

dbSNP VCF Submission Format Guidelines

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Introduction

dbSNP Submissions

dbSNP is a public database of short genetic variations. The data can be from any species, and from any part of a genome. dbSNP has been designed to include a broad collection of simple genetic variations such as single-base nucleotide substitutions, small-scale multi-base deletions or insertions, retrotransposable element insertions, and microsatellite repeats. Submissions can include genotype and allele frequency data if those data are available. dbSNP accepts submissions for all classes of simple molecular variation, including common variations as well as rare variations of germline or somatic origin that are clinically significant. Large-scale insertion/deletion, inversion and translocation data that are over 50bp long should be submitted to [dbVar](#), the NCBI database of genomic structural variation.

Submission Related Email accounts

- Submissions and questions to snp-sub@ncbi.nlm.nih.gov
- Updates to snp-update@ncbi.nlm.nih.gov

Variation Size and Data Submission Limitations

- Submit variations >50 nucleotides in length to the Database of Genomic Structural Variation (<http://www.ncbi.nlm.nih.gov/dbvar>)
- dbSNP does not accept synthetic mutations
- dbSNP does not accept variations ascertained from cross-species alignments and analysis
- dbSNP does not accept personal human data due to current NIH policy unless the participant is enrolled in a study with institutional oversight

There are a few things you should do before you begin your submission.

- *(Optional)* Raw sequence data must be deposited in a public database, e.g., [GenBank](#), [dbGaP](#), [SRA](#) or [Trace](#).
- *(Optional)* Biological samples should be registered in a [BioSamples](#).
- *(Optional)* If your data contains clinically sensitive information or is from patients who have not fully consented to having their genetic information displayed on a public website, you must first submit it to NCBI's Genotypes and Phenotypes database, dbSNP. There, sensitive information will be kept behind controlled access while anonymized variation data will be forwarded directly to dbSNP.
- *(Optional)* If you do not already have one, you may request a [PDA Login](#). The PDA login system enables NCBI to better keep track of your data and make it accessible via additional NCBI resources. Obtaining a PDA login is not required for submission to dbSNP, but it is strongly encouraged.
- *(Optional)* If your submission is part of a larger project or initiative that has been assigned an NCBI Entrez Project ID, please use the project ID (ie. [PRJNA192955](#)) as the HANDLE ID in your dbSNP submission. If you do not have one, we encourage you to request an [NCBI Entrez Project ID](#) to include in your submission. This will facilitate searching and retrieval of your data across all NCBI databases.

The Variant Call Format (VCF)

The Variant Call Format, or VCF, was developed for the [1000 Genomes Project](#) as a standardized format for storing large quantities of sequence variation data (SNPs, indels, larger structural variants, etc.) and any accompanying genotype data and annotation. A VCF file contains a header section and a data table section. Since the metadata lines in the header section can be altered to fit the requirements of the data to be submitted, you can use VCF to submit many different kinds of common variations (as well as their associated genotypes and annotation) that are contained within one reference sequence. VCF files are compressed (using bgzip), and easily accessed. See [Danecek, et. al.](#) for a concise overview of VCF, and the official 1000 Genomes site for a [detailed description of the VCF format](#). Submissions to dbSNP currently use VCF format [version 4.1](#).

NOTE: Please do not use the VCF format if you have human mutations or variations with clinical significance or phenotype. They should be submitted to ClinVar (<http://www.ncbi.nlm.nih.gov/clinvar/>) or contact dbSNP (snp-sub@ncbi.nlm.nih.gov) if you have any questions.

When should I use the VCF format for dbSNP Submissions?

Use dbSNP's Variant Call Format (VCF) to submit large or small numbers of short genetic variations that have asserted positions on genome or reference sequences^a. Large scale submitters especially will find dbSNP's VCF submission format a very useful submission tool since it allows for the submission of numerous variations generated by high-throughput sequencing (HTS) projects over multiple populations, as well as a wide variety of associated data. The VCF file for dbSNP submissions, as opposed to the standard VCF format as defined by the 1000 Genomes project, includes additional fields and attributes that describe dbSNP-specific submission and variation properties, and may include tags that are different than those used in standard VCF.

