

# AnnotSV Manual

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Version 1.2

AnnotSV is a program for annotating structural variations from the human genome.

<http://lbgf.fr/AnnotSV/>

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Please feel free to contact me for any suggestions or bug reports  
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## LEXIQUE

1000g: 1000 Genomes Project (phase 3)

BED: Browser Extensible Data

bp: base pair

CDS: CoDing Sequence

CNV: Copy Number Variation

DDD: Deciphering Developmental Disorders

DECIPHER: DatabasE of genomiC variation and Phenotype in Humans using Ensembl Resources

DEL: Deletion

DGV: Database of Genomic Variants

DNA: DesoxyriboNucleic Acid

DUP: Duplication

ENCODE: Encyclopedia of DNA Elements

ExAC: Exome Aggregation Consortium

GRCh37: Genome Reference Consortium Human Build 37

GRCh38: Genome Reference Consortium Human Build 38

HI: Haploinsufficiency

hom: homozygous

htz: heterozygous

ID: Identifier

indel: Insertion/deletion

LoF: Loss of Function

misZ = Z score indicating gene intolerance to missense variation

NAHR: Non-Allelic Homologous Recombination

NM: RefSeq identifiers

OMIM: Online Mendelian Inheritance in Man

pLI = score computed by the ExAc consortium to indicate gene intolerance to a loss of function variation

SNV : Single Nucleotide Variation

SV: Structural Variations

synZ = Z score indicating gene intolerance to synonymous variation

TAD: Topologically Associating Domains

Tcl: Tool Command Language

Tx: transcript

VCF: Variant Call Format

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## 1. INTRODUCTION

AnnotSV is a program designed for annotating Structural Variations (SV). This tool compiles functionally, regulatory and clinically relevant information and aims at providing annotations useful to i) **interpret SV potential pathogenicity** and ii) **filter out SV potential false positives**.

Different types of SV exist including deletions, duplications, insertions, inversions, translocations or more complex rearrangements. They can be either balanced or unbalanced. When unbalanced and resulting in a gain or loss of material, they are called Copy Number Variations (CNV). CNV can be described by coordinates on one chromosome, with the start and end positions of the SV (deletions, insertions, duplications). Complex rearrangements with several breakends can arbitrary be summarized as a set of novel adjacencies, as described in the Variant Call Format Specification [VCFv4.3](#) (Jul 2017).

AnnotSV takes as an input file a classical bed or VCF file describing the SV coordinates. The output file contains the overlaps of the SV with relevant genomic features where the genes refer to NCBI RefSeq genes. In addition to the gene annotations, we provide numerous additional relevant annotations (OMIM, DGV frequencies, compound heterozygosity ...).

## 2. INSTALLATION/REQUIREMENTS/UPDATE

### 2.1. Tcl (Required)

The AnnotSV program is written in the Tcl language. Modern Unix systems have this scripting language already installed (otherwise it can be downloaded from <http://www.tcl.tk/>).

AnnotSV requires **the latest release of the Tcl distribution starting with version 8.6** as well as the following 2 packages "tar" and "csv" (used only when data sources are updated).

### 2.2. AnnotSV source code (Required)

"AnnotSV sources" can be download at <http://lbgf.fr/AnnotSV/downloads> (under the GNU GPL license).

#### Install:

The sources .tar.gz should be extracted and uncompressed to any directory.

```
tar -xvf AnnotSV_latest.tar.gz
```

The installation requires simply to set the following environment variable:

```
- $ANNOTSV : "AnnotSV installation directory"
```

Make sure the program correctly finds the Tcl interpreter. By default, the best way to make a Tcl script executable is to put the following as the first line of the main script (which is already done in AnnotSV-main.tcl):

```
#!/usr/bin/env tclsh
```

It can be changed to any other path like:

```
#!/usr/local/ActiveTcl/bin tclsh
```

Typically, you can create an alias of the main Tcl script “sources/AnnotSV-main.tcl” for example to “AnnotSV”, place it in the “/bin” directory”(this is done by default already) and add the path to this in your \$PATH.

### **AnnotSV installation directory:**

By default the AnnotSV installation directory looks like this:

```
AnnotSV                #the program installation directory
|
|---- bin/              #where an alias is set to the main .tcl script
|
|---- changeLogs.txt   #description of AnnotSV changes
|
|---- configfile       #a configfile example that can be edited for modification purpose
|
|---- Example/         #command/input/output example
|
|---- Annotations/    #where external annotation files are stored (RefGene, OMIM, DGV...)
|
|---- License.txt     #GNU GPL license
|
|---- README.AnnotSV_*.pdf #this file
|
|---- Sources/        #where the source .tcl files are stored
```

### 2.3. bedtools (Required)

The “[bedtools](#)” toolset (developed by Quinlan AR) needs to be locally installed. Configuration requires to set the path to the bedtools executable in the AnnotSV configfile located in: \$ANNOTSV/configfile.

### 2.4. Annotation sources (Provided)

AnnotSV requires different data sources for the annotation of SV. **In order to provide a ready to start installation of AnnotSV, each annotation source listed below (that do not require a commercial license) is already provided with the AnnotSV sources.** The aim and update of each of these sources are explained below.

Annotation can be performed using either the GRCh37 or GRCh38 build version of the human genome (user defined, see USAGE/OPTIONS), but depending on the availability of some data sources there might be some limitations.

Some of the annotations are linked to the gene name and thus provided independently of the genome build.

#### a) GENE ANNOTATIONS

The “Gene annotation” aims at providing information for the overlapping known genes with the SV in order to list the genes from the well annotated [RefSeq](#) database. These annotations include the definition of the genes and corresponding transcripts (RefSeq), the length of the CoDing Sequence (CDS) and of the transcript, the location of the SV in the gene (e.g. « txStart-exon3 ») and the coordinates of the intersection between the SV and the transcript.

**Annotation columns:**

Adds 7 annotation columns: "Gene name", "NM", "CDS length", "tx length", "location", "intersectStart", "intersectEnd".

**Method:**

For each gene, only a single transcript from all transcripts available in RefSeq for this gene is reported. In case of transcripts with different CDS length (considering the overlapping region with the SV), the transcript with the longest CDS is reported. Otherwise, if there is no differences in CDS length, the longest transcript is reported.

**Updating the data source (if needed):**

- Remove all the files in the "\$ANNOTSV/Annotations/RefGene/GRCh37" and/or "\$ANNOTSV/Annotations/RefGene/GRCh38" directories.
- Download and place the "refGene.txt.gz" file in the "\$ANNOTSV/Annotations/RefGene/GRCh37" and/or "\$ANNOTSV/Annotations/RefGene/GRCh38" directories. The latest update of this file is available for free download at:

*Genome build GRCh37:*

<http://hgdownload.cse.ucsc.edu/goldenPath/hg19/database/refGene.txt.gz>

*Genome build GRCh38:*

<http://hgdownload.cse.ucsc.edu/goldenPath/hg38/database/refGene.txt.gz>

After the update, this refGene.txt.gz file will be processed by AnnotSV during the first run (it will take longer than usual AnnotSV runtime).

It is to notice that the **promoter's annotations update** will be done at the same time (without supplementary update command).

b) [PROMOTERS ANNOTATIONS](#)

**Aim:**

The contribution of SV overlapping with promoters to disease etiology is well established, affecting gene expression, although understanding the consequences of these regulatory variants on the human transcriptome remains a major challenge. AnnotSV reports the list of the genes whose promoters are overlapped by the SV.

