The Plant Journal (2023)

RESOURCE

Multi-environment association study highlights candidate genes for robust agronomic quantitative trait loci in a novel worldwide *Capsicum* core collection

Louis McLeod^{1,2} (b), Lorenzo Barchi^{3,*} (b), Giorgio Tumino⁴ (b), Pasquale Tripodi⁵ (b), Jérémy Salinier^{1,†} (b), Christophe Gros¹, Hatice Filiz Boyaci^{6,‡} (b), Ramazan Ozalp⁶, Yelena Borovsky⁷, Roland Schafleitner⁸ (b), Derek Barchenger⁸ (b), Richard Finkers^{4,8} (b), Matthijs Brouwer⁴ (b), Nils Stein^{9,10} (b), Mark Timothy Rabanus-Wallace^{9,1} (b), Giovanni Giuliano¹¹ (b), Roeland Voorrips⁴ (c), Ilan Paran⁷ and Véronique Lefebvre^{1,*} (c) ¹INRAE, GAFL, Montfavet, France, ²INRAE, A2M, Montfavet, France, ³Department of Agricultural, Forest and Food Sciences (DISAFA), Plant Genetics, University of Torino, Grugliasco, Italy, ⁴Plant Breeding, Wageningen University and Research (WUR), Wageningen, The Netherlands, ⁵Research Centre for Vegetable and Ornamental Crops, Council for Agricultural Research and Economics (CREA), Pontecagnano Faiano, Italy, ⁶Bati Akdeniz Agricultural Research Institute (BATEM), Antalya, Türkiye, ⁷The Volcani Center, Institute of Plant Sciences, Agricultural Research Organization (ARO), Rishon LeZion, Israel, ⁸Vegetable Diversity and Improvement, World Vegetable Center, Shanhua, Taiwan,

⁹Leibniz Institute of Plant Genetics and Crop Plant Research (IPK), Seeland,Corre, Gatersleben, Germany,

¹⁰Department of Crop Sciences, Center for Integrated Breeding Research, Georg-August-University, Göttingen, Germany,

¹¹Casaccia Research Centre, Italian National Agency for New Technologies, Energy, and Sustainable Economic Development (ENEA), Rome, Italy

Received 27 April 2023; revised 13 July 2023; accepted 4 August 2023.

*For correspondence (e-mail veronique.lefebvre@inrae.fr; lorenzo.barchi@unito.it).

[†]Present address: CIRAD, UMR PVBMT, Saint-Pierre, La Réunion, France

[‡]Present address: Department of Horticulture, Faculty of Agriculture, University of Recep Tayyip Erdogan, Rize, Türkiye

[§]Present address: GenNovation B.V., Wageningen, The Netherlands

¹Present address: Faculty of Science, University of Melbourne, Parkville, Australia

SUMMARY

Investigating crop diversity through genome-wide association studies (GWAS) on core collections helps in deciphering the genetic determinants of complex quantitative traits. Using the G2P-SOL project world collection of 10 038 wild and cultivated *Capsicum* accessions from 10 major genebanks, we assembled a core collection of 423 accessions representing the known genetic diversity. Since complex traits are often highly dependent upon environmental variables and genotype-by-environment ($G \times E$) interactions, multi-environment GWAS with a 10 195-marker genotypic matrix were conducted on a highly diverse subset of 350 *Capsicum annuum* accessions, extensively phenotyped in up to six independent trials from five climatically differing countries. Environment-specific and multi-environment quantitative trait loci (QTLs) were detected for 23 diverse agronomic traits. We identified 97 candidate genes potentially implicated in 53 of the most robust and high-confidence QTLs for fruit flavor, color, size, and shape traits, and for plant productivity, vigor, and earliness traits. Investigating the genetic architecture of agronomic traits in this way will assist the development of genetic markers and pave the way for marker-assisted selection. The G2P-SOL pepper core collection will be available upon request as a unique and universal resource for further exploitation in future gene discovery and marker-assisted breeding efforts by the pepper community.

Keywords: multi-environment trials, association study, core collection, pepper, *Capsicum annuum* L., agronomic traits, candidate genes.

