prefix

The prefix should be random i.e. the only requirement is to contribute to a euphonious and distinctive name.

Second word

If the product is radiolabelled or conjugated to another chemical, identification of this conjugate is accomplished by use of a separate, second word or acceptable chemical designation. For instance, for mAbs conjugated to a toxin, the suffix -tox can be used in the second word.

If the monoclonal antibody is used as a carrier for a radioisotope, the latter will be listed first in the INN, e.g. technetium ($^{99m}$Tc) nofetumomab merpentan (81).

The prefix peg- can be used for pegylated mAbs, but this should be avoided if it leads to over-long INN. In most cases, it is best to adopt two-word INN for pegylated mAbs, with the first word describing the mAb and the second being pegol or a related designation.
3.7. **General policies for non-glycosylated compounds** (8)

**For proteins / peptides:**
- Identification of the group with a stem, e.g. for hirudin analogues: -irudin, and indication of differences in the amino acid chain by using a random prefix (e.g. bivalirudin (72)).
- Identification of the group with a word, e.g. insulin, and indication of differences in the composition of the amino acid chain as a second element of the name (e.g. insulin argine (58)).

3.8. **General policies for skin substitutes** (4)

The products within this system are made of cells within a matrix, and skin substitutes can be considered to be engineered tissue and thus fall outside the scope of the INN system.

3.9. **General policies for transgenic products** (4)

- If an INN already exists, the same name should be used for the transgenic product, qualified in some way to identify that this product is transgenic.
- A similar system to that used for glycosylated recombinant products is suggested to differentiate new or additional sources of the same substance, and the source of the substance should be included in the definition of the INN.

3.10. **General policies for vaccines** (4) (5) (6) (7)

- At present, vaccines are not included within the INN system, but names are assigned through recommendations of the Expert Committee on Biological Standardization and through the pharmacopoeial monograph.
- During the INN Consultation in 1993, it was agreed that the prerequisite for an INN application for a recombinant vaccine

---

4 The definition of recombinant vaccines is given in item 4.24
would be fulfilled if the manufacturer was able to provide all information outlined in the guidelines entitled Definition of INNs for Substances Prepared by Biotechnology (WHO / Pharm S / Nom 1348\textsuperscript{(12)}).

- During the INN Consultation in 1998, following discussion on recombinant viruses, the experts agreed not to attempt to name live viruses.

- Another approach in vaccine technology seems to be the development of peptide vaccines\textsuperscript{5} (epitopes involved in immune response formation): since these peptides are chemically well-defined, their naming will be less problematic.

\textsuperscript{5} The definition of peptide vaccines is given in item 4.24
4. SUMMARY OF INN ASSIGNED TO BIOLOGICAL AND BIOTECHNOLOGICAL SUBSTANCES

4.1. Antimicrobial, bactericidal permeability increasing polypeptides

The pre-stem for antimicrobial, bactericidal permeability increasing polypeptides is -ganan.

iseganan (85), omiganan (89), pexiganan (78).

4.2. Antisense oligonucleotides

The common stem for antisense oligonucleotides is -rsen.

afovirsen (97), aganirsen (103), alicaforsen (97), aprinocarsen (97), beclanorsen (101), cenersen (97), custirsen (99), eteplirsen (103), fomivirsen (97), gataparsen (103), mipomersen (100), miravirsen (101), oblimersen (97), trabedersen (98), trecovirsen (97).

4.3. Antithrombins

antithrombin III (60), antithrombin alfa (93) (Rec. Glycoprotein (432aa) from transgenic goats).

---

6 The numbers in parentheses indicate the Proposed list number
4.4. **Blood coagulation cascade inhibitors**

The common stem for blood coagulation cascade inhibitors is *-cogin*.

*drotrecogin alfa (activated)* (86), *taneptacogin alfa* (90), *tifacogin* (78).

4.5. **Blood coagulation factors**

The common stem for blood coagulation factors is *-cog*.

The sub-stems *-eptacog*, *-octocog* and *-nonacog*, have been selected up to date for recombinant blood coagulation factors.

A prefix will be necessary if the amino acid sequence does not match that of the naturally occurring material.

In accordance with the general policy, *alfa*, *beta*, etc, will be added for the glycoproteins (see item 3.4 - general policies for glycosylated compounds).

When the additional statement "activated" is needed, e.g. for the blood coagulation factor VIIa, it should be spelt out in full and added in parentheses after the name.

**blood coagulation factor VII**: *-eptacog*

*eptacog alfa (activated)* (77), *eptacog alfa pegol (activated)* (101), *vatreptacog alfa (activated)* (98)

**blood factor VIII**: *-octocog*

*beroctocog alfa* (98), *moroctocog alfa* (72), *octocog alfa* (73)

**blood factor IX**: *-nonacog*

*nonacog alfa* (77), *nonacog beta pegol* (103)

**blood coagulation factor XIII**: *tridecacog*

*catridecacog* (99)

**recombinant von Willebrand factor (vWF)**: *-vonicog*

*vonicog alfa* (102).
4.6. Colony stimulating factors

The common stem for colony stimulating factors is \textit{-stim}.

\textit{ancestim} (79) (cell growth factor), \textit{garnocestim} (86) (immunomodulator), \textit{pegacaristim} (80) (megakaryocyte growth factor), \textit{romiplostim} (97) (platelet stimulating factor (through Mpl receptor))

combination of two different types of colony stimulating factors: \textit{-distim}

\textit{leridistim} (80), \textit{milodistim} (75)

granulocyte colony stimulating factor (G-CSF) type substances: \textit{-grastim}

\textit{filgrastim} (64), \textit{lenograstim} (64), \textit{nartograstim} (66), \textit{pegfilgrastim} (86), \textit{pegnartograstim} (80)

granulocyte macrophage colony stimulating factor (GM-CSF) types substances: \textit{-gramostim}

\textit{ecogramostim} (62), \textit{molgramostim} (64), \textit{gramostim} (65), \textit{sargramostim} (66)

macrophage stimulating factors (M-CSF) type substances: \textit{-mostim}

\textit{cilmostim} (71), \textit{lanimostim} (91), \textit{mirimostim} (65)

interleukin-3 analogues and derivatives: \textit{-plestim}

\textit{daniplestim} (76), \textit{muplestim} (74).

4.7. Enzymes

The common stem for enzymes, in general, is \textit{-ase}.

