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世界知识产权组织标准委员会(CWS)

第七届会议 2019年7月1日至5日,日内瓦

修订产权组织标准 ST. 26

国际局编拟的文件

导 言

- 1.. 在 2018 年于日内瓦举行的产权组织标准委员会(CWS)第六届会议上,序列表工作队(SEQL TF)对产权组织标准 ST. 26 提出了一系列修正(见文件 CWS/6/16)。这些修正包括对标准主体的修改、对产权组织 ST. 26 附件一、二、三、四和六的修正,以及增加新的附件七,提供产权组织标准 ST. 25 和产权组织标准 ST. 26 的转变映射。标准委员会通过了标准的新版本 1.2 版,其中收入了文件 CWS/6/16 中所记载的建议修改,但还增加了以下内容:
 - 将三处"legal"改为"permitted";
 - 将附件六"指导文件"15处"portion(s)"改为"regions";
 - 在附件七"将序列表从 ST.25 转至 ST.26 的建议"的场景 9 第一句后增加一句: "The nucleotide sequence feature key "modified base" is also present in both WIPO ST.25 and ST.26; however, Scenario 7 contains appropriate recommendations"。
- 2. 标准委员会第六届会议还同意将第44号任务的说明修改为:
 - "为国际局提供支持,提供用户对 ST. 26 编著和验证软件工具的要求和反馈意见;在对《PCT 行政规程》进行相应修订的工作上,为国际局提供支持;并且为产权组织标准 ST. 26 编制必要的修订。"
- 2.. 由此,2019年2月发布了产权组织ST.26的最新版1.2版。

3.. 讨论产权组织 ST. 26 修订的工作队会议召开了两次:一次在 2018 年 10 月标准委员会第六届会议期间,一次于 2019 年 4 月通过在线会议进行。

拟议修订总结

文字修改

- 4.. 在第 44 号任务的框架中,序列表工作队对产权组织 ST. 26 目前发布版本 1.2 版进行了审查。审查中,他们发现了一系列需要进行的文字修改,如多余空格或拼写错误。为确保该文件正确反映产权组织《风格指南》,也需要进行一些更正,例如用于替换"for example (例如)"的拉丁文缩写一致写为"e. g.,"。
- 5.. 为供标准委员会在第七届会议上审议,这些更正在本文件附件(产权组织 ST. 26 附件一)中用绿色高亮文本标出。工作队在产权组织 ST. 26 附件一至七中都找出了文字修改,但仅附件一转录于本文件中。

实质修改

- 6. 序列表工作队提议了下列修改,以进一步修订产权组织 ST. 26 的附件一和七,这些在附件中用黄色高亮标明增加的内容,用紫色高亮标明删除的内容。未对产权组织 ST. 26 的主体提出实质修改:
 - (a) 更新附件一表 9, 收入 INSDC 特征表 10.8 版中的更新;
 - (b) 附件一第 5.27 节中,收入下列新增的可选限定词:
 - function
 - gene
 - gene synonym
 - map
 - (c) 附件一第 5.33 节中,收入下列新增的可选限定词:
 - allele
 - direction
 - gene
 - gene synonym
 - map
 - note
 - standard name
 - (d) 附件一第 5.43 节中, 收入下列新增的可选限定词:
 - operon
 - (e) 附件一第 6.16 节中, 收入下列新增行:
 - "Example" 行: <INSDQualifier value>1.1.2.n1</INSDQualifier value>;
 - "Comment"行,增加下列案文: "Symbols including an "n", e.g. "n", "n1" and so on."

- (f) 更新附件七第三段 "Recommendations for potential added or deleted subject matter"第一句: 把"conversion"改为"transformation"。"conversion"一词暗示组件之间有一对一的映射,技术上不正确。
- (g) 更新附件七场景 8 第 24 号: 把 "SITE"一词改为 "REGION"。这项修改是开发方提出的,他们注意到概述 ST. 25 序列导入的功能规范和此例存在冲突。
- 7... 序列表工作队进一步建议,参照对其他产权组织标准提出的修正,把产权组织 ST. 26 附件三和产权组织 ST. 26 附件六附录(两者都是 XML 实例)的内容作为单独文件提供,而标准中则收入这些文件的链接。预计这将让这些例子更易被标准的读者使用。
- 8.. 有关这些修改的进一步信息,请参见附件。

9... 请标准委员会:

- (a) 注意本文件的内容;
- (b) 审议并决定是否批准上文第 4 段至第 6 段中所述以及本文件附件一和二中转录的产权组织 ST. 26 的拟议修订;
- (c) 审议并批准上文第7段中关于 把产权组织 ST. 26 附件三和附件六附录的 内容作为单独文件提供、在标准中提供链 接的建议。

[后接附件(ST. 26 附件一)]

ST.26 - ANNEX I

CONTROLLED VOCABULARY



Revision approved by the Committee on WIPO Standards (CWS) at its sixth session on October 19, 2018 Proposal presented by the SEQL Task Force for consideration and approval at the CWS/7

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SECTION 1: LIST OF NUCLEOTIDES

The nucleotide base codes to be used in sequence listings are presented in Table 1. The symbol "t" will be construed as thymine in DNA and uracil in RNA when it is used with no further description. Where an ambiguity symbol (representing two or more bases in the alternative) is appropriate, the most restrictive symbol should be used. For example, if a base in a given position could be "a or g," then "r" should be used, rather than "n". The symbol "n" will be construed as "a or c or g or t/u" when it is used with no further description.

Table 1: List of nucleotides

Symbol	Nucleotide
а	adenine
С	cytosine
g	guanine
t	thymine in DNA/uracil in RNA (t/u)
m	a or c
r	a or g
W	a or t/u
S	c or g
У	c or t/u
k	g or t/u
V	a or c or g; not t/u
h	a or c or t/u; not g
d	a or g or t/u; not c
b	c or g or t/u; not a
n	a or c or g or t/u; "unknown" or "other"

SECTION 2: LIST OF MODIFIED NUCLEOTIDES

The abbreviations listed in Table 2 are the only permitted values for the mod_base qualifier. Where a specific modified nucleotide is not present in the table below, then the abbreviation "OTHER" must be used as its value. If the abbreviation is "OTHER", then the complete unabbreviated name of the modified base must be provided in a note qualifier. The abbreviations provided in Table 2 must not be used in the sequence itself.

Table 2: List of modified nucleotides

Abbreviation	Modified Nucleotide
ac4c	4-acetylcytidine
chm5u	5-(carboxyhydroxylmethyl)uridine
cm	2'-O-methylcytidine
cmnm5s2u	5-carboxymethylaminomethyl-2-thiouridine
cmnm5u	5-carboxymethylaminomethyluridine
dhu	dihydrouridine
fm	2'-O-methylpseudouridine
gal q	beta-D-galactosylqueuosine
gm	2'-O-methylguanosine
i	inosine
i6a	N6-isopentenyladenosine
m1a	1-methyladenosine
m1f	1-methylpseudouridine
m1g	1-methylguanosine
m1i	1-methylinosine
m22g	2,2-dimethylguanosine
m2a	2-methyladenosine
m2g	2-methylguanosine
m3c	3-methylcytidine
m4c	N4-methylcytosine
m5c	5-methylcytidine
m6a	N6-methyladenosine

Abbreviation	Modified Nucleotide
m7g	7-methylguanosine
mam5u	5-methylaminomethyluridine
mam5s2u	5-methylaminomethyl-2-thiouridine
man q	beta-D-mannosylqueuosine
mcm5s2u	5-methoxycarbonylmethyl-2-thiouridine
mcm5u	5-methoxycarbonylmethyluridine
mo5u	5-methoxyuridine
ms2i6a	2-methylthio-N6-isopentenyladenosine
ms2t6a	N-((9-beta-D-ribofuranosyl-2-methylthiopurine-6-yl)carbamoyl)threonine
mt6a	N-((9-beta-D-ribofuranosylpurine-6-yl)N-methyl-carbamoyl)threonine
mv	uridine-5-oxoacetic acid-methylester
o5u	uridine-5-oxyacetic acid (v)
osyw	wybutoxosine
р	pseudouridine
q	queuosine
s2c	2-thiocytidine
s2t	5-methyl-2-thiouridine
s2u	2-thiouridine
s4u	4-thiouridine
m5u	5-methyluridine
t6a	N-((9-beta-D-ribofuranosylpurine-6-yl)carbamoyl)threonine
tm	2'-O-methyl-5-methyluridine
um	2'-O-methyluridine
yw	wybutosine
х	3-(3-amino-3-carboxypropyl)uridine, (acp3)u
OTHER	(requires note qualifier)

SECTION 3: LIST OF AMINO ACIDS

The amino acid codes to be used in sequence are presented in Table 3. Where an ambiguity symbol (representing two or more amino acids in the alternative) is appropriate, the most restrictive symbol should be used. For example, if an amino acid in a given position could be aspartic acid or asparagine, the symbol "B" should be used, rather than "X". The symbol "X" will be construed as any one of "A", "R", "N", "D", "C", "Q", "E", "G", "H", "I", "L", "K", "M", "F", "P", "O", "S", "U", "T", "W", "Y", or "V", when it is used with no further description.

Table 3: List of amino acids

Symbol	Amino acid
Α	Alanine
R	Arginine
N	Asparagine
D	Aspartic acid (Aspartate)
С	Cysteine
Q	Glutamine
Е	Glutamic acid (Glutamate)
G	Glycine
Н	Histidine
I	Isoleucine
L	Leucine
K	Lysine
М	Methionine
F	Phenylalanine
Р	Proline
0	Pyrrolysine
S	Serine
U	Selenocysteine
Т	Threonine
W	Tryptophan
Υ	Tyrosine
V	Valine
В	Aspartic acid or Asparagine
Z	Glutamine or Glutamic acid
J	Leucine or Isoleucine
Х	A or R or N or D or C or Q or E or G or H or I or L or K or M or F or P or O or S or U or T or W or Y or V; "unknown" or "other"

SECTION 4: LIST OF MODIFIED AMINO ACIDS

Table 4 lists the only permitted abbreviations for a modified amino acid in the mandatory qualifier "NOTE" for feature keys "MOD_RES" or "SITE". The value for the qualifier "NOTE" must be either an abbreviation from this table, where appropriate, or the complete, unabbreviated name of the modified amino acid. The abbreviations (or full names) provided in this table must not be used in the sequence itself.

Table 4: List of modified amino acids

Abbreviation	Modified Amino acid
Aad	2-Aminoadipic acid
bAad	3-Aminoadipic acid
bAla	beta-Alanine, beta-Aminoproprionic acid
Abu	2-Aminobutyric acid
4Abu	4-Aminobutyric acid, piperidinic acid
Acp	6-Aminocaproic acid
Ahe	2-Aminoheptanoic acid
Aib	2-Aminoisobutyric acid
bAib	3-Aminoisobutyric acid
Apm	2-Aminopimelic acid
Dbu	2,4-Diaminobutyric acid
Des	Desmosine
Dpm	2,2'-Diaminopimelic acid
Dpr	2,3-Diaminoproprionic acid
EtGly	N-Ethylglycine
EtAsn	N-Ethylasparagine
Hyl	Hydroxylysine
aHyl	allo-Hydroxylysine
ЗНур	3-Hydroxyproline
4Нур	4-Hydroxyproline
Ide	Isodesmosine
alle	allo-Isoleucine
MeGly	N-Methylglycine, sarcosine
Melle	N-Methylisoleucine
MeLys	6-N-Methyllysine
MeVal	N-Methylvaline
Nva	Norvaline
Nle	Norleucine
Orn	Ornithine

SECTION 5: FEATURE KEYS FOR NUCLEOTIDE SEQUENCES

This section contains the list of allowed feature keys to be used for nucleotide sequences, and lists mandatory and optional qualifiers. The feature keys are listed in alphabetic order. The feature keys can be used for either DNA or RNA unless otherwise indicated under "Molecule scope". Certain Feature Keys may be appropriate for use with artificial sequences in addition to the specified "organism scope".