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SOCIETY FOR EXPERIMENTAL BIOLOGY

doi: 10.1111/tpj.16425

INTRODUCTION

Most yield- and quality-related traits are inherited quantitatively and governed by multiple genes (Bertin et al., 2010; Rao et al., 2003). Genetic dissection of quantitative traits has mostly been performed by mapping quantitative trait loci (QTLs) underlying their variation (Phan & Sim, 2017). Traditional QTL analysis approaches are based on dedicated mapping progenies segregating for the traits of interest, constructed from controlled crosses between two or more parental lines exhibiting diverse phenotypes (Korte & Farlow, 2013). These mapping studies are therefore timeand resource-intensive and are limited to the genetic and phenotypic variation existing between the founder lines (Phan & Sim, 2017). The emergence of genome-wide association studies (GWAS) based on collections of ideally distantly-related accessions and the development of cutting-edge next-generation sequencing technologies are now ways to circumvent these limitations (Phan & Sim, 2017). Additionally, GWAS increases the accuracy of QTL localization by tracking a higher number of historically accumulated recombinations, as compared to methods that rely on the meiosis of parental lines only (Perez-de-Castro et al., 2012). However, relatedness between accessions and population stratification can lead to spurious associations (Price et al., 2006). To correct this, GWAS models often incorporate principal components (PCs), a kinship matrix, or an estimation of global ancestry, all derived from large sets of markers (Price et al., 2010; Zhu et al., 2008).

The concept of core collection was first introduced by Frankel (1984). It refers to limited sets of accessions representing the greatest possible portion of genetic variation found in a crop species, including its wild progenitors. They are typically subsets of larger collections, designed for use in experiments and analyses requiring a survey of genetic diversity in situations where working on the entire set would be prohibitively laborious or computationally demanding. Owing to their compactness and diversity, core collections can be phenotyped for a wider range of traits than traditional mapping progenies, and contribute to the conservation and accessibility of plant genetic resources (Brown & Spillane, 1999).

The pepper fruit is a seeded fleshy berry grown in all populated continents, mainly between the two tropics. Over 40 *Capsicum* species have been recognized so far (Barboza et al., 2022). Five of these species are domesticated, including the widely-cultivated and comparatively well-studied *Capsicum annuum* (Pereira-Dias et al., 2019). Two major types dominate the diversity in *C. annuum*: the small-fruited and pungent chili peppers and the large-fruited non-pungent sweet peppers (Pereira-Dias et al., 2019; Tripodi et al., 2021). While domestication and breeding have been mainly focused on yield improvement of

these types, numerous additional traits contribute to the production of desirable cultivated varieties, including shoot architecture, phenology, flowering time, fruit morphology, fruit quality, metabolite composition, resistance to biotic stresses and tolerance to abiotic stresses (IPGRI et al., 1995; Junior e Silva et al., 2013). The diverse natural variation observed within Capsicum has enabled the characterization of inherited beneficial variants for such traits, using a range of mapping approaches and genetic marker technologies (e.g., Lefebvre et al., 2002; Lippert et al., 1965; Paran & Fallik, 2011; Wang & Bosland, 2006; see Table S1 for an up-to-date summary). Large numbers of QTLs to be implemented for pepper improvement were located throughout the genome (Arpaci & Karatas, 2020; Jeong et al., 2015: Lin et al., 2015: Parsons et al., 2012: Rani et al., 2021; Thabuis et al., 2004).

High-throughput single nucleotide polymorphism (SNP) technologies such as genotyping-by-sequencing (GBS) have recently been used to describe the genetic structure of the pepper germplasm (Colonna et al., 2019; Hill et al., 2013; Lee, Ro, et al., 2020; Lozada et al., 2022; Ro et al., 2022; Tamisier et al., 2020; Taranto et al., 2016; Wu et al., 2019). Most recently, the G2P-SOL project produced a comprehensive survey of pepper diversity, describing the genetic structure of over 10 000 wild and cultivated accessions housed in 10 genebanks (Tripodi et al., 2021). Several core collections and GWAS have been reported in C. annuum (for an up-to-date summary, see Table S2). However, none has yet leveraged this extent of diversity to define a core collection suitable to perform a maximally powerful GWAS. Moreover, GWAS to date has relied upon phenotypic data gathered in a single or a limited number of trials within a given country (Table S2). Complex traits, however, are generally modulated by the environment and the interaction between genotype (sensu accession) and environment $(G \times E)$ (Kang, 2004). The latter especially impact crop research and breeding and necessitate replication of the trials in different locations in order to efficiently and reliably develop locally adapted varieties (Kang, 2020).

We report the compilation and analysis of a novel pepper core collection of 423 accessions, representing the accessible global pepper diversity. Twenty-three plant growth and fruit traits were phenotyped in 350 *C. annuum* accessions of this core collection in multiple environments. GWAS led to the identification of novel and previously reported QTLs, which may be used for breeding and for functional genomics studies identifying the causal genes underlying trait variation. These results demonstrated the usefulness and universality of the core collection as a worldwide research and breeding resource designed for mining the pepper genome and germplasm.