**Annotation columns:**

Adds 1 annotation column: "promoters"

**Method:**

Promoters are defined by default as 500 bp upstream from the transcription start sites (using the RefGene data). Nevertheless, the user can define a different bp size with the "promoterSize" option (see USAGE/OPTIONS). A promoter is reported only if the SV overlaps at least 70% of this promoter (user defined, see the "FeaturesOverlap" option in USAGE/OPTIONS).

**Update:**

The promoters' annotations update will be done at the same time as the Gene annotations update.

c) [DGV GOLD STANDARD ANNOTATIONS](#)

**Aim:**

The Database of Genomic Variants ([DGV](#), (MacDonald, et al., 2014)) provides SV defined as DNA elements with a size >50 bp. The content of DGV is only representing SV identified in healthy control samples from large cohorts

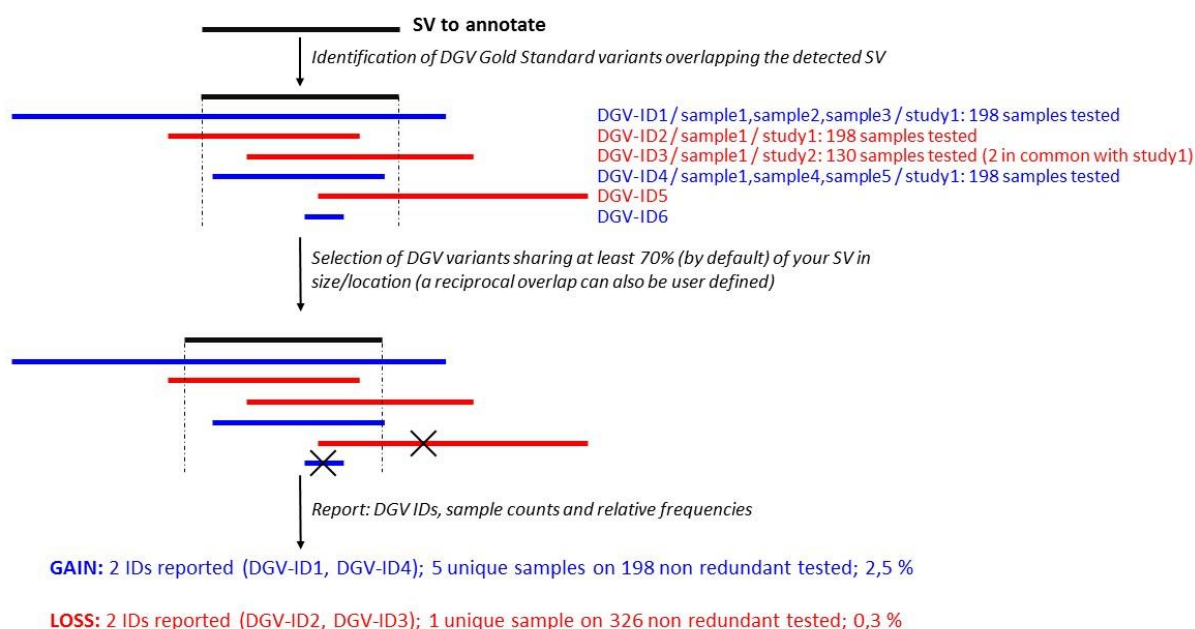
published and integrated by the DGV team. The annotations will give information about whether your SV is a rare or a common variant

**Annotation columns:**

Adds 8 annotation columns: respectively for GAIN and LOSS: “DGV\_IDs”, “n\_samples\_with\_SV”, “n\_samples\_tested” and “Frequency”.

**Method:**

First, AnnotSV searches for DGV Gold Standard variants overlapping the SV to annotate. Second, only the DGV variants overlapping at least 70% of your SV in size/location are selected (default value, a different percentage or a reciprocal overlap can also be user defined with the "SVfromDBoverlap" and "SVtoAnnOverlap" options). Third, the DGV IDs are reported. Then, all DGV samples information are merged: the counts of unique samples with gains and losses, the number of samples tested in the related studies (without redundancy) and subsequent relative frequencies are calculated and reported (genotype data are not considered).



**Warning:**

- Exceptional overestimation of the relative frequencies:

In DGV Gold Standard (March 2016), ~10% of the supporting variants are not released with sample information preventing AnnotSV to properly differentiate whether some variation are redundant or not. Consequently, some relative frequencies can be exceptionally overestimated by AnnotSV.

- Gain/Loss:

A SV call in DGV can be relative to a specific reference sample, a pool of reference samples or relative to the reference assembly. Since different reference samples may have been used in different studies, what is called as a gain in one study may actually be called a loss in another.

**Updating the data source (if needed):**

- Remove all the files in the “\$ANNOTS/Annotations/DGV/GRCh37” and/or “\$ANNOTS/Annotations/DGV/GRCh38” directories.



- Download and place the 2 following DGV files in the “\$ANNO SV/Annotations/DGV/GRCh37” and/or “\$ANNO SV/Annotations/DGV/GRCh38” directories.

*Genome build GRCh37:*

The latest update of these 2 files are available for free download at <http://dgv.tcag.ca/dgv/app/downloads>

- **DGV.GS.March2016.50percent.GainLossSep.Final.hg19.gff3** (see DGV Gold Standard Variants section)
- **GRCh37\_hg19\_supportingvariants\_2016-05-15.txt** (see Supporting Variants section)

*Genome build GRCh38:*

**The dataset is not yet available from the DGV team.**

These 2 files will be computed the first time AnnotSV will be executed after the update.

#### d) [dbVAR PATHOGENIC NR SV ANNOTATIONS](#)

##### **Aim:**

By selecting medium (< 1000000 bp) pathogenic SV records from the dbVar NR SV database, AnnotSV obtained a clinically-relevant human SV dataset. It may be used to determine genome locations where pathogenic SV are found, and so determine overlaps between these pathogenic SV and the SV to annotate.

##### **Method:**

By default, a pathogenic NR SV is reported only if it overlaps at least 70% of the SV to annotate. Nevertheless, the user can modify the default behaviour by either use a different percentage or a reciprocal overlap (see "SVfromDBoverlap" and "SVtoAnnOverlap" options in USAGE/OPTIONS).

##### **Annotation columns:**

Adds 3 annotation columns: “dbvar\_event”, “dbVar\_variant” and “dbVar\_status”.

##### **Updating the data source (if needed):**

- Remove all the “\*\_dbVar\_pathogenic\_NR\_SV\*” files in the “\$ANNO SV/Annotations/Users/GRCh37” and/or “\$ANNO SV/Annotations/Users/GRCh38” directories.
- Remove all the files in the “\$ANNO SV/Annotations/dbVar/GRCh37” and/or “\$ANNO SV/Annotations/dbVar/GRCh38” directories.
- Download and place the 2 following dbVar files in the “\$ANNO SV/Annotations/dbVar/GRCh37” and/or “\$ANNO SV/Annotations/dbVar/GRCh38” directories.