Feature key names must be used in the XML instance of the sequence listing exactly as they appear following "Feature key" in the descriptions below, except for the feature keys 3'UTR and 5'UTR. See "Comment" in the description for the 3'UTR and 5'UTR feature keys.

5.1.	Feature Key	C_regi on
	Definition	constant region of immunoglobulin light and heavy chains, and T-cell receptor alpha, beta, and gamma chains; includes one or more exons depending on the particular chain
	Optional qualifiers	allele gene gene_synonym map note product pseudo pseudogene standard_name
	Organism scope	eukaryotes
5.2.	Feature Key	CDS
	Definition	coding sequence; sequence of nucleotides that corresponds with the sequence of amino acids in a protein (location includes stop codon); feature may include amino acid conceptual translation
	Optional qualifiers	allele codon_start EC_number exception function gene gene_synonym map note number operon product protein_id pseudo pseudogene ribosomal_slippage standard_name translation transl_except trans_splicing

Comment

codon_start qualifier has valid value of 1 or 2 or 3, indicating the offset at which the first complete codon of a coding feature can be found, relative to the first base of that feature; transl_table defines the genetic code table used if other than the Standard or universal genetic code table; genetic code exceptions outside the range of the specified tables are reported in transl_except qualifier; only one of the qualifiers translation, pseugogene or pseudo are permitted with a CDS feature key; when the translation qualifier is used, the protein_id qualifier is mandatory if the translation product contains four or more specifically defined amino acids

5.3.	Feature Key	centromere
	Definition	region of biological interest identified as a centromere and which has been
	berringeron	experimentally characterized
	Optional qualifiers	note
		standard_name
	Comment	the centromere feature describes the interval of DNA that corresponds to a region where chromatids are held and a kinetochore is formed $ \frac{1}{2} \left(\frac{1}{2} \right) = \frac{1}{2} \left(\frac{1}{2} \right) \left(\frac{1}{2} \right)$
5.4.	Feature Key	D-1 oop
	Definition	displacement loop; a region within mitochondrial DNA in which a short stretch of RNA is paired with one strand of DNA, displacing the original partner DNA strand in this region; also used to describe the displacement of a region of one strand of duplex DNA by a single stranded invader in the reaction catalyzed by RecA protein
	Optional qualifiers	allele
		gene
		gene_synonym map
		note
	Molecule scope	DNA
5.5.	Feature Key	D_segment
	Definition	Diversity segment of immunoglobulin heavy chain, and T-cell receptor beta chain
	berrinteron	prversity segment of riminanogrobatrii neavy charii, and r-cerr receptor beta charii
	Optional qualifiers	allele
		gene gene_synonym
		map
		note
		product pseudo
		pseudogene
		standard_name
	Organism scope	eukaryotes
5.6.	Feature Key	exon
	Definition	region of genome that codes for portion of spliced mRNA, rRNA and tRNA; may contain 5'UTR, all CDSs and 3' UTR
	Optional qualifiers	allele
		EC_number
		function
		gene gene_synonym
		map
		note
		number product
		pseudo
		pseudogene
		standard_name trans_splicing

Definition region of biological interest identified as a gene and for which a name has been assi gned Optional qualifiers al l el e function gene gene_synonym map $not\,e$ operon product pseudo pseudogene phenotype standard_name trans_splicing the gene feature describes the interval of DNA that corresponds to a genetic trait Comment or phenotype; the feature is, by definition, not strictly bound to its positions at the ends; it is meant to represent a region where the gene is located. 5.8. Feature Key Definition intervening DNA; DNA which is eliminated through any of several kinds of recombi nati on Optional qualifiers al l el e function gene gene_synonym map note number standard_name Molecule scope DNA Comment e.g. \blacksquare in the somatic processing of immunoglobulin genes. 5.9. Feature Key intron a segment of DNA that is transcribed, but removed from within the transcript by Definition splicing together the sequences (exons) on either side of it Optional qualifiers al l el e function gene gene_synonym map note number pseudo pseudogene standard_name trans_splicing

5.7.

Feature Key

gene

5.10. Feature Key J_segment Definition joining segment of immunoglobulin light and heavy chains, and T-cell receptor alpha, beta, and gamma chains Optional qualifiers allele gene gene_synonym map note product pseudo pseudogene standard_name Organism scope eukaryotes 5.11. Feature Key mat_peptide Definition mature peptide or protein coding sequence; coding sequence for the mature or final $pepti\,de\,\,or\,\,protein\,\,product\,\,following\,\,post\text{-translational}\,\,\,modification;\,\,the\,\,location$ does not include the stop codon (unlike the corresponding CDS) Optional qualifiers allele EC_number function gene gene_synonym map note product pseudo pseudogene standard_name 5.12. Feature Key mi sc_bi ndi ng Definition site in nucleic acid which covalently or non-covalently binds another moiety that cannot be described by any other binding key (primer_bind or protein_bind) Mandatory qualifiers $bound_moiety$ Optional qualifiers allele function gene gene_synonym map note

note that the regulatory feature key and regulatory_class qualifier with the value "ribosome_binding_site" must be used for describing ribosome binding sites

Comment

5.13. Feature Key misc_difference

Definition featured sequence differs from the presented sequence at this location and cannot

be described by any other ${\tt Difference\ key\ (variation,\ or\ modified_base)}$

Optional qualifiers allele

clone compare gene gene_synonym map

note
phenotype
replace
standard name

Comment the misc_difference feature key must be used to describe variability introduced

artificially, e.g. by genetic manipulation or by chemical synthesis; use the replace qualifier to annotate a deletion, insertion, or substitution. The variation feature key must be used to describe naturally occurring genetic variability.

5.14. Feature Key misc_feature

Definition region of biological interest which cannot be described by any other feature key; a

new or rare feature

Optional qualifiers allele

function gene gene synon

gene_synonym map

map note number phenotype product pseudo pseudogene standard_name

Comment this key should not be used when the need is merely to mark a region in order to

comment on it or to use it in another feature's location

5.15. Feature Key misc_recomb

Definition site of any generalized, site-specific or replicative recombination event where

there is a breakage and reunion of $\operatorname{dupl}\operatorname{ex}\operatorname{DNA}$ that cannot be described by other

recombination keys or qualifiers of source key (proviral)

Optional qualifiers allele

gene

gene_synonym

map

note

recombination_class

standard_name

Mol ecul e scope DNA

5.16. Feature Key misc_RNA

Definition any transcript or RNA product that cannot be defined by other RNA keys

 $(prim_transcript,\ precursor_RNA,\ mRNA,\ 5'UTR,\ 3'UTR,\ exon,\ CDS,\ sig_peptide,\ transit_peptide,\ mat_peptide,\ intron,\ pol\ yA_site,\ ncRNA,\ rRNA\ and\ tRNA)$

Optional qualifiers allele

function gene

gene_synonym

note
operon
product
pseudo
pseudogene
standard_name
trans_splicing

5.17. Feature Key misc_structure

Definition any secondary or tertiary nucleotide structure or conformation that cannot be

described by other Structure keys (stem_loop and D-loop)

Optional qualifiers allele

function gene gene_synonym map note

standard_name

5.18. Feature Key mobile_element

Optional qualifiers

Definition region of genome containing mobile elements

Mandatory qualifiers mobile_element_type

allele function gene gene_synonym map

note
rpt_family
rpt_type
standard name

5.19. Feature Key modified_base

Definition the indicated nucleotide is a modified nucleotide and should be substituted for by

the indicated molecule (given in the mod_base qualifier value)

Mandatory qualifiers mod_base

Optional qualifiers allele

frequency gene gene_synonym map note

Comment value for the mandatory mod_base qualifier is limited to the restricted vocabulary

for modified base abbreviations in Section 2 of this Annex.

5.20. Feature Key mRNA

> Definition messenger RNA; includes 5' untranslated region (5'UTR), coding sequences (CDS,

exon) and 3' untranslated region (3'UTR)

Optional qualifiers al l el e

function gene

gene_synonym map $not\,e$ operon product pseudo

pseudogene

standard_name trans_splicing

5.21. Feature Key ncRNA

> a non-protein-coding gene, other than ribosomal RNA and transfer RNA, the Definition

functional molecule of which is the RNA transcript

Mandatory qualifiers ncRNA_class

Optional qualifiers allele

function gene

gene_synonym

map note operon product pseudo pseudogene standard_name trans_splicing

the ncRNA feature must not be used for ribosomal and transfer RNA annotation, for Comment

which the rRNA and tRNA feature keys must be used, respectively

5.22. Feature Key N_regi on

> Definition extra nucleotides inserted between rearranged immunoglobulin segments

Optional qualifiers al l el e

gene

gene_synonym

map note product pseudo pseudogene standard_name

Organism scope eukaryotes 5.23. Feature Key operon

Definition region containing polycistronic transcript including a cluster of genes that are

under the control of the same regulatory sequences/promoter and in the same $\,$

biological pathway

Mandatory qualifiers operon

Optional qualifiers allele

function
map
note
phenotype
pseudo
pseudogene
standard_name

5.24. Feature Key ori T

Definition origin of transfer; region of a DNA molecule where transfer is initiated during the

process of conjugation or mobilization

Optional qualifiers allele

bound_moi ety
di recti on
gene
gene_synonym
map
note
rpt_family
rpt_type
rpt_unit_range
rpt_unit_seq
standard_name

Mol ecul e Scope DNA

Comment rep_origin must be used to describe origins of replication; direction qualifier has

permitted values left, right, and both, however only left and right are valid when used in conjunction with the oriT feature; origins of transfer can be present in

the chromosome; plasmids can contain multiple origins of transfer

5.25. Feature Key polyA_site

Definition site on an RNA transcript to which will be added adenine residues by post-

 $transcriptional\ polyadenylation$

Optional qualifiers allele

gene gene_synonym

map

Organism scope eukaryotes and eukaryotic viruses

5.26. Feature Key precursor_RNA

Definition any RNA species that is not yet the mature RNA product; may include ncRNA, rRNA,

 $tRNA, \ 5' \ untranslated \ region \ (5' \ UTR), \ coding \ sequences \ (CDS, \ exon), \ intervening$

sequences (intron) and 3' untranslated region (3'UTR)

Optional qualifiers allele

function gene

gene_synonym

map
note
operon
product
standard_name
trans_splicing

Comment used for RNA which may be the result of post-transcriptional processing; if the RNA

in question is known not to have been processed, use the $prim_transcript$ key

5.27. Feature Key prim_transcript

Definition primary (initial, unprocessed) transcript; may include ncRNA, rRNA, tRNA, 5'

untranslated region (5'UTR), coding sequences (CDS, exon), intervening sequences

(intron) and 3' untranslated region (3'UTR)

Optional qualifiers allele

function
gene
gene_synonym
map
note
operon

standard_name

5.28. Feature Key primer_bind

Definition non-covalent primer binding site for initiation of replication, transcription, or

reverse transcription; includes ${\tt site}(s)$ for ${\tt synthetic}$ e.g. \blacksquare PCR primer elements

Optional qualifiers allele

gene gene_synonym map note

standard_name

Comment used to annotate the site on a given sequence to which a primer molecule binds -

not intended to represent the sequence of the primer molecule itself; since PCR reactions most often involve pairs of primers, a single primer_bind key may use the order(location, location) operator with two locations, or a pair of primer_bind keys

may be used

5.29. Feature Key propeptide

Definition propeptide coding sequence; coding sequence for the domain of a proprotein that is

cleaved to form the mature protein product.