RESULTS

The G2P-SOL core collection captures the diversity of the worldwide germplasm

The reference G2P-SOL pepper core collection

A core collection of *Capsicum* spp. consisting of 423 accessions was designed to capture the genetic variability present in the *Capsicum* panel of 10 038 accessions analyzed by Tripodi et al. (2021) by minimizing a repetitiveness criterion (Table S3). These 423 accessions comprise 391 *C. annuum* accessions and 32 accessions from other cultivated *Capsicum* species, including several likely derived from interspecific hybridizations. Principal component analyses (PCA), either based on the 10038 *Capsicum* accessions panel or on its 7848 *C. annuum* accessions subset, show that the core collection and its subset of 391 *C. annuum* accessions spanned the whole *Capsicum* and *C. annuum* germplasms, respectively (Figure 1a,b).

C. annuum accessions and markers used for GWAS

The SNP calling and filtering parameters applied to the core collection and the GBS markers resulted in 350 *C. annuum* accessions and 10 195 markers (9546 SNPs and 649 Indels) exploitable for GWAS (Table S3). The post-filtering marker set had an average density of 3.2 markers/ Mb with a mean minor allele frequency (MAF) of 12.8%, a 1.5% rate of heterozygosity, and 4.9% missing data at the marker level (Figure S1). The accessions also had on average a 1.5% rate of heterozygosity and 4.9% missing data. On average, the linkage disequilibrium (LD) estimated for each chromosome decayed below $r^2 = 0.2$ at 0.4 Mb (Figure S2).

The 350 C. annuum accessions used for GWAS were, like the entire set of 391 C. annuum accessions, representative of the whole C. annuum germplasm (Figure 1b), and there was no evidence of strong stratification in the PCA conducted solely on these 350 accessions (Figure 2). However, the negative correlation between the multienvironment best linear unbiased predictions (BLUPs) for fruit weight (FWe) and pungency (FP) confirms the existence of two major types (large-sweet versus small-spicy) dominating the C. annuum diversity (Figure 2a,b). Investigation of the population structure by Admixture revealed that K = 11 was a sensible model choice, according to the cross-validation (CV) error (Figure S3). The PCA colored according to the geographical origin and to the observed K subpopulations highlighted that the 350 accessions were partly structured according to their geographical origin (Figure 2c,d), with clusters 7 and 11 roughly corresponding to the European accessions, clusters 3 and 6 to the Eastern Asian and Middle Eastern accessions, respectively, clusters 2 and 4 to part of the Eastern European accessions, clusters 8 and 9 to part of the Southern-Southeastern Asian and Central American accessions, respectively, and finally

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cluster 5 to part of the Southern-Southeastern Asian and Middle Eastern accessions.

GWAS identified QTLs for 23 plant and fruit development traits

Field traits variation and correlations

The 350 C. annuum accessions were phenotyped for 23 traits in six independent trials performed in five countries, thus showing a rich phenotypic variability suitable for GWAS (Table 1; Tables S4 and S5). Linear models were fitted to assess the contributions of environment, genotype (sensu accession), and $G \times E$ interactions to the measured trait values. The residual variances were comprised between 2% and 13% of the total variance for all traits except locule number (LN, 32% of the total variance), indicating a sound fit (Figure 3). For most traits, the amount of total variance attributed to the genotype (accession), was largely predominant (from 27 to 93% of the total variance), while for Brix (BX) and for fruit load (FLo), it was lower than the proportion of variance attributed to the environment and to the G × E interactions, respectively. Fruit maximum width (FWi) was the least influenced by the environment and G × E interactions, while FLo was the most influenced. Broad-sense heritability ranged from 0.75 for total fruit weight (TFWe) to 0.99 for fruit maximum width (FWi), apart from FLo and total fruit number (TFN) with low heritability values of 0.40 and 0.25, respectively (Figure 3). Overall, heritability values were above 0.90 for 15 out of 23 traits.

A PCA plot based on the accession means of the 23 traits illustrates a high level of agreement between trial results (Figure S4). The average correlation values between pairs of trials (here referred to as environments) were high for most traits, especially for fruit maximum width (FWi, 0.95), fruit weight (FWe, 0.94), fruit shape index (FShI, 0.93), fruit maximum length (FLe, 0.92) and pericarp thickness (PTh, 0.88) (Figure S5). Some trials with lower correlations were omitted from downstream analyses for five traits (FF, fruit fasciation; LN, locule number; FIT, flowering time; BX, Brix; FLo, fruit load) (Figure 3).