*Genome build GRCh37:*

[https://ftp.ncbi.nlm.nih.gov/pub/dbVar/sandbox/sv\\_datasets/nonredundant/deletions/GRCh37.nr\\_deletions.tsv.gz](https://ftp.ncbi.nlm.nih.gov/pub/dbVar/sandbox/sv_datasets/nonredundant/deletions/GRCh37.nr_deletions.tsv.gz)

[https://ftp.ncbi.nlm.nih.gov/pub/dbVar/sandbox/sv\\_datasets/nonredundant/duplications/GRCh37.nr\\_duplications.tsv.gz](https://ftp.ncbi.nlm.nih.gov/pub/dbVar/sandbox/sv_datasets/nonredundant/duplications/GRCh37.nr_duplications.tsv.gz)

*Genome build GRCh38:*

[https://ftp.ncbi.nlm.nih.gov/pub/dbVar/sandbox/sv\\_datasets/nonredundant/deletions/GRCh38.nr\\_deletions.tsv.gz](https://ftp.ncbi.nlm.nih.gov/pub/dbVar/sandbox/sv_datasets/nonredundant/deletions/GRCh38.nr_deletions.tsv.gz)

[https://ftp.ncbi.nlm.nih.gov/pub/dbVar/sandbox/sv\\_datasets/nonredundant/duplications/GRCh38.nr\\_duplications.tsv.gz](https://ftp.ncbi.nlm.nih.gov/pub/dbVar/sandbox/sv_datasets/nonredundant/duplications/GRCh38.nr_duplications.tsv.gz)

These 2 files will be computed then removed the first time AnnotSV will be executed after the update.

#### e) [DECIPHER GENE ANNOTATIONS](#)

##### **Aim:**

The [Deciphering Developmental Disorders \(DDD\) Study](#) (Firth, et al., 2011) has recruited nearly 14,000 children with severe undiagnosed developmental disorders, and their parents from around the UK and Ireland. The patients have been deeply phenotyped by their referring clinician via DECIPHER using the Human Phenotype Ontology. The DNA from these children have been explored using high resolution exon-arrayCGH and exome sequencing (trio) to investigate the genetic causes of their abnormal development. These annotations give additional information on each gene overlapped by a SV (independently of the genome build version).

##### **Annotation columns:**

Adds 5 annotation columns: "DDD\_status", "DDD\_mode", "DDD\_consequence", "DDD\_disease", "DDD\_pmids".

##### **Updating the data source (if needed):**

- Remove all the **DDG2P** files in the "\$ANNOTSV/Annotations/DDD" directory.
- Download and place the "**DDG2P.csv.gz**" DECIPHER file in the "\$ANNOTSV/Annotations/DDD" directory. The latest update of this file is available for free download at:  
<http://www.ebi.ac.uk/gene2phenotype/downloads/>

This file will be computed the first time AnnotSV will be executed after the update.

##### **Warning:**

This update requires the "csv" Tcl package.

#### f) [DECIPHER FREQUENCY ANNOTATIONS](#)

##### **Aim:**

AnnotSV takes advantage of the DDD study (national blood service controls + generation Scotland controls), representing the 845 samples currently available (an update is planned in the near future).

##### **Method:**

By default, a DDD CNV is reported only if it overlaps at least 70% of the SV to annotate. Nevertheless, the user can modify the default behaviour by either use a different percentage or a reciprocal overlap (see "SVfromDBoverlap" and "SVtoAnnOverlap" options in USAGE/OPTIONS).

##### **Annotation columns:**

Adds 5 annotation columns: "DDD\_SV", "DDD\_DUP\_n\_samples\_with\_SV", "DDD\_DUP\_Frequency", "DDD\_DEL\_n\_samples\_with\_SV", "DDD\_DEL\_Frequency".

##### **Updating the data source (if needed):**

- Remove all the files in the "\$ANNOTSV/Annotations/DDD/GRCh37" directory.
- Download and place the "**population\_cnv.txt.gz**" DECIPHER files in the "\$ANNOTSV/Annotations/DDD/GRCh37" directory.  
*Genome build GRCh37:*  
The latest update of this file is available for free download at:

[https://decipher.sanger.ac.uk/files/downloads/population\\_cnv.txt.gz](https://decipher.sanger.ac.uk/files/downloads/population_cnv.txt.gz)

*Genome build GRCh38:*

**The dataset is not yet available from the DGV team.**

This file will be computed the first time AnnotSV will be executed after the update.

#### g) [1000 GENOMES ANNOTATIONS](#)

##### **Aim:**

The goal of the [1000 Genomes Project](#) (Sudmant, et al., 2015) was to find most genetic variants with frequencies of at least 1% in the populations studied. Analyses were conducted looking at both the short variations (up to 50 base pairs in length) and also the SV. These annotations give additional information on the SV allele frequencies from the 1000 genomes database overlapped by a SV to annotate.

##### **Method:**

By default, a 1000g SV is reported only if it overlaps at least 70% of the SV to annotate. Nevertheless, the user can modify the default behaviour by either use a different percentage or a reciprocal overlap (see "SVfromDBoverlap" and "SVtoAnnOverlap" options in USAGE/OPTIONS).

##### **Annotation columns:**

Adds 3 annotation columns: "1000g\_event", "1000g\_AF" and "1000g\_max\_AF".

##### **Updating the data source (if needed):**

- Remove all the **1000g** files in the "\$ANNOTSV/Annotations/1000g/GRCh37" and/or "\$ANNOTSV/Annotations/1000g/GRCh38" directories.
- Download and place the VCF files in the "\$ANNOTSV/RefSeq/GRCh37" and/or "\$ANNOTSV/RefSeq/GRCh38" directories. The latest updates of these files are available for free download at:

*Genome build GRCh37:*

[ftp://ftp.1000genomes.ebi.ac.uk/vol1/ftp/phase3/integrated\\_sv\\_map/ALL.wgs.mergedSV.v8.20130502.svs.genotypes.vcf.gz](ftp://ftp.1000genomes.ebi.ac.uk/vol1/ftp/phase3/integrated_sv_map/ALL.wgs.mergedSV.v8.20130502.svs.genotypes.vcf.gz)

[2.svs.genotypes.vcf.gz](#)

*Genome build GRCh38:*

[http://ftp.1000genomes.ebi.ac.uk/vol1/ftp/phase3/integrated\\_sv\\_map/supporting/GRCh38\\_positions](http://ftp.1000genomes.ebi.ac.uk/vol1/ftp/phase3/integrated_sv_map/supporting/GRCh38_positions)

[/ALL.wgs.mergedSV.v8.20130502.svs.genotypes.GRCh38.vcf.gz](#)

This file will be computed the first time AnnotSV will be executed after the update.

#### h) [GC CONTENT ANNOTATIONS](#)

##### **Aim:**

GC content (as well as repeated sequences, DNA sequence identity and concentration of the PRDM9 homologous recombination hotspot motif 5'-CCNCCNTNNCCNC-3') is positively correlated with the frequency of nonallelic homologous recombination (NAHR). Indeed, NAHR hot spots have a significantly higher GC content (Dittwald, et al., 2013). This information with others could help identifying a novel locus for recurrent NAHR-mediated SV.

##### **Method:**

The GC content is calculated using bedtools around each SV breakpoint (+/- 100bp) then reported.

**Annotation columns:**

Adds 2 annotation columns: "GCcontent\_left", "GCcontent\_right"

**Updating the data source (if needed):**

AnnotSV needs the human reference genome FASTA file to run the "bedtools nuc" command.