Optional qualifiers allele

function gene

gene_synonym

map note product pseudo pseudogene standard_name

5.30. Feature Key protein_bind

Definition non-covalent protein binding site on nucleic acid

Mandatory qualifiers bound_moiety

Optional qualifiers allele

function gene

gene_synonym

map note operon standard_name

Comment note that the regulatory feature key and regulatory_class qualifier with the value

 $"ri\,bosome_bi\,ndi\,ng_si\,te"\ must\ be\ used\ to\ describe\ ri\,bosome\ bi\,ndi\,ng\ si\,tes$

5.31. Feature Key regulatory

Definition any region of a sequence that functions in the regulation of transcription,

translation, replication or chromatin structure;

Mandatory qualifiers regulatory_class

Optional qualifiers allele

bound_moiety function gene

gene_synonym

map
note
operon
phenotype
pseudo
pseudogene
standard_name

5.32. Feature Key repeat_regi on Definition region of genome containing repeating units Optional qualifiers allele functiongene gene_synonym map note $rpt_fami\,l\,y$ rpt_type rpt_uni t_range rpt_uni t_seq satellite standard_name 5.33. Feature Key rep_ori gi n Definition origin of replication; starting site for duplication of nucleic acid to give two identical copies Optional Qualifiers al l el e di recti on function gene gene_synonym <u>map</u> <u>not e</u> $standard_name$ direction qualifier has valid values: left, right, or both Comment 5.34. Feature Key rRNA $mature\ \verb|ri| bosomal|\ RNA;\ RNA\ component\ of\ the\ \verb|ri| bonucleoprotein|\ particle\ (\verb|ri| bosome|)$ Definition which assembles amino acids into proteins Optional qualifiers al l el e function gene gene_synonym map note operon product pseudo pseudogene

rRNA sizes should be annotated with the product qualifier

standard_name

Comment

5.35. Feature Key S_region

Definition switch region of immunoglobulin heavy chains; involved in the rearrangement of

heavy chain DNA leading to the expression of a different immunoglobulin class from

the same B-cell

Optional qualifiers allele

gene

 $gene_synonym$

map note product pseudo pseudogene standard_name

Organism scope eukaryotes

5.36. Feature Key sig_peptide

Definition signal peptide coding sequence; coding sequence for an N-terminal domain of a

secreted protein; this domain is involved in attaching mascent polypeptide to the

membrane leader sequence

Optional qualifiers allele

function gene

gene_synonym map note

note product pseudo pseudogene standard_name 5.37. Feature Key source Definition identifies the source of the sequence; this key is mandatory; every sequence will have a single source key spanning the entire sequence Mandatory qualifiers organi sm mol_type Optional qualifiers $cel\,l_l\,i\,ne$ $cel\,l_type$ chromosome cl one cl one_l i b $col\,l\,ect\,ed_by$ collection_date cul ti var dev_stage ecotype environmental_sample germl i ne hapl ogroup $hapl\,otype$ host i dent i fi ed_by i sol ate i sol ati on_source l ab_host lat_lon macronuclear map mating_type note organel l e PCR_pri mers pl asmi d pop_vari ant provi ral rearranged segment serotype serovar sex strain sub_cl one ${\tt sub_species}$ sub_strain tissue_lib tissue_type vari ety Molecule scope any 5.38. Feature Key stem_loop Definition hairpin; a double-helical region formed by base-pairing between adjacent (inverted) complementary sequences in a single strand of RNA or DNA al l el e Optional qualifiers functiongene gene_synonym map

> note operon standard_name

5.39.	Feature Key	STS
	Definition	sequence tagged site; short, single-copy DNA sequence that characterizes a mapping landmark on the genome and can be detected by PCR; a region of the genome can be
		mapped by determining the order of a series of STSs
	Optional qualifiers	allele
	•	gene
		gene_synonym
		тар
		note
		standard_name
	Molecule scope	DNA
	Comment	STS location to include primer(s) in primer_bind key or primers
5.40.	Feature Key	telomere
	D - C11 +-1	
	Definition	region of biological interest identified as a telomere and which has been experimentally characterized
	Optional qualifiers	note
		rpt_type
		rpt_uni t_range
		rpt_uni t_seq
		standard_name
	Comment	the telomere feature describes the interval of DNA that corresponds to a specific
		structure at the end of the linear eukaryotic chromosome which is required for the
		integrity and maintenance of the end; this region is unique compared to the rest of the chromosome and represents the physical end of the chromosome
F 41	F	. DV4
5.41.	Feature Key	tmRNA
	Definition	transfer messenger RNA; tmRNA acts as a tRNA first, and then as an mRNA that encodes a peptide tag; the ribosome translates this mRNA region of tmRNA and
		attaches the encoded peptide tag to the C-terminus of the unfinished protein; this
		attached tag targets the protein for destruction or proteolysis
	Optional qualifiers	al l el e
		function
		gene
		gene_synonym
		map
		note
		product
		pseudo pseudogene
		standard_name
		tag_pepti de

5.42. Feature Key transit_peptide

Definition transit peptide coding sequence; coding sequence for an N-terminal domain of a

 $nuclear \hbox{-} encoded \ organellar \ protein; \ this \ domain \ is \ involved \ in \ post-translational$

import of the protein into the organelle

Optional qualifiers allele

function gene

gene_synonym

map note product pseudo pseudogene standard_name

5.43. Feature Key tRNA

Definition mature transfer RNA, a small RNA molecule (75-85 bases long) that mediates the

 $translation \ of \ a \ nucleic \ acid \ sequence \ into \ an \ amino \ acid \ sequence$

Optional qualifiers allele

anticodon function gene

gene_synonym

map
note
operon
product
pseudo
pseudogene
standard_name
trans_splicing

5.44. Feature Key unsure Definition a small region of sequenced bases, generally 10 or fewer in its length, which could not be confidently identified. Such a region \min ght contain called bases (a, t, g, or c), or a mixture of called-bases and uncalled-bases ('n'). Optional qualifiers al l el e compare gene gene_synonym map note repl ace Comment use the replace qualifier to annotate a deletion, insertion, or substitution. 5.45. Feature Key V_regi on Definition variable region of immunoglobulin light and heavy chains, and T-cell receptor alpha, beta, and gamma chains; codes for the variable amino terminal portion; can be composed of V_segments, D_segments, N_regions, and J_segments Optional qualifiers al l el e gene gene_synonym map note product pseudo pseudogene standard_name Organism scope eukaryotes 5.46. Feature Key V_segment Definition variable segment of immunoglobulin light and heavy chains, and T-cell receptor alpha, beta, and gamma chains; codes for most of the variable region ($V_{\rm region}$) and the last few amino acids of the leader peptide al l el e Optional qualifiers gene gene_synonym map note product pseudo pseudogene standard_name Organism scope eukaryotes

Definition a related strain contains stable mutations from the same gene (e.g. RFLPs, polymorphisms, etc.) which differ from the presented sequence at this location (and possibly others) Optional qualifiers al l el e compare frequency gene gene_synonym map note phenotype product repl ace standard name Comment used to describe alleles, RFLP's, and other naturally occurring mutations and polymorphisms; use the replace qualifier to annotate a deletion, insertion, or substitution; variability arising as a result of genetic manipulation (e.g. 2 site $\ directed\ mutagenesis)\ must\ be\ described\ with\ the\ misc_difference\ feature$ 5.48. Feature Key 3' UTR Definition 1) region at the 3' end of a mature transcript (following the stop codon) that is not translated into a protein; 2) region at the 3° end of an RNA virus (following the last stop codon) that is not translated into a protein; Optional qualifiers al l el e function gene gene_synonym map note standard_name trans_splicing Comment The apostrophe character has special meaning in XML, and must be substituted with "'" in the value of an element. Thus "3'UTR" must be represented as $"3\' \verb|UTR"| in the XML file, i.e., < INSDFeature_key > 3\' \verb|UTR < /INSDFeature_key > . |$ 5' UTR 5.49. Feature Key Definition 1) region at the 5' end of a mature transcript (preceding the initiation codon) that is not translated into a protein; 2) region at the 5' end of an RNA virus (preceding the first initiation codon) that is not translated into a protein; Optional qualifiers allele function gene gene_synonym map note standard_name trans_splicing The apostrophe character has special meaning in XML, and must be substituted with Comment "'" in the value of an element. Thus "5'UTR" must be represented as

5.47. Feature Key

vari ati on

SECTION 6: QUALIFIERS FOR NUCLEOTIDE SEQUENCES

This section contains the list of qualifiers to be used for features in nucleotide sequences. The qualifiers are listed in alphabetic order.

Where a Value format of "none" is indicated in the description of a qualifier (e.g. germline), the INSDQualifier_value element must not be used.

PLEASE NOTE: Any qualifier value provided for a qualifier with a "free text" value format may require translation for National/Regional procedures.

6. 1.	Qual i fi er	allele
	Definition	name of the allele for the given gene
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>adh1-1</insdqualifier_value>
	Comment	all gene-related features (exon, CDS etc) for a given gene should share the same allele qualifier value; the allele qualifier value must, by definition, be different from the gene qualifier value; when used with the variation feature key, the allele qualifier value should be that of the variant.
6. 2.	Qualifier	anticodon
	Definition	location of the anticodon of tRNA and the amino acid for which it codes
	Value format	(pos: <location>, aa: <amino_acid>, seq: <text>) where <location> is the position of the anticodon and <amino_acid> is the three letter abbreviation for the amino acid encoded and <text> is the sequence of the anticodon</text></amino_acid></location></text></amino_acid></location>
	Exampl e	<pre><insdqualifier_value>(pos: 3436, aa: Phe, seq: aaa) </insdqualifier_value> <insdqualifier_value>(pos: join(5, 495496), aa: Leu, seq: taa) </insdqualifier_value> <insdqualifier_value>(pos: complement(41564158), aa: Glu, seq: ttg) </insdqualifier_value> ue></pre>
6. 3.	Qualifici on	hound malata
6. 3.	Qualifier Definition	bound_moiety
	berrin tron	name of the molecule/complex that may bind to the given feature
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>GAL4</insdqualifier_value>
	Comment	A single bound_moiety qualifier is permitted on the "misc_binding", "oriT" and "protein_bind" features.
6. 4.	Qualifier	cell_line
	Definition	cell line from which the sequence was obtained
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>MCF7</insdqualifier_value>

6. 5.	Qual i fi er	cell_type
	Definition	cell type from which the sequence was obtained
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>leukocyte</insdqualifier_value>
6. 6.	Qualifier	chromosome
	Definition	chromosome (e.g. Chromosome number) from which the sequence was obtained
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqual er_val="" fi="" i="" ue="">1</insdqual> <insdqual er_val="" fi="" i="" ue="">X</insdqual>
6. 7.	Qualifier	clone
	Definition	clone from which the sequence was obtained
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>l ambda-hIL7.3</insdqualifier_value>
	Comment	a source feature must not contain more than one clone qualifier; where the sequence was obtained from multiple clones it may be further described in the feature table using the feature key misc_feature and a note qualifier to specify the multiple clones.
6. 8.	Qual i fi er	cl one_l i b
	Definition	clone library from which the sequence was obtained
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>lambda-hIL7</insdqualifier_value>
6. 9.	Qualifier	codon_start
	Definition	indicates the offset at which the first complete codon of a coding feature can be found, relative to the first base of that feature.
	Value format	1 or 2 or 3
	Exampl e	<i er_val="" fi="" i="" nsdqual="" ue="">2</i>
6. 10.	Qualifier	collected_by
	Definition	name of persons or institute who collected the specimen
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>Dan Janzen</insdqualifier_value>

6. 11.	Qual i fi er	collection_date
	Definition	date that the specimen was collected.
	Value format	YYYY-MM-DD, YYYY-MM or YYYY
	Exampl e	<insdqualifier_value>1952-10-21</insdqualifier_value> <insdqualifier_value>1952-10</insdqualifier_value> <insdqualifier_value>1952</insdqualifier_value>
	Comment	'YYYY' is a four-digit value representing the year. 'MM' is a two-digit value representing the month. 'DD' is a two-digit value representing the day of the month.
6. 12.	Qualifier	compare
	Definition	Reference details of an existing public INSD entry to which a comparison is made
	Value format	[accession-number.sequence-version]
	Exampl e	<i nsdqualifier_value="">AJ634337. 1</i>
	Comment	This qualifier may be used on the following features: misc_difference, unsure, and variation. Multiple compare qualifiers with different contents are allowed within a single feature. This qualifier is not intended for large-scale annotation of variations, such as SNPs.
0.10	Overld Ct and	
6. 13.	Qual i fi er	cul ti var
	Definition	cultivar (cultivated variety) of plant from which sequence was obtained
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<pre><insdqualifier_value>Nipponbare</insdqualifier_value> <insdqualifier_value>Tenuifolius</insdqualifier_value> <insdqualifier_value>Candy Cane</insdqualifier_value> <insdqualifier_value>IR36</insdqualifier_value></pre>
	Comment	'cultivar' is applied solely to products of artificial selection; use the variety qualifier for natural, named plant and fungal varieties.
6. 14.	Qual i fi er	dev_stage
	Definition	if the sequence was obtained from an organism in a specific developmental stage, it is specified with this qualifier
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<pre><insdqualifier_value>fourth instar larva</insdqualifier_value></pre>