^adbSNP prefers that all variant asserted positions submitted using the VCF format are submitted either on a sequence accession that is part of an assembly housed in the [NCBI Assembly Resource](#) or as an asserted location on an [INSDC](#) sequence housed in DDBJ, ENA, and GenBank.

Submission Overview

1. Check to see if your lab already has a handle assignment from NCBI. If it does not, request a handle using the [dbSNP online handle request form](#) using the BioProject ID as the handle or forward your BioProject ID to snp-sub@ncbi.nlm.nih.gov and we'll create a dbSNP handle for you from the project ID.
2. Request a FTP account from NCBI for uploading your submission files by sending your handle confirmation information to snp-sub@ncbi.nlm.nih.gov.
3. Prepare your submission:
 - a. The VCF file format required for dbSNP submissions is based on the 1000 Genomes Project VCF format guidelines with the addition of dbSNP specific fields. These additional fields describe dbSNP submission and variation properties.
 - b. Create required [metadata \(meta\) files](#) for the publication, method, population, and assay information associated with the submission.
 - c. Create a VCF Submission file for your data. Include:
 - a properly formatted [dbSNP VCF file header](#)
 - a [data table](#) that contains the [required INFO tag](#) for the variants you are submitting
 - [optional INFO tags](#) that will describe your data more fully.

- d. Upload your submission files to your assigned FTP account and notify snp-sub@ncbi.nlm.nih.gov when the upload is complete.

See the [appendix](#) of this document for an example of a VCF formatted dbSNP submission.

Required Metadata Files

In addition to VCF formatted variation files, dbSNP also requires VCF submissions to include separate Meta file(s).

- The required Meta files are: [Publication](#), [Method](#), [Population](#), and [Assay](#).
- You can submit these Meta files separately or combine them into a single text file for submission.
- Specifications for each Meta file is available in the “[How to Submit](#)” documentation for dbSNP. Links to the specific sections of the document that provide the required specifications are provided above.

Below is an example of a Meta file that combines all four Meta file types into a single file:

```
TYPE:    CONT
HANDLE:  PRJNA192955
NAME:    Jim Johnson
FAX:     111 111 1111
TEL:     222 222 2222
EMAIL:   jj@nih.gov
LAB:     NCBI
INST:    NCBI, NIH
ADDR:    9600 Rockville Pike, Bethesda, MD 20892
||
TYPE:    METHOD
HANDLE:  PRJNA192955
ID:      AgilentWholeExome
METHOD_CLASS:  Sequence
TEMPLATE_TYPE:  DIPLOID
METHOD:
Solution hybridization exome capture was carried out using the Human All Exon
System. The captured regions totaled approximately 38 or 50 Mb depending on
the kit used. Flow cell preparation and paired end read sequencing were
carried out on GAIIX and HiSeq2000 sequencers (Illumina Inc, San Diego CA).
Sequence reads were aligned with the diagCM aligner and genotypes were called
with bam2mpg (Teer et al, Systematic comparison of three genomic enrichment
methods for massively parallel DNA sequencing, Genome Res. 2010
Oct;20(10):1420-31).
||
```

```

TYPE:      POPULATION
HANDLE:    PRJNA192955
ID:       EUROPEAN
POPULATION: This population includes 712 participants of European descent.
||
TYPE:      POPULATION
HANDLE:    PRJNA192955
ID:       AFRICAN
POPULATION: This population includes 600 participants of African descent.
||
TYPE:      SNPASSAY
HANDLE:    PRJNA192955
BATCH:    Exome_SNP_Discovery
MOLTYPE:   Genomic
METHOD:    AgilentWholeExome
ORGANISM:  Homo sapiens
||
TYPE:      SNPPOPUSE
HANDLE:    PRJNA192955
BATCH:    Exome_SNP_MAF
METHOD:    AgilentWholeExome
||