- Remove all the files in the "\$ANNOTSV/Annotations/GCcontent/GRCh37" and/or "\$ANNOTSV/Annotations/GCcontent/GRCh38" directories.
- Download and place the human reference genome FASTA file in the "\$ANNOTSV/Annotations/GCcontent/GRCh37" and/or "\$ANNOTSV/Annotations/GCcontent/GRCh38" directories. The latest update of this file is available for free download at:

*Genome build GRCh37:*

<http://hgdownload.cse.ucsc.edu/goldenPath/hg19/bigZips/chromFa.tar.gz>

*Genome build GRCh38:*

<http://hgdownload.cse.ucsc.edu/goldenPath/hg38/bigZips/hg38.chromFa.tar.gz>

This FASTA file will be reprocessed during the first time AnnotSV will be executed after the update.

**Warning:**

This update requires the "tar" Tcl package.

i) [REPEATED SEQUENCES ANNOTATIONS](#)

**Aim:**

Repeated sequences (as well as GC content, DNA sequence identity and presence of the PRDM9 homologous recombination hotspot motif 5'-CCNCCNTNCCNC-3') play a major role in the formation of structural variants.

**Method:**

The overlapping repeats are identified using bedtools at the SV breakpoint (+/- 100bp) and reported (coordinates and type).

**Annotation columns:**

Adds 2 annotation columns: "Repeats\_coord" and "Repeats\_type"

**Updating the data source (if needed):**

AnnotSV needs a UCSC Repeat BED file.

- Remove all the files in the "\$ANNOTSV/Annotations/Repeat/GRCh37" and/or "\$ANNOTSV/Annotations/Repeat/GRCh38" directories.
- You can freely download the BED file from the "http://genome.ucsc.edu/cgi-bin/hgTables". There are many output options, here are the changes that you'll need to make:

"GRCh37" or "GRCh38" assembly, "Repeats" group and "Repeatmasker" track. Select output format as BED. Choose the following output filename: Repeat.bed. Then, click the get output button.

- Download and place the BED file in the "\$ANNOTSV/Annotations/Repeat/GRCh37" and/or "\$ANNOTSV/Annotations/Repeat/GRCh38" directories.

This BED file will be reprocessed during the first time AnnotSV will be executed after the update.

#### j) TAD BOUNDARIES ANNOTATIONS

##### **Aim:**

The spatial organization of the human genome helps to accommodate the DNA in the nucleus of a cell and plays an important role in the control of the gene expression. In this nonrandom organization, topologically associating domains (TAD) emerge as a fundamental structural unit able to separate domains and define boundaries. Disruption of these structures especially by SV can result in gene misexpression (Lupianez, et al., 2016).

##### **Method:**

A TAD boundary is reported only if the SV overlaps at least 70% of this TAD boundary (user defined, see the "FeaturesOverlap" option in USAGE/OPTIONS).

##### **Annotation columns:**

Adds 2 annotation columns ("TADcoordinates", "ENCODEexperiments"), containing i) the overlapping TAD coordinates with a SV and ii) the ENCODE experiments from which the TAD have been defined.

Very large SV (e.g. 30Mb) can sometime overlap too many TAD locations (e.g. more than 2600). It appears that depending on the visualisation program used (spreadsheet programs mostly) this annotation can be truncated. In order to avoid such embarrassing glitch and maybe also because overlapping so many TAD is already a problem, AnnotSV restrict the number of overlapping reported TAD to 20 (including their associated ENCODE experiments).

##### **Updating the data source (if needed):**

AnnotSV needs ENCODE experiments in BED format for the TAD annotations.

- Remove all the files in the "\$ANNOTSV/Annotations/TAD/GRCh37" and/or "\$ANNOTSV/Annotations/TAD/GRCh38" directories.
- Download and place your ENCODE BED files in the "\$ANNOTSV/Annotations/TAD/GRCh37" and/or "\$ANNOTSV/Annotations/TAD/GRCh38" directories.

These files (GRCh37 and GRCh38) are available for free download at:

[https://www.encodeproject.org/search/?type=Experiment&assay\\_title=Hi-C&files.file\\_type=bed+bed3%2B](https://www.encodeproject.org/search/?type=Experiment&assay_title=Hi-C&files.file_type=bed+bed3%2B)

Click the "bed bed3+" button on your link (else the "file.txt" is blank). Then, click the "Download" button to download a "files.txt" file that contains a list of URLs. Keep only the \*.bed URLs in your "files.txt". Then use the following command to download all the BED files in the list:

```
xargs -n 1 curl -O -L < files.txt
```

Finally, dispatch the downloaded files in either the GRCh37 or the GRCh38 directory.

These BED files will be reprocessed during the first time AnnotSV will be executed.

#### k) OMIM ANNOTATIONS

##### **Aim:**

[OMIM \(Online Mendelian Inheritance in Man\)](#) (Hamosh, et al., 2000) focuses on the relationship between phenotype and genotype. These annotations give additional information on each gene overlapped by a SV (independently of the genome build version).

**Annotation columns:**

Add 3 annotation columns: “Mim Number”, “Phenotypes”, “Inheritance”.

**Update:**

- Remove all the files in the “\$ANNOTSV/Annotations/OMIM” directory.
- Download and place the “**genemap2.txt**” OMIM file in the “\$ANNOTSV/Annotations/OMIM” directory. The latest update of this file is available for download following a registration and review process (<https://omim.org/downloads/>). It is a tab-delimited file containing OMIM's synopsis of the Human gene map including additional information such as genomic coordinates and inheritance.

I) [GENE INTOLERANCE ANNOTATIONS](#)

**Aim:**

Gene intolerance annotations from the [ExAC](#) (Lek, et al., 2016) give the significance deviation from the observed and the expected number of variants for each gene:

synZ = synonymous Z score

misZ = missense Z score

*Positive Z scores indicate gene intolerance to variation.*

pLI = score computed by the ExAC consortium

*pLI indicates the probability that a gene is intolerant to a loss of function mutation (Nonsense, splice acceptor and splice donor variants caused by SNV). ExAC consider  $pLI \geq 0.9$  as an extremely LoF intolerant set of genes.*

These annotations give additional information on each gene overlapped by a SV (independently of the genome build version).

**Annotation columns:**

Adds 3 annotation columns: “synZ”, “misZ” and “pLI”.

**Updating the data source (if needed):**

- Remove all the files in the “\$ANNOTSV/Annotations/GeneIntolerance” directory.
- Download and place the “**fordist\_cleaned\_nonpsych\_z\_pli\_rec\_null\_data.txt**” ExAC file in the “\$ANNOTSV/Annotations/GeneIntolerance” directory. The latest update of this file is available for free download at:

*Genome build GRCh37:*

[ftp://ftp.broadinstitute.org/pub/ExAC\\_release/release0.3.1/functional\\_gene\\_constraint/](ftp://ftp.broadinstitute.org/pub/ExAC_release/release0.3.1/functional_gene_constraint/)

*Genome build GRCh38:*

**The dataset is not yet available.**

This file will be reprocessed the first time AnnotSV will be executed after the update.

### m) HAPLOINSUFFICIENCY ANNOTATIONS

#### **Aim:**

Haploinsufficiency, wherein a single functional copy of a gene is insufficient to maintain normal function, is a major cause of dominant disease. As detailed in [DECIPHER](#), over 17,000 protein coding genes have been scored according to their predicted probability of exhibiting haploinsufficiency:

- High ranks (e.g. 0-10%) indicate a gene is more likely to exhibit haploinsufficiency
- Low ranks (e.g. 90-100%) indicate a gene is more likely to NOT exhibit haploinsufficiency.

