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► **To cite this version:**

Christopher Sauvage, Vincent Segura, Guillaume Bauchet, Rebecca Stevens, P.T. Do, et al.. A multilocus mixed model for GWAS reveals associations for metabolic traits in the tomato, [i]*Solanum lycopersicum*[/i]. XVIIIth Eucarpia Meeting, Vegetable section, Tomato Working group, Apr 2014, Avignon, France. 1 p. hal-01268525

HAL Id: hal-01268525

<https://hal.science/hal-01268525>

Submitted on 5 Jun 2020

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P015 – Christopher Sauvage

A multilocus mixed model for GWAS reveals associations for metabolic traits in the tomato, *Solanum lycopersicum*

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Genome-wide association studies have been successful in identifying genes involved in polygenic traits notably in crops, and can be a useful way towards crop improvement. We have applied in a major crop, tomato, a recently developed multi-locus mixed model as a general method for mapping complex traits in structured populations (Segura et al., 2012). Among major crops worldwide, tomato (*Solanum lycopersicum*) is a highly valuable fruit with excellent nutritional value (Causse et al., 2010). SNP beadchips (Hamilton et al., 2012) are available and enable GWAS for traits of interest. Recently, a pilot study defined the optimal conditions for GWAS by using cherry tomato accessions (Ranc et al., 2012). In our study, we examined a core collection of 180 tomato varieties composed of 20 wild accessions (*S. pimpinellifolium*), 130 admixed accessions (*S. cerasiforme*) and 30 domesticated accessions (*S. lycopersicum*). Multi-locus GWAS analysis was conducted using the MLM package (Segura et al., 2012) with 7700 SNP markers and a set of sugar-related, vitamin C-related and morphological traits as well as a broad range of metabolites involved in central carbon metabolism. The present study is the first one in tomato reporting associations for a large set of traits at the genome scale. We found significant associations for 89 loci with a total of 19 traits including fresh weight, sucrose, ascorbate, malate or citrate notably. Identified loci were also concordant with published quantitative trait loci (eg malate), while new loci were identified (for tocopherol). Moreover, several related metabolites, such as citrate and malate (both involved in the Krebs cycle) displayed two identical associations. These results (1) provide a list of candidate loci to be functionally validated and (2) provide a powerful analytical approach for finding genetic variants that can be directly used for crop improvement and deciphering the genetic architecture of complex traits.