```

dbSNP VCF Submission Format

VCF Submission File Header

Required VCF Header Metadata

The VCF file header for a dbSNP submission should start with the following metadata:

```

##fileformat=      {The current VCF version ID: i.e. VCF v4.1}
##fileDate=        {The date that the file was generated or the date when the
file was
                    updated. Use YYYYMMDD
                    format:i.e.20120201}
##handle=          {Your registered dbSNP submission handle}
##batch=           {A unique local batch ID. Use the same value placed in
                    The BATCH field of the Meta file SNPASSAY section; dbSNP
                    use the local batch ID to associate the VCF submission with
                    the ASSAY, PUBLICATION, and METHOD meta data}
##bioproject_id=   {A registered BioProject ID if available}
##biosample_id=    {A comma separated list of registered BioSample IDs
                    (http://www.ncbi.nlm.nih.gov/biosample/).
                    In this example of two Biosample records, the ID numbers
                    are 423 and 1595}
##reference=       {The RefSeq Assembly accession.version on which the
                    variation position is based: i.e. GCF_000001405.12. You
                    can find this ID by accessing NCBI's Genome Assembly

```

Resource (<http://www.ncbi.nlm.nih.gov/assembly/>) and search for the record of the specific assembly. You can use the organism or assembly name (e.g. GRCh37) as your search term: the assembly record for GRCh37 shows the RefSeq ID is GCF_000001405.12. Only accession.version for fully assembled genome can be reported here. For unassembled and unplaced contigs you leave this blank and use the reporting method for ISDN sequence coordinates as shown in the example below for CHROM column.}

Example of dbSNP Metadata in a VCF formatted file:

```
##fileformat=VCFv4.1
##fileDate=20120215
##handle=PRJNA192955
##batch=Exome_SNP_Discovery
##bioproject_id=60153
##biosample_id=423, 1595
##reference=GCF_000001405.12
```

INFO Tag Descriptions

The VCF header continues with tag/value descriptions for required and optional dbSNP INFO tags. These descriptions should be placed in the header following the required metadata.

The INFO tag/value descriptions you provide in the VCF header will serve to define the data you place in the INFO column of the data table. These descriptions are an important part of the VCF header as they will allow users viewing your data in VCF format to identify a tag you placed in the INFO column and see definitions for values of that tag. The data you present in the INFO column of the data table will be meaningless to some users without the inclusion of the tag/value descriptions in the VCF header for those data.

Descriptions for Required INFO Tag

Currently, the only required INFO tag for a dbSNP submission is the [Variation Type \(VRT\) tag](#). Place the VRT tag description in the VCF file header after the required metadata. The VRT tag is required for each variant submitted in VCF format. **Failure to include this required INFO tag will result in the delay of your submission.**

See the dbSNP [INFO Tag Descriptions and Examples](#) section of this document for example tag descriptions you can cut and paste into the VCF file header for both the required INFO tags and the optional INFO tags.

Descriptions for Optional INFO Tags

Place descriptions for the optional INFO tags in the VCF file header after the required metadata. These descriptions identify and define the optional INFO tags you have elected to use in the data table portion of the file.

See the dbSNP [INFO Tag Descriptions and Examples](#) section of this document for example tag descriptions you can cut and paste into the VCF file header for both the required INFO tags and the optional INFO tags.

Submission Data Table

Create a tab-delimited table to house your variations and variation data for your submission. The table header should include these six fixed, mandatory columns (in order):

```
#CHROM POS ID REF ALT INFO
```

The above columns represent six fixed fields that must be filled out for each submitted variant. If you do not have data for a particular field, use a dot (".") to represent the missing value.

VCF Data Table Examples

A) **Reporting** positions using *chromosome* coordinates (*please provide the 'reference' tag in the header if the assembly and version is known*)

<i>#CHROM</i>	<i>POS</i>	<i>ID</i>	<i>REF</i>	<i>ALT</i>	<i>QUAL</i>	<i>FILTER</i>	<i>INFO</i>
23	135498962	NG_021219.1:g.120841A>G	A	G	29	PASS	VRT=1
23	135499109	NG_021219.1:g.120988G>A	G	A	40	PASS	VRT=1
23	135499270	NG_021219.1:g.121149C>T	C	T	51	PASS	VRT=1
23	135499419	NG_021219.1:g.121298G>C	G	C	68	PASS	VRT=1

B) Reporting positions using ISDN sequence coordinates if assembly is not known.