This annotation give additional information on each gene overlapped by a SV (independently of the genome build version).

#### **Annotation columns:**

Add 1 annotation column: "HI\_percent".

#### **Update:**

- Remove all the files in the "\$ANNOTSV/Annotations/HI\_Predictions" directory.
- Download and place the "**HI\_Predictions\_Version3.bed.gz**" DECIPHER file in the "\$ANNOTSV/Annotations/HI\_Predictions" directory. The latest update of this file is available for free download at:

<https://decipher.sanger.ac.uk/about#downloads/data>

This file will be computed the first time AnnotSV will be executed after the update.

## 3. INPUT

AnnotSV takes several arguments as input to the command line including options that are detailed in section 5 ("USAGE / OPTIONS"). The different arguments can be passed either on the command line or using a specific file named "configfile". The configfile file needs to be located in the AnnotSV installation directory. Four types of INPUT files are detailed below:

### 3.1. SV input file (Required)

AnnotSV supports either the [VCF](#) (Variant Call Format) or the [BED](#) (Browser Extensible Data) input format to describe the SV to annotate. It allows the program to be easily integrated into any bioinformatics pipeline dedicated to NGS analysis.

- VCF is a text file format. It contains meta-information lines (prefixed with "##"), a header line (prefixed with "#"), and data lines each containing information about a position in the genome and genotype information on samples for each position (text fields separated by tabs). The specification are described at <https://samtools.github.io/hts-specs/VCFv4.3.pdf>  
AnnotSV supports either native or gzipped VCF file.

#### **WARNING:**

By default, AnnotSV extracts and reports only some informations from the VCF input file:

- The REF, ALT, FORMAT and samples columns
  - The SVTYPE value from the INFO column and only this one
- All other columns (QUAL, FILTER and INFO) can be reported by setting the "-SVinputInfo" option to 1.

- BED is a text file format. Every single line of the BED file define a SV including the obligatory 3 first fields to describe its coordinates:

1. *chrom* - The name of the chromosome (e.g. 3, Y, ...) - Preferred without "chr".
2. *chromStart* - The starting position of the SV on the chromosome. According to the format, the base count starts at base "0".
3. *chromEnd* - The ending position of the SV on the chromosome. The *chromEnd* base is not included in the display of the feature. For example, the first 100 bases of a chromosome are defined as *chromStart*=0, *chromEnd*=100, and span the bases numbered 0-99.

Additional fields from the BED file are optional and can be reported in the AnnotSV output file (user defined). It can be used to store quality, read depth or other metrics produced by the SV caller.

**WARNING:**

By default, AnnotSV does not report the additional fields from the BED input file.

All fields can be reported by setting the "-SVinputInfo" option to 1.

### 3.2. SNV/indel input files - for DELETION filtering (Optional)

AnnotSV can take VCF file(s) with SNV/indel as input to the command line.

These annotations report the counts of homozygous and heterozygous SNV/indel identified from the patients NGS data (user defined samples) and presents in the interval of the SV to annotate.

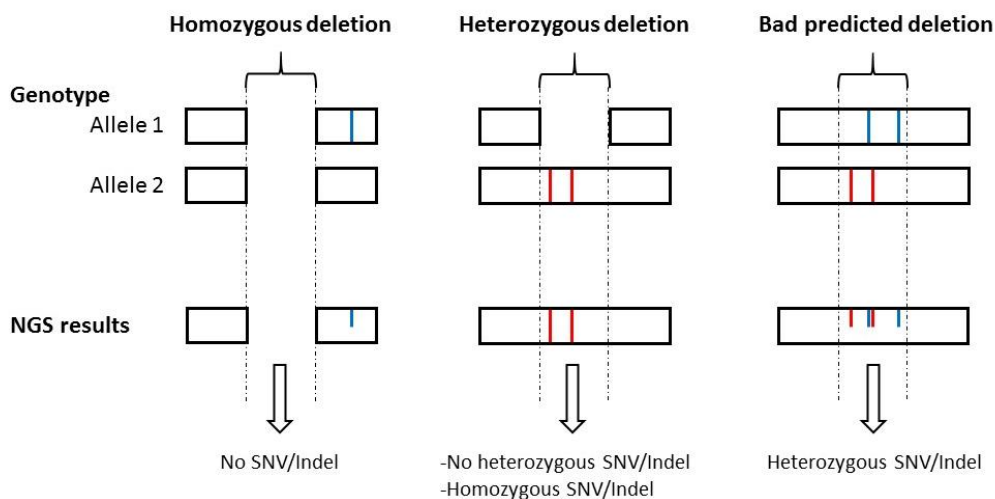
**Annotation columns/Usage:**

Add the "hom(sample)" and "htz(sample)" annotation columns.

The command line can be completed with the 2 following options: "-vcfFiles" and "-vcfSamples" (cf USAGE/OPTIONS).

**Aim:**

These annotations can be used by the user to filter out false positive SV calls or to confirm events as following:



**-Homozygous deletion** can be identified as a false positive by noting the presence of SNV/indel called at the predicted locus of the deletion in a sample.

**-Heterozygous deletion** can be identified as a false positive by noting the presence of heterozygous SNV/indel called at the predicted locus of the deletion in a sample. If no heterozygous SNV/indel are presents, the



heterozygous deletion can be confirmed by reporting the presence of homozygous SNV/indel at that locus in the sample.

**WARNING:**

In the VCF file(s), the genotype should be indicated in the format field as “GT”.

### 3.3. Filtered SNV/indel input files - for compound heterozygosity analysis (Optional)

AnnotSV can take a VCF file(s) with SNV/indel as input to the command line that is already filtered for genotype frequency and effects on protein level.

AnnotSV can report the heterozygous SNV/indel presents in the gene overlapped by the SV to annotate, as well in ‘healthy’ and ‘affected’ samples (user defined samples). This would be really useful for the user to identify compound heterozygotes with one SNV/indel and one SV.

**Usage:**

To add the “**compound-htz**” annotation column, the command line can be completed with the 2 following options: “-filteredVCFfiles” and “-filteredVCFsamples” (cf USAGE/OPTIONS).

**Background:**

In recessive genetic disorders, both copies of a certain gene are malfunctioning. This means that the maternally as well as the paternally inherited copy of an autosomal gene harbors a pathogenic mutation. And if the parents are non-consanguineous, compound heterozygosity is the best explanation for a recessive disease.

**Aim:**

AnnotSV offers an efficient filter to highlight compound heterozygous variants composed of one SV and one SNV/indel in the same gene.

In this way AnnotSV takes in input a VCF file(s) that is already filtered for genotype frequency and effects on protein level. Then, the software extracts from the input VCF file(s) the heterozygous variants (SNV/indel) presents in the gene overlapped by the SV, as well in ‘healthy’ and ‘affected’ samples (user defined samples).

**User challenge:**

The user challenge in filtering variants for compound heterozygotes is to know whether the two heterozygous variants (the SNV/indel and the SV) are in *cis* or in *trans*. And when sequencing data of more than one family member is available, one can exclude certain variants based on rules of Mendelian inheritance (transmitted in a compound heterozygous mode from parents to the patient(s)).

**WARNING:**

In the VCF file(s), the genotype should be indicated in the format field as “GT”.

