## The DDBJ/ENA/GenBank Feature Table Definition

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DDBJ/ENA/GenBank Feature Table Definition
Version 11.0 October 2020
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## 1 Introduction

Nucleic acid sequences provide the fundamental starting point for describing and understanding the structure, function, and development of genetically diverse organisms. The GenBank, EMBL, and DDBJ nucleic acid sequence data banks have from their inception used tables of sites and features to describe the roles and locations of higher order sequence domains and elements within the genome of an organism.

In February, 1986, GenBank and EMBL began a collaborative effort (joined by DDBJ in 1987) to devise a common feature table format and common standards for annotation practice.

## 2 Overview of the Feature Table format

The overall goal of the feature table design is to provide an extensive vocabulary for describing features in a flexible framework for manipulating them. The Feature Table documentation represents the shared rules that allow the three databases to exchange data on a daily basis.

The range of features to be represented is diverse, including regions which:

- \* perform a biological function,
- \* affect or are the result of the expression of a biological function,
- \* interact with other molecules,
- \* affect replication of a sequence,
- \* affect or are the result of recombination of different sequences,
- \* are a recognizable repeated unit,
- \* have secondary or tertiary structure,
- \* exhibit variation, or have been revised or corrected.

#### 2.1 Format Design

The format design is based on a tabular approach and consists of the following items:

Feature key - a single word or abbreviation indicating functional group Location - instructions for finding the feature Qualifiers - auxiliary information about a feature

## 2.2 Key aspects of this feature table design

- \* Feature keys allow specific annotation of important sequence features.
- \* Related features can be easily specified and retrieved.

  Feature keys are arranged hierarchically, allowing complex and compound features to be expressed. Both location operators and the feature keys show feature relationships even when the features are not contiguous. The hierarchy of feature keys allows broad categories of biological functionality, such as rRNAs, to be easily retrieved.
- \* Generic feature keys provide a means for entering new or undefined features. A number of "generic" or miscellaneous feature keys have been added to permit annotation of features that cannot be adequately described by existing feature keys. These generic feature keys will serve as an intermediate step in the identification and addition of new feature keys. The syntax has been designed to allow the addition of new feature keys as they are required.
- \* More complex locations (fuzzy and alternate ends, for example) can be specified. Each end point of a feature may be specified as a single point, an alternate set of possible end points, a base number beyond which the end point lies, or a region which contains the end point.
- \* Features can be combined and manipulated in many different ways.

  The location field can contain operators or functional descriptors specifying what must be done to the sequence to reproduce the feature. For example, a series of exons may be "join"ed into a full coding sequence.
- \* Standardized qualifiers provide precision and parsibility of descriptive details
  A combination of standardized qualifiers and their controlled-vocabulary
  values enable free-text descriptions to be avoided.
- \* The nature of supporting evidence for a feature can be explicitly indicated. Features, such as open reading frames or sequences showing sequence similarity to consensus sequences, for which there is no direct experimental evidence can be annotated. Therefore, the feature table can incorporate contributions from researchers doing computational analysis of the sequence databases. However, all features that are supported by experimental data will be clearly marked as such.
- \* The table syntax has been designed to be machine parsible.

  A consistent syntax allows machine extraction and manipulation of sequences coding for all features in the table.

## 2.3 Feature Table Terminology

The format and wording in the feature table use common biological research terminology whenever possible. For example, an item in the feature table such as:

Key Location/Qualifiers

CDS 23..400

/product="alcohol dehydrogenase"

/gene="adhI"

might be read as:

The feature CDS is a coding sequence beginning at base 23 and ending at base 400, has a product called 'alcohol dehydrogenase' and is coded for by a gene called "adhI".

A more complex description:

Key Location/Qualifiers
CDS join(544..589,688..>1032)

/product="T-cell receptor beta-chain"

which might be read as:

This feature, which is a partial coding sequence, is formed by joining elements indicated to form one contiguous sequence encoding a product called T-cell receptor beta-chain.

The following sections contain detailed explanations of the feature table design showing conventions for each component of the feature table, examples of how the format might be implemented, a description of the exact column placement of all the data items and examples of complete sequence entries that have been annotated using the new format. The last section of this document describes known limitations of the current feature table design.

Appendix I gives an example database entry for the DDBJ, GenBank and EMBL formats.

Appendices II and III provide reference manuals for the feature table keys and qualifiers, respectively.

Appendix IV includes controlled vocabularies such as nucleotide base codes, modified base abbreviations, genetic code tables etc.

This document defines the syntax and vocabulary of the feature table. The syntax is sufficiently flexible to allow expression of a single biological entity in numerous ways. In such cases, the annotation staffs at the databases will propose conventions for standard means of denoting the entities. This feature table format is shared by GenBank, EMBL and DDBJ. Comments, corrections, and suggestions may be submitted to any of the database staffs. New format specifications will be added as needed.

# 3 Feature table components and format

## 3.1 Naming conventions

Feature table components, including feature keys, qualifiers, accession numbers, database name abbreviations, and location operators, are all named following the same conventions. Component names may be no more than 20 characters long (Feature keys 15, Feature qualifiers 20) and must contain at least one letter. The following characters are permitted to occur in feature table component names:

```
* Uppercase letters (A-Z)
* Lowercase letters (a-z) Numbers (0-9)
* Underscore (_)
* Hyphen (-)
* Single quotation mark or apostrophe (')
* Asterisk (*)
```

## 3.2 Feature keys

## 3.2.1 Purpose

Feature keys indicate

- (1) the biological nature of the annotated feature or
- (2) information about changes to or other versions of the sequence.

  The feature key permits a user to quickly find or retrieve similar features or

features with related functions.

### 3.2.2 Format and conventions

There is a defined list of allowable feature keys, which is shown in Appendix II. Each feature must contain a feature key.

### 3.2.3 Key groups and hierarchy

The feature keys fall into families which are in some sense similar in function and which are annotated in a similar manner. A functional family may have a "generic" or miscellaneous key, which can be recognized by the 'misc.' prefix, that can used for instances not covered by the other defined keys of that group.

The feature key groups are listed below with a short definition and an annotation example:

1. Difference and change features

Indicate ways in which a sequence should be changed to produce a different
"version":

2. Transcript features

Indicate products made by a region:
misc\_RNA location

### 3. Binding features

Indicate that a sequence or nucleotide is covalently, non-covalently, or otherwise bound to something else:

misc\_binding location

/bound\_moiety="bound molecule"

## 4. Repeat features

Indicate repetitive sequence elements:

repeat\_region location

#### 5. Recombination features

Indicate regions that have been either inserted or deleted by recombination:

misc recomb location

#### 6. Structure features

Indicate sequence for which there is secondary or tertiary structural information:

misc\_structure location

### 3.2.4 Feature key examples

Key Description

CDS Protein-coding sequence rep\_origin Origin of replication

protein\_bind
Protein binding site on DNA

tRNA mature transfer RNA

See Appendix II for descriptions of all feature keys.

### 3.3 Qualifiers

## 3.3.1 Purpose

Qualifiers provide a general mechanism for supplying information about features in addition to that conveyed by the key and location.

#### 3.3.2 Format and conventions

Qualifiers take the form of a slash (/) followed by the qualifier name and, if applicable, an equal sign (=) and a value. Each qualifier should have a single value; if multiple values are necessary, these should be represented by iterating the same qualifier, eg:

Key Location/Qualifiers

source 1..1000

/culture\_collection="ATCC:11775"
/culture\_collection="CECT:515"

If the location descriptor does not need a continuation line, the first qualifier begins a new line in the feature location column. If the location descriptor requires a continuation line, the first qualifier may follow immediately after the location. Any necessary continuation lines begin in the same column. See Section 4 for a complete description of data item positions.

### 3.3.3 Qualifier values

Since qualifiers convey many different types of information, there are several value formats:

- 1. Free text
- 2. Controlled vocabulary or enumerated values
- 3. Citation or reference numbers
- 4. Sequences

#### 3.3.3.1 Free text

Most qualifier values will be a descriptive text phrase which must be enclosed in double quotation marks. When the text occupies more than one line, a single set of quotation marks is required at the beginning and at the end of the text. The text itself may be composed of any printable characters (ASCII values 32-126 decimal). If double quotation marks are used within a free text string, each set (") must be 'escaped' by placing a second double quotation mark immediately before it (""). For example:

/note="This is an example of ""escaped"" quotation marks"

#### 3.3.3.2 Controlled vocabulary or enumerated values

Some qualifiers require values from a controlled vocabulary and are entered without quotation marks. For example, the '/direction' qualifier has only three values: 'left', 'right' or 'both'. Qualifier value controlled vocabularies, like feature table component names, must be treated as completely case insensitive: they may be entered and displayed in any combination of upper and lower case ('/direction=Left' '/direction=left' and '/direction=LEFT' are all legal and all convey the same meaning). The database staffs reserve the right to regularize the case of qualifier values. Qualifier value controlled vocabularies will be maintained by the cooperating database staffs. Examples of controlled vocabularies can be found in Appendices IV and V. The database staff should be contacted for the current lists.

#### 3.3.3.3 Citation or reference numbers

The citation or published reference number (as enumerated in the entry 'REFERENCE' or 'RN' data item) should be enclosed in square brackets (e.g., [3]) to distinguish it from other numbers.

### 3.3.3.4 Sequences

Literal sequence of nucleotide bases e.g., join(12..45,"atgcatt",988..1050) in location descriptors has become illegal starting from implementation of version 2.1 of the Feature Table Definition Document (December 15, 1998)

## 3.3.4 Qualifier examples

Key Location/Qualifiers

source 1..1509

/organism="Mus musculus"

/strain="CD1"

/mol\_type="genomic DNA"

regulatory <1..9

/gene="ubc42"

/regulatory\_class="promoter"

mRNA join(10..567,789..1320)

/gene="ubc42"

CDS join(54..567,789..1254)

/gene="ubc42"

/product="ubiquitin conjugating enzyme"

/function="cell division control"

#### 3.4 Location

## 3.4.1 Purpose

The location indicates the region of the presented sequence which corresponds to a feature.

#### 3.4.2 Format and conventions

The location contains at least one sequence location descriptor and may contain one or more operators with one or more sequence location descriptors. Base numbers refer to the numbering in the entry. This numbering designates the first base (5' end) of the presented sequence as base 1.

Base locations beyond the range of the presented sequence may not be used in location descriptors, the only exception being location in a remote entry (see 3.4.2.1, e).

Location operators and descriptors are discussed in more detail below.

### 3.4.2.1 Location descriptors

The location descriptor can be one of the following:

- (a) a single base number
- (b) a site between two indicated adjoining bases
- (c) a single base chosen from within a specified range of bases (not allowed for new entries)
- (d) the base numbers delimiting a sequence span
- (e) a remote entry identifier followed by a local location descriptor(i.e., a-d)

A site between two adjoining nucleotides, such as endonucleolytic cleavage site, is indicated by listing the two points separated by a carat (^). The permitted formats for this descriptor are n^n+1 (for example 55^56), or, for circular molecules, n^1, where "n" is the full length of the molecule, ie 1000^1 for circular molecule with length 1000.

A single base chosen from a range of bases is indicated by the first base

number and the last base number of the range separated by a single period (e.g., '12.21' indicates a single base taken from between the indicated points). From October 2006 the usage of this descriptor is restricted: it is illegal to use "a single base from a range" (c) either on its own or in combination with the "sequence span" (d) descriptor for newly created entries. The existing entries where such descriptors exist are going to be retrofitted.

Sequence spans are indicated by the starting base number and the ending base number separated by two periods (e.g., '34..456'). The '<' and '>' symbols may be used with the starting and ending base numbers to indicate that an end point is beyond the specified base number. The starting and ending base positions can be represented as distinct base numbers ('34..456') or a site between two indicated adjoining bases.

A location in a remote entry (not the entry to which the feature table belongs) can be specified by giving the accession-number and sequence version of the remote entry, followed by a colon ":", followed by a location descriptor which applies to that entry's sequence (i.e. J12345.1:1..15, see also examples below)

#### 3.4.2.2 Operators

The location operator is a prefix that specifies what must be done to the indicated sequence to find or construct the location corresponding to the feature. A list of operators is given below with their definitions and most common format.

complement(location)

Find the complement of the presented sequence in the span specified by "location" (i.e., read the complement of the presented strand in its 5'-to-3' direction)

join(location, location, ... location)

The indicated elements should be joined (placed end-to-end) to form one contiguous sequence

order(location,location, ... location)

The elements can be found in the

specified order (5' to 3' direction), but nothing is implied about the reasonableness about joining them

Note: location operator "complement" can be used in combination with either "join" or "order" within the same location; combinations of "join" and "order" within the same location (nested operators) are illegal.