Sub-stems are referring to the activity of the substances.
proteinase:

with -ase suffix:
brinase (22), kallidinogenase (22), ocrase (28), pegaspargase (64), promelase (47), rasburicase (82), serrapeptase (31), sfericase (40), streptokinase (6), urokinase (48), urokinase alfa (77)

without -ase suffix:
batroxobin (29), bromelains (18), chymopapain (26), chymotrypsin (10), defibrotide (44), fibrinolysin (human) (10), sutilains (18)

Lipase: -lipase

bucelipase alfa (95), rizolipase (22)

enzymes with superoxide dismutase activity: -dismase

- ledismase (70), sudismase (58)
- isomerase (belongs to this group but in which the preferred stem has not been used)
  orgotein (31), pegorgotein (72)

plasminogen activator combined with another enzyme: -diplase

amediplase (79)

tissue-type-plasminogen activators: -teplase

alteplase (73), desmoteplase (80), dutiplase (62), lanoteplase (76), monteplase (72), nateplase (73), pamiteplase (78), retiplase (69), silteplase (65), tenecteplase (79)

anistreplase (59) (belongs to this group but in which the preferred stem has not been used)

urokinase-type-plasminogen activators: -uplase

nasaruplase (76), nasaruplase beta (86), saruplase (76)
others:

*agalsidase alfa (84):* human alpha-galactosidase isoenzyme A, isolated from human cell line, clone RAG 001, glycoform α

*agalsidase beta (84):* α-galactosidase (human clone λAG18 isoenzyme A subunit protein moiety reduced), glycoform β

*alfimeprase (85):* [3-L-serine]fibrolase-(3-203)-peptide (fibrolase : fibrinolytic enzyme isolated from *Agkistrodon contrix contrix* venom)

*alg glucerase (68):* glucosylceramidase (human placenta isoenzyme protein moiety reduced)

*alg glucosidase alfa (91):* human lysosomal prepro-α-glucosidase-(57-952)-peptide 199-arginine-223-histidine variant

*condoliase (102):* endolyase, chondroitin ABC (C-ABC), glycosaminoglycan lyase chondroitin ABC endolyase 1 (chondroitinase ABC) *Proteus vulgaris*

*dornase alfa (70):* deoxyribonuclease (human clone 18-1 protein moiety)

*epafipase (85):* 2-acetyl-1-alkyl-sn-glycero-3-phosphocholine deacetylase-(6-400)-peptide(human)

*eufauserase (84):* broad spectrum serine-protease enzyme, extracted from the Antartic krill (*Euphausia superba*)

*galsulfase (92):* N-acetylgalactosamine 4-sulfatase (human CSL4S-342 cell)

*glucarpidase (92):* recombinant glutamate carboxypeptidase (carboxypeptidase G2)

*hyalosidase (50):* hyaluronoglucosaminidase or E.C. 3.2.1.35

*hyaluronidase (1):* enzymes of various origins which depolymerize hyaluronic acid

*idursulfase (90):* α-L-iduronate sulfate sulfatase

*imiglucerase (72):* 495-L-histidineglucosylceramidase (human placenta isoenzyme protein moiety)

*laronidase (86):* 8-L-histidine-α-L-iduronidase (human)
**pegademase (63):** adenosine deaminase, reaction product with succinic anhydride, esters with polyethylene glycol monomethyl ether
   The source of the product should be indicated

**pegloticase (98):** tetramer $\alpha_4$ of des-(1-5)-(6-threonine,45-threonine,290-lysine, 300-serine]uricase (EC 1.7.3.3, urate oxidase) from *Sus scrofa* (porcine), non acetylated, of which some of the lysine 6-amine residues are engaged in a carbamate linkage with a monomethyl ether of polyoxyethylene (macrogol)

**pegsiticase (103):** pegylated Urate Oxidase from *Candida utilis*, [198-threonine(S>T)]uricase (EC 1.7.3.3, urate oxidase) *Pichia jadinii* (Yeast) (*Candida utilis*) tetramer, 6-amino group of an average of 3 lysine residues, mostly in position 16, 19, and 85 of each monomer, are amidified with $\alpha$-(3-carboxypropanoyl)-$\omega$-methoxy poly(oxyethylene)

**penicillinaspe (10):** an enzyme obtained by fermentation from cultures of *B.Cereus*

**ranpirnase (81):** ribonuclease (*Rana pipiens*)

**streptodornase (6):** enzyme obtained from cultures of various strains of *Streptococcus hemolyticus* and capable of hydrolysing deoxyribonucleoproteins

**taliglucerase alfa (101):** L-glutamyl-L-phenylalanyl-[495(497)-L-histidine(R>H)]human glucosylceramidase (beta-glucocerebrosidase) peptide with L-aspartyl-L-leucyl-L-leucyl-L-valyl-L-aspartyl-L-threonyl-L-methionine,glycosylated peptide 1-506

**tilactase (50):** $\beta$-D-galactosidase or EC 3.2.123

**velaglucerase alfa (98):** human glucosylceramidase (EC 3.2.1.45 or beta-glucocerebrosidase), glycoform $\alpha$.

### 4.8. Erythropoietin type blood factors

The common stem for erythropoietin type blood factors is *-poetin.*

In the case of erythropoietins, it was decided to select *epoetin* together with a Greek letter to differentiate between compounds of the same amino acid sequence as human erythropoietin which vary in the
glycosylation pattern (see item 3.4 - general policies for glycosylated compounds).

Substances with different amino acid sequences will be named using the -poetin stem and a random prefix.

darbepoetin alfa (85), epoetin alfa (66), epoetin beta (62), epoetin gamma (67), epoetin delta (85), epoetin epsilon (72), epoetin zeta (95), epoetin theta (95), epoetin kappa (97), epoetin omega (73).

4.9. Gene therapy products

alferminogene tadenovec (95), alipogene tiparvovec (99), amolimogene bepiplasmid (98), beperminogene perplasmid (95), contusugene ladenovec (97), golnerminogene pradenovec (101), riferminogene pecaplasmid (100), sitimagene ceradenovec (97), taberminogene vadenovec (100), tipapkinogene sovacivec (102), velimogene aliplasmid (97).

4.10. Growth factors

The common stem for growth factors is -ermin.
Sub-stems allow distinction between the various types of growth factors.
INNs for tumour necrosis factors (TNF) are also classified under the stem -ermin.

vascular endothelial growth factors: -bermin

telbermin (85)

epidermal growth factors: -dermin

murodermin (63), nepidermin (97)

fibroblast growth factors: -fermin

ersofermin (66), palifermin (88), repifermin (82), trafermin (74), velafermin (94)
leukaemia-inhibiting factors: -filermin

emfilermin (82)

tumour necrosis factors: -nermin

ardenermin (88), dulanermin (99), plusonermin (73), sonermin (68),
tasonermin (78)

platelet-derived growth factors: -plermin

becaplermin (74)

insulin-like growth factors: -sermin

mecasermin (66), mecasermin rinfabate (92)

transforming growth factors: -termin

cetermin (74), liatermin (81)

bone morphogenetic proteins: -otermin

avotermin (77), dibotermin alfa (89), eptotermin alfa (92),
radotermin (92)

others:

dapiclermin (93) (modified ciliary neurotrophic factor (CNTF)).

4.11. Growth hormone (GH) derivatives

The common stem for growth hormone derivatives is som-.
human growth hormone derivatives:
somatrem (54), somatropin (74), somatropin pegol (103)
For substances other than human, suffixes are added to indicate the species specificity of the structure.

bovine-type substances: -bove
somagrebove (63), somavubove (63), sometribove (74), somidobove (58)

porcine-type substances: -por
somalapor (62), somenopor (62), somfasepor (66), sometripor (75)

salmon-type substances: -salm
somatosalm (69)

others (growth hormone related peptides):
somatorelin (57) (pituitary hormone-release stimulating peptides, see item 4.26), somatostatin (46) (growth hormone release inhibitor).