#CHROM	POS	ID	REF	ALT	QUAL	FILTER	INFO
NG_021219.1	140860	SNV1	T	C	29	PASS	VRT=1
NG_021219.1	140879	SNV2	A	G	40	PASS	VRT=1
NG_021219.1	140921	SNV3	T	C	51	PASS	VRT=1
NG_021219.1	140939	SNV4	C	T	68	PASS	VRT=1

VCF Data Table Field Values

#CHROM

This field contains the chromosome identifier from the reference genome where the variant is located or an angle-bracketed ID String ("*<ID>*") pointing to a contig in the assembly file. (cf. the *##assembly* line in the header). Entries for a specific CHROM should form a contiguous block within the VCF file. Alternative, the sequence accession and version can be used for this field if the variation position is based on a non-chromosomal sequence (see example B above). Do not use the colon symbol (:) in a chromosome name.

#POS

This field contains the reference position of the variant, which is the 1st base of the variation event. Positions are sorted numerically within each reference sequence chromosome (CHROM) in increasing order. When there are multiple variation allele starting at the same POS, submit one variation type(VRT) per line. Telomeres are indicated by using positions 0 or N+1, where N is the length of the corresponding chromosome or contig.

Note: For short, simple insertions and deletions in which the REF or one of the ALT alleles would otherwise be null/empty, the POS field must contain the coordinates of the base preceding the indel event. See the [Submission Data Table Special Case Examples](#) section of this document for instruction on reporting insertion/deletion POS values.

Large indels and structural variants must be submitted to [dbVAR](#)

ID

This field contains the unique local ID (AD) of the variant, **and is a required value (cannot be NULL)**.The AD provided here combined with the handle must be unique for a particular submitter. You can use an

HGVS expression (<http://www.hgvs.org/mutnomen/recs.html>) for the variant ID if you do not have a unique identifier of your own.

REF

This field contains the reference allele of the variant. The bases representing the reference allele can be any of the following: A, C, G, T, N (case insensitive).

Note: In order for the variant to be included in dbSNP, the maximum length for the REF allele is 51bp.

Note: For short, simple insertions and deletions in which the REF or one of the ALT alleles would otherwise be null/empty, the REF and ALT Strings must include the base preceding the indel event. See the [Submission Data Table Special Case Examples](#) section of this document for instruction on reporting indel reference (REF) alleles.

ALT

This field contains a comma separated list of alternate, non-reference alleles that you have called in at least one sample. You can use A, C, G, T or N (case insensitive) or you can use an angle-bracketed ID String ("`<ID>`"). If there are no alternative alleles, put a dot (".") placeholder the ALT column. **Note: In order for the variant to be included in dbSNP, the maximum length of each REF allele is 51bp.**

Note: For short, simple insertions and deletions in which the REF or one of the ALT alleles would otherwise be null/empty, the REF and ALT Strings must include the base preceding the indel event. See the [Submission Data Table Special Case Examples](#) section of this document for instruction on reporting indel alternate (ALT) alleles.

QUAL

This field contains the quality score for the assertion if available.

FILTER

This field contains the filter status if available.

INFO

This field contains additional information for the reported variation. INFO fields are encoded as a semicolon-separated series of short keys with optional values in the format: `<key>=<data>[,data]` See the [INFO Tag Descriptions and Examples](#) section of this document for examples of the required and optional INFO Tags that dbSNP supports.

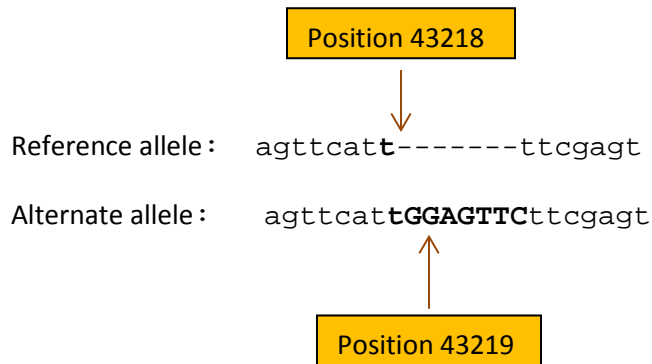
Submission Data Table Special Case Examples: Reporting POS, REF and ALT for insertion/deletion variants

For simple insertions and deletions where either the REF or one of the ALT alleles would otherwise be null/empty, include the base preceding the variation event (a “padding base”) in the REF and ALT allele Strings, and report the coordinates of this “padding base” in POS.

