Gene-centered View of Human Structural Variants -GeneVar

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Methods

Module 1: SV<-> gene

In: dbVar + Gencode Out TSV: **var_id**, gene_id

Module 2: Gene impact

In: dbVar + Gencode

Out TSV: *var_id*, *gene_id*, *elt_type*

Module 3: SV<-> clinical SNV/indels

In: dbVar + (ClinVar or custom variant set)
Out TSV: var_id, clinvar_pathogenic_snv

Module 4: Allele frequency

In: dbVar + gnomAD-SV Out TSV: **var_id**, af

Module 5: SV<-> clinical SVs

In: dbVar + (ClinVar or ClinGen)
Out TSV: var_id, clinvar_pathogenic_sv

Module 6: External resources

In: gene_id + OMIM + ... Out TSV: **gene_id**, omim_url App locally or online Status: *prototype*

Module N: Awesome info

In: awesome DB

Out TSV: *variant_id/gene_id*, awesome_info

Tools

- SnpEff
- AnnotSV
- bedtools
- Custom R or python scripts
- R-Shiny

Resources

- dbVar
- Gencode
- gnomAD-SV
- OMIM
- ClinVar
- ClinGen
- QC tracks from UCSC
- GWAS catalog SNVs
- ...

App prototype