4.12. Growth hormone antagonists

pegvisomant (82).

4.13. Heparin derivatives including low molecular mass heparins

The common stem for heparin derivatives including low molecular mass heparins is -parin.

ardeparin sodium (68), bemiparin sodium (75), certoparin sodium (70),
dalteparin sodium (77), deligoparin sodium (89), enoxaparin sodium (77),
heparin sodium (54), livaraparin calcium (86), minolteparin sodium (74),
nadroparin calcium (78), parnaparin sodium (77), reviparin sodium (78),
semuloparin sodium (99), tafoxiparin sodium (102), tinzaparin sodium (77).


The common stem for hirudin derivatives is -irudin.
bivalirudin (72), desirudin (76), lepirudin (76), pegmusirudin (77).

4.15. Pituitary hormone-release inhibiting peptides

The common stem for pituitary hormone-release inhibiting peptides is -relix.

abarelix (78), cetrorelix (66), degarelix (86), detirelix (56), ganirelix (65), iturelix (79), ozarelix (94), prazarelix (81), ramorelix (69), teverelix (78).

4.16. Human papilloma virus

verpasep caltespen (95) (heat-shock protein HSP 65 (Mycobacterium bovis strain BCG) fusion protein with transcription factor E7 (human papilloma virus 16)).

The suffix -tespen is the indicator of heat shock protein.

4.17. Insulins

Up to now, the insulin derivatives have been named using the two-word approach. The compounds named represent a structure with an additional amino acid, such as insulin argine (58), or represent modifications of the amino acid sequence, i.e. insulin aspart (76).

biphasic insulin injection (16), compound insulin zinc suspension (06), dalanated insulin (14), globin zinc insulin injection (06), insulin argine (58), insulin aspart (76), insulin defalan (37), insulin degludec (101), insulin detemir (80), insulin glargine (76), insulin glulisine (84), insulin human (48), insulin lispro (72), insulin tregopil (103), insulin zinc suspension (amorphous) (04), insulin zinc suspension (crystalline) (04), isophane insulin (04), neutral insulin injection (15), protamine zinc insulin injection (06)

argine: B30-yl-l-arginyll-l-arginine
aspart: [B28-L-aspartic acid]
dalanated: des-B30-alanine
defalan: des-B1-phenylalanine
degludec: $N^6$, B29-$[N-(15\text{-carboxypentadecanoyl})-L-\gamma\text{-glutamyl}]-\text{des-30B-L-threonine}$
detemir: $N^6$, B29-tetradecanoyl-des-B30-L-threonine
glargine: [A21-glycine], B30-yl-L-arginyl-L-arginine
glulisine: [B3-lysine, B29-glutamic acid]
lispro: [B28-yl-lysine, B29-L-proline]
tregopil: $N^6$, B29-(4,7,10,13-tetraoxatetradecanoyl).

4.18. Interferons

Interferon was published as an INN in 1962 with a general definition based on the origin and activity, e.g. "a protein formed by the interaction of animal cells with viruses capable of conferring on animal cells resistance to virus infection".

The name was revised in the 1980s when human interferon and its variations alfa, beta and gamma were produced by recombinant biotechnology. The INN Expert Group would have preferred to replace the old INN interferon by alfaferon, betaferon and gammaferon; however, this approach was barred as these names had already been registered as trade marks. The system adopted was thus to take interferon alfa, interferon beta and interferon gamma, and to provide, when necessary, for further distinction by additional numbers, or in the case of mixtures, by additional codes.

albinterferon alfa-2b (99), interferon alfa (73), interferon alfacon-1 (77), interferon beta (73), interferon gamma (73), peginterferon alfa-2a (84), peginterferon alfa-2b (84).
4.19. **Interleukin receptor antagonists**

The common stem for interleukin receptor antagonists is -kinra.

interleukin-1 (IL-1) receptor antagonists: -nakinra

*anakinra* (72)

interleukin-4 (IL-4) receptor antagonists: -trakinra

*pitrakinra* (87).

4.20. **Interleukin type substances**

The common stem for interleukin type substances is -kin.

In accordance with general policy for naming glycosylated proteins (see item 3.4), it was agreed to publish the INNs for glycosylated interleukins with alfa, beta.

interleukin-1 (IL-1) analogues and derivatives: -nakin

interleukin-1\(\alpha\) analogues and derivatives: -onakin

*pifonakin* (77)

interleukin-1\(\beta\) analogues and derivatives: -benakin

*mobenakin* (72)

interleukin-2 (IL-2) analogues and derivatives: -leukin

*adargileukin alfa* (89), *aldesleukin* (63), *celmoleukin* (65), *denileukin difitox* (78), *pegaldesleukin* (74), *teceleukin* (67), *tucotuzumab celmoleukin* (95)

interleukin-3 (IL-3) analogues and derivatives: -plestim (belongs to this group but in which the preferred stem has not been used)

*daniplestim* (76), *muplestim* (74)

interleukin-4 (IL-4) analogues and derivatives: -trakin

*binetrakin* (82)
interleukin-6 (IL-6) analogues and derivatives: -exakin
atexakin alfa (72)
interleukin-8 (IL-8) analogues and derivatives: -octakin
emoctakin (74)
interleukin-10 (IL-10) analogues and derivatives: -decakin
ilodecakin (81)
interleukin-11 (IL-11) analogues and derivatives: -elvekin
oprelvekin (76)
interleukin-12 (IL-12) analogues and derivatives: -dodekin
edodekin alfa (79)
interleukin-13 (IL-13) analogues and derivatives: -tredekin
cintredekin besudotox (92)
a recombinant human interleukin-18 (IL-18) with 157 amino acids:
iboctadekin (92)
a recombinant human interleukin-21: -enicokin
denenicokin (99)
neurotrophins (interleukin-78, brain derived neurotropic factor): -neurin (pre-stem)
abrineurin (84).

4.21. Monoclonal antibodies

The common stem for monoclonal antibodies is -mab.

INNs for monoclonal antibodies alphabetically by origin:
-axomab (pre-sub-stem, rat-murine hybrid)
catumaxomab (93), ertuxaxomab (93)

-omab (mouse origin)

abagovomab (95), afelimomab (80), altumomab (80), anatumomab
mafenatox (86), arcitumomab (74), bectumomab (81), besilesomab (92),
biciromab (66), blinatumomab (100), capromab (80), detumomab (80),
dorlimomab aritox (66), edobacomab (80), edrecolomab (74), elsilimomab
(89), enlimomab pegol (77), epitumomab (97), epitumomab
cituxetan (89), faralimomab (81), gavilimomab (84), ibrutumomab ituxetan
(86), igovomab (86), inciromab (66), inolimomab (80), lenalesomab (86),
maslimomab (66), minretumomab (80), mitumomab (82), moxetumomab
pasudotox (102), nacolomab tafenatox (80), naptumomab estafenatox (96),
nerelimomab (81), odulimomab (81), oregovomab (86), racotumomab (100),
saturnomab (81), sulesomab (86), taplitumomab paptopx (84), technetium
(99mTc) fanolesomab (86), technetium (99mTc) nofetumomab merpentan (81),
technetium (99mTc) pintumomab (86), telimomab aritox (66), tenatumomab
(99), tositumomab (80), vepalimomab (80), zolimomab aritox (80)