To correct for environment and $G \times E$ interaction effects, adjusted means per accession were estimated using a BLUP model. These BLUPs, used as GWAS phenotypic input, were calculated both for single environments and across highly correlated environments. Since the correlations between single- and multi-environment BLUPs were high for all traits, with an average by trait ranging from 0.82 to 0.98 (Table 2), we focused on the multienvironment BLUPs to investigate the correlation between traits (Figure S6a). Based on the observed correlations, all traits except IFCG (external immature fruit color green) could be grouped into two negatively correlated clusters. On the one hand, fruit size and shape traits (FWe, fruit weight; FWi, fruit maximum width; PTh, pericarp thickness;

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The Plant Journal, (2023), doi: 10.1111/tpj.16425

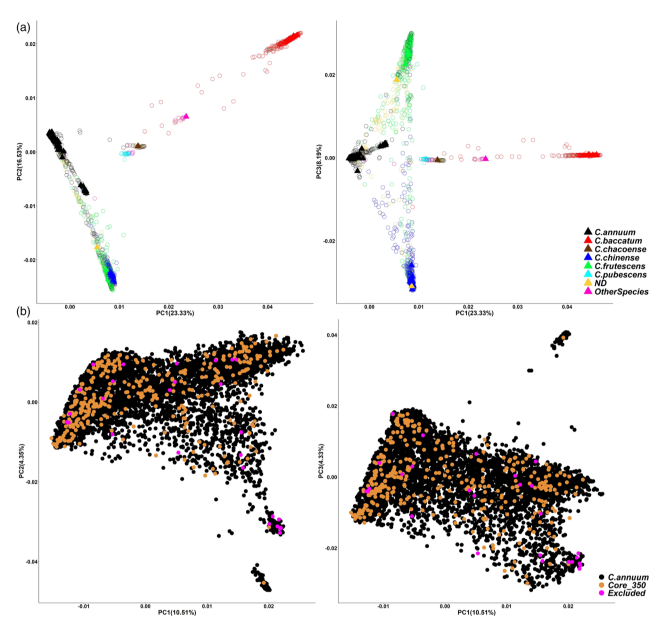


Figure 1. The worldwide genetic diversity of pepper based on single nucleotide polymorphisms.

(a) Plots of the first three components of the principal component analysis of the 10 038 *Capsicum* accessions from Tripodi et al. (2021), with the whole core collection as triangles.

(b) Plots of the first three components of the principal component analysis of the 7848 *Capsicum annuum* accessions, using 1073 linkage disequilibrium-pruned markers with a minor allele frequency superior to 0.05. The 350 *C. annuum* accessions of the core collection that were used for genome-wide association studies are colored in orange, and the rest of the *C. annuum* accessions of the core collection that were filtered out are colored in pink.

FF, fruit fasciation; LN, locule number; FLe, fruit maximum length; FShO, fruit predominant shape oblate) and plant productivity traits (TFWe, total fruit weight; Flo, fruit load) were positively correlated. On the other hand, fruit external color L, a and b traits (FCL, FCa, and FCb), flavor traits (BX, Brix; FP, fruit pungency), a fruit shape trait (FShI, fruit shape index), plant vigor and earliness traits (FIT, flowering time; AL, axis length; PHe, total plant height) and a plant productivity trait (TFN, total fruit number) were positively correlated.

Marker-phenotype associations and QTLs

We scanned for marker-phenotype associations using three models from the GAPIT R package: Mixed Linear Model (MLM), Multi-Locus Mixed-Model (MLMM), and Bayesian-information and Linkage-disequilibrium Iteratively Nested Keyway (BLINK). We identified a total of 704 significant marker-phenotype associations corresponding to 453 distinct quantitative trait nucleotides (QTNs) for the 23 studied traits. Associations differed between environments including the multi-environment (Figure 4a). They

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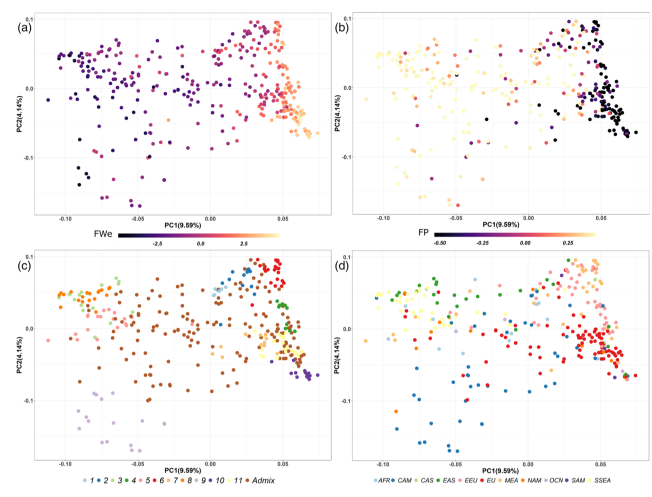


Figure 2. Population structure of the 350 Capsicum annuum accessions used for genome-wide association studies. Plot of the first and second component of a principal component analysis based on 2357 pruned single nucleotide polymorphism markers.