6. 15.	Qual i fi er	direction
	Definition	direction of DNA replication
	Value format	left, right, or both where left indicates toward the 5' end of the sequence (as presented) and right indicates toward the 3' end
	Exampl e	<insdqualifier_value>left</insdqualifier_value>
	Comment	The values left, right, and both are permitted when the direction qualifier is used to annotate a rep_origin feature key. However, only left and right values are permitted when the direction qualifier is used to annotate an oriT feature key.
6. 16.	Qual i fi er	EC_number
	Definition	Enzyme Commission number for enzyme product of sequence
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<pre><insdqualifier_value>1.1.2.4</insdqualifier_value> <insdqualifier_value>1.1.2</insdqualifier_value> <insdqualifier_value>1.1.2.n</insdqualifier_value> <insdqualifier_value>1.1.2.n</insdqualifier_value></pre>
	Comment	valid values for EC numbers are defined in the list prepared by the Nomenclature Committee of the International Union of Biochemistry and Molecular Biology (NC-IUBMB) (published in Enzyme Nomenclature 1992, Academic Press, San Diego, or a more recent revision thereof). The format represents a string of four numbers separated by full stops; up to three numbers starting from the end of the string may be replaced by dash "-" to indicate uncertain assignment. Symbols including an symbol "n", e.g., "n", "n1" and so on, may be used in the last position instead of a number where the EC number is awaiting assignment. Please note that such incomplete EC numbers are not approved by NC-IUBMB.
6. 17.	Qual i fi er	ecotype
	Definition	a population within a given species displaying genetically based, phenotypic traits that reflect adaptation to a local habitat $ \frac{1}{2} \int_{\mathbb{R}^{n}} \frac{1}{2} \left(\frac{1}{2} \int_{\mathbb{R}^{n}} \frac{1}{2}$
	Value Format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>Columbia</insdqualifier_value>
	Comment	an example of such a population is one that has adapted hairier than normal leaves as a response to an especially sunny habitat. 'Ecotype' is often applied to standard genetic stocks of Arabidopsis thaliana, but it can be applied to any sessile organism.

6. 18.	Qualifier	environmental_sample
0. 10.	•	
	Definition	identifies sequences derived by direct molecular isolation from a bulk environmental DNA sample (by PCR with or without subsequent cloning of the product, DGCE, or other anonymous methods) with no reliable identification of the source organism. Environmental samples include clinical samples, gut contents, and other sequences from anonymous organisms that may be associated with a particular host. They do not include endosymbionts that can be reliably recovered from a particular host, organisms from a readily identifiable but uncultured field sample (e.g. many cyanobacteria), or phytoplasmas that can be reliably recovered from diseased plants (even though these cannot be grown in axenic culture)
	Value format	none
	Comment	used only with the source feature key; source feature keys containing the environmental_sample qualifier should also contain the isolation_source qualifier; a source feature including the environmental_sample qualifier must not include the strain qualifier.
6. 19.	Qualifier	excepti on
	Definition	indicates that the coding region cannot be translated using standard biological rules
	Value format	One of the following controlled vocabulary phrases: RNA editing rearrangement required for product annotated by transcript or proteomic data
	Exampl e	<pre><insdqualifier_value>RNA editing</insdqualifier_value> <insdqualifier_value>rearrangement required for product</insdqualifier_value></pre>
	Comment	only to be used to describe biological mechanisms such as RNA editing; protein translation of a CDS with an exception qualifier will be different from the corresponding conceptual translation; must not be used where transl_except qualifier would be adequate, e.g. in case of stop codon completion use.
6. 20.	Qual i fi er	frequency
	Definition	frequency of the occurrence of a feature
	Value format	free text representing the proportion of a population carrying the feature expressed as a fraction (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>23/108</insdqualifier_value> <insdqualifier_value>1 in 12</insdqualifier_value> <insdqualifier_value>0.85</insdqualifier_value>
6. 21.	Qualifier	function
	Definition	function attributed to a sequence
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<pre><insdqualifier_value>essential for recognition of cofactor </insdqualifier_value></pre>
	Comment	The function qualifier is used when the gene name and/or product name do not convey the function attributable to a sequence.

6. 22.	Qualifier	gene
	Definition	symbol of the gene corresponding to a sequence region
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<i er_val="" fi="" i="" nsdqual="" ue="">i l vE</i>
	Comment	Use gene qualifier to provide the gene symbol; use standard_name qualifier to provide the full gene name.
6. 23.	Qualifier	gene_synonym
	Definition	synonymous, replaced, obsolete or former gene symbol
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<pre><insdqualifier_value>Hox-3.3</insdqualifier_value> in a feature where the gene qualifier value is Hoxc6</pre>
	Comment	used where it is helpful to indicate a gene symbol synonym; when the gene_synonym qualifier is used, a primary gene symbol must always be indicated in a gene qualifier
6. 24.	Qualifier	germl i ne
	Definition	the sequence presented has not undergone somatic rearrangement as part of an adaptive immune response; it is the unrearranged sequence that was inherited from the parental germline
	Value format	none
	Comment	germline qualifier must not be used to indicate that the source of the sequence is a gamete or germ cell; germline and rearranged qualifiers must not be used in the same source feature; germline and rearranged qualifiers must only be used for molecules that can undergo somatic rearrangements as part of an adaptive immune response; these are the T-cell receptor (TCR) and immunoglobulin loci in the jawed vertebrates, and the unrelated variable lymphocyte receptor (VLR) locus in the jawless fish (lampreys and hagfish); germline and rearranged qualifiers should not be used outside of the Craniata (taxid=89593)
6. 25.	Qual i fi er	hapl ogroup
	Definition	name for a group of similar haplotypes that share some sequence variation. Haplogroups are often used to track migration of population groups.
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>H*</insdqualifier_value>
6. 26.	Qual i fi er	haplotype
	Definition	name for a specific set of alleles that are linked together on the same physical chromosome. In the absence of recombination, each haplotype is inherited as a unit, and may be used to track gene flow in populations.
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>Dw3 B5 Cw1 A1</insdqualifier_value>

6. 27.	Qual i fi er	host
	Definition	natural (as opposed to laboratory) host to the organism from which sequenced molecule was obtained
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<pre><insdqualifier_value>Homo sapiens</insdqualifier_value> <insdqualifier_value>Homo sapiens 12 year old girl</insdqualifier_value> <insdqualifier_value>Rhizobium NGR234</insdqualifier_value></pre>
6. 28.	Qual i fi er	i dentifi ed_by
	Definition	name of the expert who identified the specimen taxonomically
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>John Burns</insdqualifier_value>
6. 29.	Qual i fi er	i sol ate
	Definition	individual isolate from which the sequence was obtained
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<pre><insdqualifier_value>Patient #152</insdqualifier_value> <insdqualifier_value>DGGE band PSBAC-13</insdqualifier_value></pre>
6. 30.	Qual i fi er	i sol ati on_source
	Definition	describes the physical, environmental and/or local geographical source of the biological sample from which the sequence was derived
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl es	<pre><insdqualifier_value>rumen isolates from standard Pelleted ration-fed steer #67</insdqualifier_value> <insdqualifier_value>permanent Antarctic sea ice</insdqualifier_value> <insdqualifier_value>denitrifying activated sludge from carbon_limited continuous reactor</insdqualifier_value></pre>
	Comment	used only with the source feature key; source feature keys containing an environmental_sample qualifier should also contain an isolation_source qualifier
6. 31.	Qual i fi er	l ab_host
	Definition	scientific name of the laboratory host used to propagate the source organism from which the sequenced molecule was obtained
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<pre><insdqualifier_value>Gallus gallus</insdqualifier_value> <insdqualifier_value>Gallus gallus embryo</insdqualifier_value> <insdqualifier_value>Escherichia coli strain DH5 alpha</insdqualifier_value> <insdqualifier_value>Homo sapiens HeLa cells</insdqualifier_value></pre>
	Comment	the full binomial scientific name of the host organism should be used when known; extra conditional information relating to the host may also be included

6. 32.	Qual i fi er	lat_lon
	Definition	granuplical accordington of the leasting whom the arctimen was called a
	Del i i i i i i i i i i i i i i i i i i i	geographical coordinates of the location where the specimen was collected
	Value format	free text - degrees latitude and longitude in format "d[d.dddd] N S d[dd.dddd] W E" (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	$<$ INSDQualifier_value>47.94 N 28.12 W $<$ INSDQualifier_value>45.0123 S 4.1234 E
6. 33.	Qual i fi er	macronuclear
	Definition	if the sequence shown is DNA and from an organism which undergoes chromosomal differentiation between macronuclear and micronuclear stages, this qualifier is used to denote that the sequence is from macronuclear DNA
	Value format	none
6. 34.	Qualifier	тар
	Definition	genomic map position of feature
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>8q12-q13</insdqualifier_value>
6. 35.	Qualifier	mating_type
	Definition	mating type of the organism from which the sequence was obtained; mating type is used for prokaryotes, and for eukaryotes that undergo meiosis without sexually dimorphic gametes
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Examples	<pre><insdqualifier_value>MAT-1</insdqualifier_value> <insdqualifier_value>plus</insdqualifier_value> <insdqualifier_value>-</insdqualifier_value> <insdqualifier_value>odd</insdqualifier_value> <insdqualifier_value>even</insdqualifier_value>"</pre>
	Comment	mating_type qualifier values male and female are valid in the prokaryotes, but not in the eukaryotes; for more information, see the entry for the sex qualifier.

6. 36. Qualifier mobile_element_type Definition type and name or identifier of the mobile element which is described by the parent feature Value format <mobile_element_type>[:<mobile_element_name>] where <mobile_element_type> is one of the following: transposon retrotransposon integron insertion sequence non-LTR retrotransposon SINE MI TE LINE other Exampl e $<\!I\,NSDQual\,i\,fi\,er_val\,ue\!>\!t\,ransposon:\,Tnp9<\!/I\,NSDQual\,i\,fi\,er_val\,ue\!>$ Comment $mobile_element_type~is~permitted~on~mobile_element~feature~key~only.~Mobile~element$ should be used to represent both elements which are currently mobile, and those which were mobile in the past. Value "other" for <mobile_element_type> requires a <mobile_element_name> 6.37. Qualifier mod base Definition abbreviation for a modified nucleotide base modified base abbreviation chosen from this Annex Section 2 Value format <INSDQualifier_value>m5c</INSDQualifier_value> Example <INSDQualifier_value>OTHER</INSDQualifier_value> Comment specific modified nucleotides not found in Section 2 of this Annex are annotated by entering OTHER as the value for the mod_base qualifier and including a note qualifier with the full name of the modified base as its value 6. 38. Qual i fi er mol_type Definition molecule type of sequence Value format One chosen from the following: genomic DNA genomic RNA mRNA t RNA rRNA other RNA other DNA transcribed RNA viral cRNA unassigned DNA unassigned RNA <INSDQualifier_value>genomic DNA</INSDQualifier_value> Exampl e <INSDQualifier_value>other RNA</INSDQualifier_value> Comment mol_type qualifier is mandatory on the source feature key; the value "genomic DNA" does not imply that the molecule is nuclear (e.g. \blacksquare organelle and plasmid DNA must be described using "genomic DNA"); ribosomal RNA genes must be described using "genomic DNA"; "rRNA" must only be used if the ribosomal RNA molecule itself has been sequenced; values "other RNA" and "other DNA" must be applied to synthetic molecules, values "unassigned DNA", "unassigned RNA" must be applied where in vivo molecule is unknown.

