# genewiseR FORMATICS

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#### **ABSTRACT**

#### Motivation:

R is the language of choice for statistical analysis in genomic research. This is because of its open source nature and the advance collection of R packages for statistical tests and visualisation of data. In GWAS studies, pooled sequencing is gaining popularity. Many tools exist to work with next-gen sequencing data in a gwas study; Bioconductor and Python are the go to places. Very few packages and tools exist for pooled seq gwas in R. This is mainly because of slow speed and memory constraints for R when working with large datasets. New high performance computing techniques in R have enabled the fullfilment of this gap. Works well on desktop and cluster configurations. Open source. Fast and intuitive methods for biology scientist to perform gwas analysis.

**Results:** The availablity of genewiseR on github makes it easy for scientists to install the package in their development environment is perform gwas analysis with ease. Because of addition on boostrapping feature for p-values of snps, this package allow scientist to indentify noise in their data and reduce the dataset early on in their gewas analysis.

**Availability:** genewiseR is open source and easily downloadable at github here. The

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1 INTRODUCTION

Genome

- what is gwas?
- why pool seq and why new skills and large datasets?
- why snps to genes?
- study design to make use of genewiseR

Text. Figure 1 shows that the above method Text Text Text Text Text Text Text. (Bag *et al.*, 2001) wants to know about text follows.

$$\sum x + y = Z \tag{1}$$

Table 1. This is table caption

head1	head2	head3	head4
row1	row1	row1	row1
row2	row2	row2	row2
row3	row3	row3	row3
row4	row4	row4	row4

This is a footnote

Fig. 1. Caption, caption.

Fig. 2. Caption, caption.

## 2 DESCRIPTION

#### 2.1 Input Data

The primary input data for this package is filtered sync files of Pooled Sequencing Data. The reads of pool-seq experiment are mapped to reference genome and then converted to multi-pileup(mpileup) format using 'Samtools' (?). The mpileup format is converted to sync using Popoolation2 (?). For detailed description refer [online documentation link here].

- Hypothesis testing, QC with bootstrapping
- Multiple testing correction
- Snps to Genes
- · Calc gene level statistics
- LD assesment
- Utilities required by biologist
- Pathway analysis with Revigo/David

2.1.1 Parallelisation and Performance Sequenning pool of individuals are cheaper than individuals but Bofelli et al., 2000

# 3 EXAMPLE

### 4 VALIDATION OR CONCLUSION

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Text Text Text Text Text Text Text. Bofelli et al., 2000 might want to know about text text text text

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