The “padding base” is not required for complex substitutions or other events where all alleles have at least one base represented in their Strings.

Insertion Example

Sequence: TCAGTCTCACCATGAAAGTTCATT [-/GGAGTTC]TTCGAGTAAATGGTTCACGCGGG

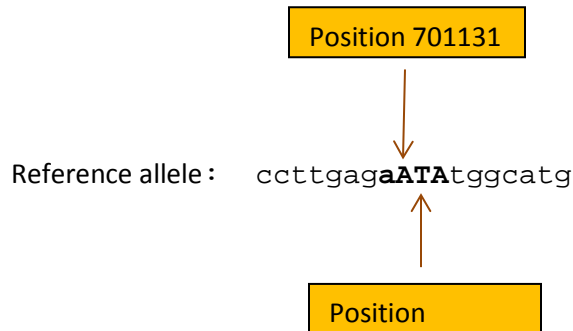


If the coordinates of first base of the insertion event (“G” at position 43219) in the above sequence were used as the reference position (POS) of this event, the REF field would have no value since the inserted bases are only present in the ALT allele. In such a case, report the coordinates of the base that precedes the insertion event— the “t” at position 43218 — for POS and include this “padding base” in the REF and ALT Strings:

#CHROM	POS	ID	REF	ALT	QUAL	FILTER	INFO
10	43218	NC_000010.10:g.43218_43219insGGAGTTC	T	TGGA GTTC	.	.	VRT=2;AA=T;NIO=12;SSR=0;SAO=0;SCS=0

Deletion Example

Sequence : AGAGATTCACAGCCTCACCTTGAGA[ATA/-]TGGCATGGAGAATATTTTGGATAAT



Alternate allele : ccttgaga---tgcatg

If the coordinates of first base of this deletion event (“A” at position 701132) in the above sequence were used as the reference position (POS) of this variant, the ALT field would have no value since the deleted bases are only present in the reference (REF) allele. In such a case, report the coordinates of the base that precedes the deletion event— the “a” at position 701131 — for POS and include this “padding base” in the REF and ALT Strings:

#CHROM	POS	ID	REF	ALT	QUAL	FILTER	INFO
15	701131	NC_000015.9:g.701132_701134delATA	AA TA	A	.	.	VRT=2;AA=A;NIO=5;SSR=0;SAO=0;SCS=0

INFO Tag Descriptions and Examples

Required dbSNP VCF INFO Tag

Place the required tag in the INFO column of the [data table](#) and place the corresponding tag description in the [file header](#).

Variation Type (VRT) INFO Tag

The required “VRT” INFO tag allows you to define the kind of variation you are submitting to dbSNP. We use this information to verify position and that the reported alleles are consistent with reported variation type. **Failure to include this required INFO tag will result in the delay of your submission.** Also only one variation type (VRT) can be reported per row. For instance if you have both a deletion

variation and SNV at the same location, they should be reported two separate rows with the corresponding VRT value.

VRT Tag/Value Description

```
##INFO=<ID=VRT,Number=1,Type=Integer,Description="Variation type,1 - SNV:
single nucleotide variation,2 - DIV: deletion/insertion variation,3 -
HETEROZYGOUS: variable, but undefined at nucleotide level,4 - STR: short
tandem repeat (microsatellite) variation, 5 - NAMED: insertion/deletion
variation of named repetitive element,6 - NO VARIATON: sequence scanned for
variation, but none observed,7 - MIXED: cluster contains submissions from 2
or more allelic classes (not used),8 - MNV: multiple nucleotide variation
with alleles of common length greater than 1,9 - Exception">
```

VRT Data Format Example

<i>#CHROM</i>	<i>POS</i>	<i>ID</i>	<i>REF</i>	<i>ALT</i>	<i>QUAL</i>	<i>FILTER</i>	<i>INFO</i>
1	140860	NC_000001.10:g140860T>C	T	C	.	.	VRT=1

Optional dbSNP VCF INFO Tags

The following INFO tags are optional and need only be used if they describe available data. If you want to include any of the following INFO tags with your submitted data, place the tag in the INFO column of the [data table](#) and place the corresponding tag description in the [file header](#). Optional VCF INFO tags for dbSNP submissions include:

[Alternate Designations](#)

[Ancestral Allele](#)

[Free Text for Comment](#)

[LinkOut](#)

[Mutation](#)

[Number of Independent Observations](#)

[OMIM/OMIA Record](#)

[Population Frequency](#)

[Population ID](#)

[PubMed ID](#)

[Variant Allele Origin](#)

[Variant Suspect Reason](#)

Alternate Designations (AD) or Names

The optional "AD" INFO tag allows you to provide dbSNP with a (comma separated) set of alternative names or common names used to describe the same submitted variant.