## 3.4.3 Location examples

The following is a list of common location descriptors with their meanings:

Location Description

467 Points to a single base in the presented sequence

021	DDB3 - Home
340565	Points to a continuous range of bases bounded by and including the starting and ending bases
<345500	Indicates that the exact lower boundary point of a feature is unknown. The location begins at some base previous to the first base specified (which need not be contained in the presented sequence) and continues to and includes the ending base
<1888	The feature starts before the first sequenced base and continues to and includes base 888
1>888	The feature starts at the first sequenced base and continues beyond base 888
102.110	Indicates that the exact location is unknown but that it is one of the bases between bases 102 and 110, inclusive
123^124	Points to a site between bases 123 and 124
join(1278,134202)	Regions 12 to 78 and 134 to 202 should be joined to form one contiguous sequence
complement(34126)	Start at the base complementary to 126 and finish at the base complementary to base 34 (the feature is on the strand complementary to the presented strand)
complement(join(269145	71 4019 [162]
complement(join(209145)	Joins regions 2691 to 4571 and 4918 to 5163, then complements the joined segments (the feature is on the strand complementary to the presented strand)
join(complement(4918516	Complement(26914571))  Complements regions 4918 to 5163 and 2691 to 4571, then joins the complemented segments (the feature is on the strand complementary to the presented strand)
J00194.1:100202	Points to bases 100 to 202, inclusive, in the entry (in this database) with primary accession number 'J00194'
join(1100,J00194.1:100	Joins region 1100 of the existing entry with the region 100202 of remote entry J00194

# **4 Feature table Format**

The examples below show the preferred sequence annotations for a number of commonly occurring sequence types. These examples may not be appropriate in all cases but should be used as a guide whenever possible. This section

describes the columnar format used to write this feature table in "flat-file" form for distributions of the database.

## 4.1 Format examples

```
Feature table format example (EMBL):
                  1..1859
    source
                  /db_xref="taxon:3899"
FT
                  /organism="Trifolium repens"
FΤ
FT
                  /tissue_type="leaves"
                  /clone_lib="lambda gt10"
                  /clone="TRE361"
FT
                  /mol_type="genomic DNA"
FΤ
    CDS
                  14..1495
FΤ
                  /db xref="MENDEL:11000"
FT
                  /db_xref="UniProtKB/Swiss-Prot:P26204"
FT
                  /note="non-cyanogenic"
FΤ
                  /EC_number="3.2.1.21"
FT
FΤ
                  /product="beta-glucosidase"
FΤ
                  /protein_id="CAA40058.1"
                  /translation="MDFIVAIFALFVISSFTITSTNAVEASTLLDIGNLSR......
30
                                 40
                                          50
Feature table format example (GenBank):
    source
                  1..8959
                  /organism="Homo sapiens"
                  /db_xref="taxon:9606"
                  /mol_type="genomic DNA"
                  212..8668
    gene
                  /gene="NF1"
    CDS
                  212..8668
                  /gene="NF1"
                  /note="putative"
                  /codon_start=1
                  /product="GAP-related protein"
                  /protein_id="AAA59924.1"
                  /translation="MAAHRPVEWVQAVVSRFDEQLPIKTGQQNTHTKVSTE......
  .-----
                         30
                                          50
                                                   60
                                                            70
       10
                20
                                 40
                                                                    79
Feature table format example (DDBJ):
    source
                  1..2136
                  /clone="pK28"
                  /organism="Rattus norvegicus"
                  /strain="Sprague-Dawley"
                  /tissue_type="kidney"
                  /mol_type="genomic DNA"
```

```
mRNA
                19..2128
   CDS
                31..1212
                /codon_start=1
                /function="Dual specificity protein tyrosine/threonine
                kinase"
                /product="MAP kinase kinase"
                /protein id="BAA02603.1"
                /translation="MPKKKPTPIQLNPAPDGSAVNGTSSAETNLEALQKKL......
-----
             20
                     30
                             40
                                    50
                                            60
                                                   70
                                                           79
     10
```

## 4.2 Definition of line types

The feature table consists of a header line, which contains the column titles for the table, and the individual feature entries. Each feature entry is composed of a feature descriptor line and qualifier and continuation lines, if needed. The feature descriptor line contains the feature's name, key, and location. If the location cannot be contained on the first line of the feature descriptor, it is continued on a continuation line immediately following the descriptor line. If the feature requires further attributes, feature qualifier lines immediately follow the corresponding feature descriptor line (or its continuation). Qualifier information that cannot be contained on one line continues on the following continuation lines as necessary.

Thus, there are 4 types of feature table lines:

Line type	Content	#/entry	#/teature
Header	Column titles	1*	N/A
Feature descriptor	Key and location	1 to many*	1
Feature qualifiers	Qualifiers and values	N/A	0 to many
Continuation lines	Feature descriptor or	0 to many	0 to many
	qualifier continuation		

## 4.3 Data item positions

column position

The position of the data items within the feature descriptor line is as follows:

·	
1-5	blank
6-20	feature key
21	blank
22-80	location

data item

Data on the qualifier and continuation lines begins in column position 22 (the first 21 columns contain blanks). The EMBL format for all lines differs from the GenBank / DDBJ formats that it includes a line type abbreviation in columns 1 and 2.

### 4.4 Use of blanks

Blanks (spaces) may, in general, be used within the feature location and qualifier values to make the construction more readable. The following rules should be observed:

- \* Names of feature table components may not contain blanks (see Section 3.1)
- \* Operator names may not be separated from the following open parenthesis (the beginning of the operand list) by blanks.
- \* Qualifiers may not be separated from the preceding slash or the following equals sign (if one) by blanks

## 5 Examples of sequence annotation

The examples below show the preferred sequence annotations for a number of commonly occurring sequence types. These examples may not be appropriate in all cases but should be used as a guide whenever possible.

## 5.1 Eukaryotic gene

```
source
                1..1509
                /organism="Mus musculus"
                /strain="CD1"
                /mol_type="genomic DNA"
regulatory
                <1..9
                /gene="ubc42"
                /regulatory_class="promoter"
mRNA
                join(10..567,789..1320)
                /gene="ubc42"
CDS
                join(54..567,789..1254)
                /gene="ubc42"
                /product="ubiquitin conjugating enzyme"
                /function="cell division control"
                /translation="MVSSFLLAEYKNLIVNPSEHFKISVNEDNLTEGPPDTLY
                QKIDTVLLSVISLLNEPNPDSPANVDAAKSYRKYLYKEDLESYPMEKSLDECS
                AEDIEYFKNVPVNVLPVPSDDYEDEEMEDGTYILTYDDEDEEEDEEMDDE"
exon
                10..567
                /gene="ubc42"
                /number=1
                568..788
intron
                /gene="ubc42"
                /number=1
                789..1320
exon
                /gene="ubc42"
                /number=2
regulatory
                1310..1317
                /gene="ubc42"
                /regulatory_class="polyA_signal_sequence"
```

### 5.2 Bacterial operon

source 1..9430

/organism="Lactococcus sp."

/strain="MG1234"

/mol\_type="genomic DNA"

operon 160..6865

/operon="gal"

regulatory 160..165

/operon="gal"

/regulatory\_class="minus\_35\_signal"

regulatory 179..184

/operon="gal"

/regulatory\_class="minus\_10\_signal"

CDS 405..1934

/operon="gal"
/gene="galA"

/product="galactose permease"
/function="galactose transporter"

CDS 2003..3001

/operon="gal"
/gene="galM"

/product="aldose 1-epimerase"

/EC\_number="5.1.3.3"
/function="mutarotase"

CDS 3235..4537

/operon="gal"
/gene="galK"

/product="galactokinase"
/EC\_number="2.7.1.6"

mRNA 189..6865

/operon="gal"

## 5.3 Artificial cloning vector (circular)

source 1..5300

/organism="Cloning vector pABC"
/lab\_host="Escherichia coli"

/mol\_type="other DNA"

/focus

source 1..5138

/organism="Escherichia coli"

/mol\_type="other DNA"

/strain="K12"

source 5139..5247

/organism="Aequorea victoria"

/mol\_type="other DNA"
/dev\_stage="adult"

source 5248..5300

/organism="Escherichia coli"

/mol\_type="other DNA"

/strain="K12"

cds join(complement(1..799),complement(5080..5120))
/gene="mob1"

/product="mobilization protein 1"

CDS complement(1697..2512)

/gene="Km"

/product="kanamycin resistance protein"

CDS 3037..3711

/gene="rep1"

/product="replication protein 1"

CDS complement(4170..4829)

/gene="Cm"

/product="chloramphenicol resistance protein"

CDS 5139..5247

/gene="GFP"

/product="green fluorescent protein"

## 5.4 Plasmid

source 1..2245

/organism="Escherichia coli"

/plasmid="Plasmid XYZ"

/strain="K12"

/mol\_type="genomic DNA"

rep\_origin 6

/direction=LEFT

/note="ori"

CDS complement(join(21..349,567..795))

/gene="trbC"

/product="transfer protein C"

CDS 803..1344

/gene="traN"

/product="transfer protein N"

CDS 1559..1985

/gene="incA"

/product="incompatability protein A"

CDS join(2004..2195,3..20)

/gene="finP"

/product="fertility inhibition protein P"

## 5.5 Repeat element

source 1..1011

/organism="Homo sapiens"
/clone="pha281u/1D0"
/mol type="genomic DNA"

repeat\_region 80..401

/rpt\_type=DISPERSED
/rpt\_family="Alu-J"

## 5.6 Immunoglobulin heavy chain

source 1..321

/organism="Mus musculus "

/strain="BALB/c2

/cell\_line="hybridoma 1A4"

/rearranged
/mol\_type="mRNA"

CDS <1..>321

/codon\_start=1

/gene="VFM1-DFL16.1-JH4"

/product="immunoglobulin heavy chain"

V\_region 1..277

/gene="VFM1"

/product="immunoglobulin heavy chain variable region"

## 5.7 T-cell receptor

source 1..402

/organism="Homo sapiens"

/sex="male"

/cell\_type="CD4+ T-lymphocyte"

/rearranged
/clone="TCR1A.12"
/mol\_type="mRNA"

sig\_peptide 1..54

/gene="TCR1A"

CDS 1..402

/gene="TCR1A"

/product="T-cell receptor alpha chain"

mat\_peptide 55..399

/gene="TCR1A"

/product="T-cell receptor alpha chain"

V\_region 55..327

/gene="TCR1A"

J\_segment 328..393

/gene="TCR1A"

C\_region 394..399

/gene="TCR1A"

## 5.8 Transfer RNA

source 1..2345

/organism="Yersinia sp."

/strain="IP134"

/mol\_type="genomic DNA"

regulatory 644..650

/gene="tRNA-Leu(UUR)"

/regulatory\_class="minus\_35\_signal"

tRNA 655..730

/gene="tRNA-Leu(UUR)"

```
/anticodon=(pos:678..680,aa:Leu,seq:taa)
/product="transfer RNA-Leu(UUR)"
```

## 6 Limitations of this feature table design

During the development of the feature table design numerous choices between simplicity and representational power had to be made. In order to create a design which was capable of representing the most common features of biological significance, a certain degree of complexity in the syntax was guaranteed. However, to limit that level of complexity, certain limitations of the design syntax have been accepted.

# 7 Appendices

## 7.1 Appendix I EMBL, GenBank and DDBJ entries

## 7.1.1 EMBL Format

```
X64011; SV 1; linear; genomic DNA; STD; PRO; 756 BP.
XX
AC
     X64011; S78972;
XX
     X64011.1
SV
XX
     28-APR-1992 (Rel. 31, Created)
DT
DT
     30-JUN-1993 (Rel. 36, Last updated, Version 6)
XX
DE
     Listeria ivanovii sod gene for superoxide dismutase
XX
     sod gene; superoxide dismutase.
ΚW
XX
     Listeria ivanovii
0S
     Bacteria; Firmicutes; Bacillus/Clostridium group;
OC.
     Bacillus/Staphylococcus group; Listeria.
OC
XX
RN
     [1]
     MEDLINE; 92140371.
RX
RA
     Haas A., Goebel W.;
     "Cloning of a superoxide dismutase gene from Listeria ivanovii by
RT
RT
     functional complementation in Escherichia coli and characterization of the
RT
     gene product.";
     Mol. Gen. Genet. 231:313-322(1992).
RL
XX
     [2]
RN
     1-756
RP
     Kreft J.;
RA
RT
     Submitted (21-APR-1992) to the EMBL/GenBank/DDBJ databases.
RL
     J. Kreft, Institut f. Mikrobiologie, Universitaet Wuerzburg, Biozentrum Am
RL
     Hubland, 8700 Wuerzburg, FRG
RL
XX
```

```
FΗ
     Key
                     Location/Qualifiers
FΗ
                     1..756
FΤ
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FT
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FΤ
FΤ
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                     /mol_type="genomic DNA"
FΤ
FΤ
     regulatory
                     95..100
FΤ
                     /gene="sod"
FΤ
                     /regulatory_class="ribosome_binding_site"
FΤ
     regulatory
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                     /gene="sod"
FT
                     /regulatory_class="terminator"
FΤ
     CDS
                     109..717
FT
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FΤ
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FΤ
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                     /db_xref="HSSP:P00448"
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FT
FΤ
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FΤ
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FT
FΤ
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FT
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FΤ
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XX
SQ
     Sequence 756 BP; 247 A; 136 C; 151 G; 222 T; 0 other;
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                                                                          60
     gtaatttctt .....
                                                                          120
//
7.1.2 GenBank Format
            LISOD
LOCUS
                                     756 bp
                                               DNA
                                                        linear
                                                                 BCT 30-JUN-1993
DEFINITION Listeria ivanovii sod gene for superoxide dismutase.
           X64011 S78972
ACCESSION
VERSION
            X64011.1 GI:44010
KEYWORDS
            sod gene; superoxide dismutase.
SOURCE
            Listeria ivanovii
 ORGANISM Listeria ivanovii
            Bacteria; Firmicutes; Bacillales; Listeriaceae; Listeria.
REFERENCE
           1 (bases 1 to 756)
 AUTHORS
           Haas,A. and Goebel,W.
 TITLE
            Cloning of a superoxide dismutase gene from Listeria ivanovii by
            functional complementation in Escherichia coli and characterization
            of the gene product
           Mol. Gen. Genet. 231 (2), 313-322 (1992)
  JOURNAL
            92140371
 MEDLINE
```