### 3.4. External BED and/or TSV annotation files (Optional)

**Aim:**

Several users might want to add their own private annotations to the one already provided by AnnotSV.

**Inputs:**

AnnotSV can integrate external annotations for specific regions that will be imported from a BED and/or a TSV file into the output file.

Each external BED or TSV annotation file should be **copy or linked** in:

*Genome build GRCh37:*

→ the “\$ANNO SV/Annotations/Users/GRCh37/” directory

*Genome build GRCh38:*

→ the “\$ANNO SV/Annotations/Users/GRCh38/” directory

**WARNING:** The copy and/or linked users file(s) will be deleted the first time AnnotSV will be executed after an update.

#### **BED user file:**

Each external BED annotation file (e.g. ‘User’.bed) should be completed with a tab separated values file (e.g. ‘User’.header.tsv.) describing the header of these new annotations.

The following example has been set to provide the SV overlap with Regions of Homozygosity (RoH) of 2 individuals (sample1 and sample2):

‘User’.header.tsv file contains:

Chrom	Start	End	RoH
-------	-------	-----	-----

‘User’.bed file contains:

1	2806107	107058351	sample1, sample2
12	25687536	25699754	sample2

"RoH" annotation column is then available in the output file.

#### **TSV user file:**

Each external TSV annotation file (e.g. ‘User’.tsv) should contain a header beginning with “#”.

‘User’.tsv file contains:

#Chrom	Start	End	RoH
1	2806107	107058351	sample1, sample2
12	25687536	25699754	sample2

"RoH" annotation column is then available in the output file.

#### **Method:**

By default, a user region is reported only if it overlaps at least 70% of the SV to annotate. Nevertheless, the user can modify the default behaviour by either use a different percentage or a reciprocal overlap (see “SVfromDBoverlap” and “SVtoAnnOverlap” options in USAGE/OPTIONS).

### [\*3.5.External Gene annotation files \(Optional\)\*](#)

In order to further enrich the annotation for each SV gene, AnnotSV can integrate external annotations imported from tab separated values file(s) into the output file. The first line should be a header including a column entitled "genes".

The following example has been set to provide annotation for the interacting partners of a gene.

genes	Interacting genes
BBS1	BBS7, TTC8, BBS5, BBS4, BBS9, ARL6, BBS2, RAB3IP, BBS12, BBS10

"Interacting genes" annotation column is then available in the output file.

Each external gene annotation file (\*.tsv ) should be located in the "\$ANNOTSV/Annotations/Users/" directory. It is to notice that these files should not contain any of these 2 specific characters "{" and "}" (that would be replaced by "(" and ")").

AnnotSV supports either native or gzipped tsv file.

## 4. OUTPUT

### **Format:**

Giving a SV input file, AnnotSV produces a tab-separated values file that can be easily integrated in bioinformatics pipelines or directly read in a spreadsheet program.

### **Output file path and name:**

Two options (-outputDir and -outputFile) can be used to specify the output directory and/or file name. The output file extension should be ".tsv" (tab separated values).

By default, an output directory is created where AnnotSV is run ('YYYYMMDD'\_AnnotSV). As an example, an input SV file named "mySVinputFile.vcf" will produce by default an output file named "20180320\_AnnotSV/mySVinputFile.annotated.tsv".

### **Output lines:**

There are 2 types of lines produced by AnnotSV (cf the "AnnotSV type" output column):

- An annotation on the "full" length of the SV. Every SV are reported, even those not covering a gene. This type of annotation gives an estimate of the SV event itself.

- An annotation of the SV "split" by gene. This type of annotation gives an estimate of the gene composition of the corresponding SV and is meant to analyse the consequences more deeply. Thus, in some cases, when a SV spans over several genes, the output will contain as many annotations lines as covered genes (cf example in FAQ). This latter annotation is extremely powerful to shorten the identification of mutation in a implicating a specific gene.

-typeOfAnnotation

### 4.1.Annotation columns available in the output file

In the following table, we describe the annotations that are available in the AnnotSV output file. It is to notice that, since AnnotSV can be configured to output the annotations using 2 different modes (full or split), in some cases specific gene annotations are only present while using one of the two modes.

Column name	Annotation	Full	Split
<b>SV chrom</b>	Name of the chromosome	X	X
<b>SV start</b>	Starting position of the SV in the chromosome	X	X
<b>SV end</b>	Ending position of the SV in the chromosome	X	X
<b>REF</b>	Nucleotide sequence in the reference genome (extracted only from a VCF input file)	X	X
<b>ALT</b>	Alternate nucleotide sequence (extracted only from a VCF input file)	X	X
<b>SVTYPE</b>	Type of the SV (extracted only from a VCF input file)	X	X

<b>FORMAT</b>	The FORMAT column from a VCF file	X	X
<b>Sample ID</b>	The sample ID column from a VCF file	X	X
<b>AnnotSV type</b>	Indicate the type of annotation generated: - annotation on the SV full length ("full") - annotation on each gene overlapped by the SV ("split")	X	X
<b>Gene name</b>	Gene symbol	X	X
<b>NM</b>	Transcript symbol <sup>1</sup>		X
<b>CDS length</b>	Length of the CoDing Sequence (CDS) (bp) overlapping with the SV		X
<b>tx length</b>	Length of the transcript (bp) overlapping with the SV		X
<b>location</b>	SV location in the gene (e.g. « txStart-exon1 », « intron3-exon7 »)		X
<b>intersectStart</b>	Start position of the intersection between the SV and the transcript		X
<b>intersectEnd</b>	End position of the intersection between the SV and the transcript		X
<b>promoters</b>	List of the genes whose promoters are overlapped by the SV	X	X
<b>DGV_GAIN_IDs</b>	DGV Gold Standard GAIN IDs overlapped with the annotated SV	X	X
<b>DGV_GAIN_n_samples_with_SV</b>	Number of individuals with a shared DGV_GAIN_ID	X	X
<b>DGV_GAIN_n_samples_tested</b>	Number of individuals tested	X	X
<b>DGV_GAIN_Frequency</b>	Relative GAIN Frequency=DGV_GAIN_n_samples_with_SV/DGV_GAIN_n_samples_tested	X	X
<b>DGV_LOSS_IDs</b>	DGV Gold Standard LOSS IDs overlapped with the annotated SV	X	X
<b>DGV_LOSS_n_samples_with_SV</b>	Number of individuals with a shared DGV_LOSS_ID	X	X
<b>DGV_LOSS_n_samples_tested</b>	Number of individuals tested	X	X
<b>DGV_LOSS_Frequency</b>	Relative LOSS Frequency=DGV_LOSS_n_samples_with_SV/DGV_LOSS_n_samples_tested	X	X
<b>DDD_SV</b>	Deciphering Developmental Disorders (DDD) SV coordinates from the DDD study (data control sets) overlapped with the annotated SV	X	X
<b>DDD_DUP_n_samples_with_SV</b>	Number of individuals with a shared DDD_DUP	X	X
<b>DDD_DUP_Frequency</b>	DUP Frequency	X	X
<b>DDD_DEL_n_samples_with_SV</b>	Number of individuals with a shared DDD_DEL	X	X
<b>DDD_DEL_Frequency</b>	DEL Frequency	X	X
<b>DDD_status</b>	Deciphering Developmental Disorders (DDD) category e.g. confirmed, probable, possible, ...	X	X
<b>DDD_mode</b>	Deciphering Developmental Disorders (DDD) allelic requirement e.g. biallelic, hemizygous, ...	X	X
<b>DDD_consequence</b>	Deciphering Developmental Disorders (DDD) mutation consequence e.g. "loss of function", uncertain, ...	X	X
<b>DDD_disease</b>	Deciphering Developmental Disorders (DDD) disease name e.g. "OCULOauricular syndrome"	X	X
<b>DDD_pmids</b>	Deciphering Developmental Disorders (DDD) pmids	X	X
<b>1000g_event</b>	1000 genomes event types (e.g. DEL, DUP, ALU, <CN3>...)	X	X
<b>1000g_AF</b>	1000 genomes allele frequency	X	X
<b>1000g_max_AF</b>	Maximum observed allele frequency across the 1000 genomes populations	X	X
<b>dbVar_event</b>	dbVar NR SV event types (e.g. deletion, duplication...)	X	X
<b>dbVar_variant</b>	dbVar NR SV accession (e.g. nssv1415016)	X	X
<b>dbVar_status</b>	dbVar NR SV clinical assertion (e.g. pathogenic, likely pathogenic)	X	X
<b>synZ</b>	Positive synZ (Z score) indicate gene intolerance to synonymous variation		X

