

Tomato variation, origin and domestication

Jose Miguel Blanca, J. Monterau-Pau, Christopher Sauvage, Guillaume Bauchet, E. Illa, Monique Diez, Mathilde M. Causse, D. Francis, Joaquin Canizares, E. van Der Knaap

▶ To cite this version:

Jose Miguel Blanca, J. Monterau-Pau, Christopher Sauvage, Guillaume Bauchet, E. Illa, et al.. Tomato variation, origin and domestication. XVIII. Eucarpia Meeting, Vegetable section, Tomato Working group, Apr 2014, Avignon, France. 1 p. hal-02796249

HAL Id: hal-02796249 https://hal.inrae.fr/hal-02796249

Submitted on 5 Jun 2020

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



T001 - José Blanca

Tomato variation, origin and domestication

Blanca, J. (1), Montero-Pau, J. (1), Sauvage, C. (2), Bauchet, G. (2), Illa, E. (3), Díez, M. J. (1), Francis, D. (3), Causse, M. (2), van der Knaap, E. (3), Cañizares, J. (1)

- (1) Institute for the Conservation and Improvement of Agricultural Biodiversity (COMAV), Polytechnic University of Valencia, Spain
- (2) INRA, Unité de Génétique et Amélioration des Fruits et Légumes, Montfavet, France
- (3) Department of Horticulture and Crop Science, The Ohio State University/Ohio Agricultural Research and Development Center, USA

A collection of 949 accessions composed by *S. lycopersicum* var. lycopersicum (SLL; 514), S. l. var. cerasiforme (SLC; 291), S. pimpinellifolium (SP; 128), S. galapagense (SG;3), S. neorickii (SN; 1), S. chmielewski (Schm; 1) and 11 crosses between SP, SLC, SLL and S. pennellii was analyzed. The genotyping was carried out using the 7720 markers of the SolCAP genotyping platform and some key genes involved in the domestication process (CNR/FW2.2, SIKLUH/FW3.2, LC, FAS, OVATE and SUN). Using Principal Component Analyses (PCAs) based on the genetic data a new genetic-based classification divided the accessions in SP (121), SLC (236), SLL (492) and mixture (100). Within SP and Andean SLC a complex geographical structure correlated with the climate was found. Ecuadorian SP was remarkably differentiated from the rest of SP. In non-Andean SLC the structure found was not as marked. In a neighbor phylogenetic network SLC appeared intermediate between SP and SLC. Thus SLC was found to be a semi-domesticate with a key role in the tomato domestication. Andean SLC had a diversity comparable to SP while the non-Andean SLC had a lower diversity comparable to vintage SLL, indicating that the main bottleneck in the tomato domestication happened in the migration from the Andean region to Mesoamerica. Most SLC accessions were reported to have cherrysized fruit, but interestingly some South American SLC were collected in markets as fullsized vintage tomatoes. Thus the vintage tomato in South America included SLC as well as SLL materials.