

## k-mer-Based GWAS Unlock Causal Variant Discovery in Soybean

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The occurrence of short sequence motifs, pinpointed from the re- sults of a GWAS, distinguished

The occurrence of short sequence motifs, pinpointed from the re- sults of a GWAS, distinguished between white- and purple-flowered soybean accessions. Figure reproduced with permission from the following article by Sundaramoorthy et al. published in the Journal of the Korean Society for Applied Biological Chemistry: https://doi.org/10.1007/s13765-015-0077-z.

GenomeIwide association studies (GWAS) allow the discovery of loci associated with traits of interest in crops by identifying statistical associations between molecular markers and traits. Many studies use singleIhucleotide polymorphisms (SNPs) as molecular markers, but observed variation in traits can be due to other types of genetic changes such as structural variants (SVs). GWAS frequently fail to identify them. Researchers tested if short sequence motifs, known as *k*Imers, could identify causal variants regardless of if they are SNPs or SVs. Using a population of 363 cultivated soybean lines, they applied SNPI SVI and *k*ImerIbased GWAS to 13 traits. The researchers found that *k*Imers could pinpoint known causal variants at four loci, while identifying promising causal genes for several other traits. These analyses can assess all types of variants in a single analytic framework.

Such analyses hold promise for speeding up the application of genomics to plant breeding. The framework and computational tools developed by the authors for downstream analysis of *k*ImerIbased GWAS will facilitate the adoption of this method.

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