

Genome-Wide Association Study

[Genome-Wide Association Study](#) (GWAS) is a hypothesis-free method for identifying associations between genetic regions and traits. GWAS analysis are usually used to identify genes involved in human disease. By applying GWAS analysis to variant data we will be able to identify a given variant (or a set of variants) involved in a given phenotype or disorder. Based on a statistical test, GWAS analysis will provide a level of significance (or p-value) for each variant. OpenCGA implements GWAS analysis based on the statistical tests: chi-square and Fisher.

Implementation

OpenCGA GWAS analysis extends Oskar GWAS implementation. GWAS is implemented using Hadoop MapReduce over HBase.

Input

Parameters

OpenCGA support different input parameters:

- Variant data with sample genotypes
- Two list of samples (case-control study)
- Statistical test: chi square or fisher.

Instead of providing two lists of samples, users can provide:

- A phenotype, and
- A pedigree for those variant samples

Configuration

OpenCGA implementation supports different configuration variables. This can be setup in the OpenCGA installation folder or specified during execution.

Output

Files

GWAS analysis result includes a text file with score and plot image (WIP)

Text file that consists of a header line (starting with #), and then one line per variant with the following 12-13 columns:

chromosome	Chromosome code
start	Start base-pair coordinate
end	End base-pair coordinate
strand	Strand
reference	Reference allele
alternate	Alternate allele
dbSNP	Variant identifier
gene	Gene name
biotype	Biotype
conseq. types	List of consequence types
<i>chi square</i>	Allelic test chi-square statistic. <i>Not present with 'fisher' test.</i>
p-value	Allelic test p-value
odd ratio	Odd ratio: odds(allele 1 case) / odds(allele 1 control)

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Useful Links

- [GWAS Study](#)

Next, it shows the first lines of a result file after executing a GWAS analysis using the chi square test:

```
#chromosome      start      end      strand      reference
alternate      dbSNP      gene      biotype      consequ.
types      chi square      p-value      odd ratio
22      16054454      16054454      +      C      T
rs373998521      intergenic_variant
2.47272727272727      0.11583677431831574      0.0
22      16065809      16065809      +      T
C      ENSG00000233866      lincRNA
downstream_gene_variant      0.053968253968253915
0.8162967146689325      0.8
22      16065809      16065809      +      T
C      regulatory_region_variant
0.053968253968253915      0.8162967146689325      0.8
22      16077310      16077310      +      T
A      ENSG00000229286      unprocessed_pseudogene
2KB_upstream_variant      0.9714285714285711
0.3243241555798487      3.0
22      16077310      16077310      +      T
A      regulatory_region_variant
0.9714285714285711      0.3243241555798487      3.0
22      16080499      16080499      +      A      G
rs200119791      ENSG00000229286      unprocessed_pseudogene
upstream_gene_variant      1.8888888888888886
0.16932729721206297      Infinity
22      16080499      16080499      +      A      G
rs200119791      ENSG00000235265      unprocessed_pseudogene
downstream_gene_variant      1.8888888888888886
0.16932729721206297      Infinity
22      16084621      16084621      +      T
C      ENSG00000235265      unprocessed_pseudogene
non_coding_transcript_exon_variant,non_coding_transcript_variant
2.4425287356321843      0.11808572685033702      Infinity
```

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Score can be indexed.

Useful Links

- https://en.wikipedia.org/wiki/Genetic_association
- https://en.wikipedia.org/wiki/Genome-wide_association_study
- <https://www.cog-genomics.org/plink/1.9/assoc>