<b>misZ</b>	Positive misZ (Z score) indicate gene intolerance to missense variation		X
<b>pLI</b>	Score computed in the ExAC database indicating the probability that a gene is intolerant to a loss of function variation (Nonsense, splice acceptor and donor variants caused by SNV). ExAC consider pLI >= 0.9 as an extremely LoF intolerant set of genes		X
<b>HI_percent</b>	Haploinsufficiency ranks		X
<b>Mim Number</b>	OMIM unique six-digit identifier		X
<b>Phenotypes</b>	e.g. Charcot-Marie-Tooth disease		X
<b>Inheritance</b>	e.g. AD (= "Autosomal dominant") <sup>2</sup>		X
<b>GCcontent_left</b>	GC content around the left SV breakpoint (+/- 100bp)	X	
<b>GCcontent_right</b>	GC content around the right SV breakpoint (+/- 100bp)	X	
<b>Repeats_coord_left</b>	Repeats coordinates around the left SV breakpoint (+/- 100bp)	X	
<b>Repeats_type_left</b>	Repeats type around the left SV breakpoint (+/- 100bp) e.g. AluSp, L2b, L1PA2, LTR12C, SVA_D, ...	X	
<b>Repeats_coord_right</b>	Repeats coordinates around the right SV breakpoint (+/- 100bp)	X	
<b>Repeats_type_right</b>	Repeats type around the right SV breakpoint (+/- 100bp) e.g. AluSp, L2b, L1PA2, LTR12C, SVA_D, ...	X	
<b>TADcoordinates</b>	Coordinates of the TAD whose boundaries overlapped with the annotated SV (boundaries included in the coordinates)	X	
<b>ENCODEexperiments</b>	ENCODE experiments from where the TAD have been defined	X	
<b>compound-htz(sample)</b>	List of heterozygous SNV/indel (reported with "chrom_position") presents in the gene overlapped by the annotated SV	X	X
<b>hom(sample)</b>	Number of homozygous variants in the individual "sample" which are presents: - in the SV for the "full" annotation - between intersectStart and intersectEnd for the "split" annotation. Values are extracted from the input VCF file(s)	X	X
<b>htz(sample)</b>	Number of heterozygous variants in the individual "sample" which are presents: - in the SV for the "full" annotation - between intersectStart and intersectEnd for the "split" annotation. Values are extracted from the input VCF file(s)	X	X

<sup>1</sup>Given one gene, only a single transcript from all transcripts available in RefSeq is reported. In case of transcripts with different CDS length (considering the overlapping region with the SV), the transcript with the longest CDS is reported. Otherwise, if there is no differences in CDS length, the longest transcript is reported.

<sup>2</sup>Detailed in the FAQ

## 5. USAGE / OPTIONS

To run AnnotSV, the default command line is the following:

```
$ANNOTSV/bin/AnnotSV -SVinputFile '/Path/Of/Your/VCF/or/BED/Input/File' >& AnnotSV.log &
```

The command line can be completed by the list of options described below or modified in the configfile. To show the options simply type:

```
$ANNOTSV/bin/AnnotSV -help or $ANNOTSV/bin/AnnotSV
```

OPTIONS:

-----

-SVinputFile: Path of the input file (VCF or BED) with SV coordinates

Gzipped VCF file is supported

- SVinputInfo:** To extract the additional SV input fields and insert the data in the output file  
Range values: 0 (default) or 1
- bedtools:** Path of the bedtools local installation
- FeaturesOverlap:** Minimum overlap (%) of the features (promoter, TAD...) with the annotated SV to be reported  
Range values: [0-100], default = 70
- filteredVCFfiles:** Path of the filtered VCF input file(s) with SNV/indel coordinates for compound heterozygotes report (optional)  
Gzipped VCF files are supported as well as regular expression
- filteredVCFsamples:** To specify the sample names from the VCF files defined from the -filterVCFfiles option  
Default: use all samples from the filtered VCF files
- genomeBuild:** Genome build used  
Values: GRCh37 (default) or GRCh38
- help:** More information on the arguments
- outputDir:** Output path name
- outputFile:** Output path and file name
- promoterSize:** Number of bases upstream from the transcription start site  
Default = 500
- SVfromDBoverlap:** Minimum overlap (%) of the SV from external databases (DGV, DDD) with the annotated SV to report the features  
Range values: [0-100], default = 0
- SVminSize:** SV minimum size (in bp)  
Default = 50
- SVtoAnnOverlap:** Minimum overlap (%) of the annotated SV with the SV from external databases (DGV, DDD...) to report the features  
Range values: [0-100], default = 70
- typeOfAnnotation:** Description of the types of lines produced by AnnotSV  
Values: both (default), full or split
- vcfFiles:** Path of the VCF input file(s) with SNV/indel coordinates used for false positive discovery  
Use counts of the homozygous and heterozygous variants  
Gzipped VCF files are supported as well as regular expression
- vcfPASS:** Boolean. To only use variants from VCF input files that passed all filters during the calling (FILTER column value equal to PASS)  
Range values: 0 (default) or 1

-vcfSamples: To specify the sample names from the VCF files defined from the -vcfFiles option  
Default: use all samples from the VCF files

## 6. [Test](#)

In order to validate the AnnotSV installation and its functioning, an example is available in the “\$ANNOTSV/Example” directory. Command lines examples are available in the following file “\$ANNOTSV/Example/commands.README”.

Moreover, an input/output example (the HG00096 individual from the 1000 Genomes project) is available on the [AnnotSV website](#).

## 7. [FAQ](#)

### **Q: What are Structural Variations (SV)?**

SV are generally defined as variation in a DNA region that vary in length from ~50 base pairs to many megabases and include several classes such as translocations, inversions, insertions, deletions.

### **Q: What are Copy Number Variations (CNV)?**

CNV are deletions and duplications in the genome (unbalanced SV) that vary in length from ~50 base pairs to many megabases.

### **Q: What are the differences between SV and CNV?**

CNV are unbalanced SV with gain or loss of genomic material. For example, a heterozygous duplication as a CNV will be characterized with the start and end coordinates and the number of copies which is 3.