2 (bases 1 to 756)

REFERENCE

```
AUTHORS
            Kreft,J.
 TITLE
            Direct Submission
            Submitted (21-APR-1992) J. Kreft, Institut f. Mikrobiologie,
  JOURNAL
            Universitaet Wuerzburg, Biozentrum Am Hubland, 8700 Wuerzburg, FRG
FEATURES
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     source
                     1..756
                     /organism="Listeria ivanovii"
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                     /gene="sod"
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                     95..746
     gene
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                     109..717
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                     /protein_id="CAA45406.1"
                     /translation="MTYELPKLPYTYDALEPNFDKETMEIHYTKHHNIYVTKLNEAVS
                     {\sf GHAELASKPGEELVANLDSVPEEIRGAVRNHGGGHANHTLFWSSLSPNGGGAPTGNLK}
                     AAIESEFGTFDEFKEKFNAAAAARFGSGWAWLVVNNGKLEIVSTANODSPLSEGKTPV
                     LGLDVWEHAYYLKFQNRRPEYIDTFWNVINWDERNKRFDAAK"
                     723..746
     regulatory
                     /gene="sod"
                     /regulatory class="terminator"
ORIGIN
        1 cgttatttaa ggtgttacat agttctatgg aaatagggtc tatacctttc gccttacaat
       61 gtaatttctt .....
//
7.1.3 DDBJ Format
LOCUS
                                     756 bp
                                                       linear
                                                                BCT 30-JUN-1993
DEFINITION Listeria ivanovii sod gene for superoxide dismutase.
ACCESSION
          X64011 S78972
VERSION
           X64011.1 GI:44010
KEYWORDS
            sod gene; superoxide dismutase.
SOURCE
            Listeria ivanovii
 ORGANISM Listeria ivanovii
            Bacteria; Firmicutes; Bacillales; Listeriaceae; Listeria.
REFERENCE
           1 (bases 1 to 756)
 AUTHORS
            Haas, A. and Goebel, W.
 TITLE
            Cloning of a superoxide dismutase gene from Listeria ivanovii by
```

```
functional complementation in Escherichia coli and characterization
            of the gene product
           Mol. Gen. Genet. 231 (2), 313-322 (1992)
 JOURNAL
 MEDLINE
           92140371
REFERENCE 2 (bases 1 to 756)
 AUTHORS
           Kreft,J.
 TITLE
           Direct Submission
 JOURNAL
            Submitted (21-APR-1992) J. Kreft, Institut f. Mikrobiologie,
            Universitaet Wuerzburg, Biozentrum Am Hubland, 8700 Wuerzburg, FRG
FEATURES
                     Location/Oualifiers
                     1..756
    source
                     /organism="Listeria ivanovii"
                     /strain="ATCC 19119"
                     /db_xref="taxon:1638"
                     /mol_type="genomic DNA"
    regulatory
                     95..100
                     /gene="sod"
                     /regulatory_class="ribosome_binding_site"
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    gene
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    CDS
                     /gene="sod"
                     /EC_number="1.15.1.1"
                     /codon_start=1
                     /transl_table=11
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                     /db_xref="HSSP:P00448"
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                     /db xref="UniProtKB/Swiss-Prot:P28763"
                     /protein_id="CAA45406.1"
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                     AAIESEFGTFDEFKEKFNAAAAARFGSGWAWLVVNNGKLEIVSTANQDSPLSEGKTPV
                     LGLDVWEHAYYLKFQNRRPEYIDTFWNVINWDERNKRFDAAK"
                     723..746
    regulatory
                     /gene="sod"
                     /regulatory_class="terminator"
BASE COUNT
                    247 a
                                                                 222 t
                                   136 c
                                                  151 g
ORIGIN
       1 cgttatttaa ggtgttacat agttctatgg aaatagggtc tatacctttc gccttacaat
       61 gtaatttctt .....
//
```

### 7.2 Appendix II: Feature keys reference

The following has been organized according to the following format:

Feature Key the feature key name

Definition the definition of the key

Mandatory qualifiers qualifiers required with the key; if there are no

mandatory qualifiers, this field is omitted.

Optional qualifiers optional

optional qualifiers associated with the key

Organism scope valid organisms for the key; if the scope is any

organism, this field is omitted.

Molecule scope valid molecule types; if the scope is any molecule

type, this field is omitted.

References citations of published reports, usually supporting the

feature consensus sequence

Comment comments and clarifications

Abbreviations:

accnum an entry primary accession number

<amino\_acid> abbreviation for amino acid

<base\_range> location descriptor for a simple range of bases
<br/>
<bool> Boolean truth value. Valid values are yes and no

<integer> unsigned integer value

<location> general feature location descriptor

<modified\_base> abbreviation for modified nucleotide base

[number] integer representing number of citation in entry's

reference list

<repeat\_type> value indicating the organization of a repeated

sequence.

"text" any text or character string. Since the string is

delimited by double quotes, double quotes may only appear as part of the string if they appear in pairs.

For example, the sentence:

The "label" qualifier is no longer legal.

would be formatted thus:

"The ""label"" qualifier is no longer legal."

Feature Key assembly\_gap

Definition gap between two components of a genome or transcriptome assembly;

Mandatory qualifiers /estimated\_length=unknown or <integer>

/gap\_type="TYPE"

/linkage\_evidence="TYPE" (Note: Mandatory only if the

/gap\_type is "within scaffold", "repeat within scaffold" or "contamination". If there are multiple types of linkage\_evidence they will appear as multiple

/linkage\_evidence="TYPE" qualifiers. For all other types of assembly\_gap features,

use of the /linkage\_evidence qualifier is invalid.)

Mandatory qualifiers under assembly\_gap feature for transcriptome

shotgun assemblies (TSA):
/estimated\_length=<integer>

/gap\_type="within scaffold" and /linkage\_evidence="TYPE"

where TYPE can not be "unspecified";

Comment the location span of the assembly\_gap feature for an unknown gap has

to be specified by the submitter; the specified gap length has to be

reasonable (less or = 1000) and will be indicated as "n"'s in sequence. However, the value for the estimated\_length of assembly\_gap features within a single (non-CON) transcriptome record must be an integer and can not be "unknown";

```
Feature Key
                      C_region
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
                      /allele="text"
Optional qualifiers
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /product="text"
                      /pseudo
                      /pseudogene="TYPE"
                      /standard_name="text"
Organism scope
                      eukaryotes
                      CDS
Feature Key
Definition
                      coding sequence; sequence of nucleotides that
                      corresponds with the sequence of amino acids in a
                      protein (location includes stop codon);
                      feature includes amino acid conceptual translation.
Optional qualifiers
                      /allele="text"
                      /artificial_location="[artificial_location_value]"
                      /circular_RNA
                      /citation=[number]
                      /codon_start=<1 or 2 or 3>
                      /db_xref="<database>:<identifier>"
                      /EC number="text"
                      /exception="[exception_value]"
                      /experiment="[CATEGORY:]text"
                      /function="text"
                      /gene="text"
                      /gene_synonym="text"
```

/inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE\_BASIS]"

```
/locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /number=unquoted text (single token)
                      /old_locus_tag="text" (single token)
                      /operon="text"
                      /product="text"
                      /protein_id="<identifier>"
                      /pseudo
                      /pseudogene="TYPE"
                      /ribosomal_slippage
                      /standard_name="text"
                      /translation="text"
                      /transl_except=(pos:<location>,aa:<amino_acid>)
                      /transl_table=<integer>
                      /trans_splicing
Comment
                      /codon_start 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 universal genetic code table;
                      genetic code exceptions outside the range of the specified
                      tables is reported in /transl_except qualifier;
                      /protein_id consists of a stable ID portion (from the end of 2018
                      new accessions may be extended to a 3+7 accession format with
                      3 position letters and 7 numbers; existing data before the end of
                      2018 uses a 3+5 format) plus a version number after the decimal point;
                      when the protein sequence encoded by the CDS changes, only the version
                      number of the /protein_id value is incremented; the
                      stable part of the /protein_id remains unchanged and as
                      a result will permanently be associated with a given protein;
```

Feature Key	centromere
Definition	region of biological interest identified as a centromere and which has been experimentally characterized;
Optional qualifiers	<pre>/citation=[number] /db_xref="<database>:<identifier>" /experiment="[CATEGORY:]text" /inference="[CATEGORY:]TYPE[(same species)][:EVIDENCE_BASIS]" /note="text" /standard_name="text"</identifier></database></pre>
Comment	the centromere feature describes the interval of DNA that corresponds to a region where chromatids are held and a kinetochore is formed

```
Feature Key
                      D-loop
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="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
Molecule scope
                      DNA
Feature Key
                      D segment
Definition
                      Diversity segment of immunoglobulin heavy chain, and
                      T-cell receptor beta chain;
                      /allele="text"
Optional qualifiers
                      /citation=[number]
                      /db xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /product="text"
                      /pseudo
                      /pseudogene="TYPE"
                      /standard_name="text"
Organism scope
                      eukaryotes
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="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /EC_number="text"
                      /experiment="[CATEGORY:]text"
                      /function="text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /number=unquoted text (single token)
                      /old_locus_tag="text" (single token)
                      /product="text"
                      /pseudo
                      /pseudogene="TYPE"
                      /standard_name="text"
                      /trans_splicing
Feature Key
                      gap
Definition
                      gap in the sequence
Mandatory qualifiers /estimated_length=unknown or <integer>
Optional qualifiers
                      /experiment="[CATEGORY:]text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /map="text"
                      /note="text"
Comment
                      the location span of the gap feature for an unknown
                      gap is 100 bp, with the 100 bp indicated as 100 "n"'s in
                      the sequence. Where estimated length is indicated by
                      an integer, this is indicated by the same number of
                      "n"'s in the sequence.
                      No upper or lower limit is set on the size of the gap.
Feature Key
                      gene
Definition
                      region of biological interest identified as a gene
                      and for which a name has been assigned;
                      /allele="text"
Optional qualifiers
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /function="text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
```

```
/locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /operon="text"
                      /product="text"
                      /pseudo
                      /pseudogene="TYPE"
                      /phenotype="text"
                      /standard_name="text"
                      /trans_splicing
Comment
                      the gene feature describes the interval of DNA that
                      corresponds to a genetic trait or phenotype; the feature is,
                      by definition, not strictly bound to it's positions at the
                      ends; it is meant to represent a region where the gene is
                      located.
Feature Key
                      iDNA
Definition
                      intervening DNA; DNA which is eliminated through any of
                      several kinds of recombination;
Optional qualifiers
                      /allele="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /function="text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /number=unquoted text (single token)
                      /old_locus_tag="text" (single token)
                      /standard_name="text"
Molecule scope
                      DNA
Comment
                      e.g., in the somatic processing of immunoglobulin genes.
Feature Key
                      intron
                      a segment of DNA that is transcribed, but removed from
Definition
                      within the transcript by splicing together the sequences
                      (exons) on either side of it;
Optional qualifiers
                      /allele="text"
                      /citation=[number]
```

```
/db_xref="<database>:<identifier>"
/experiment="[CATEGORY:]text"
/function="text"
/gene="text"
/gene_synonym="text"
/inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
/locus_tag="text" (single token)
/map="text"
/note="text"
/number=unquoted text (single token)
/old_locus_tag="text" (single token)
/pseudo
/pseudogene="TYPE"
/standard_name="text"
/trans_splicing
J_segment
```

```
Feature Key
Definition
                      joining segment of immunoglobulin light and heavy
                      chains, and T-cell receptor alpha, beta, and gamma
                      chains;
Optional qualifiers
                      /allele="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /product="text"
                      /pseudo
                      /pseudogene="TYPE"
```

/standard\_name="text"

Organism scope eukaryotes

## Feature Key mat\_peptide

Definition mature peptide or protein coding sequence; coding sequence for the mature or final peptide or protein product following post-translational modification; the location does not include the stop codon (unlike the

corresponding CDS);

Optional qualifiers /allele="text"

/citation=[number]