(a) Accessions are colored according to their fruit weight (FWe) multi-environment BLUP values.

(b) Accessions are colored according to their fruit pungency (FP) multi-environment BLUP values. (c) Accessions are colored according to their cluster of structure identified with ADMIXTURE, from K1 to K11.

(d) Accessions are colored according to their geographical origin (AFR, Africa; CAM, Central America; CAS, Central Asia; EAS, Eastern Asia; EEU, Eastern Europe; EU, Europe; MEA, Middle East; NAM, Northern America; OCN, Oceania; SAM, South America; SSEA, Southern-Southeastern Asia).

were primarily found with the BLINK model (499 associations), considerably less with the MLM model (110 associations), and with the MLMM model (95 associations) (Table S6). The 453 QTNs were grouped into 423 QTLs, the chromosome-specific LD extent using value (Figure S2): 332 QTLs for the quantitative traits and 91 QTLs for the ordinal and binary categorical traits (Table S7; Figure S7). Overall, QTL size varied from 600 to 1819 kb (mean: 821 kb). They explained up to 45% of the multienvironment BLUPs variation (R^2) , with 19 QTLs explaining more than 20%, 61 QTLs explaining between 10 and 20%, 106 QTLs explaining between 5 and 10%, and 237 QTLs explaining less than 5% (Table S7).

Some of these QTLs were particularly robust, i.e., detected in several environments (Figure 4a; Figure S7). Among the most robust QTLs, FLe-P10.1 (fruit maximum length), FShO-P5.6 (fruit predominant shape oblate), FShI-P10.1 (fruit shape index), and IFCG-P1.2 (external immature fruit color green) were detected in all investigated environments (FShI-P10.1 and IFCG-P1.2 by all three GWAS models in each environment), FWe-P2.4 (fruit weight) in six of the seven investigated environments, FShO-P5.3 and FShO-P5.4 in five of the six investigated environments, and FShO-P3.3, FShO-P6.4, FShO-P9.1, FShO-P10.1 in four of the six investigated environments. Overall, 411 out of the 423 QTLs controlled traits phenotyped in several environments, of which 74 were robust (18%), 73 (18%) were detected with the multi-environment BLUPs only, and the remaining 264 (64%) were detected in one trial. The proportion of investigated environments in which a QTL was detected was only slightly correlated to the effect of the QTL (R^2) on the multi-environment BLUPs (Pearson correlation = 0.12, P = 0.011), meaning that the QTLs having a major effect on a trait were not systematically robust.

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Table 1 Descriptive statistics on the accession means of the 23 traits

Trait			Descriptive statistics ^a						
Code	Name	Unit	Nb. val	Min	Max	Median	Mean	SEM	SD
AL	Axis length	cm	350	8.54	76.54	28.50	30.04	0.47	8.78
BX	Brix	°Bx	350	5.27	13.33	7.75	7.91	0.06	1.21
FCa	Fruit external color a	-	349	-2.25	44.63	34.08	33.36	0.34	6.43
FCb	Fruit external color b	-	349	2.28	70.64	20.40	22.23	0.46	8.64
FCL	Fruit external color L	-	349	26.67	78.25	38.15	39.25	0.30	5.60
FLe	Fruit maximum length	cm	350	1.07	21.08	8.25	8.59	0.23	4.37
FIT	Flowering time	days	350	13.71	63.39	27.16	28.64	0.43	8.11
FShl	Fruit shape index	-	350	0.56	19.53	3.04	3.59	0.14	2.54
FWe	Fruit weight	g	350	0.30	201.55	17.89	37.39	2.38	44.44
FWi	Fruit maximum width	cm	350	0.43	8.55	2.53	3.22	0.11	2.04
IFCa	Immature fruit external color a	-	329	-20.68	5.81	-13.66	-13.43	0.19	3.49
IFCb	Immature fruit external color b	-	329	3.65	49.46	20.53	22.46	0.44	7.90
IFCL	Immature fruit external color L	-	329	28.78	82.60	39.46	42.84	0.53	9.68
LN	Locule number	count	350	1.94	4.31	2.61	2.72	0.03	0.50
PHe	Total plant height	cm	350	43.98	154.22	91.61	92.91	1.11	20.75
PTh	Pericarp thickness	mm	350	0.72	5.97	2.26	2.61	0.07	1.39
TFN	Total fruit number	count	346	0.00	226.25	17.54	24.15	1.32	24.52
TFWe	Total fruit weight	g	347	0.00	1273.33	413.21	431.07	13.86	258.18
FF	Fruit fasciation	1–4	350	1.00	3.77	1.79	1.85	0.03	0.60
FLo	Fruit load	1–3	350	1.40	3.00	2.31	2.30	0.02	0.31
FP	Fruit pungency	0 versus 1	348	0.00	1.00	0.73	0.57	0.02	0.43
FShO	Fruit predominant shape oblate	0 versus 1	350	0.00	1.00	0.00	0.03	0.01	0.13
IFCG	External immature fruit color green	0-4	349	0.58	4.00	3.07	2.80	0.04	0.68