-umab (human origin)

adalimumab (85), adecatumumab (90), atorolimumab (80), belimumab (89),
bertilimumab (88), briakinumab (101), canakinumab (97), cixutumumab
(100), conatumumab (99), daratumumab (101), denosumab (94),
drozitumab (103), efungumab(95), exbivirumab (91), fezakinumab (101),
figitumumab (100), foralumab (103), foravirumab (100), fresolimumab
(101), ganitumab (103), gantenerumab (97), glembatumumab (102),
golimumab (91), intetumumab (101), ipilimumab (94), iratumumab (94),
lerdelimumab (86), lexatumumab (95), libivirumab (91), lucatumumab (98),
mapatumumab (93), mavrilimumab (102), metelimumab (88),
morolimumab (79), nebacumab (66), necitumumab (100), ofatumumab (93),
olaratumab (103), oxelumab (103), panatumumab (96), panobacumab (100),
pritumumab (89), rafivirumab (100), ramucirumab (100), raxibacumab
(92), regavirumab (80), rilotumumab (101), robatumumab (100), roledumab
(103), secukinumab (102), sevirumab (66), sifalimumab (101), stomatumab
(95), teprotumumab (101), tralokinumab (102), tremelimumab (97),
tuvirumab (66), ustekinumab (99), votumumab (80), zalutumumab (93),
zanolimumab (92), ziralimumab (84)
-ximab (chimeric origin)

abciximab (80), basiliximab (81), bavituximab (95), brentuximab vedotin (103), cetuximab (82), clenoliximab (77), ecromeximab (87), ensiuximab (103), galiximab (89), girentuximab (101), infliximab (77), iodine (124I), girentuximab (101), keliximab (81), lumiliximab (90), pagibaximab (93), priliximab (80), rituximab (77), siluximab (100), teneliximab (87), vapaliximab (87), volociximab (93)

-xizumab (chimeric/humanized origin, under discussion)

otelixizumab (99)

-zumab (humanized origin)

alacizumab pegol (98), alemtuzumab (83), anruxizumab (98), apolizumab (87), aselizumab (88), bapineuzumab (93), benralizumab (102), bevacizumab (86), bivatuzumab (86), cantuzumab mertansine (89), cedelizumab (81), certolizumab pegol (97), citatuzumab bogatox (99), yttrium (90Y) clivatuzumab tetraxetan (102), dacetuzumab (98), daclizumab (78), dalotuzumab (10), ecilizumab (87), efalizumab (85), eoliximab (100), epratuzumab (82), etaracizumab (99), erlizumab (84), farletuzumab (100), felvizumab (77), fontolizumab (87), gemtuzumab (83), ibalizumab (97), inotuzumab ozogamicin (92), itolizumab (103), labetuzumab (85), lebrikizumab (101), lintuzumab (86), lorivotuzumab mertansine (103), matuzumab (88), mepolizumab (81), milatuzumab (98), motavizumab (95), natalizumab (79), nimotuzumab (94), obinutuzumab (101), ocrelizumab (95), olokizumab (103), oportuzumab monatox (100), oxelumab (103), palivizumab (79), pascolizumab (87), pertuzumab (89), pexelizumab (86), ranibizumab (90), restiluzumab (85), rontalizumab (101), roveluzumab (81), ruplizumab (83), samalizumab (103), sibrotuzumab (86), siplizumab (87), solanezumab (100), sotuzumab (94), suviximab (102), tadocizumab (94), talizumab (89), tanezumab (99), tefizumab (92), teplizumab (97), tigatuzumab (98), tocilizumab (90), toralizumab (87), trastuzumab (78), trastuzumab emtansine (103), tucotuzumab celmoleukin (95), urtoxazumab (90), vedolizumab (100), veltuzumab (98), visilizumab (84), yttrium (90Y) tacatuzumab tetraxetan (93)

Others: muromonab-CD3 (59) (belongs to this group but in which the preferred stem has not been used).
4.22. Oxytocin derivatives

The common stem for oxytocin derivatives is \textit{-tocin}.

\begin{itemize}
\item argiprestocin (13), aspartocin (11), carbetocin (45), cargutocin (35), demoxytocin (22), nacartocin (51), oxytocin (13).
\end{itemize}

4.23. Peptides and glycopeptides

for special groups of peptides see \textit{-actide} (see item 4.28), \textit{-pressin} (see item 4.31), \textit{-relin} (see item 4.26), \textit{-tocin} (see item 4.22)

The common stem for peptides and glycopeptides is \textit{-tide}.

analgesic: leconotide (86), ziconotide (78)

angiogenesis inhibitor: cilengitide (81)

angiotensin converting-enzyme inhibitor: teprotide (36)

antianaemic : peginesatide (103)

anti-inflammator: icrocaptide (89)

antiarrythmic: rotigaptide (94), danegaptide (101)

antidepressant: nemifitide (87)

antidiabetic: albighitude (97), amlintide (76), davalintide (101), dulaglutide (103), exenatide (89), liraglutide (87), lixisenatide (99), pramlintide (74), seglitide (57), semaglutide (101), taspoglutide (99)

antidiarrhoeal: lagatide (75)

antiobesity drug: obinepitide (96)

antithrombotic: eptifibatide (78) (-fiba- is a pre-substem for platelet aggregation inhibitor (GPIIb/IIIa receptor antagonist))

antiviral: enfuvirtide (85), tifuvirtide (91)

atrial natriuretic factor type substance: anaritide (57), neseritide (80), ularitide (69)
autoimmune disorders: dirucotide (100)

cardiac stimulant: carperitide (65)

diagnostic: betiatide (58), bibapcitide (78), ceruletide (34), depreotide (80), fluciclatide (18F) (103), maraciclatide (103), mertiatide (60), pendetide (70), technetium (99mTc) apcitide (86), teriparatide (50)

expectorant (in cystic fibrosis): lancovutide (99)

gastro-intestinal bleeding / antineoplastic: edotreotide (84), ilatreotide (68), lanreotide (64), octreotide (52), pentetreotide (66), vapreotide (62)

gastro-intestinal functions normalizing agent: teduglutide (90), linaclotide (97)

growth stimulant-veterinary: nosiheptide (35)

gut motility increasing: ociltide (52)

hormone analogue: semparatide (80)

immunological agents - antineoplastics: almurtide (74), delmitide (92), disomotide (94), edratide (89), elpamotide (103), goralatide (72), mifamurtide (95), murabutide (49), ovemotide (94), pentigetide (60), pimelautide (53), prezatide copper acetate (67), rolipoltide (94), romurtide (61), tabilautide (60), temurtide (60), tertomotide (98), tigapotide (95), tiplimotide (82)

inhibition of growth hormone release: pasireotide (90)

kallicrein inhibitor: ecallantide (93)

melanocortin receptor agonist: bremelanotide (95), afamelanotide (99)

neuromodulator / neuroprotective agent: ebiratide (56), davunetide (100), vanutide cridificar (100)
peptic ulcer: sulglicotide (29), triletide (50)
pulmonary surfactant: lusupultide (80), sinapultide (78)
sedative: emideltide (70)
transforming growth factor beta-1 inhibitor: disitertide (99)
treatment of Parkinson's disease: doreptide (59), pareptide (38)
wound healing agent: rusalatide (96)
zonulin antagonist (in celiac disease): larazotide (99)
other: defibrotide (44) (nucleotide).