AD Tag/Value Description

##INFO=<ID=AD,Number=1,Type=String,Description="Alternate designations; a set of (comma separated)alternative names used to describe the same variant">

AD Tag/Value Example

<i>#CHR OM</i>	<i>POS</i>	<i>ID</i>	<i>REF</i>	<i>ALT</i>	<i>QUAL</i>	<i>FILTER</i>	<i>INFO</i>
10	1340172 95	NC_000010.10:g.134017 295A>G	A	G	.	.	VRT=1;AA=T; NIO=12;AD=SNP- 12313,chr10:1340172 95A>G;

Ancestral Allele (AA)

The optional "AA" INFO tag allows you to provide dbSNP with the ancestral allele (if you know it) for a variant.

AA Tag/Value Description

##INFO=<ID=AA,Number=1,Type=String,Description="Provide Ancestral Allele if known">

AA Tag/Value Example

<i>#CHROM</i>	<i>POS</i>	<i>ID</i>	<i>REF</i>	<i>ALT</i>	<i>QUAL</i>	<i>FILTER</i>	<i>INFO</i>
8	19863	NC_000008.10:g.19863G>C	G	C	.	.	VRT=1;AA=T;

Free Text for Comment (CMT)

The optional "CMT" INFO tag allows you to provide dbSNP with text about any additional important information that cannot be described (e.g. phenotypic information) using the other available INFO tags

CMT Tag/Value Description

```
##INFO=<ID=CMT=1,Type=String,Description="Comment">
```

CMT Data Format Example

<i>#CHROM</i>	<i>POS</i>	<i>ID</i>	<i>REF</i>	<i>ALT</i>	<i>QUAL</i>	<i>FILTER</i>	<i>INFO</i>
8	19863	NC_000008.10:g.19863G>C	G	C	.	.	VRT=1;CMT="A variant identified in SLC10A1 gene with possible correlation to disease susceptibilities(PM ID: 12436193)";

LinkOut (LKO)

The optional "[LKO](#)" INFO tag allows you to point to this variant on your organization's web site or to other relevant online information about your submission.

LKO Tag/Value Description

```
##INFO=<ID=LKO,Number=1,Type=String,Description="A link out URL for this variant on the submitting organization's website">
```

LKO Data Format Example:

#C HR OM	PO S	ID	R E F	A L T	Q U A L	F I L T E R	INFO
8	19 86 3	NC_000008 .10:g.198 63G>C	G	C	.	.	VRT=1;AA=T;LKO= http://variantgps.nci.nih.gov/cgfseq/pages/sequenceSubmit.do?method=sequence&SNP500ID=LPL-08 ;

Number of Independent Observations (NIO)

The optional “NIO” INFO tag allows you to provide dbSNP with the number of times you observed this variant occur independently in your experimental analysis.

NIO Tag/Value Description

##INFO=<ID=NIO,Number=1,Type=Integer,Description="Number of Independent Observations;the number of times the submitter observed this variant occurring independently">

NIO Tag/Value Example

#CHRO M	POS	ID	R E F	A L T	Q U A L	F I L T E R	INFO
8	1986 3	NC_000008.10:g.19863G >C	G	C	.	.	VRT=1;AA=T;NIO=12 ;

OMIM and OMIA (OMIM/OMIA) Records

The optional “OMIM” and “OMIA” INFO tags allow you to provide dbSNP with any available [OMIM](#) or [OMIA](#) record and variant ID (if available) associated with a variant.

