

Genetics and population analysis

MultiGWAS: A tool for GWAS analysis on tetraploid organisms by integrating results of four GWAS software

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Abstract

Summary: At present, genome-wide association studies (GWAS) are increasingly common to analyze non-model organisms that are important for agriculture as polyploid crops. A critical aspect in these studies is the importance to replicate the analysis, and one way to do this task is by using different tools to validate the accuracy of the associations. Currently, software for GWAS in polyploids is scarce, but recent advances in this area along with widely used diploid software can be used to replicate GWAS analyses. However, each software has its own characteristics (interface, inputs, outputs, and arguments) which can may cost time and effort to successfully use them. Here, we present MultiGWAS, a tool to do GWAS analysis in tetraploid organisms by executing in parallel and integrating the results from four existing GWAS software: two for polyploids (GWASpoly and SHEsis) and the other two for diploids (Plink and Tassel). The tool deals with all the matters of the GWAS process in the four software, uses different control quality filters for the genomic data, and allows the execution of two GWAS models: Full and Naive, the first with control for population structure and individual relatedness, and the second without any control. The summary reports generated by MultiGWAS provide the user with tables and plots describing intuitively the markers found by each tool and by more than one tool, which help users to check for potential true or false associations.

Availability and implementation: Source code, examples, documentation and installations instructions are available at <https://github.com/agrosavia/multiGWAS>

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1 Introduction

Genome-wide association studies (GWAS) allows to analyze genomic data to identify the set of variants across different individuals of a species that are associated with a phenotypic characteristic of research interest. Due to the advances in the next-gen sequencing technology, GWAS analysis are currently more frequently used in non-model species as plants which include polyploid crops, that are important for agriculture and the food security of different developing countries.

One of the main challenges in these GWAS analysis is to identify true from false associations, which can be validated by replicating

the analysis using different tools. However, each tool has its own characteristics with different user interfaces (GUI or command line based), genotype/phenotype formats, models and algorithm assumptions, and different outputs, all of which take great effort to consider and makes it difficult for researches to do the replication.

The two most strong tools to perform GWAS across different organisms are Plink (Purcell *et al.*, 2007) and Tassel (Bradbury *et al.*, 2007). However, both are limited to diploid species and to be used for polyploids, the genomic data need to be "diploidized" (Lindqvist-Kreuzer *et al.*, 2014; Schulz *et al.*, 2016). Fortunately, in 2016 were published two software

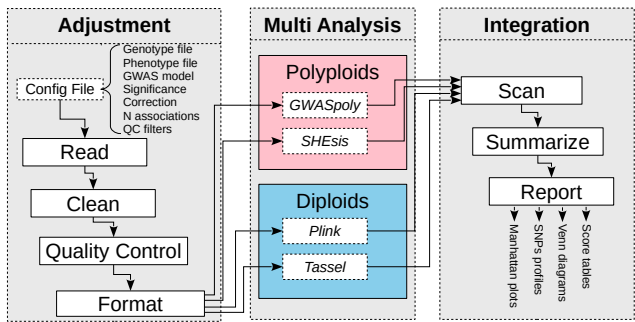


Fig. 1. Flowchart of the central steps in the MultiGWAS tool. The analysis is carried out by in three stages: adjustment, multi analysis, and integration. In the first stage, inputs are read from a configuration file (see below) and the genotype and phenotype are cleaned and filtered using the quality control (QC) filters. In the second stage, the new filtered genotype/phenotype are formatted for each GWAS tool which are executed in parallel. In the last stage, the output files generated from each tool are scanned, and their results are summarized and reported through score tables, Venn diagrams, SNP profiles, and Manhattan plots. The configuration file includes: the genotype/phenotype filenames, genome-wide significance threshold, multiple testing correction method, GWAS model, number of associations to be reported, and TRUE or FALSE whether to use QC filters or not. The QC filters are: minor allele frequency, individual missing rate, SNP missing rate, Hardy-Weinberg threshold.

explicitly designed for GWAS in polyploid species. They are the R package GWASpoly (Rosyara *et al.*, 2016) and the SHESIS tool (Shen *et al.*, 2016).

With these considerations in mind, we developed the MultiGWAS tool that performs GWAS analyses by using four different GWAS software. The tool deals with all matters of the GWAS process in the four software: preprocessing genomic data using different quality control filters, transforming it to particular tool formats, executing each GWAS software in parallel, postprocessing outputs and creating reports. The reports provides the user with tables and plots showing the significant and best ranked markers, also the markers found by one tool and by more than one tool, all of this with the objective to help users to decide in a more intuitively way the possible true or false associations.

2 Methods and Implementation

A flowchart of the main central steps involved in the three stages of the MultiGWAS tool is outlined in figure 1.

2.1 Adjustment stage

MultiGWAS takes as input a configuration file where the user specifies the genomic data along with the parameters that will be used by the four tools to perform their particular GWAS analysis (see figure 1). It starts by preprocessing the genomic data by selecting individuals present in both genotype and phenotype, and excluding individuals and SNPs that are likely to be of poor-quality.