6. 39. Qual i fi er ncRNA class Definition a structured description of the classification of the non-coding RNA described by the ncRNA parent key Value format TYPE where TYPE is one of the following controlled vocabulary terms or phrases: antisense_RNA $autocatal\,yti\,cal\,l\,y_spl\,i\,ced_i\,ntron$ ri bozyme hammerhead_ri bozyme l ncRNA RNase_P_RNA RNase_MRP_RNA telomerase_RNA gui de_RNA segRNA rasi RNA SCRNA scaRNA si RNA pre_mi RNA mi RNA pi RNA snoRNA snRNA SRP_RNA vault_RNA Y_RNA other <INSDQualifier_value>autocatalytically_spliced_intron </INSDQualifier_value> Example <INSDQualifier_value>siRNA</INSDQualifier_value> <INSDQualifier_value>scRNA</INSDQualifier_value> $<\!I\,NSDQual\,i\,fi\,er_val\,ue\!>\!other<\!/I\,NSDQual\,i\,fi\,er_val\,ue\!>$ specific ncRNA types not yet in the ncRNA_class controlled vocabulary must be Comment annotated by entering "other" as the ncRNA_class qualifier value, and providing a brief explanation of novel $ncRNA_class$ in a note qualifier 6. 40. Qual i fi er note Definition any comment or additional information Value format free text (NOTE: this value may require translation for National/Regional procedures) <INSDQualifier_value>A comment about the feature</INSDQualifier_value> Exampl e Qual i fi er number Definition a number to indicate the order of genetic elements (e.g. exons or introns) in the 5' to 3' direction Value format free text (with no whitespace characters) (NOTE: this value may require translation for National/Regional procedures) Exampl e <INSDQualifier_value>4</INSDQualifier_value> $<\!I\,NSDQual\,i\,fi\,er_val\,ue\!>\!6B\!<\!/I\,NSDQual\,i\,fi\,er_val\,ue\!>$ text limited to integers, letters or combination of integers and/or letters Comment represented as a data value that contains no whitespace characters; any additional terms should be included in a standard_name qualifier. Example: a number qualifier with a value of 2A and a standard_name qualifier with a value of "long"

6. 42.	Qual i fi er	operon
	Definition	name of the group of contiguous genes transcribed into a single transcript to which that feature belongs ${\sf S}$
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>lac</insdqualifier_value>
6. 43.	Qualifier	organel l e
	Definition	type of membrane-bound intracellular structure from which the sequence was obtained
	Value format	One of the following controlled vocabulary terms and phrases: chromatophore hydrogenosome mitochondri on nucleomorph plastid mitochondri on: kinetoplast plastid: chloroplast plastid: api coplast plastid: chromoplast plastid: chromoplast
		pl astid: l eucopl ast pl astid: propl astid
	Exampl es	<pre><insdqualifier_value>chromatophore</insdqualifier_value> <insdqualifier_value>hydrogenosome</insdqualifier_value> <insdqualifier_value>mitochondrion</insdqualifier_value> <insdqualifier_value>nucleomorph</insdqualifier_value> <insdqualifier_value>plastid</insdqualifier_value> <insdqualifier_value>mitochondrion: kinetoplast</insdqualifier_value> <insdqualifier_value>plastid: chloroplast</insdqualifier_value> <insdqualifier_value>plastid: apicoplast</insdqualifier_value> <insdqualifier_value>plastid: chromoplast</insdqualifier_value> <insdqualifier_value>plastid: chromoplast</insdqualifier_value> <insdqualifier_value>plastid: cyanelle</insdqualifier_value> <insdqualifier_value>plastid: proplastid: leucoplast</insdqualifier_value> <insdqualifier_value>plastid: proplastid</insdqualifier_value></pre> <pre> <insdqualifier_value>plastid: proplastid</insdqualifier_value></pre> <pre> <insdqualifier_value>plastid: proplastid</insdqualifier_value></pre>
6. 44.	Qualifier	organism
	Definition	scientific name of the organism that provided the sequenced genetic material, if known, or the available taxonomic information if the organism is unclassified; or an indication that the sequence is a synthetic construct
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<pre><insdqualifier_value>Homo sapiens</insdqualifier_value></pre>

6. 45.	Qual i fi er	PCR_pri mers
	Definition	PCR primers that were used to amplify the sequence. A single PCR_primers qualifier should contain all the primers used for a single PCR reaction. If multiple forward or reverse primers are present in a single PCR reaction, multiple sets of
		fwd_name/fwd_seq or rev_name/rev_seq values will be present
	Value format	[fwd_name: XXX1,]fwd_seq: xxxxx1, [fwd_name: XXX2,]fwd_seq: xxxxx2, [rev_name: YYY1,]rev_seq: yyyyy1, [rev_name: YYY2,]rev_seq: yyyyy2
	Exampl e	<pre><insdqualifier_value>fwd_name: C01P1, fwd_seq: ttgattttttggtcayccwgaagt,rev_name: C01R4, rev_seq: ccwvytardcctarraartgttg</insdqualifier_value> <insdqualifier_value>fwd_name: hoge1, fwd_seq: cgkgtgtatcttact, rev_name: hoge2, rev_seq: cg<i>gtgtatcttact</insdqualifier_value> <insdqualifier_value>fwd_name: C01P1, fwd_seq: ttgattttttggtcayccwgaagt, fwd_name: C01P2, fwd_seq: gatacacaggtcayccwgaagt, rev_name: C01R4, rev_seq: ccwvytardcctarraartgttg</insdqualifier_value></pre>
	Comment	fwd_seq and rev_seq are both mandatory; fwd_name and rev_name are both optional. Both sequences must be presented in 5'>3' order. The sequences must be given in the symbols from Section 1 of this Annex, except for the modified bases, which must be enclosed within angle brackets < >. In XML, the angle brackets < and > must be substituted with < and > since they are reserved characters in XML.
6. 46.	Qualifier	phenotype
	Definition	phenotype conferred by the feature, where phenotype is defined as a physical, biochemical or behavioural characteristic or set of characteristics
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<pre><insdqualifier_value>erythromycin resistance</insdqualifier_value></pre>
6. 47.	Qual i fi er	pl asmi d
	Definition	name of naturally occurring plasmid from which the sequence was obtained, where plasmid is defined as an independently replicating genetic unit that cannot be described by chromosome or segment qualifiers
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>pC589</insdqualifier_value>
6. 48.	Qual i fi er	pop_vari ant
	Definition	name of subpopulation or phenotype of the sample from which the sequence was derived $% \left(1\right) =\left(1\right) \left($
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>pop1</insdqualifier_value> <insdqualifier_value>Bear Paw</insdqualifier_value>

6. 49.	Qual i fi er	product
	Definition	name of the product associated with the feature, e.g. \blacksquare the mRNA of an mRNA feature, the polypeptide of a CDS, the mature peptide of a mat_peptide, etc.
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<pre><insdqualifier_value>trypsinogen</insdqualifier_value> (when qualifier appears in CDS feature) <insdqualifier_value>trypsin</insdqualifier_value> (when qualifier appears in mat_peptide feature) <insdqualifier_value>XYZ neural-specific transcript</insdqualifier_value> (when qualifier appears in mRNA feature)</pre>
6. 50.	Qual i fi er	protei n_i d
	Definition	protein sequence identification number, an integer used in a sequence listing to designate the protein sequence encoded by the coding sequence identified in the corresponding CDS feature key and translation qualifier
	Value format	an integer greater than zero
	Exampl e	<insdqualifier_value>89</insdqualifier_value>
6. 51.	Qual i fi er	provi ral
	Definition	this qualifier is used to flag sequence obtained from a virus or phage that is integrated into the genome of another organism
	Value format	none
6. 52.	Qual i fi er	pseudo
	Definition	indicates that this feature is a non-functional version of the element named by the feature \ensuremath{key}
	Value format	none
	Comment	The qualifier pseudo should be used to describe non-functional genes that are not formally described as pseudogenes, e.g. CDS has no translation due to other reasons than pseudogenization events. Other reasons may include sequencing or assembly errors. In order to annotate pseudogenes the qualifier pseudogene must be used, indicating the TYPE of pseudogene.

6. 53. Qual i fi er pseudogene Definition indicates that this feature is a pseudogene of the element named by the feature key Value format where TYPE is one of the following controlled vocabulary terms or phrases: processed unprocessed uni tary allelic unknown Example <INSDQualifier value>processed</INSDQualifier value> <INSDQualifier_value>unprocessed</INSDQualifier_value> <INSDQualifier_value>unitary</INSDQualifier_value> <INSDQualifier_value>allelic</INSDQualifier_value> <INSDQualifier_value>unknown</INSDQualifier_value> Comment Definitions of TYPE values: processed - the pseudogene has arisen by reverse transcription of a mRNA into cDNA, followed by reintegration into the genome. Therefore, it has lost any intron/exon structure, and it might have a pseudo-polyA-tail. unprocessed - the pseudogene has arisen from a copy of the parent gene by duplication followed by accumulation of random mutations. The changes, compared to their functional homolog, include insertions, deletions, premature stop codons, frameshifts and a higher proportion of non-synonymous versus synonymous substitutions. unitary - the pseudogene has no parent. It is the original gene, which is functional is some species but disrupted in some way (indels, mutation, recombination) in another species or strain. allelic - a (unitary) pseudogene that is stable in the population but importantly it has a functional alternative allele also in the population. i.e., one strain may have the gene, another strain may have the pseudogene. MHC haplotypes have allelic pseudogenes. $unknown \ \hbox{- the submitter does not know the method of pseudogenization}.$

6.54. Qualifier

rearranged

Definition

the sequence presented in the entry has undergone somatic rearrangement as part of an adaptive immune response; it is not the unrearranged sequence that was inherited from the parental germline

Value format

none

Comment

The rearranged qualifier must not be used to annotate chromosome rearrangements that are not involved in an adaptive immune response; germline and rearranged qualifiers must not be used in the same source feature; germline and rearranged qualifiers must only be used for molecules that can undergo somatic rearrangements as part of an adaptive immune response; these are the T-cell receptor (TCR) and immunoglobulin loci in the jawed vertebrates, and the unrelated variable lymphocyte receptor (VLR) locus in the jawless fish (lampreys and hagfish); germline and rearranged qualifiers should not be used outside of the Craniata (taxid=89593)

6. 55. Qual i fi er recombination class Definition a structured description of the classification of recombination hotspot region within a sequence Value format TYPE where TYPE is one of the following controlled vocabulary terms or phrases: meiotic mitotic non_allelic_homologous chromosome_breakpoint other Exampl e <INSDQualifier_value>meiotic</INSDQualifier_value> <INSDQualifier_value>chromosome_breakpoint</INSDQualifier_value> Comment specific recombination classes not yet in the recombination_class controlled vocabulary must be annotated by entering "other" as the recombination_class qualifier value and providing a brief explanation of the novel recombination_class in a note qualifier 6. 56. Qual i fi er regul atory_cl ass a structured description of the classification of transcriptional, translational, Definition replicational and chromatin structure related regulatory elements in a sequence Value format TYPE where TYPE is one of the following controlled vocabulary terms or phrases: attenuator CAAT_si gnal ${\tt DNase_I_hypersensitive_site}$ enhancer_bl ocki ng_el ement GC_si gnal imprinting_control_region insulator locus_control_region matrix_attachment_region mi nus_35_si gnal mi nus_10_si gnal pol yA_si gnal _sequence recoding_stimulatory_region replication_regulatory_region response element ri bosome_bi ndi ng_si te ri boswi tch silencer TATA_box termi nator transcriptional_cis_regulatory_region Exampl e <INSDQualifier_value>promoter</INSDQualifier_value> $<\!I\,NSDQual\,i\,fi\,er_val\,ue\!>\!enhancer<\!/I\,NSDQual\,i\,fi\,er_val\,ue\!>$ <INSDQualifier_value>ribosome_binding_site</INSDQualifier_value> Comment specific regulatory classes not yet in the regulatory_class controlled vocabulary must be annotated by entering "other" as the regulatory_class qualifier value and providing a brief explanation of the novel regulatory_class in a note qualifier