4.24. Peptide vaccines / recombinant vaccines

Definition of peptide vaccines: vaccine in which antigens are produced from synthetic peptides and transported through the bloodstream by an adjuvant, in order to stimulate an immune response.

Definition of recombinant vaccines: vaccine produced from a cloned gene.

Description of recombinant vaccines: there are certain antigens on viruses and bacteria which are better at stimulating an antibody response by the animal than others. The genes for these antigens can be isolated, and made to produce large quantities of the antigens they code for. A recombinant vaccine contains these antigens, not the whole organism. Compare with "modified live vaccine" and "killed vaccine".

The following substances are peptide vaccines: disomotide (94), elpamotide (103), ovemotide (94), tertomotide (98), tiplimotide (82).

4.25. Pituitary / placental glycoprotein hormones

The names selected by the International Union of Pure and Applied Chemistry–International Union of Biochemistry (IUPAC-IUB) have, to date, been chosen for compounds with an amino acid sequence identical
to that of the naturally occurring human hormones. Addition of a 
Greek letter as the second part of the name will allow differentiation of 
different glycosylation patterns for compounds produced by 
biochemistry (see item 3.4 - general policies for glycosylated 
compounds).

follicle stimulating hormones: ending in (-)follitropin

corifollitropin alfa (80), follitropin alfa (71), follitropin beta (75),
urofollitropin (57), varfollitropin alfa (101)

gonadotropin: ending in -gonadotropin

choriogonadotropin alfa (76), chorionic gonadotrophin (01): chorionic 
gonadotropins, obtained from human serum and urine during pregnancy and 
has both lutropin and follitropin activity

serum gonadotrophin (01): used for the follicle stimulating hormone (FSH, 
follitropin) from serum of pregnant mares

luteinizing hormones: ending in (-)lutropin

lutropin alfa (71).

4.26. Pituitary hormone-release stimulating peptides

The common stem for pituitary hormone-release stimulating peptides is 
-relin.

LHRH-release-stimulating peptides:

avorelin (74), buserelin (36), deslorelin (61), fertirelin (42), gonadorelin 
(32), goserelin (55), histrelin (53), leuprolelin (47), lutrelin (51), nafarelin 
(50), peforelin (93), triptorelin (58)

growth hormone release-stimulating peptides: -morelin

anamorelin (97), capromorelin (83), dumorelin (59), examorelin (72), 
ipamorelin (78), macimorelin (100), pralmorelin (77), rismorelin (74), 
sermorelin (56), somatotropin (57), tabimorelin (86), tesamorelin (96), 
ulimorelin (103)
thyrotropin releasing hormone analogues: -tirelin

azetirelin (60), montirelin (58), orotirelin (58), posatirelin (60), protirelin (31), taltirelin (75)

thyrotropin alfa (78) (thyrotropin releasing hormone (TRH) analog, belongs to this group but in which the preferred stem has not been used)

other: corticorelin (66) (diagnostic agent).

4.27. Receptor molecules, native or modified

The stem for receptor molecules, native or modified is -cept.

A preceding infix should designate the target.

vascular endothelial growth factor receptors: -ber-
aflibercept (96)

complement receptors: -co-
mirococept (91)

subgroup of interferon receptors: -far-
bifarcept (86)

lymphocyte function-associated antigen 3 receptors: -efa-
alefacept (84)

interleukin-1 receptors: -na-
rilonacept (95)

transforming growth factor receptors: -ter-
sotatercept (102)

tumour necrosis factor receptors: -nercept.
baminercept (99), etanercept (81), lenercept (72), onercept (86), pegsunercept (95).

cytotoxic T lymphocyte-associated antigen 4 (CTLA-4) receptors: -ta-
abatacept (91), belatacept (93)

antiviral receptors: -vir-
alvircept sudotox (69)
other: atacicept (95), briobacept (98).

4.28. Synthetic polypeptides with a corticotropin-like action

The common stem for synthetic polypeptides with a corticotropin-like action is -actide.

alsactide (45), codactide (24), giractide (29), norleusactide (18), seractide (31), tetracosactide (18), tosactide (24), tricosactide (44), tridecactide (97).

4.29. Thrombomodulins

sorthrombomodulin alfa (101), thrombomodulin alfa (94).

4.30. Toxins

toxin ML-1 (mistletoe lectin I) (Viscum album): aviscumine (86).

4.31. Vasoconstrictors, vasopressin derivatives

The common stem for vasoconstrictors, vasopressin derivatives is -pressin.

argipressin (13), desmopressin (33), felypressin (13), lypressin (13), ornipressin (22), terlipressin (46), vasopressin injection (16).

4.32. Various

- alisporivir (100): [8-(N-methyl-D-alanine),9-(N-ethyl-L-valine)]cyclosporine
• agatolimod (98): DNA, d(P-thio)(T-C-G-T-C-G-T-T-G-T-C-G-T-T-G-T-C-G-T-T)

• angiotensin II (65): 5-L-isoleucineangiotensin II (the source of the material should be indicated)

• angiotensinamide (12): N-{1-{N-{N-[N-(N²-asparaginylarginyl)valyl]tyrosyl}valyl}histidyl}prolyl]-3-phenylalanine

• bevasiranib (99): siRNA inhibitor of Vascular Endothelial Growth Factor (VEGF) production

• calcitonin (80): a polypeptide hormone that lowers the calcium concentration in blood (the species specificity should be indicated in brackets behind the name)

• conestat alfa (98): human plasma protease C1 inhibitor (C1 esterase inhibitor) (N,O-glycosylated recombinant protein expressed in the mammary gland of transgenic rabbits), glycoform α

• epelestat (92): human recombinant neutrophil elastase inhibitor, bovine pancreatic trypsin inhibitor (BPTI) homologue

• edifoligide (89): oligonucleotide

• hemoglobin glutamer (80): the species specificity should be indicated in brackets behind the name, "(bovine)"; the average mass of the polymer is given as e.g. haemoglobin glutamer-250 for 250kD

• hemoglobin crosfumaril (76): hemoglobin A₀ (human α₂β₂ tetrameric subunit), α-chain 99,99'-diamide with fumaric acid

• hemoglobin raffimer (89)