/db\_xref="<database>:<identifier>"

```
/EC_number="text"
                      /experiment="[CATEGORY:]text"
                      /function="text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /product="text"
                      /pseudo
                      /pseudogene="TYPE"
                      /standard_name="text"
Feature Key
                      misc_binding
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="text"
Optional qualifiers
                     /allele="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /function="text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      note that feature key regulatory with /regulatory_class="ribosome_binding_site"
Comment
                      should be used for ribosome binding sites.
Feature Key
                      misc difference
Definition
                      feature sequence is different from that presented
                      in the entry and cannot be described by any other
                      difference key (old_sequence, variation, or modified_base);
Optional qualifiers
                      /allele="text"
                      /citation=[number]
                      /clone="text"
                      /compare=[accession-number.sequence-version]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
```

```
/gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /phenotype="text"
                      /replace="text"
                      /standard_name="text"
Comment
                      the misc_difference feature key should be used to
                      describe variability that arises as a result of
                      genetic manipulation (e.g. site directed mutagenesis);
                      use /replace="" to annotate deletion, e.g.
                      misc_difference 412..433
                                      /replace=""
                      misc_feature
Feature Key
Definition
                      region of biological interest which cannot be described
                      by any other feature key; a new or rare feature;
Optional qualifiers
                      /allele="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /function="text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /number=unquoted text (single token)
                      /old_locus_tag="text" (single token)
                      /phenotype="text"
                      /product="text"
                      /pseudo
                      /pseudogene="TYPE"
                      /standard_name="text"
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
Feature Key
                      misc_recomb
Definition
                      site of any generalized, site-specific or replicative
                      recombination event where there is a breakage and
                      reunion of duplex DNA that cannot be described by other
```

```
recombination keys or qualifiers of source key
                      (/proviral);
Optional qualifiers
                      /allele="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /recombination_class="TYPE"
                      /standard name="text"
Molecule scope
                      DNA
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, polyA_site, ncRNA, rRNA and tRNA);
                      /allele="text"
Optional qualifiers
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /function="text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /operon="text"
                      /product="text"
                      /pseudo
                      /pseudogene="TYPE"
                      /standard_name="text"
                      /trans_splicing
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="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /function="text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /standard_name="text"
Feature Key
                      mobile_element
Definition
                      region of genome containing mobile elements;
Mandatory qualifiers /mobile_element_type="<mobile_element_type>
                      [:<mobile_element_name>]"
                      /allele="text"
Optional qualifiers
                      /citation=[number]
                      /db xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /function="text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /rpt_family="text"
                      /rpt_type=<repeat_type>
                      /standard name="text"
                      modified_base
Feature Key
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=<modified_base>
Optional qualifiers
                      /allele="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /frequency="text"
```

/gene="text"

```
/gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
Comment
                      value is limited to the restricted vocabulary for
                      modified base abbreviations;
Feature Key
                      mRNA
Definition
                      messenger RNA; includes 5'untranslated region (5'UTR),
                      coding sequences (CDS, exon) and 3'untranslated region
                      (3'UTR);
Optional qualifiers
                      /allele="text"
                      /artificial_location="[artificial_location_value]"
                      /circular_RNA
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /function="text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /operon="text"
                      /product="text"
                      /pseudo
                      /pseudogene="TYPE"
                      /standard_name="text"
                      /trans_splicing
                      ncRNA
Feature Key
Definition
                      a non-protein-coding gene, other than ribosomal RNA and
                      transfer RNA, the functional molecule of which is the RNA
                      transcript;
Mandatory qualifiers /ncRNA_class="TYPE"
Optional qualifiers
                      /allele="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /function="text"
                      /gene="text"
```

```
/gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /operon="text"
                      /product="text"
                      /pseudo
                      /pseudogene="TYPE"
                      /standard_name="text"
                      /trans_splicing
Example
                      /ncRNA_class="miRNA"
                      /ncRNA_class="siRNA"
                      /ncRNA_class="scRNA"
Comment
                      the ncRNA feature is not used for ribosomal and transfer
                      RNA annotation, for which the rRNA and tRNA feature keys
                      should be used, respectively;
Feature Key
                      N region
Definition
                      extra nucleotides inserted between rearranged
                      immunoglobulin segments.
Optional qualifiers
                      /allele="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /product="text"
                      /pseudo
                      /pseudogene="TYPE"
                      /standard_name="text"
Organism scope
                      eukaryotes
Feature Key
                      old_sequence
Definition
                      the presented sequence revises a previous version of the
                      sequence at this location;
Mandatory qualifiers /citation=[number]
```

```
/compare=[accession-number.sequence-version]
                      /allele="text"
Optional qualifiers
                      /db xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /replace="text"
Comment
                      /replace="" is used to annotate deletion, e.g.
                      old_sequence 12..15
                                   /replace=""
                      NOTE: This feature key is not valid in entries/records
                      created from 15-Oct-2007.
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="text"
Optional qualifiers
                      /allele="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /function="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /map="text"
                      /note="text"
                      /phenotype="text"
                      /pseudo
                      /pseudogene="TYPE"
                      /standard_name="text"
Feature Key
                      oriT
Definition
                      origin of transfer; region of a DNA molecule where transfer is
                      initiated during the process of conjugation or mobilization
                      /allele="text"
Optional qualifiers
                      /bound_moiety="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
```

```
/direction=value
                      /experiment="[CATEGORY:]text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /rpt_family="text"
                      /rpt_type=<repeat_type>
                      /rpt_unit_range=<base_range>
                      /rpt_unit_seq="text"
                      /standard_name="text"
Molecule Scope
                      DNA
Comment
                      rep_origin should be used for origins of replication;
                      /direction has legal values RIGHT, LEFT and BOTH, however only
                      RIGHT and LEFT 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
Feature Key
                      polyA_site
Definition
                      site on an RNA transcript to which will be added adenine
                      residues by post-transcriptional polyadenylation;
Optional qualifiers
                      /allele="text"
                      /citation=[number]
                      /db xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
Organism scope
                      eukaryotes and eukaryotic viruses
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);
                      /allele="text"
Optional qualifiers
```

```
/citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /function="text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /operon="text"
                      /product="text"
                      /standard_name="text"
                      /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.
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="text"
                      /citation=[number]
                      /db xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /function="text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /operon="text"
                      /standard_name="text"
Feature Key
                      primer_bind
Definition
                      non-covalent primer binding site for initiation of
                      replication, transcription, or reverse transcription;
                      includes site(s) for synthetic e.g., PCR primer elements;
Optional qualifiers
                      /allele="text"
                      /citation=[number]
```

```
/db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /standard_name="text"
                      /PCR_conditions="text"
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; PCR components and reaction times may
                      be stored under the "/PCR_conditions" qualifier;
                      since PCR reactions most often involve pairs of primers,
                      a single primer_bind key may use the order() operator
                      with two locations, or a pair of primer_bind keys may be
                      used.
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="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /function="text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /product="text"
                      /pseudo
                      /pseudogene="TYPE"
                      /standard_name="text"
Feature Key
                      protein_bind
Definition
                      non-covalent protein binding site on nucleic acid;
Mandatory qualifiers /bound_moiety="text"
Optional qualifiers
                      /allele="text"
```

/citation=[number]

/db\_xref="<database>:<identifier>"

```
/experiment="[CATEGORY:]text"
                      /function="text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /operon="text"
                      /standard_name="text"
Comment
                      note that feature key regulatory with /regulatory_class="ribosome_binding_site"
                      should be used for ribosome binding sites.
Feature Key
                      regulatory
Definition
                      any region of sequence that functions in the regulation of
                      transcription, translation, replication or chromatin structure;
Mandatory qualifiers /regulatory class="TYPE"
Optional qualifiers
                     /allele="text"
                      /bound_moiety="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /function="text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /operon="text"
                      /phenotype="text"
                      /pseudo
                      /pseudogene="TYPE"
                      /standard_name="text"
Comment
                      This feature has replaced the following Feature Keys on 15-DEC-2014:
                      enhancer, promoter, CAAT_signal, TATA_signal, -35_signal, -10_signal,
                      RBS, GC_signal, polyA_signal, attenuator, terminator, misc_signal.
Feature Key
                      repeat_region
Definition
                      region of genome containing repeating units;
Optional qualifiers
                      /allele="text"
```

```
/citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /function="text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /rpt_family="text"
                      /rpt_type=<repeat_type>
                      /rpt_unit_range=<base_range>
                      /rpt_unit_seq="text"
                      /satellite="<satellite_type>[:<class>][ <identifier>]"
                      /standard_name="text"
Feature Key
                      rep_origin
Definition
                      origin of replication; starting site for duplication of
                      nucleic acid to give two identical copies;
Optional Qualifiers
                      /allele="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /direction=value
                      /experiment="[CATEGORY:]text"
                      /function="text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /standard_name="text"
                      /direction has valid values: RIGHT, LEFT, or BOTH.
Comment
Feature Key
                      rRNA
Definition
                      mature ribosomal RNA; RNA component of the
                      ribonucleoprotein particle (ribosome) which assembles
                      amino acids into proteins.
Optional qualifiers
                     /allele="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /function="text"
```

```
/gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /operon="text"
                      /product="text"
                      /pseudo
                      /pseudogene="TYPE"
                      /standard_name="text"
Comment
                      rRNA sizes should be annotated with the /product
                      qualifier.
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="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /product="text"
                      /pseudo
                      /pseudogene="TYPE"
                      /standard_name="text"
Organism scope
                      eukaryotes
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 nascent polypeptide to the
                      membrane leader sequence;
Optional qualifiers
                      /allele="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
```

```
/experiment="[CATEGORY:]text"
/function="text"
/gene="text"
/gene_synonym="text"
/inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
/locus_tag="text" (single token)
/map="text"
/note="text"
/old_locus_tag="text" (single token)
/product="text"
/pseudo
/pseudogene="TYPE"
/standard_name="text"
```

### Feature Key

#### source

Definition

identifies the biological source of the specified span of the sequence; this key is mandatory; more than one source key per sequence is allowed; every entry/record will have, as a minimum, either a single source key spanning the entire sequence or multiple source keys, which together, span the entire sequence.

```
Mandatory qualifiers /organism="text"
```

```
/mol_type="genomic DNA", "genomic RNA", "mRNA", "tRNA",

"rRNA", "other RNA", "other DNA", "transcribed RNA",

"viral cRNA", "unassigned DNA", "unassigned RNA"
```

Optional qualifiers

```
/altitude="text"
/bio_material="[<institution-code>:[<collection-code>:]]<material_id>"
/cell line="text"
/cell type="text"
/chromosome="text"
/citation=[number]
/clone="text"
/clone lib="text"
/collected_by="text"
/collection_date="text"
/country="<country_value>[:<region>][, <locality>]"
/cultivar="text"
/culture_collection="<institution-code>:[<collection-code>:]<culture_id>"
/db_xref="<database>:<identifier>"
/dev_stage="text"
/ecotype="text"
/environmental sample
/focus
/germline
/haplogroup="text"
/haplotype="text"
```

/host="text"

```
/identified_by="text"
                      /isolate="text"
                      /isolation_source="text"
                      /lab host="text"
                      /lat_lon="text"
                      /macronuclear
                      /map="text"
                      /mating_type="text"
                      /metagenome_source="text"
                      /note="text"
                      /organelle=<organelle_value>
                      /PCR_primers="[fwd_name: XXX, ]fwd_seq: xxxxx,
                      [rev_name: YYY, ]rev_seq: yyyyy"
                      /plasmid="text"
                      /pop_variant="text"
                      /proviral
                      /rearranged
                      /segment="text"
                      /serotype="text"
                      /serovar="text"
                      /sex="text"
                      /specimen_voucher="[<institution-code>:[<collection-code>:]]<specimen_id>"
                      /strain="text"
                      /sub_clone="text"
                      /submitter_seqid="text"
                      /sub_species="text"
                      /sub strain="text"
                      /tissue_lib="text"
                      /tissue_type="text"
                      /transgenic
                      /type_material="<type-of-type> of <organism name>"
                      /variety="text"
Molecule scope
                      any
                      transgenic sequences must have at least two source feature
Comment
                      keys; in a transgenic sequence the source feature key
                      describing the organism that is the recipient of the DNA
                      must span the entire sequence;
                      see Appendix III /organelle for a list of <organelle_value>
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.
Optional qualifiers
                     /allele="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
```

```
/experiment="[CATEGORY:]text"
                      /function="text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /operon="text"
                      /standard_name="text"
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;
                      /allele="text"
Optional qualifiers
                      /citation=[number]
                      /db xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /standard name="text"
Molecule scope
                      DNA
                      STS location to include primer(s) in primer_bind key or
Comment
                      primers.
                      telomere
Feature Key
Definition
                      region of biological interest identified as a telomere
                      and which has been experimentally characterized;
Optional qualifiers
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"/note="text"
                      /inference="[CATEGORY:]TYPE[(same species)][:EVIDENCE_BASIS]"
                      /note="text"
                      /rpt_type=<repeat_type>
                      /rpt_unit_range=<base_range>
                      /rpt_unit_seq="text"
                      /standard_name="text"
```

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 represent the physical end of the chromosome;

#### 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

```
/allele="text"
/citation=[number]
/db_xref="<database>:<identifier>"
/experiment="[CATEGORY:]text"
/function="text"
/gene="text"
/gene_synonym="text"
/inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
/locus_tag="text" (single token)
/map="text"
/note="text"
/old_locus_tag="text" (single token)
/product="text"
/pseudo
/pseudogene="TYPE"
/standard_name="text"
```

# Feature Key

# transit\_peptide

/tag\_peptide=<base\_range>

/locus\_tag="text" (single token)

Definition

transit peptide coding sequence; coding sequence for an N-terminal domain of a nuclear-encoded organellar protein; this domain is involved in post-translational import of the protein into the organelle;

Optional qualifiers

```
/allele="text"
/citation=[number]
/db_xref="<database>:<identifier>"
/experiment="[CATEGORY:]text"
/function="text"
/gene="text"
/gene_synonym="text"
/inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
```

```
/map="text"
/note="text"
/old_locus_tag="text" (single token)
/product="text"
/pseudo
/pseudogene="TYPE"
/standard_name="text"
```