The allowed format for the marker data is the “ACGT” (e.g. AAAT, ... ,CCCC), suitable for the polyploid software GWASpoly and SHESIS, but what is needed to “diploidize” for Plink and Tassel (e.g. AT, ... ,CG). MultiGWAS does this by coding each marker in one of two possible ways: all possible homozygous genotypes are coded with two nucleotides (AAAA→AA, CCCC→CC, GGGG→GG, TTTT→TT); and all possible heterozygous genotypes are coded with the combination of their reference and alternate alleles calculated from the tetraploid marker (e.g. AAAT→AT, ... ,CCCC→CG). After that, the new filtered genotype and phenotype are transformed to the specific formats required for each tool using different own functions from MultiGWAS.

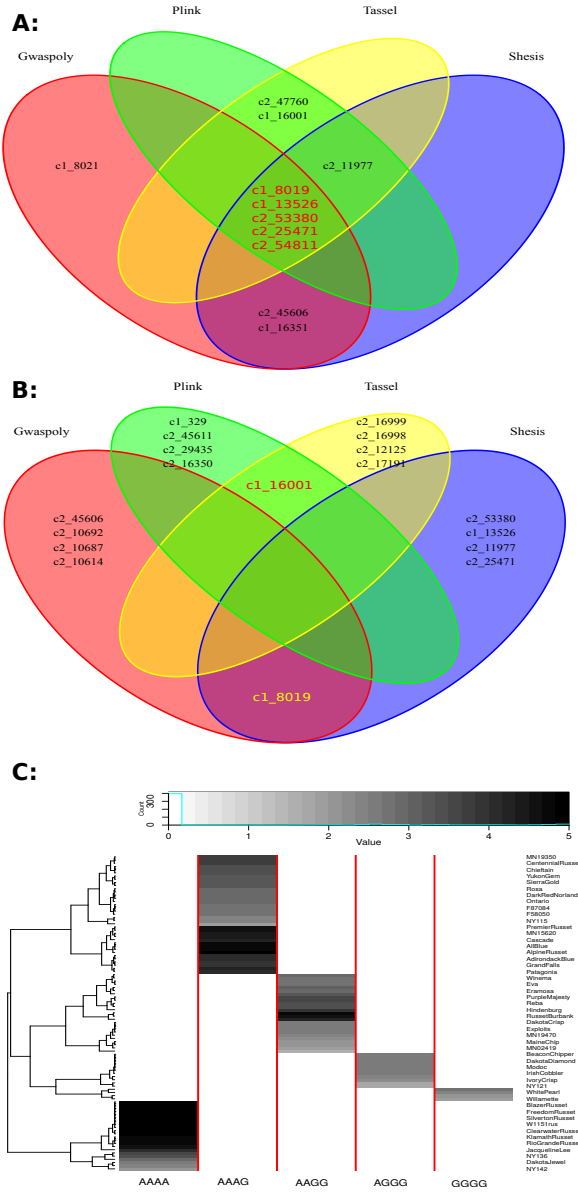


Fig. 2. Venn diagrams and SNP profile generated by the MultiGWAS tool for the SolCAP potato panel. A: GWAS with Naive model in which the four software found the same set of five markers (center area, red text). B: GWAS with Full model in which the two diploid software found one common marker (upper-central area, red text), but the other two polyploid tools found a different common marker (lower-central area, yellow text). In both cases, other marker were found either by two or by one software.

2.2 Multi analysis stage

The four tools are executed in parallel by MultiGWAS using for each one a particular script parameterized with the transformed genotype/phenotype and parameters set in the configuration file (see figure 1). One of these parameters is the type of statistical GWAS model to be conducted, where MultiGWAS allows two of them: Full (Q+K) and Naive (Sharma *et al.*, 2018). The first with control for population structure and individual relatedness; and the second without any of these controls. Controlling of population structure is based on the principal components (PCs) where each tool calculates the PCs and uses the top ten as covariates in their GWAS analysis. Controlling of individual relatedness is based on kinship matrices, where Tassel and GWASpoly make their own calculations,

but Plink and SHEsis calculate them externally by the king software (Manichaikul *et al.*, 2010).

2.3 Integration stage

The outputs resulting from the four tools are scanned and processed to identify the SNPs with both: significant and best ranked associations. This is done by correcting the reported *p-values*, and calculating their threshold value using the correction method and significance level α set in configuration file. These values are calculated taking into account only the number of valid genotype calls (nonmissing genotype, phenotype, and covariates). All this information is summarized to report them as tables and Venn diagrams for significative and best ranked associations, SNPs profiles, and Manhattan plots.

3 Results and Discussion

Figure 2 presents two marker plots generated by MultiGWAS in the analysis of the genomic data for the Solanaceae Coordinated Agricultural Project (SolCAP) potato diversity panel, used to test the GWASpoly software for polyploid species (Rosyara *et al.*, 2016). In the first case (figure 2.A), it shows that the four software agreed on a large set of common markers (text in red, figure 2.A). It can be explained as the statistical GWAS model used was Naive, without any controls for population structure or individual relatedness. However, two markers from this set, c1_8019 and c2_25471, were also reported by Rosayra *et al.*, the first as the most significative association, and second as the second best ranked association. So, the four tools agreed

In the second case (figure 2.B), it shows that the set of common markers found by the tools was reduced, which can be explained as the statistical GWAS model used was Full, with controls for population structure (10 PCs) and individual relatedness (kinship matrix). Now, the two polyploid tools GWASpoly and SHEsis found the common marker c1_8019, as in the Naive GWAS, which help us to consider with more confidence that this marker is a true association. But, the two diploid tools found a different common marker, the c1_16001, which was not found in the first case (figure 2.A) and may be a false association.

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