OMIM and OMIA Tag/Value Descriptions

OMIM:

##INFO=<ID=OMIM,Number=1,Type=String,Description="Provide OMIM (http://omim.org)record and variant ID if available i.e. 300746.0001">

OMIA:

```
##INFO=<ID=OMIA,Number=1,Type=String,Description="Provide OMIA  
(http://www.ncbi.nlm.nih.gov/omia)record and variant ID if available i.e.  
000011-9615">
```

OMIM and OMIA Data Format Example

#CHR OM	POS	ID	RE F	AL T	QUA L	FILT ER	INFO
16	9199 82	NC_000016.9:g.9199 82G>C	G	C	.	.	VRT=1;AA=T;OMIM=300746 .0001;

Population IDs (for Allele Frequency, Genotype Frequency, or Observed Heterozygosity data submissions)

If you intend to report allele frequency, genotype frequency, or observed heterozygosity in your VCF formatted dbSNP submission, place the population ID for each assayed population in the VCF header after the INFO Tag/Value descriptions, and before your data table. The POP IDs you will provide in the VCF header are the same ones you placed in the ID field of the Meta File.

Population_ID Tag/Value Description

```
##population_id=<A unique local population ID e.g. "HapMap", "Case",  
"Control", "Healthy Blood Donors"> Use the same value placed in the ID field  
of the Meta file POPULATION section where the population details are  
described.
```

Population_ID Example

```
##INFO=<ID=GEN_FRQ,Number=1,Type=string,Description="Report population,  
sample size (number of distinct chromosomes assayed), and frequency for each  
genotype  
##population_id=EUROPEAN  
##population_id=AFRICAN
```

Allele Frequency Format Examples

The format for reporting allele frequency follows the convention for reporting for genotype.

- Add a reporting FORMAT column to specify the data type and order. Suggested data types are listed below.


```
##FORMAT=<ID=NA,Number=1,Type=Integer,Description="Number of alleles for
the population."
##FORMAT=<ID=NS,Number=1,Type=Integer,Description="Number of samples for
the population."
##FORMAT=<ID=FRQ,Number=.,Type=Float,Description="Frequency of each
alternate allele."
##FORMAT=<ID=AC,Number=.,Type=Integer,Description="Allele count for each
alternate allele."
```

- Add additional column for each population
- Report under the population column the total allele count (NA) or population samples (NS) follow by the allele frequency (FRQ) or allele count (AC) separated by a colon ‘:’

#CHR OM	POS	I D	RE F	AL T	QU AL	FILT ER	INFO	FORM AT	EUROPEA N	AFRICA N
X	1408 79	.	A	G	.	.	VRT=1;AD=SNV:chrX: 140879;	NA:F RQ	1424:0. 001	1200:0 .05

PubMed ID (PMID) INFO Tag

The optional “PMID” INFO tag allows you to provide dbSNP with the PubMed ID (if available) for an original publication associated with a variant. If multiple PubMed IDs (PMID) are available for a single variant, report them using a comma separated list (see example below). Report PMIDs for multiple variants as a batch in the [ASSAY](#) and [PUBLICATION](#) meta files.

PMID Tag/Value Description

```
##INFO=<ID=PMID,Number=.,Type= Integer,Description="PubMed ID linked to
variation if available">
```

PMID Data Format Example

#CHRO M	POS	ID	RE F	AL T	QUA L	FILTE R	INFO
16	91998 2	NC_000016.9:g.91998 2G>C	G	C	.	.	VRT=1;AA=T; PMID=21840 003;

Variant Allele Origin (SAO) INFO Tag

The optional “SAO” or “Variant Allele Origin” INFO tag allows you to provide dbSNP with the source of the sample from which the variant was derived.

Note: Although the name we use to refer to Allele Origin has changed from “SNP Allele Origin” (SAO) to “Variant Allele Origin” to emphasize that the dbSNP database contains both rare and polymorphic variants, the database itself still uses the acronym “SAO”.