• imetelstat (101): oligonucleotide telomerase inhibitor; 3'-amino-3'-deoxy-P-thiothymidylyl-(3'→5')-3'-amino-2',3'-dideoxy-P-thiodenyllyl-(3'→5')-3'-amino-2',3'-dideoxy-P-thioguanlyyl-(3'→5')-3'-amino-
2',3'-dideoxy-\textit{P}-thioguanylyl-(3'\rightarrow5')-3'-amino-2',3'-dideoxy-\textit{P}-thioguanylyl-(3'\rightarrow5')-3'-amino-3'-deoxy-\textit{P}-thiothymidylyl-(3'\rightarrow5')-3'-amino-2',3'-dideoxy-\textit{P}-thioadenylyl-(3'\rightarrow5')-3'-amino-2',3'-dideoxy-\textit{P}-thioguanylyl-(3'\rightarrow5')-3'-amino-2',3'-dideoxy-\textit{P}-thioadenylyl-(3'\rightarrow5')-3'-amino-2',3'-dideoxy-\textit{P}-thiocytidylyl-(3'\rightarrow5')-3'-amino-2',3'-dideoxy-\textit{P}-thioadenylyl-(3'\rightarrow5')-3'-amino-2',3'-dideoxyadenosine 5'-\{O-[2-hydroxy-3-(hexadecanoylamino)propyl] hydrogen phosphorothioate\}

- **iodinated \(^{125}\text{I}\) human serum albumin (24):** human serum albumin iodinated with radioactive iodine \(^{125}\text{I}\)

- **iodinated \(^{131}\text{I}\) human serum albumin (24):** human serum albumin iodinated with radioactive iodine \(^{131}\text{I}\)

- **iroplact (74):** \(N\)-L-methionyl blood platelet factor 4 (human subunit)

- **ismomultin alfa (91):** 47-261-Glycoprotein gp 39 (human clone CDM8-gp39 reduced)


- **macrosalb \(^{131}\text{I}\) (33):** macroaggregated iodinated \(^{131}\text{I}\) human albumin

- **macrosalb \(^{99m}\text{Tc}\) (33):** technetium \(^{99m}\text{Tc}\) labelled macroaggregated human serum albumin

- **metenkefalin (97):** L-tyrosylglycylglycyl-L-phenylalanyl-L-methionine \(\beta\)-endorphin human-(1-5)-peptide

- **metreleptin (82):** \(N\)-methionylleptin (human)

- **mirostipen (85):** [23-methionine] human myeloid progenitor inhibitory factor 1-(23-99)-peptide

- **muromonab-CD3 (59):** a biochemically purified IgG\(_{2\alpha}\) immunoglobulin consisting of a heavy chain of approx. 50,000 daltons and a light chain of approx. 25,000 daltons. It is manufactured by a process involving the fusion of mouse myeloma
cells to lymphocytes from immunized animals to produce a hybridoma which
secretes antigen-specific antibodies to the T3 antigen of human T-lymphocytes.

- **nagrestipen (76)**: 26-L-alaninelymphokine MIP 1α (human clone pAT464
macrophage inflammatory)

- **ocriplasmin (101)**: truncated human plasmin:
  human plasmin heavy chain A-(543-561)-peptide (548-666;558-566)-bisdisulfide
  with human plasmin light chain B

- **opebacan (83)**: 132-L-alanine-1-193-bactericidal / permeability-increasing protein
  (human)

- **orgotein (31)**: a group of soluble metalloproteins isolated from liver, red blood
cells, and other mammalian tissues

- **ovandrotone albumin (52)**: 3-[(3,17-dio xoandrost-4-en-7α-yl)thio]propionic acid,
  serum albumin conjugate

- **parathyroid hormone (90)**: non glycosylated human parathyroid hormone, the
  origin should be indicated between brackets after the INN, for example (r. *E. coli*)
  for recombinant produced by *Escherichia coli*

- **pegaptanib (88)**: 5’-ester of (2’-deoxy-2’-fluoro)C-Gm-Gm-A-A-(2’-deoxy-2’-
  fluoro)U-(2’-deoxy-2’-fluoro)C-(2’-deoxy-2’-fluoro)C-Gm-(3’→3’)-dT with α,α-
  [[(1S)-1-[[5-(phosphonooxy)pentyl]carbamoyl]pentane-1,5-
diyl]bis(iminocarbonyl)]bis[ω-methoxypoly(oxyethane-1,2-diyl)]

- **pegdinetanib (103)**: 94 residues protein derived from human fibronectin 10th type
  III domain, pegylated:
glycyl[1438-L-arginine(D>R),1439-L-histidine(A>H),1441-L-
histidine(A>H),1442-L-phenylalanine(V>F),1443-L-proline(T>P),1444-L-threonine(V>T),1467-L-leucine(G>L),1468-L-glutamine(S>Q),1469-L-
proline(K>P),1470-L-proline(S>P),1492-L-aspartic acid(G>D),1493-glycine(R>G),1494-L-arginine(G>R),1495-L-asparagine(D>N),1496-glycine(S>G),1497-L-arginine(P>R),1498-L-leucine(A>L),1499-L-
leucine(S>L),1501-L-isoleucine(K>I),1515-S-[(3RS)-1-(1-[[α-
methylpoly(oxyethylene)carbamoyl]-3-[(α-methylpoly(oxyethylene)carbamoyl)oxy]methyl]-8,13-dioxo-1,4-dioxao-9,12-diazapentadecan-15-yl]-2,5-dioxopyrrolidin-3-yl]-L-cysteine(S>C)][human fibronectin-(1424-1516)-peptide

- **rintatolimod (102):** poly[5'-inosinylyl-(3'→)] duplex with poly[dodecakis[3']-cytidylyl-(5'→)3'-uridylyl-(5'→)]

- **secretin (01):** hormone of the duodenal mucosa which activates the pancreatic secretion and lowers the blood-sugar level

- **talactoferrin alfa (93):** recombinant human lactoferrin

- **tadekinig alfa (90):** interleukin-18 binding protein (human gene IL 18BP isoform a precursor)

- **thrombin alfa (97):** human thrombin (recombinant, glycoform α)

- **tiprelestat (103):** human elafin (elastase-specific inhibitor, skin-derived antileukoproteinase, peptidase inhibitor 3)

- **torapsel (91):** 42-89-glycoprotein (human clone PMT21:PL85 P-selectin glycoprotein ligand 1) fusion protein with immunoglobulin (human constant region)

- **tremacamra (78):** 1-453-glycoprotein ICAM-I (human reduced)

- **troplasminogen alfa (99):** thrombin-activable plasminogen; endo-[((558a(559)-558h(365))-human coagulation factor XI-(363-370)-peptide]-des-(559-562)-(606(610)-lysine,623(627)-lysine]human plasminogen, glycoform α

- **votucalis (96):** methionyl[145-leucine]FS-HBP2 (*Rhipicephalus appendiculatus* (Brown ear tick) Female-Specific Histamine-Binding Protein 2).
5. CURRENT CHALLENGES

The challenges currently faced include the following:

- The INN Expert Group, when selecting names for recombinant proteins, has to deal not with substances with well-defined structures but with products of highly complex composition or even with mixtures of such products.

- It is not only modified proteins that might differ from their naturally occurring counterparts, products derived by expression of the natural gene in foreign host cells may also differ structurally, biologically or immunologically from the natural protein.

- Glycoproteins particularly may occur in forms that differ in the structure of one or more of their carbohydrate units, a phenomenon known as microheterogeneity and resulting in a heterogeneous population of molecules. Such differences may affect both the size and the charge of individual glycoproteins.

- A variety of novel biotechnology-derived products are under development, all of which will require specific policies on how to deal with such products.