6. 57.	Qual i fi er	repl ace
	Definition	indicates that the sequence identified in a feature's location is replaced by the sequence shown in the qualifier's value; if no sequence (i.e., no value) is contained within the qualifier, this indicates a deletion
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<pre><insdqualifier_value>a</insdqualifier_value> <insdqualifier_value></insdqualifier_value> - for a deletion</pre>
6. 58.	Qual i fi er	ri bosomal_sli ppage
	Definition	during protein translation, certain sequences can program ribosomes to change to an alternative reading frame by a mechanism known as ribosomal slippage
	Value format	none
	Comment	a join operator, e.g. \blacksquare : [join(4861784,17874810)] must be used in the CDS feature location to indicate the location of ribosomal_slippage
6. 59.	Qual i fi er	rpt_family
	Definition	type of repeated sequence; "Alu" or "Kpn", for example
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>Alu</insdqualifier_value>

6. 60. Qual i fi er rpt_type Definition structure and distribution of repeated sequence Value format One of the following controlled vocabulary terms or phrases: tandem di rect inverted fl anki ng nested <u>termi nal</u> di spersed long_terminal_repeat $non_l\,tr_retrotransposon_pol\,ymeri\,c_tract$ centromeri c_repeat tel omeri c_repeat x_el ement_combi natori al _repeat y_pri me_el ement other Example <INSDQualifier_value>inverted</INSDQualifier_value> $<\!I\,NSDQual\,i\,fi\,er_val\,ue>\!l\,ong_t\,ermi\,nal\,_repeat<\!/I\,NSDQual\,i\,fi\,er_val\,ue>$ Comment Definitions of the values: tandem - a repeat that exists adjacent to another in the same orientation; direct - a repeat that exists not always adjacent but is in the same orientation; inverted – a repeat pair occurring in reverse orientation to one another on the same molecule; $flanking \ \hbox{-} \ a \ repeat \ lying \ outside \ the \ sequence \ for \ which \ it \ has \ functional$ significance (eg. transposon insertion target sites); nested - a repeat that is disrupted by the insertion of another element; dispersed - a repeat that is found dispersed throughout the genome; terminal - a repeat at the ends of and within the sequence for which it has functional significance (eg. transposon LTRs); long_terminal_repeat - a sequence directly repeated at both ends of a defined sequence, of the sort typically found in retroviruses; non_ltr_retrotransposon_polymeric_tract - a polymeric tract, such as poly(dA), within a non LTR retrotransposon; $centromeri\,c_repeat\ \hbox{-}\ a\ repeat\ region\ found\ within\ the\ modul\,ar\ centromere;}$ telomeric_repeat - a repeat region found within the telomere; x_{el} ement_combinatorial_repeat - a repeat region located between the X element and the telomere or adjacent Y' element; $y_prime_element$ - a repeat region located adjacent to telomeric repeats or Xelement combinatorial repeats, either as a single copy or tandem repeat of two to four copies; other - a repeat exhibiting important attributes that cannot be described by other

6. 61.	Qualifier	rpt unit range
0. 01.	quarrirei	r pc_unit c_r ange
	Definition	location of a repeating unit expressed as a range
	Value format	<pre><base_range> - where <base_range> is the first and last base (separated by two dots) of a repeating unit</base_range></base_range></pre>
	Exampl e	<i er_val="" fi="" i="" nsdqual="" ue="">202 245</i>
	Comment	used to indicate the base range of the sequence that constitutes a repeating unit within the region specified by the feature keys oriT and repeat_region.

values.

6. 62.	Qual i fi er	rpt_uni t_seq
	Definition	identity of a repeat sequence
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<i fier_val="" i="" nsdqual="" ue="">aagggc</i> <i fier_val="" i="" nsdqual="" ue="">ag(5) tg(8) </i> <i fier_val="" i="" nsdqual="" ue="">(AAAGA) 6(AAAA) 1(AAAGA) 12</i>
	Comment	used to indicate the literal sequence that constitutes a repeating unit within the region specified by the feature keys oriT and repeat_region
6. 63.	Qualifier	satellite
	Definition	identifier for a satellite DNA marker, compose of many tandem repeats (identical or related) of a short basic repeated unit
	Value format	<pre><satellite_type>[:<class>][<identifier>] - where <satellite_type> is one of the following: satellite; microsatellite; minisatellite</satellite_type></identifier></class></satellite_type></pre>
	Exampl e	<insdqualifier_value>satellite: S1a</insdqualifier_value> <insdqualifier_value>satellite: alpha</insdqualifier_value> <insdqualifier_value>satellite: gamma III</insdqualifier_value> <insdqualifier_value>microsatellite: DC130</insdqualifier_value>
	Comment	many satellites have base composition or other properties that differ from those of the rest of the genome that allows them to be identified.
6. 64.	Qual i fi er	segment
	Definition	name of viral or phage segment sequenced
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>6</insdqualifier_value>
6. 65.	Qualifier	serotype
	Definition	serological variety of a species characterized by its antigenic properties
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>B1</insdqualifier_value>
	Comment	used only with the source feature key; the Bacteriological Code recommends the use of the term 'serovar' instead of 'serotype' for the prokaryotes; see the International Code of Nomenclature of Bacteria (1990 Revision) Appendix 10.B "Infraspecific Terms".

6. 66.	Qual i fi er	serovar
	Definition	serological variety of a species (usually a prokaryote) characterized by its antigenic properties $% \left(1\right) =\left(1\right) \left(1\right) +\left(1\right) \left(1\right) \left(1\right) +\left(1\right) \left(1\right) \left(1\right) \left(1\right) +\left(1\right) \left(1\right) \left$
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqual er_val="" fi="" i="" ue="">0157: H7</insdqual>
	Comment	used only with the source feature key; the Bacteriological Code recommends the use of the term 'serovar' instead of 'serotype' for prokaryotes; see the International Code of Nomenclature of Bacteria (1990 Revision) Appendix 10. B "Infraspecific Terms".
6. 67.	Qualifier	sex
	Definition	sex of the organism from which the sequence was obtained; sex is used for eukaryotic organisms that undergo meiosis and have sexually dimorphic gametes
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Examples	<pre><insdqualifier_value>female</insdqualifier_value> <insdqualifier_value>male</insdqualifier_value> <insdqualifier_value>hermaphrodite</insdqualifier_value> <insdqualifier_value>hermaphrodifier_value> <insdqualifier_value>sexual</insdqualifier_value> <insdqualifier_value>bisexual</insdqualifier_value> <insdqualifier_value>asexual</insdqualifier_value> <insdqualifier_value>monoecious</insdqualifier_value> [or monecious] <insdqualifier_value>dioecious</insdqualifier_value> [or diecious]</insdqualifier_value></pre>
	Comment	The sex qualifier should be used (instead of mating_type qualifier) in the Metazoa, Embryophyta, Rhodophyta & Phaeophyceae; mating_type qualifier should be used (instead of sex qualifier) in the Bacteria, Archaea & Fungi; neither sex nor mating_type qualifiers should be used in the viruses; outside of the taxa listed above, mating_type qualifier should be used unless the value of the qualifier is taken from the vocabulary given in the examples above
6. 68.	Qual i fi er	standard_name
	Definition	accepted standard name for this feature
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<pre><insdqualifier_value>dotted</insdqualifier_value></pre>
	Comment	use standard_name qualifier to give full gene name, but use gene qualifier to give gene symbol (in the above example gene qualifier value is Dt).
6. 69.	Qual i fi er	strain
	Definition	strain from which sequence was obtained
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>BALB/c</insdqualifier_value>
	Comment	feature entries including a strain qualifier must not include the environmental_sample qualifier

6. 70.	Qualifier	sub_cl one
	Definition	sub-clone from which sequence was obtained
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>l ambda-hIL7. 20g</insdqualifier_value>
	Comment	a source feature must not contain more than one sub_clone qualifier; to indicate that the sequence was obtained from multiple sub_clones, multiple sources may be further described using the feature key "misc_feature" and the qualifier "note"
6. 71.	Qual i fi er	sub_speci es
	Definition	name of sub-species of organism from which sequence was obtained
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Example	<insdqualifier_value>lactis</insdqualifier_value>
6. 72.	Qual i fi er	sub_strain
	Definition	name or identifier of a genetically or otherwise modified strain from which sequence was obtained, derived from a parental strain (which should be annotated in the strain qualifier). sub_strain from which sequence was obtained
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>abis</insdqualifier_value>
	Comment	must be accompanied by a strain qualifier in a source feature; if the parental strain is not given, the modified strain should be annotated in the strain qualifier instead of sub_strain. For example, either a strain qualifier with the value K-12 and a substrain qualifier with the value MG1655 or a strain qualifier with the value MG1655
6. 73.	Qual i fi er	tag_peptide
	Definition	base location encoding the polypeptide for proteolysis tag of \ensuremath{tmRNA} and its termination codon
	Value format	<pre><base_range> - where <base_range> provides the first and last base (separated by two dots) of the location for the proteolysis tag</base_range></base_range></pre>
	Exampl e	<insdqualifier_value>90 122</insdqualifier_value>
	Comment	it is recommended that the amino acid sequence corresponding to the tag_peptide be annotated by describing a 5' partial CDS feature; e.g. \blacksquare CDS with a location of $<\!90122$
6. 74.	Qual i fi er	tissue_lib
	Definition	tissue library from which sequence was obtained
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Example	<pre><insdqualifier_value>tissue library 772</insdqualifier_value></pre>

6. 75.	Qual i fi er	tissue_type
	Definition	tissue type from which the sequence was obtained
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>liver</insdqualifier_value>
6. 76.	Qual i fi er	transl_except
	Definition	translational exception: single codon the translation of which does not conform to genetic code defined by organism or transl_table.
	Value format	(pos:location, aa: <amino_acid>) where <amino_acid> is the three letter abbreviation for the amino acid coded by the codon at the base_range position</amino_acid></amino_acid>
	Exampl e	<pre><insdqualifier_value>(pos: 213 215, aa: Trp)</insdqualifier_value></pre>
	Comment	if the amino acid is not one of the specific amino acids listed in Section 3 of this Annex, use OTHER as <amino_acid> and provide the name of the unusual amino acid in a note qualifier; for modified amino-acid selenocysteine use three letter abbreviation 'Sec' (one letter symbol 'U' in amino-acid sequence) for <amino_acid>; for modified amino-acid pyrrolysine use three letter abbreviation 'Pyl' (one letter symbol 'O' in amino-acid sequence) for <amino_acid>; for partial termination codons where TAA stop codon is completed by the addition of 3' A residues to the mRNA either a single base_position or a base_range is used for the location, see the third and fourth examples above, in conjunction with a note qualifier indicating 'stop codon completed by the addition of 3' A residues to the mRNA'.</amino_acid></amino_acid></amino_acid>
6. 77.	Qual i fi er	transl_table
	Definition	definition of genetic code table used if other than universal or standard genetic code table. Tables used are described in this Annex
	Value format	<pre><integer> where <integer> is the number assigned to the genetic code table</integer></integer></pre>
	Exampl e	$<\!$ INSDQualifier_value> $\!$ 3 $<\!$ /INSDQualifier_value> - example where the yeast mitochondrial code is to be used
	Comment	if the transl_table qualifier is not used to further annotate a CDS feature key, then the CDS is translated using the Standard Code (i.e. Universal Genetic Code). Genetic code exceptions outside the range of specified tables are reported in transl_except qualifiers.
6. 78.	Qual i fi er	trans_splicing
	Definition	indicates that exons from two RNA molecules are ligated in intermolecular reaction to form mature RNA $$
	Value format	none
	Comment	should be used on features such as CDS, mRNA and other features that are produced as a result of a trans-splicing event. This qualifier must be used only when the splice event is indicated in the "join" operator, e.g. join(complement(6961169724), 139856140087) in the feature location

6. 79.	Qual i fi er	translation
	Definition	one-letter abbreviated amino acid sequence derived from either the standard (or universal) genetic code or the table as specified in a transl_table qualifier and as determined by an exception in the transl_except qualifier
	Value format	contiguous string of one-letter amino acid abbreviations from Section 3 of this Annex, " X " is to be used for AA exceptions.
	Exampl e	<pre><insdqualifier_value>MASTFPPWYRGCASTPSLKGLIMCTW</insdqualifier_value></pre>
	Comment	to be used with CDS feature only; must be accompanied by protein_id qualifier when the translation product contains four or more specifically defined amino acids; see transl_table for definition and location of genetic code Tables; only one of the qualifiers translation, pseudo and pseudogene are permitted to further annotate a CDS feature.
6. 80.	Qualifier	variety
	Definition	variety (= varietas, a formal Linnaean rank) of organism from which sequence was derived.
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<insdqualifier_value>insularis</insdqualifier_value>
	Comment	use the cultivar qualifier for cultivated plant varieties, i.e., products of artificial selection; varieties other than plant and fungal variatas should be annotated via a note qualifier, e.g. with the value <insdqualifier_value>breed: Cukorova</insdqualifier_value>

SECTION 7: FEATURE KEYS FOR AMINO ACID SEQUENCES

This section contains the list of allowed feature keys to be used for amino acid sequences. The feature keys are listed in alphabetic order.