### 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="text"
/anticodon=(pos:<location>,aa:<amino_acid>,seq:<text>)
/circular_RNA
/citation=[number]
/db_xref="<database>:<identifier>"
/experiment="[CATEGORY:]text"
/function="text"
/gene="text"
/gene_synonym="text"
/inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
/locus_tag="text" (single token)
/map="text"
/note="text"
/old_locus_tag="text" (single token)
/operon="text"
/product="text"
/pseudo
/pseudogene="TYPE"
```

## Feature Key

### unsure

/standard\_name="text"
/trans\_splicing

Definition

a small region of sequenced bases, generally 10 or fewer in its length, which could not be confidently identified. Such a region might contain called bases (A, T, G, or C), or a mixture of called-bases and uncalled-bases ('N'). The unsure feature should not be used when annotating gaps in genome assemblies. Please refer to assembly\_gap feature for gaps within the sequence of an assembled genome. For annotation of gaps in other sequences than assembled genomes use the gap feature.

Optional qualifiers

```
/allele="text"
/citation=[number]
/compare=[accession-number.sequence-version]
/db_xref="<database>:<identifier>"
/experiment="[CATEGORY:]text"
/gene="text"
```

```
/gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /replace="text"
Comment
                      use /replace="" to annotate deletion, e.g.
                      unsure
                                  11..15
                                  /replace=""
Feature Key
                      V_region
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
                      /allele="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /product="text"
                      /pseudo
                      /pseudogene="TYPE"
                      /standard_name="text"
Organism scope
                      eukaryotes
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_region)
                      and the last few amino acids of the leader peptide;
Optional qualifiers
                      /allele="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /gene="text"
                      /gene_synonym="text"
```

```
/inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /product="text"
                      /pseudo
                      /pseudogene="TYPE"
                      /standard_name="text"
Organism scope
                      eukaryotes
Feature Key
                      variation
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);
                      /allele="text"
Optional qualifiers
                      /citation=[number]
                      /compare=[accession-number.sequence-version]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /frequency="text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /phenotype="text"
                      /product="text"
                      /replace="text"
                      /standard name="text"
Comment
                      used to describe alleles, RFLP's, and other naturally occurring
                      mutations and polymorphisms; variability arising as a result
                      of genetic manipulation (e.g. site directed mutagenesis) should
                      be described with the misc_difference feature;
                      use /replace="" to annotate deletion, e.g.
                      variation 4..5
                                  /replace=""
                      3'UTR
Feature Key
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
                     /allele="text"
                      /citation=[number]
                      /db xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /function="text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /standard_name="text"
                      /trans_splicing
                      5'UTR
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 genome (preceding the
                      first initiation codon) that is not translated into a protein;
Optional qualifiers
                     /allele="text"
                      /citation=[number]
                      /db_xref="<database>:<identifier>"
                      /experiment="[CATEGORY:]text"
                      /function="text"
                      /gene="text"
                      /gene_synonym="text"
                      /inference="[CATEGORY:]TYPE[ (same species)][:EVIDENCE_BASIS]"
                      /locus_tag="text" (single token)
                      /map="text"
                      /note="text"
                      /old_locus_tag="text" (single token)
                      /standard name="text"
                      /trans_splicing
```

# 7.3 Appendix III: Summary of qualifiers for feature keys

## 7.3.1 Qualifier List

```
The following is a list of available qualifiers for feature keys and their usage. The information is arranged as follows:
```