- Clearly, the INN nomenclature of biological medicinal products is an area of increasing complexity.
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21. International nonproprietary names (INN) for pharmaceutical substances: names for radicals, groups & others (Comprehensive list), WHO/EMP/QSM/2010.3 *

   * These documents are available on the INN Programme Website at: http://www.who.int/medicines/services/inn/en/
ANNEX 1

The list of fusion proteins published\textsuperscript{7} classified by groups

-cept

\textit{abatacept (91)}

1-25-oncostatin M (human precursor) fusion protein with CTLA-4 (antigen) (human) fusion protein with immunoglobulin G1 (human heavy chain fragment), bimolecular (146\textrightarrow146\textsuperscript{'})-disulfide

\textit{aflibercept (95)}

des-432-lysine-[human vascular endothelial growth factor receptor 1-(103-204)-peptide (containing Ig like C2 type 2 domain) fusion protein with human vascular endothelial growth factor receptor 2-(206-308)-peptide (containing Ig like C2 type 3 domain fragment) fusion protein with human immunoglobulin G1-(227 \ C-terminal residues)-peptide (Fc fragment)], (211-211\textsuperscript{'}:214-214\textsuperscript{'})-bisdisulfide dimer

\textit{atacicept (95)}

[86-serine,101-glutamic acid,196-serine,197-serine,222-aspartic acid,224-leucine][human tumor necrosis factor receptor superfamily member 13B-(30-110)-peptide (TAC1 fragment containing TNFR-Cys 1 and TNFR-Cys 2) fusion protein with human immunoglobulin G1-(232 \ C-terminal residues)-peptide (γ1-chain Fc fragment), (92-92\textsuperscript{'}:95-95\textsuperscript{'})-bisdisulfide dimer

\textit{baminercept (99)}

human tumor necrosis factor receptor superfamily member 3 (lymphotoxin-β receptor, TNF C receptor)-(2-195)-peptide (fragment of extracellular domain) fusion protein with human immunoglobulin heavy constant γ1 chain Fc fragment [227 residues, hinge (195-205) des-(1-4),C5>V, CH2 (206-315), CH3 (316-421) des-K\textsuperscript{107}]

\textsuperscript{7} It should be noted that this list may not be comprehensive and the descriptions under the names are the ones published
rilonacept (95)

[653-glycine][human interleukin-1 receptor accessory protein-(1-339)-peptide (extracellular domain fragment) fusion protein with human type 1 interleukin-1 receptor-(5-316)-peptide (extracellular domain fragment) fusion protein with human immunoglobulin G1-(229 C-terminal residues)-peptide (Fc fragment)], (659-659':662-662')-bisdisulfide dimer

sotatercept (102)

fusion protein for immune applications (FPIA) comprising ACVR2A (activin receptor type 2A, activin receptor type IIA) fragment fused with immunoglobulin G1 Fc fragment, and binding activin, a member of the TGF beta family; ACVR2A, 21-135 precursor fragment (1-115) -threonyl-triglycyl linker (116-119) -gamma1 chain H-CH2-CH3 fragment (120-344) [Homo sapiens IGHG1*03 hinge (120-127), CH2, A115>V (128-237), CH3 (238-344)]; (123-123':126-126')-bisdisulfide dimer

-cept & -tox

alvircept sudotox (69)


interferon

albinterferon alfa-2b (97)

human serum albumin (585 residues) fusion protein with human interferon α-2b (165 residues)

\textsuperscript{8} The names and the descriptions of toxins are published in Annex 4-1 of "International Nonproprietary Names (INN) for pharmaceutical substances. Names for radicals, groups & others: comprehensive list (WHO/EMP/QSM/2010.3)"
-kin & -tox

**cintredekin besudotox (92)**

toxin hIL13-PE38QQR (plasmid phuIL13-Tx)

**denileukin difitox (78)**


-mab & -kin

**tucotuzumab celmoleukin (95)**

immunoglobulin G1, anti-(tumor associated calcium signal transducer 1 (KS 1/4 antigen)) (human-mouse monoclonal huKS-IL2 heavy chain) fusion protein with interleukin 2 (human), disulfide with human-mouse monoclonal huKS-IL2 light chain, dimer

-mab & -tox

**anatumomab mafenatox (86)**

immunoglobulin G 1, anti-(human tumor-associated glycoprotein 72) (human-mouse clone pMB125 Fab fragment γ1-chain) fusion protein with enterotoxin A (227-alanine) (*Staphylococcus aureus*) complex with mouse clone pMB125 κ-chain

**citatumab bogatox (99)**

immunoglobulin Fab fusion protein, anti-*[Homo sapiens* tumor-associated calcium signal transducer 1 (TACSTD1, gastrointestinal tumor-associated protein 2, GA733-2, epithelial glycoprotein 2, EGP-2, epithelial cell adhesion molecule Ep-CAM, KSA, KS1/4 antigen, M4S, tumor antigen 17-1A, CD326)], humanized
Fab fused with *Bougainvillea spectabilis* Willd rRNA N-glycosidase [type I ribosome inactivating protein (RIP), bougainin], VB6-845; gamma1 heavy chain fragment (1-225) [hexahistidyl (1-6) -humanized VH from 4D5MOC-B (*Homo sapiens* FR/*Mus musculus* CDR, *Homo sapiens* IGHJ4*01, V124>L) [8.8.9] (7-122) -*Homo sapiens* IGHG1*01 CH1-hinge fragment EPKSC (123-225)], (225-219')-disulfide with kappa fusion chain (1'-481') [humanized V-KAPPA from clone 4D5MOC-B (*Homo sapiens* FR/*Mus musculus* CDR, *Homo sapiens* IGKJ1*01, I126>L) [11.3.9] (1'-112') -*Homo sapiens* IGKC*01 (113'-219') -12-mer furin linker (proteolytic cleavage spacer from *Pseudomonas* exotoxin A) (220'-231') -*Bougainvillea spectabilis* Willd bouganin fragment (27-276 from precursor, V354'>A, D358'>A, Y364'>N, I383'>A) (232'-481')]

dorlimomab aritox (66)

ricin A chain-antibody ST 1 F(ab')2 fragment immunotoxin

*moxetumomab pasudotox* (102)

immunoglobulin Fv fragment fused to *Pseudomonas* toxin, anti-*[Homo sapiens* CD22 (sialic acid-binding Ig-like lectin 2, Siglec-2, SIGLEC2, Leu-14, B-lymphocyte cell adhesion molecule, BL-CAM)], *Mus musculus* monoclonal antibody disulfide stabilized Fv fragment with the variable heavy VH domain fused with the truncated form PE38 of *Pseudomonas aeruginosa* exotoxin A (VH-PE38), disulfide linked with the variable kappa domain (V-KAPPA); VH-PE38 (1-476) comprising the VH domain (1-123) [methionyl-*Mus musculus* VH [[IGHV5-12-1*01 -IGHD]-IGHJ3*01] [8.8.16] (2-123)] fused with a 7-mer linker (124-130) and with the *Pseudomonas aeruginosa* exotoxin A (ETA) PE38 fragment (131-476) [277-638 precursor fragment with del 389-405>N (131-476), containing domain II (131-243) with furin proteolytic cleavage site (152-164), domain Ib (244-267), domain III (268-476)], (45-101')-disulfide with V-KAPPA (1'-108') [methionyl-*Mus musculus* V-KAPPA [[IGHKV10-96*01 -IGKJ1*01] [6.3.9] (2'-108')]