7. 1.	Feature Key	ACT_SITE
	Definition	Amino $\operatorname{acid}(s)$ involved in the activity of an enzyme
	Optional qualifiers	NOTE
	Comment	Each amino acid residue of the active site must be annotated separately with the ACT_SITE feature key. The corresponding amino acid residue number must be provided as the location descriptor in the feature location element.
7. 2.	Feature Key	BI NDI NG
	Definition	Binding site for any chemical group (co-enzyme, prosthetic group, etc.). The chemical nature of the group is indicated in the NOTE qualifier ${\cal O}(1)$
	Mandatory qualifiers	NOTE
	Comment	Examples of values for the "NOTE" qualifier: "Heme (covalent)" and "Chloride." Where appropriate, the features keys CA_BIND, DNA_BIND, METAL, and NP_BIND should be used rather than BINDING.
7. 3.	Feature Key	CA_BI ND
	Definition	Extent of a calcium-binding region
	Optional qualifiers	NOTE
7. 4.	Feature Key	CARBOHYD
	Definition	Glycosylation site
	Mandatory qualifiers	NOTE
	Comment	This key describes the occurrence of the attachment of a glycan (mono- or polysaccharide) to a residue of the protein. The type of linkage (C-, N- or O-linked) to the protein is indicated in the "NOTE" qualifier. If the nature of the reducing terminal sugar is known, its abbreviation is shown between parentheses. If three dots '' follow the abbreviation this indicates an extension of the carbohydrate chain. Conversely no dots means that a monosaccharide is linked. Examples of values used in the "NOTE" qualifier: N-linked (GlcNAc); O-linked (GlcNAc); O-linked (Glc); C-linked (Man) partial; O-linked (Ara).
7. 5.	Feature Key	CHAI N
	Definition	Extent of a polypeptide chain in the mature protein
	Optional qualifiers	NOTE
7. 6.	Feature Key	COI LED
	Definition	Extent of a coiled-coil region
	Optional qualifiers	NOTE

7. 7.	Feature Key	COMPBLAS
	Definition	Extent of a compositionally biased region
	Optional qualifiers	NOTE
7. 8.	Feature Key	CONFLICT
	Definition	Different sources report differing sequences
	Optional qualifiers	NOTE
	Comment	Examples of values for the "NOTE" qualifier: Missing; K -> Q; GSDSE -> RIRLR; V -> A.
7. 9.	Feature Key	CROSSLNK
	Definition	Post translationally formed amino acid bonds
	Mandatory qualifiers	NOTE
	Comment	Covalent linkages of various types formed between two proteins (interchain crosslinks) or between two parts of the same protein (intrachain cross-links); except for cross-links formed by disulfide bonds, for which the "DISULFID" feature key is to be used. For an interchain cross-link, the location descriptor in the feature location element is the residue number of the amino acid cross-linked to the other protein. For an intrachain cross-link, the location descriptors in the feature location element are the residue numbers of the cross-linked amino acids in conjunction with the "join" location operator, e.g. "join(42,50)." The NOTE qualifier indicates the nature of the cross-link; at least specifying the name of the conjugate and the identity of the two amino acids involved. Examples of values for the "NOTE" qualifier: "Isoglutamyl cysteine thioester (Cys-Gln);" "Betamethyllanthionine (Cys-Thr);" and "Glycyl lysine isopeptide (Lys-Gly) (interchain with G-Cter in ubiquitin)"
7. 10.	Feature Key	DI SULFI D
	Definition	Di sul fi de bond
	Mandatory qualifiers	NOTE
	Comment	For an interchain disulfide bond, the location descriptor in the feature location element is the residue number of the cysteine linked to the other protein. For an intrachain cross-link, the location descriptors in the feature location element are the residue numbers of the linked cysteines in conjunction with the "join" location operator, e.g. "join(42,50)". For interchain disulfide bonds, the NOTE qualifier indicates the nature of the cross-link, by identifying the other protein, for example, "Interchain (between A and B chains)"
7. 11.	Feature Key	DNA_BI ND
	Definition	Extent of a DNA-binding region
	Mandatory qualifiers	NOTE
	Comment	The nature of the DNA-binding region is given in the NOTE qualifier. Examples of values for the "NOTE" qualifier: "Homeobox" and "Myb 2"
7. 12.	Feature Key	DOMAI N
	Definition	Extent of a domain, which is defined as a specific combination of secondary structures organized into a characteristic three-dimensional structure or fold

	Mandatory qualifiers	NOTE
	Comment	The domain type is given in the NOTE qualifier. Where several copies of a domain are present, the domains are numbered. Examples of values for the "NOTE" qualifier: "Ras-GAP" and "Cadherin 1"
7. 13.	Feature Key	HELIX
	Definition	Secondary structure: Helices, for example, Alpha-helix; 3(10) helix; or Pi-helix
	Optional qualifiers	NOTE
	Comment	This feature is used only for proteins whose tertiary structure is known. Only three types of secondary structure are specified: helices (key HELIX), beta-strands (key STRAND) and turns (key TURN). Residues not specified in one of these classes are in a 'loop' or 'random-coil' structure.
7. 14.	Feature Key	I NI T_MET
	Definition	Initiator methionine
	Optional qualifiers	NOTE
	Comment	The location descriptor in the feature location element is "1". This feature key indicates the N-terminal methionine is cleaved off. This feature is not used when the initiator methionine is not cleaved off.
7. 15.	Feature Key	I NTRAMEM
	Definition	Extent of a region located in a membrane without crossing it
	Optional qualifiers	NOTE
7. 16.	Feature Key	LIPID
	Definition	Covalent binding of a lipid moiety
	Mandatory qualifiers	NOTE
	Comment	The chemical nature of the bound lipid moiety is given in the NOTE qualifier, indicating at least the name of the lipidated amino acid. Examples of values for the "NOTE" qualifier: "N-myristoyl glycine"; "GPI-anchor amidated serine" and "S-diacylglycerol cysteine."
7. 17.	Feature Key	METAL
	Definition	Binding site for a metal ion.
	Mandatory qualifiers	NOTE
	Comment	The NOTE qualifier indicates the nature of the metal. Examples of values for the "NOTE" qualifier: "Iron (heme axial ligand)" and "Copper".

7. 18.	Feature Key	MOD_RES
	Definition	Posttranslational modification of a residue
	Mandatory qualifiers	NOTE
	Comment	The chemical nature of the modified residue is given in the NOTE qualifier, indicating at least the name of the post-translationally modified amino acid. If the modified amino acid is listed in Section 4 of this Annex, the abbreviation may be used in place of the the full name. Examples of values for the "NOTE" qualifier: "N-acetylalanine"; "3-Hyp"; and "MeLys" or "N-6-methyllysine"
7. 19.	Feature Key	MOTI F
	Definition	Short (up to 20 amino acids) sequence motif of biological interest
	Optional qualifiers	NOTE
7. 20.	Feature Key	MUTAGEN
	Definition	Site which has been experimentally altered by mutagenesis
	Optional qualifiers	NOTE
7. 21.	Feature Key	NON_STD
	Definition	Non-standard amino acid
	Optional qualifiers	NOTE
	Comment	This key describes the occurrence of non-standard amino acids selenocysteine (U) and pyrrolysine (O) in the amino acid sequence.
7. 22.	Feature Key	NON_TER
	Definition	The residue at an extremity of the sequence is not the terminal residue
	Optional qualifiers	NOTE
	Comment	If applied to position 1, this means that the first position is not the N-terminus of the complete molecule. If applied to the last position, it means that this position is not the C-terminus of the complete molecule.
7. 23.	Feature Key	NP_BI ND
	Definition	Extent of a nucleotide phosphate-binding region
	Mandatory qualifiers	NOTE
	Comment	The nature of the nucleotide phosphate is indicated in the NOTE qualifier. Examples of values for the "NOTE" qualifier: "ATP" and "FAD".
7. 24.	Feature Key	PEPTI DE
	Definition	Extent of a released active peptide
	Optional qualifiers	NOTE

7. 25.	Feature Key	PROPEP
	Definition	Extent of a propeptide
	Optional qualifiers	NOTE
7. 26.	Feature Key	REGI ON
	Definition	Extent of a region of interest in the sequence
	Optional qualifiers	NOTE
7. 27.	Feature Key	REPEAT
	Definition	Extent of an internal sequence repetition
	Optional qualifiers	NOTE
7. 28.	Feature Key	SIGNAL
	Definition	Extent of a signal sequence (prepeptide)
	Optional qualifiers	NOTE
7. 29.	Feature Key	SITE
	Definition	Any interesting single amino-acid site on the sequence that is not defined by another feature key. It can also apply to an amino acid bond which is represented by the positions of the two flanking amino acids
	Mandatory qualifier	NOTE
	Comment	When SITE is used to annotate a modified amino acid the value for the qualifier "NOTE" must either be an abbreviation set forth in Section 4 of this Annex, or the complete, unabbreviated name of the modified amino acid.
7. 30.	Feature Key	SOURCE
	Definition	Identifies the source of the sequence; this key is mandatory; every sequence will have a single SOURCE feature spanning the entire sequence
	Mandatory qualifiers	MOL_TYPE ORGANI SM
	Optional qualifiers	NOTE
7. 31.	Feature Key	STRAND
	Definition	Secondary structure: Beta-strand; for example Hydrogen bonded beta-strand or residue in an isolated beta-bridge
	Optional qualifiers	NOTE
	Comment	This feature is used only for proteins whose tertiary structure is known. Only three types of secondary structure are specified: helices (key HELIX), beta-strands (key STRAND) and turns (key TURN). Residues not specified in one of these classes are in a 'loop' or 'random-coil' structure.

7. 32.	Feature Key	TOPO_DOM
	Definition	Topol ogi cal domai n
	Optional qualifiers	NOTE
7. 33.	Feature Key	TRANSMEM
	Definition	Extent of a transmembrane region
	Optional qualifiers	NOTE
7. 34.	Feature Key	TRANSIT
	Definition	Extent of a transit peptide (mitochondrion, chloroplast, thylakoid, cyanelle, peroxisome etc.)
	Optional qualifiers	NOTE
7. 35.	Feature Key	TURN
	Definition	Secondary structure Turns, for example, H-bonded turn (3-turn, 4-turn or 5-turn)
	Optional qualifiers	NOTE
	Comment	This feature is used only for proteins whose tertiary structure is known. Only three types of secondary structure are specified: helices (key HELIX), beta-strands (key STRAND) and turns (key TURN). Residues not specified in one of these classes are in a 'loop' or 'random-coil' structure.
7. 36.	Feature Key	UNSURE
	Definition	Uncertainties in the sequence
	Optional qualifiers	NOTE
	Comment	Used to describe $\operatorname{region}(s)$ of an amino acid sequence for which the authors are unsure about the sequence presentation.
7. 37.	Feature Key	VARI ANT
	Definition	Authors report that sequence variants exist
	Optional qualifiers	NOTE
_		
7. 38.	Feature Key	VAR_SEQ
	Definition	Description of sequence variants produced by alternative splicing, alternative promoter usage, alternative initiation and ribosomal frameshifting
	Optional qualifiers	NOTE

7. 39.	Feature Key	ZN_FI NG
	Definition	Extent of a zinc finger region
	Mandatory qualifiers	NOTE
	Comment	The type of zinc finger is indicated in the NOTE qualifier. For example: "GATA-type" and "NR C4-type"

SECTION 8: QUALIFIERS FOR AMINO ACID SEQUENCES

This section contains the list of allowed qualifiers to be used for amino acid sequences.