max, maximal value; min, minimal value; Nb. val, number of values; SD, standard deviation; SEM, standard error on the mean. ^aOutliers were removed before computing the statistics.

Highest-confidence QTLs

Among the 423 QTLs, 114 QTLs which were located on the 12 pepper chromosomes, and 5 QTLs which were located on the artefactual chromosome P0 were considered particularly promising for plant breeding, as they were either robust (i.e., identified in several environments), or identified by several GWAS models, or with a particularly high $-\log 10(P-value)$ (Table S7; Figure 5). Fifty-one of these 114 QTLs affected six of the eight traits never analyzed by GWAS (AL, BX, FCa, FCL, TFN, and FShO). Twenty-one out of the 114 most promising QTLs overlapped on eight distinct chromosomal regions. Three of these regions, on chromosome P1, P2, and P10, included the most robust QTLs for traits related to immature fruit color, to fruit vield and to fruit size and shape, respectively (Table S8; Figure 5). As an example, the fruit maximum length QTL FLe-P10.1 was detected in all environments (Figure 4a), and accessions of the AA and GG genotypes at the corresponding marker exhibited contrasting phenotypes (Student test, P = 5.632e-06;Figure 4b,c).

Candidate genes were identified for 53 of the highestconfidence QTLs

There was a total of 2790 genes in the intervals of the 114 most promising QTLs mapped on the 12 chromosomes,

with an average of 137 genes per trait and 24.5 genes per QTL. We identified 97 candidate genes likely to determine the traits for 53 of these QTLs, based on previously reported biological function (Table S9).

Among the eight candidate genes potentially affecting fruit flavor, the *Pun1* gene and homologs of two MYB transcription factors (*MYB61* and *MYB63*) underlie fruit pungency QTLs *FP-P2.1* and *FP-P7.3*, while a homolog of *SNF1-RELATED PROTEIN KINASE REGULATORY SUBUNIT BETA-2* (*KINB2*) underlies the Brix QTL *BX-P5.2*.

The 28 candidate genes potentially affecting fruit size and shape include three Ovate Family Proteins transcription factors: C. annuum OVATE FAMILY PROTEIN 20 (CaOFP20) for colocalized QTLs FLe-P10.1, FShl-P10.1, FShO-P10.1, FF-P10.1, and PTh-P10.1; a homolog of transcriptional repressor OVATE FAMILY PROTEIN 3 (OFP3) for FLe-P6.4; and a homolog of OVATE for FWe-P2.4. Among the other strong candidates, a homolog of FANTASTIC FOUR 3 (FAF3) underlies FLe-P6.2, a homolog of BEL1-LIKE HOMEODOMAIN PROTEIN 2 (BLH2) underlies FWe-P2.4, a homolog of a probable leucine-rich repeat receptor-like protein kinase of the CLAVATA family At5g63930 underlies LN-P3.1, homologs of KATANIN P60 and SUBUNIT ATPASE-CONTAINING (AAA1) A1 and PURINE PERMEASE 3 (PUP3) underlie FShl-P1.1 and FShl-P6.3, respectively.

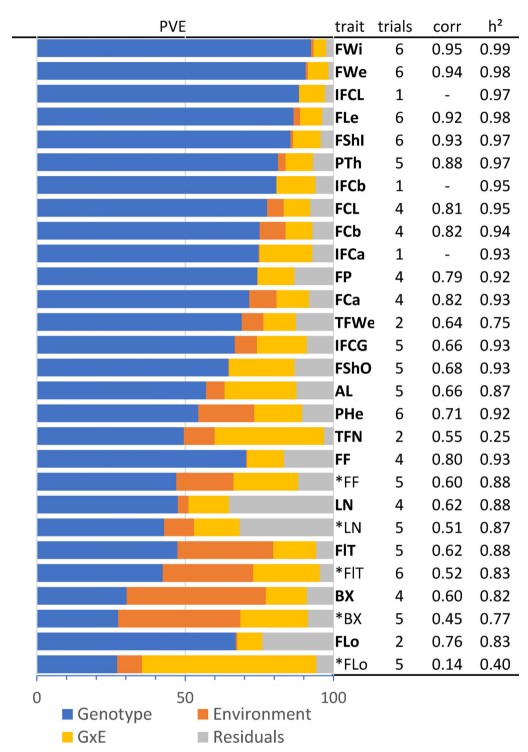


Figure 3. Proportion of phenotypic variance explained by the effects of genotype, environment and their interaction, and heritability values for the 23 agronomic traits.