```
Qualifier name of qualifier; qualifier requires a value if followed by an equal
```

sign

Definition definition of the qualifier
Value format format of value, if required

Example example of qualifier with value comments, questions and clarifications Comment

Qualifier /allele=

Definition name of the allele for the given gene

Value format "text"

Example /allele="adh1-1"

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.

Qualifier /altitude=

Definition geographical altitude of the location from which the sample

was collected

Value format "text"

Example /altitude="-256 m"

/altitude="330.12 m"

Values indicate altitudes above or below nominal sea level Comment

provided in metres

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 abbreviation

for the amino acid encoded and seq is the sequence of the anticodon

Example /anticodon=(pos:34..36,aa:Phe,seq:aaa)

/anticodon=(pos:join(5,495..496),aa:Leu,seq:taa)

/anticodon=(pos:complement(4156..4158),aa:Gln,seq:ttg)

Qualifier /artificial\_location

Definition indicates that location of the CDS or mRNA is modified to adjust

for the presence of a frameshift or internal stop codon and not

because of biological processing between the regions.

"heterogeneous population sequenced", "low-quality sequence region" Value format

/artificial\_location="heterogeneous population sequenced" Example

/artificial\_location="low-quality sequence region"

expected to be used only for genome-scale annotation. Comment

**Oualifier** /bio material=

Definition identifier for the biological material from which the nucleic

> acid sequenced was obtained, with optional institution code and collection code for the place where it is currently stored.

"[<institution-code>:[<collection-code>:]]<material\_id>" Value format

Example /bio\_material="CGC:CB3912" <- Caenorhabditis stock centre

the bio\_material qualifier should be used to annotate the Comment

identifiers of material in biological collections that are not

appropriate to annotate as either /specimen\_voucher or

/culture\_collection; these include zoos and aquaria, stock centres, seed banks, germplasm repositories and DNA banks; material\_id is mandatory, institution\_code and collection\_code are optional; institution code is mandatory where collection code is present; institution code and collection code are taken from a controlled vocabulary maintained by the INSDC.

Qualifier /bound\_moiety=

Definition name of the molecule/complex that may bind to the

given feature

Value format "text"

Example /bound\_moiety="GAL4"

Comment A single /bound\_moiety qualifier is legal on the "misc\_binding",

"oriT" and "protein\_bind" features.

Qualifier /cell\_line=

Definition cell line from which the sequence was obtained

Value format "text"

Example /cell\_line="MCF7"

Qualifier /cell\_type=

Definition cell type from which the sequence was obtained

Value format "text"

Example /cell\_type="leukocyte"

Qualifier /chromosome=

Definition chromosome (e.g. Chromosome number) from which

the sequence was obtained

Value format "text"

Example /chromosome="1"

Qualifier /circular\_RNA

Definition indicates that exons are out-of-order or overlapping

because this spliced RNA product is a circular RNA (circRNA) created by backsplicing, for example when a downstream exon in the gene is located 5' of an

upstream exon in the RNA product

Value format none

Example /circular\_RNA

Comment should be used on features such as CDS, mRNA, tRNA and other

features that are produced as a result of a backsplicing event. This qualifier should be used only when the splice event is

indicated in the "join" operator,

eg join(101627..101652,102190..102421,73380..73493)

Qualifier /citation=

Definition reference to a citation listed in the entry reference field Value format [integer-number] where integer-number is the number of the

reference as enumerated in the reference field

Example /citation=[3]

Comment used to indicate the citation providing the claim of and/or

evidence for a feature; brackets are used for conformity.

Qualifier /clone=

Definition clone from which the sequence was obtained

Value format "text"

Example /clone="lambda-hIL7.3"

Comment not more than one clone should be specified for a given source

feature; to indicate that the sequence was obtained from multiple clones, multiple source features should be given.

Qualifier /clone\_lib=

Definition clone library from which the sequence was obtained

Value format "text"

Example /clone\_lib="lambda-hIL7"

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
Example /codon\_start=2

Qualifier /collected\_by=

Definition name of persons or institute who collected the specimen

Value format "text"

Example /collected\_by="Dan Janzen"

Qualifier /collection\_date=

Definition The date on which the specimen was collected.

Date/time ranges are supported by providing two collection dates from among

the supported value formats, delimited by a forward-slash character.

Collection times are supported by adding "T", then the hour, minute and

second, after the date.

Collection times must be in Coordinated Universal Time (UTC), otherwise

known as "Zulu Time" (Z).

Value format "DD-Mmm-YYYY", "Mmm-YYYY", "YYYY"

"YYYY-MM-DDThh:mmZ", "YYYY-MM-DDThh:mm:ssZ", "YYYY-MM-DDThhZ", "YYYY-MM-DD", or "YYYY-MM"

Example /collection\_date="21-Oct-1952"

/collection\_date="Oct-1952"

/collection\_date="1952"

/collection\_date="1952-10-21T11:43Z"

/collection\_date="1952-10-21T11Z"

/collection\_date="1952-10-21"

/collection\_date="1952-10"

/collection\_date="21-Oct-1952/15-Feb-1953"

/collection\_date="Oct-1952/Feb-1953"

/collection\_date="1952/1953"

/collection\_date="1952-10-21/1953-02-15"

/collection\_date="1952-10/1953-02"

/collection\_date="1952-10-21T11:43Z/1952-10-21T17:43Z"

/collection\_date="2015-10-11T17:53:03Z"

Comment 'Mmm' represents a three-letter month abbreviation, and can be one of

the following: Jan, Feb, Mar, Apr, May, Jun, Jul, Aug, Sep, Oct, Nov, Dec

'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.

'hh' is a two-digit value representing the hour of the day (00 to 23)

'mm' is a two-digit value representing the minute of the hour (00 to 59)

'ss' is a two-digit value representing the second of the hour (00 to 59)

Within a date range, value formats that make use of 'Mmm' (month abbreviations) cannot be combined with value formats that make use of 'MM' (two-digit month number)

Collection dates that are specified to at least the month, day, and year (DD-Mmm-YYYY or YYYY-MM-DD) are strongly encouraged. If the day and/or month of the collection date are not known, Mmm-YYYY or YYYY-MM or YYYY may be used.

Within a collection date range, the first date (possibly including time) must be prior to the second date (possibly including time).

Within a collection date range for which the day, month, and year are identical, the first time value must be prior to the second time value.

## Qualifier /compare=

Definition Reference details of an existing public INSD entry

to which a comparison is made

Value format [accession-number.sequence-version]

Example /compare=AJ634337.1

Comment This qualifier may be used on the following features:

misc difference, unsure, old sequence and variation.

The feature "old\_sequence" must have either a

/citation or a /compare qualifier. 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.

## Qualifier /country=

Definition locality of isolation of the sequenced sample indicated in

terms of political names for nations, oceans or seas, followed

by regions and localities

Value format "<country\_value>[:<region>][, <locality>]" where

country\_value is any value from the controlled vocabulary at http://www.insdc.org/documents/country-qualifier-vocabulary

Example /country="Canada:Vancouver"

/country="France:Cote d'Azur, Antibes"

/country="Atlantic Ocean:Charlie Gibbs Fracture Zone"

> organism was isolated or sampled. Regions and localities should be indicated where possible. Note that the physical geography of

the isolation or sampling site should be represented in /isolation source.

Qualifier /cultivar=

Definition cultivar (cultivated variety) of plant from which sequence was

obtained.

Value format "text"

Example /cultivar="Nipponbare"

/cultivar="Tenuifolius"
/cultivar="Candy Cane"

/cultivar="IR36"

Comment 'cultivar' is applied solely to products of artificial

selection; use the variety qualifier for natural, named

plant and fungal varieties;

Qualifier /culture\_collection=

Definition institution code and identifier for the culture from which the

nucleic acid sequenced was obtained, with optional collection

code.

Value format "<institution-code>:[<collection-code>:]<culture\_id>"

Example /culture\_collection="ATCC:26370"

Comment the /culture\_collection qualifier should be used to annotate

live microbial and viral cultures, and cell lines that have been deposited in curated culture collections; microbial cultures in personal or laboratory collections should be annotated in strain

qualifiers;

annotation with a culture\_collection qualifier implies that the sequence was obtained from a sample retrieved (by the submitter or a collaborator) from the indicated culture collection, or that the sequence was obtained from a sample that was deposited (by the submitter or a collaborator) in the indicated culture collection; annotation with more than one culture\_collection qualifier indicates that the sequence was obtained from a sample that was deposited (by the submitter or a collaborator) in more

than one culture collection.

culture\_id and institution\_code are mandatory, collection\_code
is optional; institution code and collection code are taken

from a controlled vocabulary maintained by the INSDC.

http://www.insdc.org/controlled-vocabulary-culturecollection-qualifier

Qualifier /db\_xref=

Definition database cross-reference: pointer to related information in

another database.

Value format "<database:identifier>" where database is

the name of the database containing related information, and identifier is the internal identifier of the related information according to the naming conventions of the cross-referenced

database.

Example /db\_xref="UniProtKB/Swiss-Prot:P28763"

Comment the complete list of allowed database types is kept at

http://www.insdc.org/db\_xref.html

Qualifier /dev\_stage=

Definition if the sequence was obtained from an organism in a specific

developmental stage, it is specified with this qualifier

Value format "text"

Example /dev\_stage="fourth instar larva"

Qualifier /direction=

Definition direction of DNA replication

Value format left, right, or both where left indicates toward the 5' end of

the entry sequence (as presented) and right indicates toward

the 3' end

Example /direction=LEFT

Qualifier /EC\_number=

Definition Enzyme Commission number for enzyme product of sequence

Value format "text"

Example /EC\_number="1.1.2.4"

/EC\_number="1.1.2.-"
/EC\_number="1.1.2.n"
/EC number="1.1.2.n1"

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 can

be replaced by dash "." to indicate uncertain assignment. Symbols including an "n", e.g. "n", "n1" and so on, can 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.

Qualifier /ecotype=

Definition a population within a given species displaying genetically

based, phenotypic traits that reflect adaptation to a local habitat.

Value Format "text"

Example /ecotype="Columbia"

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.

Qualifier /environmental sample

Definition identifies sequences derived by direct molecular

isolation from a bulk environmental DNA sample (by PCR with or without subsequent cloning of the product, DGGE, 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

Example /environmental\_sample

Comment used only with the source feature key; source feature

> keys containing the /environmental\_sample qualifier should also contain the /isolation\_source qualifier. entries including /environmental\_sample must not include

the /strain qualifier

Qualifier /estimated\_length=

estimated length of the gap in the sequence Definition

Value format unknown or <integer> Example /estimated\_length=unknown /estimated\_length=342

**Qualifier** /exception=

Definition indicates that the coding region cannot be translated using

standard biological rules

Value format "RNA editing", "reasons given in citation",

"rearrangement required for product",

"annotated by transcript or proteomic data"

Example /exception="RNA editing"

/exception="reasons given in citation"

/exception="rearrangement required for product"

/exception="annotated by transcript or proteomic data"

Comment only to be used to describe biological mechanisms such

as RNA editing; where the exception cannot easily be described a published citation must be referred to; protein translation of /exception CDS will be different from the according conceptual translation;

- An /inference qualifier should accompany any use of /exception="annotated by transcript or proteomic data", to provide support for the existence of the transcript/protein.

- must not be used where transl\_except would be adequate,

e.g. in case of stop codon completion use:

/transl\_except=(pos:6883,aa:TERM)

/note="TAA stop codon is completed by addition of 3' A residues to mRNA".

- must not be used for ribosomal slippage, instead use join operator,

e.g.: CDS join(486..1784,1787..4810)

/note="ribosomal slip on tttt sequence at 1784..1787"

Qualifier /experiment=

Definition a brief description of the nature of the experimental

evidence that supports the feature identification or assignment.

Value format "[CATEGORY:]text"

where CATEGORY is one of the following:

"COORDINATES" support for the annotated coordinates

"DESCRIPTION" support for a broad concept of function such as that based on phenotype, genetic approach, biochemical function, pathway

information, etc.

"EXISTENCE" support for the known or inferred existence of the product

where text is free text (see examples)

Example /experiment="5' RACE"

/experiment="Northern blot [DOI: 12.3456/FT.789.1.234-567.2010]" /experiment="heterologous expression system of Xenopus laevis

oocytes [PMID: 12345678, 10101010, 987654]" /experiment="COORDINATES: 5' and 3' RACE"

Comment detailed experimental details should not be included, and would

normally be found in the cited publications; PMID, DOI and any experimental database ID is allowed to be used in /experiment qualifier; value "experimental evidence, no additional details recorded" was used to replace instances of /evidence=EXPERIMENTAL in

December 2005

Qualifier /focus

Definition identifies the source feature of primary biological

interest for records that have multiple source features originating from different organisms and that are not

transgenic.

Value format none Example /focus

Comment the source feature carrying the /focus qualifier

identifies the main organism of the entry, this determines: a) the name displayed in the organism lines, b) if no translation table is specified, the translation table, c) the DDBJ/EMBL/GenBank taxonomic division in which the entry will appear; only one source feature with /focus is allowed in an entry; the /focus and /transgenic qualifiers are mutually exclusive

in an entry.

Qualifier /frequency=

Definition frequency of the occurrence of a feature

Value format text representing the proportion of a population carrying the

feature expressed as a fraction

Example /frequency="23/108"

/frequency="1 in 12"
/frequency=".85"

Qualifier /function=

Definition function attributed to a sequence

Value format "text"

Example function="essential for recognition of cofactor"

Comment /function is used when the gene name and/or product name do not

convey the function attributable to a sequence.

Qualifier /gap\_type=

Definition type of gap connecting components in records of a genome assembly,

or the type of biological gap in a record that is part of a genome

assembly;

Value format "between scaffolds", "within scaffold", "telomere", "centromere",

"short arm", "heterochromatin", "repeat within scaffold",

"repeat between scaffolds", "contamination", "unknown"

Example /gap\_type="between scaffolds"

/gap\_type="within scaffold"

Comment This qualifier is used only for assembly\_gap features and its values

are controlled by the AGP Specification version 2.1:

https://www.ncbi.nlm.nih.gov/assembly/agp/AGP\_Specification/

Please also visit: http://www.insdc.org/controlled-vocabulary-gaptype-qualifier

Qualifier /gene=

Definition symbol of the gene corresponding to a sequence region

Value format "text"
Example /gene="ilvE"

Qualifier /gene\_synonym=

Definition synonymous, replaced, obsolete or former gene symbol

Value format "text"

Example /gene\_synonym="Hox-3.3"

in a feature where /gene="Hoxc6"

Comment used where it is helpful to indicate a gene symbol synonym;

when used, a primary gene symbol must always be indicated

in /gene or a /locus\_tag must be used.

Qualifier /germline

Definition the sequence presented in the entry 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 Example /germline

Comment /germline should not be used to indicate that the source of

the sequence is a gamete or germ cell;

/germline and /rearranged cannot be used in the same source

feature;

/germline and /rearranged should 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 should not be used outside of the

Craniata (taxid=89593)

Qualifier /haplogroup=

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

"text"

Example

/haplogroup="H\*"

Qualifier

#### /haplotype=

Definition

name for a combination 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

"text"

Example

/haplotype="Dw3 B5 Cw1 A1"

Qualifier

#### /host=

Definition

natural (as opposed to laboratory) host to the organism from

which sequenced molecule was obtained

Value format

"text"

Example

/host="Homo sapiens"

/host="Homo sapiens 12 year old girl"

/host="Rhizobium NGR234"

Qualifier

#### /identified\_by=

Definition

name of the expert who identified the specimen taxonomically

Value format

"text"

Example

/identified\_by="John Burns"

#### Qualifier

#### /inference=

Definition

a structured description of non-experimental evidence that supports  $% \left( 1\right) =\left( 1\right) \left( 1\right)$ 

the feature identification or assignment.

Value format

"[CATEGORY:]TYPE[ (same species)][:EVIDENCE\_BASIS]"

where CATEGORY is one of the following:

"COORDINATES" support for the annotated coordinates

"DESCRIPTION" support for a broad concept of function such as that based on phenotype, genetic approach, biochemical function, pathway information, etc.

"EXISTENCE" support for the known or inferred existence of the product

where TYPE is one of the following:

"non-experimental evidence, no additional details recorded"

"similar to sequence"

"similar to AA sequence"

"similar to DNA sequence"

"similar to RNA sequence"

"similar to RNA sequence, mRNA"

"similar to RNA sequence, EST"

"similar to RNA sequence, other RNA"

"profile"

"nucleotide motif"

"protein motif"

"ab initio prediction"

"alignment"

where the optional text "(same species)" is included when the inference comes from the same species as the entry.

where the optional "EVIDENCE\_BASIS" is either a reference to a database entry (including accession and version) or an algorithm (including version), eg 'INSD:AACN010222672.1', 'InterPro:IPR001900',

'ProDom:PD000600', 'Genscan:2.0', etc. and is structured "[ALGORITHM][:EVIDENCE\_DBREF[,EVIDENCE\_DBREF]\*[,...]]"

Example /inference="COORDINATES:profile:tRNAscan:2.1"

/inference="similar to DNA sequence:INSD:AY411252.1"

/inference="similar to RNA sequence, mRNA:RefSeq:NM\_000041.2"

/inference="similar to DNA sequence (same

species):INSD:AACN010222672.1"

/inference="protein motif:InterPro:IPR001900"
/inference="ab initio prediction:Genscan:2.0"

/inference="alignment:Splign:1.0"

/inference="alignment:Splign:1.26p:RefSeq:NM\_000041.2,INSD:BC003557.1"

Comment /inference="non-experimental evidence, no additional details

recorded" was used to replace instances of

/evidence=NOT\_EXPERIMENTAL in December 2005; any database ID can be used in /inference= qualifier; recommendations for choice of resource acronym for [EVIDENCE\_BASIS] are provided in the /inference qualifier vocabulary recommendation document (http://www.insdc.org/inference.html);

Qualifier /isolate=

Definition individual isolate from which the sequence was obtained

Value format "text"

Example /isolate="Patient #152"

/isolate="DGGE band PSBAC-13"

Qualifier /isolation\_source=

Definition describes the physical, environmental and/or local

geographical source of the biological sample from which

the sequence was derived

Value format "text"

Examples /isolation\_source="rumen isolates from standard

Pelleted ration-fed steer #67"

/isolation\_source="permanent Antarctic sea ice"

/isolation\_source="denitrifying activated sludge from

carbon\_limited continuous reactor"

Comment used only with the source feature key;

source feature keys containing an /environmental\_sample qualifier should also contain an /isolation\_source qualifier; the /country qualifier should be used to describe the country and major geographical sub-region.

Qualifier /lab\_host=

Definition scientific name of the laboratory host used to propagate the

source organism from which the sequenced molecule was obtained

Value format "text"

Example /lab\_host="Gallus gallus"

/lab\_host="Gallus gallus embryo"

/lab host="Escherichia coli strain DH5 alpha"

/lab\_host="Homo sapiens HeLa cells"

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

Qualifier /lat\_lon=

Definition geographical coordinates of the location where the specimen was

collected

Value format "text"

Example /lat\_lon="47.94 N 28.12 W"

/lat\_lon="45.0123 S 4.1234 E"

Comment degrees latitude and longitude in format

"d[d.dddd] N|S d[dd.dddd] W|E" (see the examples)

Qualifier /linkage\_evidence=

Definition type of evidence establishing linkage across an assembly\_gap.

Only allowed to be used with assembly\_gap features that have

a /gap\_type value of "within scaffold", "repeat within scaffold" or "contamination";

Please note if /gap\_type="contamination", /linkage\_evidence must be used and the value of

/linkage\_evidence must be "unspecified".

Value format "pcr", "paired-ends", "align genus", "align xgenus", "align trnscpt",

"within clone", "clone contig", "map", "strobe", "proximity ligation",

"unspecified"

Example /linkage\_evidence="paired-ends"

/linkage\_evidence="within clone"

Comment This qualifier is used only for assembly\_gap features and its values are

controlled by the AGP Specification version 2.1:

https://www.ncbi.nlm.nih.gov/assembly/agp/AGP\_Specification/

Please also visit: http://www.insdc.org/controlled-vocabulary-linkageevidence-qualifier

Qualifier /locus\_tag=

Definition a submitter-supplied, systematic, stable identifier for a gene

and its associated features, used for tracking purposes

Value Format "text"(single token)

but not "<1-5 letters><5-9 digit integer>[.<integer>]"

Example /locus\_tag="ABC\_0022"

/locus\_tag="A1C\_00001"

Comment /locus\_tag can be used with any feature that /gene can be used with;

identical /locus\_tag values may be used within an entry/record,

but only if the identical /locus\_tag values are associated with the same gene; in all other circumstances the /locus\_tag

value must be unique within that entry/record. Multiple /locus\_tag

values are not allowed within one feature for entries created

after 15-0CT-2004.

If a /locus\_tag needs to be re-assigned the /old\_locus\_tag qualifier  $\,$ 

should be used to store the old value. The /locus\_tag value should

not be in a format which resembles INSD accession numbers, accession.version, or /protein\_id identifiers.

Qualifier /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

Example /macronuclear

Qualifier /map=

Definition genomic map position of feature

Value format "text"

Example /map="8q12-q13"

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 "text"

Examples /mating\_type="MAT-1"

/mating\_type="plus"
/mating\_type="-"
/mating\_type="odd"
/mating\_type="even"

Comment /mating\_type="male" and /mating\_type="female" are

valid in the prokaryotes, but not in the eukaryotes;

for more information, see the entry for /sex.

Qualifier /metagenome\_source=

Definition sequences from a Metagenome Assembled Genome (MAG), i.e a single-taxon assembly

drawn from a binned metagenome, are specified with this qualifier to indicate

that the assembly is derived from a metagenomic source, rather than from an isolated organism. Where this qualifier is present it must contain the word "metagenome" and must exist in the