*nacolomab tafenatox* (80)

immunoglobulin G1, anti-(human colorectal tumor antigen C242) Fab fragment (mouse monoclonal r-C242Fab-SEA clone pkP941 γ1-chain) fusion protein with enterotoxin A (*Staphylococcus aureus*), disulfide with mouse monoclonal r-C242Fab-SEA clone pkP941 κ-chain
naptumomab estafenatox (96)


oportuzumab monatox (100)

immunoglobulin scFv fusion protein, anti-[Homo sapiens tumor-associated calcium signal transducer 1 (TACSTD1, gastrointestinal tumor-associated protein 2, GA733-2, epithelial glycoprotein 2, EGP-2, epithelial cell adhesion molecule Ep-CAM, KSA, KS1/4 antigen, M4S1, tumor antigen 17-1A, CD326)] humanized monoclonal antibody scFv fused with Pseudomonas aeruginosa exotoxin A; hexahistidyl-humanized scFv [V-KAPPA (Homo sapiens IGKV1-39*01 (78%)-IGKJ1*01, I126->L) [11.3.9] (7-118)-26-mer linker -VH (Homo sapiens IGHV7-4-1*02 -IGHD)-IGHJ4*01, V124->L) [8.8.9] (145-260)] -20-mer linker - Pseudomonas aeruginosa exotoxin A (ETA) [277-633 precursor fragment, containing domain II (281-393) with furin proteolytic cleavage site (302-313), domain Ib (394-433), domain III (434-637)] (281-637) -hexahistidyl-lysyl-aspartyl-glutamylleucyl

taplitumomab paptovx (84)

immunoglobulin G1, anti-(human antigen CD19) (mouse monoclonal B43 \(\gamma_1\)-chain), disulfide with mouse monoclonal B43 \(\kappa\)-chain, dimer, disulfide with protein PAP (pokeweed antiviral)

telimomab aritox (66)

ricin A chain-antibody T 101 Fab fragment immunotoxin

zolimomab aritox (80)

immunoglobulin G1, anti-(human CD5 (antigen) heavy chain) (mouse monoclonal H65-RTA \(\gamma_1\)-chain), disulfide with mouse monoclonal H65-RTA light chain, dimer, disulfide with ricin (castor bean A-chain)
-stim (colony stimulating factors)

**romiplostim** (97)

L-methionyl[human immunogloblin heavy constant gamma 1-(227 C-terminal residues)-peptide (Fc fragment)] fusion protein with 41 amino acids peptide, (7-7';10,10')-bisdisulfide dimer

-tide

**albiglutide** (97)


**dulaglutide** (103)


**vanutide cridificar** (100)

inactivated diphtheria toxin (carrier) covalently linked to human beta-amyloid protein 42 short fragments: pentadecakis[N^6-Lys-(sulfanylacetyl)]-[52-glutamic acid(G>E)]diphtheria toxin Corynebacterium diptheriae thioether with human beta-amyloid protein 42-(1-7)-peptidylcysteine

Others

**torapsel** (91)

42-89-glycoprotein (human clone PMT21:PL85 P-selectin glycoprotein ligand fusion protein with immunoglobulin (human constant region)
transferrin aldifitox (95)

a conjugate of the precursor of human serotransferrin (siderophillin) with a primary amine group used to form an amidine with (4-iminobutane-1,4-diy) sulfanediyl[(3RS)-2,5-dioxopyrrolidine-1,3-diy]-1,3-phenylenecarbonyl and forming an N-benzoyl derivative of a primary amine group of diphtheria [550-L-phenylalanine]toxin from Corynebacterium diphtheriae-(26-560)-peptide

verpasep caltespen (95)

60 kDa chaperonin 2 (heat shock protein 65 from Mycobacterium bovis strain BCG) fusion protein with L-histidylprotein E7 from human papillomavirus type 16
## ANNEX 2

### Transliteration of Greek letters in English, French and Spanish

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<thead>
<tr>
<th>Upper case</th>
<th>Lower case</th>
<th>English</th>
<th>French</th>
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* * letters to be avoided
ANNEX 3

The previous naming scheme for monoclonal antibodies

General policies for monoclonal antibodies

- The common stem for monoclonal antibodies is -mab.
- Sub-stems for source of product:

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<thead>
<tr>
<th>Sub-stem</th>
<th>Source</th>
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<tr>
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<td>axo (pre-sub-stem)</td>
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<td>hamster</td>
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<td>xi</td>
<td>chimeric</td>
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<td>zu</td>
<td>humanized</td>
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</table>

The distinction between chimeric and humanized antibodies is as follows:

A chimeric antibody is one that contains contiguous foreign-derived amino acids comprising the entire variable region of both heavy and light chains linked to heavy and light constant regions of human origin.

A humanized antibody has segments of foreign-derived amino acids interspersed among variable region segments of human-derived amino acid residues and the humanized heavy-variable and light-variable regions are linked to heavy and light constant regions of human origin.
- Sub-stems for disease or target class:

<table>
<thead>
<tr>
<th>Sub-stem</th>
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<td>fungal</td>
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<td>interleukin</td>
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<td>-le(s) -</td>
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</tr>
<tr>
<td>-li(m) -</td>
<td>immunomodulator</td>
</tr>
<tr>
<td>-os -</td>
<td>bone</td>
</tr>
<tr>
<td>-vi(r) -</td>
<td>viral</td>
</tr>
</tbody>
</table>

- Tumours:

<table>
<thead>
<tr>
<th>Sub-stem</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>-co(l) -</td>
<td>colon</td>
</tr>
<tr>
<td>-go(t) -</td>
<td>testis</td>
</tr>
<tr>
<td>-go(v) -</td>
<td>ovary</td>
</tr>
<tr>
<td>-ma(r) -</td>
<td>mammary</td>
</tr>
<tr>
<td>-me(l) -</td>
<td>melanoma</td>
</tr>
<tr>
<td>-pr(o) -</td>
<td>prostate</td>
</tr>
<tr>
<td>-tu(m) -</td>
<td>miscellaneous</td>
</tr>
</tbody>
</table>

Whenever there is a problem in pronunciation, the final letter of the sub-stems for diseases or targets may be deleted, e.g. -vi(r)-, -ba(c)-, -li(m)-, -co(l)-, etc.

**Prefix**

Should be random e.g. the only requirement is to contribute to a euphonious and distinctive name.
Second word

If the product is radiolabelled or conjugated to another chemical, such as toxin, identification of this conjugate is accomplished by use of a separate, second word or acceptable chemical designation.

If the monoclonal antibody is used as a carrier for a radioisotope, the latter will be listed first in the INN, e.g. technetium ($^{99m}$Tc) pintumomab (86).

-toxa- infix

For monoclonals conjugated to a toxin, the infix -toxa- can be inserted either into the first (main) name or included in the second word.