Data and text mining

ASSIsT: an automatic SNP scoring tool for in- and outbreeding species

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Abstract

ASSIsT (Automatic SNP Scoring Tool) is a user-friendly customized pipeline for efficient calling and filtering of SNPs from Illumina Infinium arrays, specifically devised for custom genotyping arrays. Illumina has developed an integrated software for SNP data visualization and inspection called GenomeStudio® (GS). ASSIsT builds on GS-derived data and identifies those markers that follow a bi-allelic genetic model and show reliable genotype calls. Moreover, ASSIsT re-edits SNP calls with null alleles or additional SNPs in the probe annealing site. ASSIsT can be employed in the analysis of different population types such as full-sib families and mating schemes used in the plant kingdom (backcross, F1, F2), and unrelated individuals. The final result can be directly exported in the format required by the most common software for genetic mapping and marker–trait association analysis. ASSIsT is developed in Python and runs in Windows and Linux.

Availability and implementation: The software, example data sets and tutorials are freely available at http://compbiotoolbox.fmach.it/assist/.

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1 Introduction

Advances in whole genome genotyping technologies enabled the investigation of several hundred thousand SNP markers simultaneously on a genome-wide scale. To date, Illumina (GoldenGate® and Infinium®) and Affimetrix (Axiom®) are the most widely used array-based genotyping platforms worldwide. Illumina has developed GenomeStudio®, a proprietary software with a graphical user interface (GUI) for SNP data visualization and filtering that enables the selection of high-quality markers showing robust performance across the examined germplasm. However, the actual filtering of such SNPs requires a deep understanding of the performance of SNP markers, genetic segregation patterns and familiarity with the many tools and parameters in GenomeStudio® (GS). ASSIsT accounts for this by offering a user-friendly, automated pipeline that builds on the results of Illumina’s GenCall algorithm (Kermani, 2006) as incorporated in GS.

In addition to filtering, ASSIsT also re-edits GS-calls in order to better explore the available information for SNPs showing null alleles or additional SNP clusters® due to additional polymorphisms at the probe annealing site. This re-editing enhances correct SNP calling and reduces unnecessary removal of potentially valuable markers.
2 Methods
The analysis and selection of SNPs performed by ASSIsT is based on the calls produced by Illumina’s GenCall algorithm (Kermani, 2006). A two tiers approach that employs a bi-allelic genetic model, and then a tri-allelic model is used to classify SNPs on the basis of their real performance on examined germplasm. The tri-allelic model is used to describe more complex segregation patterns due to null-alleles or alleles with variable signal intensity due to additional SNP, as the bi-allelic genetic model used by GS cannot account for such polymorphisms (Bassil et al., 2015; Gardner et al., 2014; Pikunova et al., 2014; Troggio et al., 2013). In this case, ASSIsT may re-edit GS-calls by applying de novo filters using the original light intensity data and the segregation patterns in the germplasm.

3 Results
ASSIsT supports the analyses of different population types, such as full-sib families (e.g. human, livestock, cross pollinating plants), mating schemes common in plants (backcross, F1, F2) and individuals with unknown genetic relationships. ASSIsT’s GUI allows easy parameter setting and provides a visual output of the SNP clustering analysis. The results produced by ASSIsT can be directly exported to the input format of the most widely used software for genetic and marker-trait association analysis (FlexQTLTM, GAPIT, JoinMap, the input format of the most widely used software for genetic and marker-trait association studies, where as ‘classical’ heterozygotes are not informative in the generation of genetic linkage maps as it is not possible to determine the parental origin of the alleles. Additional SNPs in the probe, as well as INDELS (Pikunova et al., 2014), may also give rise to null alleles, due to the lack of signal in one of the DNA templates, which results in additional clusters. GS cannot currently account for this scenario; thus, informative markers are lost. Conversely, ASSIsT succeeds in the analysis of the majority of such markers (A0 × A0, A0 × 00 and A0 × B0), allowing a more efficient marker calling.

All the above-mentioned SNP classes are suitable for the generation of genetic linkage maps or for marker–trait association studies. Discarded markers are grouped according to their performance considering absence of or severe distortion in segregation, presence of not allowed genotypes in segregating families and number of No Calls.

ASSIsT has been used to analyze SNP markers of several bi-parenatal full-sib families and germplasm of apple (Bianco et al., 2014), peach, melon and grape. For each family, ~99% of the ‘approved’ (those that passed the filtering procedure) SNPs showed to have high-quality data as they integrated smoothly in the generation of high-quality genetic linkage maps. The remaining 1% presented several types of issues, largely related to the presence of paralog loci where the AB cluster was too close or even merged to one of the two homoyzygous clusters.

ASSIsT thus proved to be an effective tool for genotyping studies as it allows to easily filter informative and well-performing SNP and to recover potentially useful SNPs from indels or regions of high-sequence divergence, feeding them directly to the most common downstream analysis tools through its easy interface.

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Conflict of Interest: none declared

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