PLEASE NOTE: Any qualifier value provided for a qualifier with a "free text" value format may require translation for National/Regional procedures.

8. 1.	Qualifier	MOL_TYPE
	Definition	In vivo molecule type of sequence
	Value format	protein
	Exampl e	<pre><insdqualifier_value>protein</insdqualifier_value></pre>
	Comment	The "MOL_TYPE" qualifier is mandatory on the SOURCE feature key.
8. 2.	Qual i fi er	NOTE
	Definition	Any comment or additional information
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<pre><insdqualifier_value>Heme (covalent)</insdqualifier_value></pre>
	Comment	The "NOTE" qualifier is mandatory for the feature keys: BINDING; CARBOHYD; CROSSLNK; DISULFID; DNA_BIND; DOMAIN; LIPID; METAL; MOD_RES; NP_BIND and ZN_FING
8. 3.	Qualifier	ORGANI SM
	Definition	Scientific name of the organism that provided the peptide
	Value format	free text (NOTE: this value may require translation for National/Regional procedures)
	Exampl e	<pre><insdqualifier_value>Homo sapiens</insdqualifier_value></pre>
	Comment	The "ORGANISM" qualifier is mandatory for the SOURCE feature key.

SECTION 9: GENETIC CODE TABLES

Table 5 reproduces Genetic Code Tables to be used for translating coding sequences. The value for the trans_table qualifier is the number assigned to the corresponding genetic code table. Where a CDS feature is described with a translation qualifier but not a transl_table qualifier, the 1 - Standard Code is used by default for translation. (Note: Genetic code tables 7, 8, 15, and 17 to 20 do not exist, therefore these numbers do not appear in Table 5.)

Table 5: Genetic Code Tables

	1 - Standard Code
AAs =	FFLLSSSSYY**CC*WLLLLPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGG
Starts =	M M M M M
Base1 =	tttttttttttttttttcccccccccccccaaaaaaaaa
Base2 =	ttttccccaaaaggggttttccccaaaaggggttttccccaaaagggg
Base3 =	t cagt cagt cagt cagt cagt cagt cagt cag
	2 - Vertebrate Mitochondrial Code
AAs =	FFLLSSSSYY**CCWWLLLLPPPPHHQQRRRRIIMMTTTTNNKKSS**VVVVAAAADDEEGGGG
Starts =	MMMM M
Base1 =	tttttttttttttttttcccccccccccccaaaaaaaaa
	ttttccccaaaaggggttttccccaaaaggggttttccccaaaaggggttttccccaaaagggg
Base3 =	t cagt cagt cagt cagt cagt cagt cagt cag
	3 - Yeast Mitochondrial Code
AAs =	FFLLSSSSYY**CCWWTTTTPPPPHHQQRRRRIIMMTTTTNNKKSSRRVVVVAAAADDEEGGGG
Starts =	
D 1	
Base1 =	ttttttttttttttttccccccccccccaaaaaaaaaaa
Base2 =	ttttccccaaaaggggttttccccaaaaggggttttccccaaaagggg
Base3 =	tcagtcagtcagtcagtcagtcagtcagtcagtcagtcag
	4 - Mold, Protozoan, Coelenterate Mitochondrial Code &
	Mycoplasma/Spiroplasma Code
	, , , ,
AAs =	FFLLSSSSYY**CCWWLLLLPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGG
Starts =	FFLLSSSSYY**CCWWLLLLPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGG
Starts = Base1 =	FFLLSSSSYY**CCWWLLLLPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGGMMM
Starts = Base1 = Base2 =	FFLLSSSSYY**CCWWLLLLPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGGMM
Starts = Base1 = Base2 = Base3 =	FFLLSSSSYY**CCWWLLLLPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGGMM
Starts = Base1 = Base2 = Base3 =	FFLLSSSSYY**CCWWLLLLPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGGMM
Starts = Base1 = Base2 = Base3 = AAs = Starts =	FFLLSSSSYY**CCWWLLLLPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGGMM
Starts = Base1 = Base2 = Base3 = AAs = Starts = Base1 =	FFLLSSSSYY**CCWWLLLLPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGGMM
Starts = Base1 = Base2 = Base3 = Starts = Base1 = Base2 =	FFLLSSSSYY**CCWWLLLLPPPPHHQQRRRRIIIMTTTNNKKSSRRVVVVAAAADDEEGGGGMM
Starts = Base1 = Base2 = Base3 = Starts = Base1 = Base2 =	FFLLSSSSYY**CCWWLLLLPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGGMM
Starts = Base1 = Base2 = Base3 = Starts = Base1 = Base2 = Base3 =	FFLLSSSSYY**CCWWLLLLPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGGMM
Starts = Base1 = Base2 = Base3 = Starts = Base1 = Base2 = Base3 = AAs =	FFLLSSSSYY**CCWWLLLLPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGGMM
Starts = Base1 = Base2 = Base3 = AAs = Starts = Base2 = Base3 = AAs = Starts =	FFLLSSSYY**CCWWLLLLPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGGMM

9 - Echinoderm and Flatworm Mitochondrial Code Base2 = ttttccccaaaaggggttttccccaaaaggggttttccccaaaaggggttttccccaaaagggg 10 - Euplotid Nuclear Code AAs = FFLLSSSSYY**CCCWLLLLPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGG Base2 = ttttccccaaaaggggttttccccaaaaggggttttccccaaaaggggttttccccaaaagggg 11 - Bacterial, Archaeal and Plant Plastid Code $AAS \ = \ FFLLSSSSYY**CC*WLLLLPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGG$ $Base 2 \quad = \quad ttttccccaaaaggggttttccccaaaaggggttttccccaaaaggggttttccccaaaagggg$ $Base 3 \quad = \quad t cag t c$ 12 - Alternative Yeast Nuclear Code $AAs \ = \ FFLLSSSSYY**CC*WLLLSPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGG$ $Base 2 \quad = \quad ttttccccaaaaggggttttccccaaaaggggttttccccaaaaggggttttccccaaaagggg$ 13 - Ascidian Mitochondrial Code $\textbf{AAS} \quad = \quad \textbf{FFLLSSSSYY**CCWWLLLPPPPHHQQRRRRIIMMTTTTNNKKSSGGVVVVAAAADDEEGGGG}$ Starts = --- M------ M------ M------ M------ $Base 2 \quad = \quad ttttccccaaaaggggttttccccaaaaggggttttccccaaaaggggttttccccaaaagggg$ 14 - Alternative Flatworm Mitochondrial Code AAS = FFLLSSSSYYY*CCWWLLLLPPPPHHQQRRRRIIIMTTTTNNNKSSSSVVVVAAAADDEEGGGG Starts = $Base 2 \quad = \quad ttttccccaaaaggggttttccccaaaaggggttttccccaaaaggggttttccccaaaagggg$ Starts tttteeccaaaaggggtttteeceaaaaggggtttteeceaaaaggggtttteeccaaaagggg 16 - Chlorophycean Mitochondrial Code AAs = FFLLSSSSYY*LCC*WLLLLPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGG ----- M------ M-----Starts = Base2 = ttttccccaaaaggggttttccccaaaaggggttttccccaaaaggggttttccccaaaagggg

21 - Trematode Mitochondrial Code $\textbf{AAS} \quad = \quad \textbf{FFLLSSSSYY**CCWWLLLLPPPPHHQQRRRRIIMMTTTTNNNKSSSSVVVVAAAADDEEGGGG}$ Starts = ttttccccaaaaggggttttccccaaaaggggttttccccaaaaggggttttccccaaaagggg22 - Scenedesmus obliquus Mitochondrial Code FFLLSS*SYY*LCC*WLLLLPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGGStarts = ----- M----- M-----Base2 = ttttccccaaaaggggttttccccaaaaggggttttccccaaaaggggttttccccaaaagggg 23 - Thraustochytrium Mitochondrial Code $AAS \ = \ FF*LSSSSYY**CC*WLLLLPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGG$ Starts = ----- M--M------ M--M-----Base2 = ttttccccaaaaggggttttccccaaaaggggttttccccaaaaggggttttccccaaaagggg 24 - Pterobranchia Mitochondrial Code FFLLSSSSYY**CCWWLLLLPPPPHHQQRRRRIIIMTTTTNNKKSSSKVVVVAAAADDEEGGGG Base2 = ttttccccaaaaggggttttccccaaaaggggttttccccaaaaggggttttccccaaaagggg 25 - Candidate Division SR1 and Gracilibacteria Code AAs = FFLLSSSSYY**CCGWLLLLPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGG Base1 Base2 = ttttccccaaaaggggttttccccaaaaggggttttccccaaaaggggttttccccaaaagggg 26 - Pachysolen tannophilus Nuclear Code = FFLLSSSSYY**CC*WLLLAPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGG Base2 = ttttccccaaaaggggttttccccaaaaggggttttccccaaaaggggttttccccaaaagggg27 - Karyorelict Nuclear Code -SSYYQQCCWWLLLAPPPPHHQQRRRRI I I MTTTTNNKKSSRRVVVVAAAADDEEGGGG<mark>FFLLSSSSYYQQCCW</mark> PPPHHQQRRRRI I I MTTTTNNKKSSRRVVVVAAAADDEEGGGG Base2 = ttttccccaaaaggggttttccccaaaaggggttttccccaaaaggggttttccccaaaagggg

28 - Condylostoma Nuclear Code AAS = FFLLSSSSYYQQCCWWLLLAPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAAADDEEGGGGFFLLSSSSYYQQCCWWLLLPPPPHHQQRRRIIIMTTTNNKKSSRRVVVVAAAAADDEEGGGG Base2 = ttttccccaaaaggggttttccccaaaaggggttttccccaaaaggggttttccccaaaagggg $Base 3 \quad = \quad t cag t c$ 29 - Mesodinium Nuclear Code AAs = FFLLSSSSYYYYCC* WLLLAPPPPHHQQRRRRI I LMTTTTNNKKSSRRVVVVAAAAADDEEGGGGWLLLLPPPPHHQQRR RRI I I MTTTTNNKKSSRRVVVVAAAADDEEGGGG --Starts = ----- M------Base2 = ttttccccaaaaggggttttccccaaaaggggttttccccaaaaggggttttccccaaaagggg 30 - Peritrich Nuclear Code AAS = FILLSSSSYYEECC*WLLLAPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAAADDEEGGGG<mark>WLLLLPPPPPHHQQRR</mark>RIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGG Base2 = ttttccccaaaaggggttttccccaaaaggggttttccccaaaaggggttttccccaaaagggg 31 - Blastocrithidia Nuclear Code AAS = FFLLSSSSYYEECCWWLLLLPPPPHHQQRRRRIIIMTTTTNNKKSSRRVVVVAAAADDEEGGGG Base2 = ttttccccaaaaggggttttccccaaaaggggttttccccaaaaggggttttccccaaaagggg 33 - Cephalodiscidae Mitochondrial UAA-Tyr Code AAS = FFLLSSSSYYY*CCWWLLLLPPPPHHQQRRRRIIIMTTTTNNKKSSSKVVVVAAAADDEEGGGG Base2 = ttttccccaaaaggggttttccccaaaaggggttttccccaaaaggggttttccccaaaagggg

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