### **Q: Can AnnotSV annotate every type of SV?**

AnnotSV supports as well VCF or BED format in input.

- VCF format supports complex rearrangements with breakends, that can arbitrary be summarized as a set of novel adjacencies, as described in the Variant Call Format Specification [VCFv4.3](#) (Jul 2017).
- BED format doesn't allow inter-chromosomal feature definitions (e.g. inter-chromosomal translocation). A new file format (BEDPE) is proposed in order to concisely describe disjoint genome features but it is not yet supported by AnnotSV.

### **Q: I would like to annotate my SV with new annotation sources but I don't know how to do that...**

No problem. AnnotSV is under active and continuous development. You can email me with a detailed request and I will answer as quickly as possible.

### **Q: I have just updated AnnotSV or the annotations sources and the annotation process is longer than usual, is it normal?**

After an update of AnnotSV sources, some files will be reprocessed and thus taking several additional time. Further use of AnnotSV will be quicker!

### **Q: How to cite AnnotSV in my work?**

If you are using AnnotSV, please cite our work using the following reference:

**AnnotSV: An integrated tool for Structural Variations annotation.** Geoffroy V, Herenger Y, Kress A, Stoetzel C, Piton A, Dolfus H, Muller J. Bioinformatics. 2018 Apr 14. doi: [10.1093/bioinformatics/bty304](https://doi.org/10.1093/bioinformatics/bty304)

**Q: What are the WARNINGS that AnnotSV mention while running?**

AnnotSV writes to the standard output progress of the analysis including warnings about issues or missing information that can be either blocking or simply informative.

**Q: Why are some values empty in the output files?**

When no information is available for a specific type of annotation, then the value is empty.

**Q: Why can we have several gene annotations for one SV?**

In some cases, one SV overlaps a large portion of the genome including several genes. In these cases, the annotation of the SV is split on several lines.

Annotation example for the deletion 1:16892807-17087595

AnnotSV keep all gene annotations, with only one transcript annotation for each gene:

1	16892807	17087595	DEL	CROCCP2	NR_026752	1	12652	txStart-txEnd
1	16892807	17087595	DEL	ESPNP	NR_026567	1	28941	txStart-txEnd
1	16892807	17087595	DEL	FAM231A	NM_001282321	511	511	txStart-txEnd
1	16892807	17087595	DEL	FAM231C	NM_001310138	511	656	txStart-txEnd
1	16892807	17087595	DEL	LOC102724562	NR_135824	1	2998	txStart-txEnd
1	16892807	17087595	DEL	MIR3675	NR_037446	1	75	txStart-txEnd
1	16892807	17087595	DEL	MST1L	NM_001271733	2015	6468	txStart-exon14
1	16892807	17087595	DEL	MST1P2	NR_027504	1	4848	txStart-txEnd
1	16892807	17087595	DEL	NBPF1	NM_017940	2912	47294	intron3-txEnd

**Q: I am confused with the "AnnotSV type" in the AnnotSV output. The program returned two (or more than two) lines for each CNV segments, each with a "AnnotSV type" coded as "full" or "split". I am not sure what does the full and split means, and why the following columns show different messages between lines of the same segment.**

AnnotSV constructs an annotation based on the full-length SV (and so the AnnotSV type column is set to "full") but also an annotation for each gene within the SV (and so the AnnotSV type column is set to "split"). You will so have access to :

- all the overlapped genes information (ID, OMIM...)
- the SV location within each overlapped gene (e.g. "exon3-intron11", "txStart-intron19", ...). You could so determine fusion, truncation events...

If a SV overlaps only 1 gene, you may have only 1 "split" line.

But in the case of a SV overlapping several genes, the "split" lines are really useful by reporting one annotation line for each overlapped gene.

Depending of your analyses, you can choose to keep only the full annotation lines thanks to the "-typeOfAnnotation" option.

**Q: Why some SV have empty gene annotation in the output file?**

If a SV is located in an intergenic region and so doesn't cover a gene, then the SV is reported in the output file but without gene annotation.

**Q: What do the OMIM Inheritance annotations mean?**

- AD = "Autosomal dominant"
- AR = "Autosomal recessive"
- XLD = "X-linked dominant"
- XLR = "X-linked recessive"
- YLD = "Y-linked dominant"
- YLR = "Y-linked recessive"
- XL = "X-linked"
- YL = "Y-linked"



**Q: What is the overlap used by AnnotSV between an annotated SV and features?**

*Concerning DGV, DDD, 1000g and user defined regions:*

By default, one of these regions is reported only if it overlaps at least 70% of the SV to annotate. Nevertheless, the user can modify the default behaviour by either use a different percentage or a reciprocal overlap (see "SVfromDBoverlap" and "SVtoAnnOverlap" options in USAGE/OPTIONS).

*Concerning TAD boundaries and promoters regions:*

By default, one of these regions is reported only if the SV overlaps at least 70% of this region (user defined, see the "FeaturesOverlap" option in USAGE/OPTIONS).

**Q: Why do I get this error message: "Feature (10:134136286-134136486) beyond the length of 10 size (133797422 bp). Skipping."**

One possibility is that you are using the bad "-genomeBuild" option.

For example, you are using a bedfile in input with the SV coordinates on GRCh37 but with the "-genomeBuild GRCh38" option.

**Q: How to interpret the presence of my SV in DGV or DDD databases?**

DGV is populated with healthy samples whereas DDD is presenting affecting patients.

The presence of a SV from your sample in DGV or DDD does not necessarily imply a disease causing event. Healthy carriers of pathogenic SV do exist in either databases. When available allele frequency can be helpful to decide on the status.

**Q: Is AnnotSV available for other organisms?**

The main objective of AnnotSV is to annotate SV information from human data. All the annotations are based on human specific databases. Nevertheless, some files can be modified with the proper dataset but this is not currently supported.

**Q: Is there an option to just generate SV "split" by gene?**

You can choose to keep only the split annotation lines thanks to the "-typeOfAnnotation" option.

**Q: I am unable to run the code on the input files provided. It crashes on the Repeat annotation step due to a bad\_alloc error. Do you have any ideas on why this is happening?**

AnnotSV needs to be run with an appropriate RAM (depending of the annotations used). Setting your system to allocate 10 Go should solve the problem.

**Q: I'm getting the error: "ANNOTSV environment variable not specified. Please define it before running AnnotSV. Exit". How can I fix this problem?**

ANNOTSV is the environment variable defining the installation path of the software.

- In csh, you can define it with the following command line:  
setenv ANNOTSV /path\_of\_AnnotSV\_installation/bin
- In bash, you can define it with the following command line:  
Export ANNOTSV=/path\_of\_AnnotSV\_installation/bin

I advise you to save the good command in your .cshrc or .bashrc file.

**Q: My annotated SV is intersecting both a benign SV and a pathogenic SV. How can I explain that?**

Here is a corresponding use-case:

	Positions	ID
SV to annotate	1:27804112-27804450	

<b>Benign SV (from DGV)</b>	1:27803915-27804641	gssvL1678
<b>Pathogenic SV (from NR SV)</b>	1:27133503-28011702	nssv577197

Several possible explanations can be considered:

- The pathogenicity can concern a recessive disease. So the pathogenic SV can be present in the heterozygous state in the healthy population (with a DGV low frequency)
- The pathogenic region of the dbVar SV is not overlapping the DGV SV

## 8. REFERENCES

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