NCBI taxonomy database: https://www.ncbi.nlm.nih.gov/Taxonomy/

Value format "text"

Examples /metagenome\_source="human gut metagenome"

/metagenome\_source="soil metagenome"

Comment the qualifier /metagenome source is mandatory when a single-taxon sequence is derived from

a metagenome; sequences with a /metagenome\_source require also an /environmental\_sample qualifier.

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", "MITE", "LINE", "other".

Example /mobile\_element\_type="transposon:Tnp9"

Comment /mobile\_element\_type is legal 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" requires a mobile\_element\_name.

Qualifier /mod\_base=

Definition abbreviation for a modified nucleotide base

Value format modified\_base Example /mod\_base=m5c

Comment modified nucleotides not found in the restricted vocabulary

list can be annotated by entering '/mod\_base=OTHER' with

'/note="name of modified base"'

Qualifier /mol\_type=

Definition in vivo molecule type of sequence

Value format "genomic DNA", "genomic RNA", "mRNA", "tRNA", "rRNA",

"other RNA", "other DNA", "transcribed RNA", "viral cRNA",

"unassigned DNA", "unassigned RNA"

Example /mol\_type="genomic DNA"

Comment all values refer to the in vivo or synthetic molecule for

primary entries and the hypothetical molecule in Third Party
Annotation entries; the value "genomic DNA" does not imply that
the molecule is nuclear (e.g. organelle and plasmid DNA should
be described using "genomic DNA"); ribosomal RNA genes should be
described using "genomic DNA"; "rRNA" should only be used if the
ribosomal RNA molecule itself has been sequenced; /mol\_type is
mandatory on every source feature key; all /mol\_type values
within one entry/record must be the same; values "other RNA" and
"other DNA" should be applied to synthetic molecules, values
"unassigned DNA", "unassigned RNA" should be applied where in

vivo molecule is unknown

Please also visit:

http://www.insdc.org/controlled-vocabulary-moltype-qualifier

Qualifier /ncRNA\_class=

Definition a structured description of the classification of the

non-coding RNA described by the ncRNA parent key

Value format "TYPE"

Example /ncRNA\_class="miRNA"

/ncRNA\_class="siRNA"
/ncRNA\_class="scRNA"

Comment TYPE is a term taken from the INSDC controlled vocabulary for ncRNA

classes. For a complete list of supported values, please see:

http://www.insdc.org/documents/ncrna-vocabulary;

ncRNA classes not yet in the INSDC /ncRNA\_class controlled

vocabulary can be annotated by entering

'/ncRNA\_class="other"' with either '/product="[name of the product]"' OR

'/note="[brief explanation of novel ncRNA\_class]"';

Qualifier /note= Definition any comment or additional information Value format "text" Example /note="This qualifier is equivalent to a comment." Qualifier /number= Definition a number to indicate the order of genetic elements (e.g., exons or introns) in the 5' to 3' direction Value format unquoted text (single token) Example /number=4 /number=6B Comment text limited to integers, letters or combination of integers and/or letters represented as an unquoted single token (e.g. 5a, XIIb); any additional terms should be included in /standard\_name.

Example: /number=2A

/standard\_name="long"

Qualifier /old\_locus\_tag=

Definition feature tag assigned for tracking purposes

Value Format "text" (single token)

Example /old\_locus\_tag="RSc0382"

/locus\_tag="YP00002"

Comment /old\_locus\_tag can be used with any feature where /gene is valid and

where a /locus\_tag qualifier is present.

Identical /old\_locus\_tag values may be used within an entry/record, but only if the identical /old\_locus\_tag values are associated with the same gene; in all other circumstances the /old\_locus\_tag

value must be unique within that entry/record.

Multiple/old\_locus\_tag qualifiers with distinct values are allowed within a single feature; /old\_locus\_tag and /locus\_tag

values must not be identical within a single feature.

Qualifier /operon=

Definition name of the group of contiguous genes transcribed into a

single transcript to which that feature belongs.

Value format "text"
Example /operon="lac"

Qualifier /organelle=

Definition type of membrane-bound intracellular structure from which the

sequence was obtained

Value format chromatophore, hydrogenosome, mitochondrion, nucleomorph,

plastid, mitochondrion:kinetoplast, plastid:chloroplast,
plastid:apicoplast, plastid:chromoplast, plastid:cyanelle,

plastid:leucoplast, plastid:proplastid

Examples /organelle="chromatophore"

/organelle="hydrogenosome"
/organelle="mitochondrion"
/organelle="nucleomorph"
/organelle="plastid"

/organelle="mitochondrion:kinetoplast"

/organelle="plastid:chloroplast"
/organelle="plastid:apicoplast"
/organelle="plastid:chromoplast"
/organelle="plastid:cyanelle"
/organelle="plastid:leucoplast"
/organelle="plastid:proplastid"

Comments modifier text limited to values from controlled list

Please also visit: http://www.insdc.org/controlled-vocabulary-organelle-qualifier

Qualifier /organism=

Definition scientific name or higher-level classification of the

organism or agent that provided the sequenced genetic

material.

Value format "text"

Examples /organism="Homo sapiens"

/organism="Lactobacillaceae bacterium"

/organism="West Nile virus"
/organism="synthetic construct"
/organism="uncultured bacterium"

Comment includes names for Prokaryotes, Eukaryotes, Viruses, synthetic

sequences, uncultured samples, and unclassified organisms. the organism name which appears on the OS or ORGANISM line will match the value of the /organism qualifier of the source key in the simplest case of a one-source sequence.

Qualifier /partial

Definition differentiates between complete regions and partial ones

Value format none Example /partial

Comment not to be used for new entries from 15-DEC-2001;

use '<' and '>' signs in the location descriptors to

indicate that the sequence is partial.

Qualifier /PCR\_conditions=

Definition description of reaction conditions and components for PCR

Value format "text"

Example /PCR\_conditions="Initial denaturation:94degC,1.5min"

Comment used with primer\_bind key

Qualifier /PCR primers=

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 /PCR\_primers="[fwd\_name: XXX1, ]fwd\_seq: xxxxx1,[fwd\_name: XXX2,]

fwd\_seq: xxxxx2, [rev\_name: YYY1, ]rev\_seq: yyyyy1,

[rev\_name: YYY2, ]rev\_seq: yyyyy2"

Example /PCR\_primers="fwd\_name: CO1P1, fwd\_seq: ttgattttttggtcayccwgaagt,

rev\_name: CO1R4, rev\_seq: ccwvytardcctarraartgttg"

/PCR\_primers=" fwd\_name: hoge1, fwd\_seq: cgkgtgtatcttact,

rev\_name: hoge2, rev\_seq: cg<i>gtgtatcttact"

/PCR\_primers="fwd\_name: CO1P1, fwd\_seq: ttgattttttggtcayccwgaagt, fwd\_name: CO1P2, fwd\_seq: gatacacaggtcayccwgaagt, rev\_name: CO1R4,

rev\_seq: ccwvytardcctarraartgttg"

Comment fwd seg and rev seg are both mandatory; fwd name and rev name are

both optional. Both sequences should be presented in 5'>3' order. The sequences should be given in the IUPAC degenerate-base alphabet, except for the modified bases; those must be enclosed within angle

brackets <>

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 "text"

Example /phenotype="erythromycin resistance"

Qualifier /plasmid=

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

Value format "text"

Example /plasmid="C-589"

Qualifier /pop\_variant=

Definition name of subpopulation or phenotype of the sample from which the sequence

was derived

Value format "text"

Example /pop\_variant="pop1"

/pop variant="Bear Paw"

Qualifier /product=

Definition name of the product associated with the feature, e.g. the mRNA of an

mRNA feature, the polypeptide of a CDS, the mature peptide of a

mat\_peptide, etc.

Value format "text"

Example /product="trypsinogen" (when qualifier appears in CDS feature)

/product="trypsin" (when qualifier appears in mat\_peptide feature)
/product="XYZ neural-specific transcript" (when qualifier appears in

mRNA feature)

Qualifier /protein\_id=

Definition protein identifier, issued by International collaborators.

this qualifier consists of a stable ID portion (accessioned data before the end of 2018 uses a 3+5 format; from the end of 2018 new accessions may be extended to a 3+7 accession format with 3 position letters and 7 numbers) plus a version number after

the decimal point.

Value format <identifier>

Example /protein\_id="AAA12345.1"

/protein\_id="AAA1234567.1"

Comment when the protein sequence encoded by the CDS changes, only

the version number of the /protein\_id value is incremented; the stable part of the /protein\_id remains unchanged and as a result will permanently be associated with a given protein; this qualifier is valid only on CDS features which translate

into a valid protein.

Qualifier /proviral

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
Example /proviral

Qualifier /pseudo

Definition indicates that this feature is a non-functional version of the

element named by the feature key

Value format none
Example /pseudo

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 pseudogenisation events.

Other reasons may include sequencing or assembly errors.

In order to annotate pseudogenes the qualifier /pseudogene= must be

used indicating the TYPE which can be taken from the INSDC controlled vocabulary

for pseudogenes.

Qualifier /pseudogene=

Definition indicates that this feature is a pseudogene of the element named

by the feature key

Value format "TYPE"

where TYPE is one of the following:

processed, unprocessed, unitary, allelic, unknown

Example /pseudogene="processed"

/pseudogene="unprocessed"
/pseudogene="unitary"
/pseudogene="allelic"
/pseudogene="unknown"

Comment TYPE is a term taken from the INSDC controlled vocabulary for pseudogenes

( http://www.insdc.org/documents/pseudogene-qualifier-vocabulary ):

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 mutation. 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: the submitter does not know the method of pseudogenisation.

## 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

Example /rearranged

Comment /rearranged should not be used to annotate chromosome

rearrangements that are not involved in an adaptive immune

response;

/germline and /rearranged cannot be used in the same source

feature;

/germline and /rearranged should 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 should not be used outside of the

Craniata (taxid=89593)

## Qualifier /recombination\_class

Definition a structured description of the classification of recombination

hotspot region within a sequence

Value format "TYPE"

Example /recombination\_class="meiotic"

/recombination\_class="chromosome\_breakpoint"

Comment TYPE is a term taken from the INSDC controlled vocabulary for recombination

classes ( http://www.insdc.org/controlled-vocabulary-recombination-class );

in DEC 2017, the following terms were valid:

"meiotic"

"mitotic"

"non\_allelic\_homologous"

"chromosome\_breakpoint"

"other"

recombination classes not yet in the INSDC /recombination\_class controlled

vocabulary can be annotated by entering /recombination\_class="other" with
/note="[brief explanation of novel /recombination\_class]";

Qualifier /regulatory\_class

Definition a structured description of the classification of transcriptional,

translational, replicational and chromatin structure related

regulatory elements in a sequence

Value format "TYPE"

Example /regulatory\_class="promoter"

/regulatory\_class="enhancer"

/regulatory\_class="ribosome\_binding\_site"

Comment TYPE is a term taken from the INSDC controlled vocabulary for regulatory

classes. For a complete list of supported values, please see:
http://www.insdc.org/controlled-vocabulary-regulatoryclass;

regulatory classes not yet in the INSDC /regulatory\_class controlled vocabulary can be annotated by entering /regulatory\_class="other" with /note="[brief explanation of novel regulatory\_class]";

Qualifier /replace=

Definition indicates that the sequence identified a feature's intervals is

replaced by the sequence shown in "text"; if no sequence is contained within the qualifier, this indicates a deletion.

Value format "text"

Example /replace="a"

/replace=""

Qualifier /ribosomal slippage

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

Example /ribosomal\_slippage

Comment a join operator, e.g.: [join(486..1784,1787..4810)] should be used

in the CDS spans to indicate the location of  $ribosomal\_slippage$ 

Qualifier /rpt family=

Definition type of repeated sequence; "Alu" or "Kpn", for example

Value format "text"

Example /rpt\_family="Alu"

Qualifier /rpt\_type=

Definition structure and distribution of repeated sequence

Value format tandem, direct, inverted, flanking, nested, dispersed, terminal,

long\_terminal\_repeat, non\_ltr\_retrotransposon\_polymeric\_tract,

centromeric\_repeat, telomeric\_repeat, x\_element\_combinatorial\_repeat,

y\_prime\_element and other

Example /rpt\_type=INVERTED

Comment the values are case-insensitive, i.e. both "INVERTED" and "inverted"

are valid; For the most current list of allowed values and their definitions please visit: http://www.insdc.org/controlled-vocabulary-rpttype-qualifier

Qualifier /rpt\_unit\_range=

Definition identity of a repeat range

Value format <base\_range>

Example /rpt\_unit\_range=202..245

Comment used to indicate the base range of the sequence that constitutes

a repeated sequence specified by the feature keys oriT and repeat\_region; qualifiers /rpt\_unit\_range and /rpt\_unit\_seq

replaced qualifier /rpt\_unit in December 2005

Qualifier /rpt\_unit\_seq=

Definition identity of a repeat sequence

Value format "text"

Example /rpt\_unit\_seq="aagggc"

/rpt\_unit\_seq="ag(5)tg(8)"

/rpt\_unit\_seq="(AAAGA)6(AAAA)1(AAAGA)12"

Comment used to indicate the literal sequence that constitutes a

repeated sequence specified by the feature keys oriT and repeat\_region; qualifiers /rpt\_unit\_range and /rpt\_unit\_seq

replaced qualifier /rpt\_unit in December 2005

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 "<satellite\_type>[:<class>][ <identifier>]"

where satellite\_type is one of the following

"satellite", "microsatellite", "minisatellite"

Example /satellite="satellite: S1a"

/satellite="satellite: alpha"
/satellite="satellite: gamma III"
/satellite="microsatellite: DC130"

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.

Please also visit: http://www.insdc.org/controlled-vocabulary-satellite-qualifier

Qualifier /segment=

Definition name of viral or phage segment sequenced

Value format "text"
Example /segment="6"

Qualifier /serotype=

Definition serological variety of a species characterized by its

antigenic properties

Value format "text"

Example /serotype="B1"

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".

Qualifier /serovar=

Definition serological variety of a species (usually a prokaryote)

characterized by its antigenic properties

Value format "text"

Example /serovar="0157:H7"

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".

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 "text"

Examples /sex="female"

/sex="male"

/sex="hermaphrodite"
/sex="unisexual"
/sex="bisexual"
/sex="asexual"

/sex="monoecious" [or monecious]
/sex="dioecious" [or diecious]

Comment /sex should be used (instead of /mating\_type)

in the Metazoa, Embryophyta, Rhodophyta & Phaeophyceae;

/mating\_type should be used (instead of /sex)

in the Bacteria, Archaea & Fungi;

neither /sex nor /mating\_type should be used

in the viruses;

outside of the taxa listed above, /mating\_type should be used unless the value of the qualifier is taken from the vocabulary given in the examples

above

Qualifier /specimen\_voucher=

Definition identifier for the specimen from which the nucleic acid

sequenced was obtained

Value format /specimen\_voucher="[<institution-code>:[<collection-code>:]]<specimen\_id>"

Example /specimen\_voucher="UAM:Mamm:52179"

/specimen\_voucher="AMCC:101706"

/specimen voucher="USNM:field series 8798"

/specimen\_voucher="personal:Dan Janzen:99-SRNP-2003"

/specimen voucher="99-SRNP-2003"

Comment the /specimen\_voucher qualifier is intended to annotate a

reference to the physical specimen that remains after the

sequence has been obtained;

if the specimen was destroyed in the process of sequencing, electronic images (e-vouchers) are an adequate substitute for a physical voucher specimen; ideally the specimens will be deposited in a curated museum, herbarium, or frozen tissue collection, but often they will remain in a personal or

laboratory collection for some time before they are deposited in

a curated collection;

there are three forms of specimen\_voucher qualifiers; if the text of the qualifier includes one or more colons it is a

'structured voucher'; structured vouchers include

institution-codes (and optional collection-codes) taken from a controlled vocabulary maintained by the INSDC that denotes the museum or herbarium collection where the specimen resides;

Please also visit: http://www.insdc.org/controlled-vocabulary-specimenvoucher-qualifier

Qualifier /standard\_name=

Definition accepted standard name for this feature

Value format "text"

Example /standard\_name="dotted"

Comment use /standard\_name to give full gene name, but use /gene to

give gene symbol (in the above example /gene="Dt").

Qualifier /strain=

Definition strain from which sequence was obtained

Value format "text"

Example /strain="BALB/c"

Comment entries including /strain must not include

the /environmental\_sample qualifier

Qualifier /sub\_clone=

Definition sub-clone from which sequence was obtained

Value format "text"

Example /sub\_clone="lambda-hIL7.20g"

Comment the comments on /clone apply to /sub\_clone

Qualifier /submitter\_seqid=

Definition identifier attributed to each sequence within an assembly. This identifier

is appropriate for WGS, TSA, TLS and CON records. The submitter\_seqid should be

unique within the context of a single set of assembled sequences.

Value format "text"

Example /submitter\_seqid="NODE\_1"

Comment The length of the value should be limited to <51 characters. Spaces, greater than (>),

left/right square brackets ([ ]) and vertical bar (|) in addition to

double quotation marks (") can not be used for the value of /submitter\_seqid qualifier.

Qualifier /sub\_species=

Definition name of sub-species of organism from which sequence was

obtained

Value format "text"

Example /sub\_species="lactis"

Qualifier /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 "text"

Example /sub strain="abis"

Comment If the parental strain is not given, this should

be annotated in the strain qualifier instead of sub\_strain.

Either:

/strain="K-12"

/sub\_strain="MG1655"

or:

/strain="MG1655"

Qualifier /tag\_peptide=

Definition base location encoding the polypeptide for proteolysis tag of

tmRNA and its termination codon;

Example /tag\_peptide=90..122

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. CDS <90..122;

Qualifier /tissue\_lib=

Definition tissue library from which sequence was obtained

Value format "text"

Example /tissue\_lib="tissue library 772"

Qualifier /tissue\_type=

Definition tissue type from which the sequence was obtained

Value format "text"

Example /tissue\_type="liver"

Qualifier /transgenic

Definition identifies the source feature of the organism which was

the recipient of transgenic DNA.

Value format none

Example /transgenic

Comment transgenic sequences must have at least two source feature keys;

the source feature key having the /transgenic qualifier must span the whole sequence; the source feature carrying the

/transgenic qualifier identifies the main organism of the entry, this determines: a) the name displayed in the organism lines, b) if no translation table is specified, the translation table; only one source feature with /transgenic is allowed in an entry; the /focus and /transgenic qualifiers are mutually exclusive in

an entry.

Qualifier /translation=

Definition automatically generated one-letter abbreviated amino acid

sequence derived from either the universal genetic code or the table as specified in /transl\_table and as determined by an

exception in the /transl\_except qualifier

Value format IUPAC one-letter amino acid abbreviation, "X" is to be used

for AA exceptions.

Example /translation="MASTFPPWYRGCASTPSLKGLIMCTW"

Comment to be used with CDS feature only; this is a mandatory qualifier

in the CDS feature key except where /pseudogene="TYPE" or /pseudo is shown; see /transl\_table for definition and location of genetic

code tables.

Qualifier /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

amino acid coded by the codon at the base\_range position

Example /transl\_except=(pos:213..215,aa:Trp)

/transl\_except=(pos:1017,aa:TERM)

/transl\_except=(pos:2000..2001,aa:TERM)

/transl\_except=(pos:X22222:15..17,aa:Ala)

Comment if the amino acid is not on the restricted vocabulary list use

e.g., '/transl\_except=(pos:213..215,aa:OTHER)' with

'/note="name of unusual amino acid"';

for modified amino-acid selenocysteine use three letter code

'Sec' (one letter code 'U' in amino-acid sequence)

/transl\_except=(pos:1002..1004,aa:Sec);

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, e.g.

if partial stop codon is a single base:

/transl\_except=(pos:1017,aa:TERM)

if partial stop codon consists of two bases:

/transl\_except=(pos:2000..2001,aa:TERM) with

 $\mbox{'/note='stop}$  codon completed by the addition of 3' A residues

to the mRNA'.

Qualifier /transl\_table=

Definition definition of genetic code table used if other than universal

genetic code table. Tables used are described at the specified URLs in appendix IV.

Value format <integer> 1=universal table 1;2=non-universal table 2;...

Example /transl\_table=4

Comment genetic code exceptions outside range of specified tables are

reported in /transl\_except qualifier.

Qualifier /trans splicing

Definition indicates that exons from two RNA molecules are ligated in

intermolecular reaction to form mature RNA

Value format none

Example /trans\_splicing

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 should be used only when the splice event is indicated in the "join" operator, eg join(complement(69611..69724),139856..140087)

Qualifier /type\_material=

Definition indicates that the organism from which this sequence was obtained is

a nomenclatural type of the species (or subspecies) corresponding with

the /organism identified in the sequence entry

Value format "<type-of-type> of <organism name>"

Example /type\_material="type strain of Escherichia coli"

/type\_material="holotype of Cercopitheus lomamiensis"
/type\_material="paratype of Cercopitheus lomamiensis"

Comment <type-of-type> is taken from the INSDC controlled vocabulary for /type\_material

at: http://www.insdc.org/controlled-vocabulary-typematerial-qualifer

<organism name> should be listed as the scientific name

(or as a synonym) at the species (or subspecies) node in the taxonomy database. INSDC will automatically populate this qualifier from the NCBI taxonomy database to

flag sequences of form type in the INSDC databases (ENA/GenBank/DDBJ).

Qualifier /variety=

Definition variety (= varietas, a formal Linnaean rank) of organism

from which sequence was derived.

Value format "text"

Example /variety="insularis"

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 /note, e.g. /note="breed:Cukorova"

## 7.4 Appendix IV: Controlled vocabularies

This appendix contains information on the restricted vocabulary fields used in the Feature Table. The information contained in this appendix is subject to change, please contact the database staff for the most recent information concerning controlled vocabularies. This appendix is organized as follows:

Authority The organization with authority to define the vocabulary

Reference Publications of (or about) the vocabulary

Contact Name of database staff responsible for maintaining

the database copy of the vocabulary

Scope Feature Table qualifiers which take members of this vocabulary

as values

Listing A listing of the current vocabulary with definitions or

explanations

This appendix includes reference lists for the following controlled vocabulary fields:

- Nucleotide base codes (IUPAC)
- Modified base abbreviations
- Amino acid abbreviations
- Modified and unusual Amino Acids

- Genetic Code Tables
- Country Names

# 7.4.1 Nucleotide base codes (IUPAC)

Authority Nomenclature Committee of the International Union of

Biochemistry

Reference Cornish-Bowden, A. Nucl Acid Res 13, 3021-3030 (1985)

Contact EMBL-EBI

Scope Location descriptors

Listing