SAO Tag/Value Description

```
##INFO=<ID=SAO,Number=.,Type=Integer,Description="Variant Allele Origin: 0 - unspecified, 1 - Germline, 2 - Somatic, 3 - Both">
```

SAO Data Format Example

<i>#CHROM</i>	<i>POS</i>	<i>ID</i>	<i>REF</i>	<i>ALT</i>	<i>QUAL</i>	<i>FILTER</i>	<i>INFO</i>
16	91998 2	NC_000016.9:g.919982G >C	G	C	.	.	VRT=1;AA=T;SAO=1 ;

Note: If you are providing more than one allele origin value, place the allele origin values in a comma separated list in the order that they appear in the submission. List the value for the reference allele first, followed by the allele origin value for the 1st alternate allele, 2nd alternate allele, etc.:

Variant Suspect Reason (SSR) INFO Tag

The optional “SSR” or “SNP Suspect Reason” INFO tag allows you to provide dbSNP with the reason you suspect that a variant is a false positive. Evidence for false positives can include information indicating the presence of a paralogous sequence in the genome ([Musumeci et al. 2010](#)) ([Sudmant et al. 2010](#)), or evidence of sequencing error or computation artifacts.

Note: Although the name we use to refer to the Suspect Reason code has changed from “SNP Suspect Reason” (SSR) to “Variant Suspect Reason” to emphasize that the dbSNP database contains both rare and polymorphic variants, the database itself still uses the acronym “SSR”.

SSR Tag Description

```
##INFO=<ID=SSR,Number=.,Type=Integer,Description="Variant Suspect Reason Code, 0 - unspecified, 1 - Paralog, 2 - byEST, 3 - Para_EST, 4 - oldAlign, 5 - other">
```

SSR Data Format Example

#CHROM	POS	ID	REF	ALT	QUAL	FILTER	INFO
16	91998 2	NC_000016.9:g.919982G >C	G	C	.	.	VRT=1;AA=T;SSR=1 ;

Appendix: Example of a VCF Formatted dbSNP Submission

```
##fileformat=VCFv4.1
##fileDate=20120215
##handle=PRJNA192955
##batch=Exome_SNP_Discovery
##reference=GCF_000001405.12
##INFO=<ID=VRT,Number=1,Type=Integer,Description="Variation type, 1 - SNV:
single nucleotide variation, 2 - DIV: deletion/insertion variation, 3 -
HETEROZYGOUS: variable, but undefined at nucleotide level, 4 - STR: short
tandem repeat (microsatellite) variation, 5 - NAMED: insertion/deletion
variation of named repetitive element, 6 - NO VARIATION: sequence scanned for
variation, but none observed, 7 - MIXED: cluster contains submissions from 2
or more allelic classes, 8 - MNV: multiple nucleotide variation with alleles
of common length greater than 1, 9 - Exception">
##INFO=<ID=AD,Number=1,Type=string,Description="Unique local alternative ID
or variation name for display. The AD provided here combined with the handle
must be unique for a particular submitter. An HGVS expression
(http://www.hgvs.org/mutnomen/recs.html) can be used here">
##FORMAT=<ID=NA,Number=1,Type=Integer,Description="Number of alleles for the
population."
##FORMAT=<ID=NS,Number=1,Type=Integer,Description="Number of samples for the
population."
##FORMAT=<ID=FRQ,Number=.,Type=Float,Description="Frequency of each alternate
allele."
##FORMAT=<ID=AC,Number=.,Type=Integer,Description="Allele count for each
alternate allele."
##population_id=EUROPEAN
##population_id=AFRICAN
```

<i>#CHR OM</i>	<i>POS</i>	<i>I D</i>	<i>RE F</i>	<i>AL T</i>	<i>QU AL</i>	<i>FILT ER</i>	<i>INFO</i>	<i>FORM AT</i>	<i>EUROPEA N</i>	<i>AFRICAN</i>
X	1408 60	.	T	C	.	.	VRT=1;AD=SNV:chrX: 140860;	NA:F RQ	1424:0. 056	.
X	1408 79	.	A	G	.	.	VRT=1;AD=SNV:chrX: 140879;	NA:F RQ	1424:0. 001	1200:0. 05
X	1409 21	.	T	C	.	.	VRT=1;AD=SNV:chrX: 140921;	NA:F RQ	1424:0. 003	1200:0. 002
X	1409 39	.	C	T	.	.	VRT=1;AD=SNV:chrX: 140939;	NA:F RQ	1424:0. 01	.

Revision History:

10/13/2013

- Replaced tag for ancestral allele (ANC) with AA.
- Replaced tag for local ID (LID) with alternative variation ID (AD).
- Added submission limitations
- Added optional submissions to other NCBI resources (SRA, BioSamples, and BioProjects).