Environment corresponds to the trial (or to the block for IFCa, IFCb and IFCL only). Bold characters indicate that only highly correlated trials were included, asterisks (*) indicate that lesser-correlated trials were included as well. PVE, Proportion of phenotypic variance explained; trials, number of trials; corr, average of the 2-by-2 correlations across trials; *h*², broad-sense heritability; G × E, genotype-by-environment interaction; AL, axis length; BX, brix; FCa, fruit external color a; FCb, fruit external color b; FCL, fruit external color L; FF, fruit fasciation; FLe, fruit maximum length; FLo, fruit load; FIT, flowering time; FP, fruit pungency; FShl, fruit shape index; FShO, fruit predominant shape oblate; FWe, fruit weight; FWi, fruit maximum width; IFCa, immature fruit external color a; IFCb, immature fruit external color b; IFCG, immature fruit color green; IFCL, immature fruit external color L; LN, locule number; PHe, total plant height; PTh, pericarp thickness; TFN, total fruit number; TFWe, total fruit weight.

Table 2 Pearson correlation between single-environment BLUPs and multi-environment BLUPs

	Trial							
Trait code	ARO Israel April 2020– August 2020	WorldVeg Taiwan October 2020– January 2021	BATEM Türkiye August 2020– March 2021	CREA Italy June 2019– December 2019	INRAE 2019 France June– November	INRAE 2020 France June– November	Average	
AL	0.68	NA	0.87	0.84	0.93	0.87	0.84	
BX	0.66	NA	0.85	NA	0.88	0.87	0.82	
FCa	0.82	NA	NA	0.93	0.96	0.96	0.92	
FCb	0.90	NA	NA	0.94	0.93	0.92	0.92	
FCL	0.89	NA	NA	0.93	0.93	0.91	0.92	
FLe	0.97	0.97	0.96	0.96	0.95	0.98	0.97	
FIT	0.77	0.87	0.88	NA	0.83	0.81	0.83	
FShI	0.96	0.97	0.96	0.97	0.97	0.98	0.97	
FWe	0.98	0.98	0.98	0.98	0.96	0.99	0.98	
FWi	0.98	0.98	0.98	0.97	0.97	0.99	0.98	
LN	0.75	NA	0.87	NA	0.89	0.90	0.85	
PHe	0.73	0.84	0.85	0.94	0.94	0.89	0.87	
PTh	0.93	NA	0.94	0.92	0.95	0.98	0.94	
TFN	NA	0.95	0.84	NA	NA	NA	0.90	
TFWe	NA	0.85	0.97	NA	NA	NA	0.91	
FF	NA	NA	0.85	0.96	0.96	0.91	0.92	
FLo	NA	NA	NA	NA	0.94	0.93	0.94	
FP	NA	NA	0.86	0.95	0.96	0.92	0.92	
FShO	0.77	NA	0.65	1.00	0.99	0.96	0.87	
IFCG	0.87	NA	0.92	0.86	0.93	0.93	0.90	
minimum	0.66	0.84	0.65	0.84	0.83	0.81	0.82	
maximum	0.98	0.98	0.98	1.00	0.99	0.99	0.98	
average	0.84	0.93	0.89	0.94	0.94	0.93	0.91	

NA indicates that the trait was not phenotyped by the partner, or was removed for multi-environment BLUPs computation. All BLUPs are used to compute correlations (no phenotypic outlier removed). All correlations are significant (P < 0.001).

We identified 32 candidate genes potentially affecting fruit color. Regarding immature fruit color, a homolog of *ARABIDOPSIS PSEUDO RESPONSE REGULATOR2 (APRR2)* underlies colocalizing QTLs *IFCa-P1.1, IFCb-P1.1* and *IFCG-P1.2,* and *CAPSANTHIN-CAPSORUBIN SYNTHASE (CCS)* underlies *IFCL-P6.1*. Regarding ripe fruit color, QTLs *FCa-P1.2, FCa-P3.1, FCa-P6.1,* and *FCa-P6.2* are underlain by two homologs of a *LICODIONE SYNTHASE (CYP93B1),* homologs of three MYB transcription factors (*MYB35, MYB59,* and *MYB98*), of two E3 ubiquitin-protein ligases (*UPL3* and a putative UPL3 LIN-like isoform X1), of a *STE-ROL 3-BETA-GLUCOSYLTRANSFERASE (UGT80B1),* of an F-box/kelch-repeat protein (*At3g18720*) and by a putative *FLAVONOL SYNTHASE/FLAVANONE 3-HYDROXYLASE (CA06g12570).*