```
Symbol Meaning
-----
а
       a; adenine
       c; cytosine
С
        g; guanine
g
       t; thymine in DNA; uracil in RNA
t
        a or c
r
        a or g
        a or t
       c or g
s
       c or t
У
k
        g or t
       a or c or g; not t
       a or c or t; not g
h
d
        a or g or t; not c
b
       c or g or t; not a
n
        a or c or g or t
```

# 7.4.2 Modified base abbreviations

Authority Sprinzl, M. and Gauss, D.H.

Reference Sprinzl, M. and Gauss, D.H. Nucl Acid Res 10, r1 (1982).

(note that in Cornish\_Bowden, A. Nucl Acid Res 13, 3021-3030 (1985) the IUPAC-IUB declined to recommend a set of abbreviations for modified nucleotides)

abbreviations for modified

Contact NCBI
Scope /mod\_base

Abbreviation	Modified base description
ac4c	4-acetylcytidine
chm5u	5-(carboxyhydroxylmethyl)uridine
cm	2'-O-methylcytidine

 ${\tt cmnm5s2u} \qquad \qquad {\tt 5-carboxymethylaminomethyl-2-thiouridine}$ 

cmnm5u 5-carboxymethylaminomethyluridine

dhu dihydrouridine

fm 2'-O-methylpseudouridine

gal q beta-D-galactosylqueuosine 2'-0-methylguanosine gm i inosine N6-isopentenyladenosine i6a 1-methyladenosine m1a m1f 1-methylpseudouridine 1-methylguanosine m1g m1i 1-methylinosine m22g 2,2-dimethylguanosine 2-methyladenosine m2a 2-methylguanosine m2g m3c 3-methylcytidine N4-methylcytosine m4c 5-methylcytidine m5c m6a N6-methyladenosine m7g 7-methylguanosine mam5u 5-methylaminomethyluridine mam5s2u 5-methylaminomethyl-2-thiouridine beta-D-mannosylqueuosine man q 5-methoxycarbonylmethyl-2-thiouridine mcm5s2u mcm5u 5-methoxycarbonylmethyluridine mo5u 5-methoxyuridine ms2i6a 2-methylthio-N6-isopentenyladenosine ms2t6a N-((9-beta-D-ribofuranosyl-2-methyltiopurin-6-yl)carbamoyl)threonine mt6a N-((9-beta-D-ribofuranosylpurine-6-yl)N-methyl-carbamoyl)threonine mν uridine-5-oxoacetic acid methylester uridine-5-oxyacetic acid (v) o5u wybutoxosine osyw pseudouridine р queuosine q 2-thiocytidine s2c s2t 5-methyl-2-thiouridine s2u 2-thiouridine 4-thiouridine s4u m5u 5-methyluridine N-((9-beta-D-ribofuranosylpurine-6-yl)carbamoyl)threonine t6a 2'-0-methyl-5-methyluridine †m 2'-O-methyluridine um wybutosine уw 3-(3-amino-3-carboxypropyl)uridine, (acp3)u х OTHER (requires /note= qualifier)

## 7.4.3 Amino acid abbreviations

Authority IUPAC-IUB Joint Commission on Biochemical Nomenclature.

Reference IUPAC-IUB Joint Commission on Biochemical Nomenclature.

Nomenclature and Symbolism for Amino Acids and Peptides.

Eur. J. Biochem. 138:9-37(1984).

IUPAC-IUBMB JCBN Newsletter, 1999

http://www.chem.qmul.ac.uk/iubmb/newsletter/1999/item3.html

Scope /anticodon, /transl\_except

Contact EMBL-EBI

Listing (note that the abbreviations are legal values for amino acids, not the full names)

Abbreviation		Amino acid name
Ala	Α	Alanine
Arg	R	Arginine
Asn	N	Asparagine
Asp	D	Aspartic acid (Aspartate)
Cys	С	Cysteine
Gln	Q	Glutamine
Glu	E	Glutamic acid (Glutamate)
Gly	G	Glycine
His	Н	Histidine
Ile	I	Isoleucine
Leu	L	Leucine
Lys	K	Lysine
Met	М	Methionine
Phe	F	Phenylalanine
Pro	Р	Proline
Pyl	0	Pyrrolysine
Ser	S	Serine
Sec	U	Selenocysteine
Thr	Т	Threonine
Trp	W	Tryptophan
Tyr	Υ	Tyrosine
Val	V	Valine
Asx	В	Aspartic acid or Asparagine
Glx	Z	Glutamine or Glutamic acid.
Xaa	X	Any amino acid.
Xle	J	Leucine or Isoleucine
TERM		termination codon

# 7.4.4 Modified and unusual Amino Acids

Abbreviation	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
Аср	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

N-Ethylglycine EtGly EtAsn N-Ethylasparagine Hyl Hydroxylysine aHy1 allo-Hydroxylysine 3-Hydroxyproline ЗНур 4Нур 4-Hydroxyproline Isodesmosine Ide aIle allo-Isoleucine

MeGly N-Methylglycine, sarcosine

MeIle N-Methylisoleucine
MeLys 6-N-Methyllysine
MeVal N-Methylvaline
Nva Norvaline
Nle Norleucine
Orn Ornithine

OTHER (requires /note=)

### 7.4.5 Genetic Code Tables

Authority International Nucleotide Sequence Database Collaboration

Contact NCBI

Scope /transl\_table qualifier

URL https://www.ncbi.nlm.nih.gov/Taxonomy/Utils/wprintgc.cgi?mode=c

URL http://www.insdc.org/genetic-code-tables

# 7.4.6 Country Names

Authority International Nucleotide Sequence Database Collaboration

Contact INSDC member databases Scope /country qualifier

URL http://www.insdc.org/country

#### 7.4.7 Announces

Additional controlled vocabulary terms for qualifier values might be added outside of the cycle of the Feature Table document release. See also www.insdc.org with controlled vocabularies in the Feature Table document.

From December 2016 a complete list of the genetic codes will also be maintained outside of the cycle of the Feature Table document release at:

http://www.insdc.org/genetic-code-tables

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