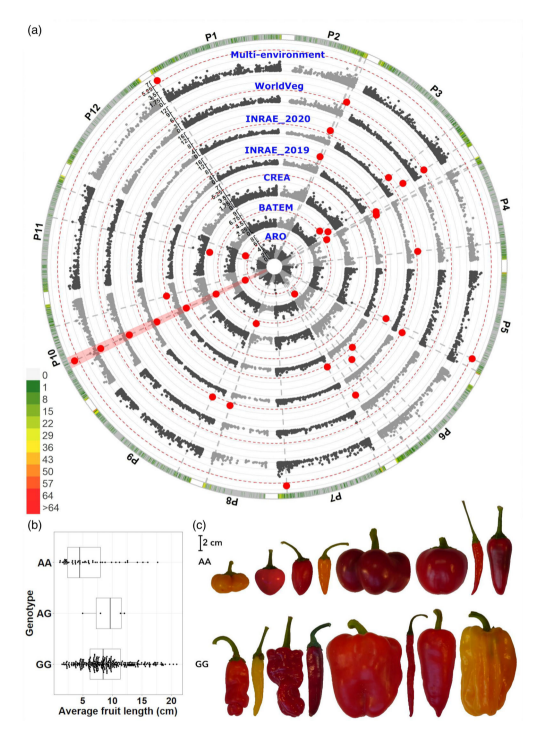
We also found 15 candidate genes potentially affecting plant growth and earliness. Homologs of *GROWTH-REGULATING FACTOR 12* (*GRF12*) and *PLAT DOMAIN-CONTAINING PROTEIN 1* (*PLAT1*) underlie axis length QTLs *AL-P1.2* and *AL-P5.4*, respectively. A homolog of *NARROW LEAF 1* (*NAL1*) underlies the total plant height QTL *PHe-P2.1*. Regarding flowering time, *FIT-P9.2* is underlain by homologs of *SNW/SKI-INTERACTING PROTEIN* (*SKIP*) and of putative *GATA TRANSCRIPTION FACTOR 22* (GATA22), and FIT-P9.2 by a homolog of Zinc finger protein CONSTANS-LIKE 6 (COL6).

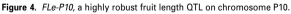
Finally, we identified 14 candidate genes potentially affecting plant productivity. Among these, a NAC transcription factor (*CA03g28100*) and a homolog of *AINTEGU-MENTA* (*ANT*) underlie total fruit number QTLs *TFN-P12.2* and *TFN-P6.1*, respectively.

DISCUSSION

The G2P-SOL core collection is a valuable new resource for pepper genetics

Until recently, no global comprehensive survey of pepper phenotypic and genotypic diversity had been reported. Association panels and core collections mainly originated from rather limited numbers of accessions from one or few genebanks (Colonna et al., 2019; Lozada et al., 2022; Nimmakayala et al., 2014, 2016; Tamisier et al., 2020; Wu et al., 2019; Zewdie et al., 2004), and were designed based on the phenotypic diversity for the traits of interest (Hanson et al., 2004; Quenouille et al., 2016; Thies & Fery, 2002), and on limited genotypic data (Fan et al., 2004; Lee et al., 2016; Nicolaï et al., 2013). This limitation has now been lifted owing largely to the G2P-SOL project's survey of the global





(a) Circular Manhattan-plot depicting the fruit maximum length QTNs detected with the multi-locus BLINK model. The seven internal circles show Manhattan plots for all single environments and the multi-environment, and the external circle displays the marker density (markers/megabase, according to the color scale). Red dashed circles represent the Bonferroni-adjusted significance threshold for each environment, and red dots the –log10(*P*-value) of significant marker-trait associations. Gray radial dashed lines indicate the projection of significant associations on all circles. QTL *FLe-P10* is highlighted in red on chromosome P10.

(b) Boxplots representing the fruit maximum length of the 350 accessions averaged over the six trials, according to their genotype at QTL *FLe-P10*. AA genotype: 49 accessions. AG genotype: 4 accessions. GG genotype: 283 accessions. AA and GG genotypes exhibited contrasting phenotypes (Student test, *P*-value = 5.632e-06).

(c) Pictures of eight accessions spanning the first to third quartiles of fruit maximum length for the AA and GG genotypes at QTL FLe-P10. QTL, quantitative trait loci; QTN, quantitative trait nucleotide.

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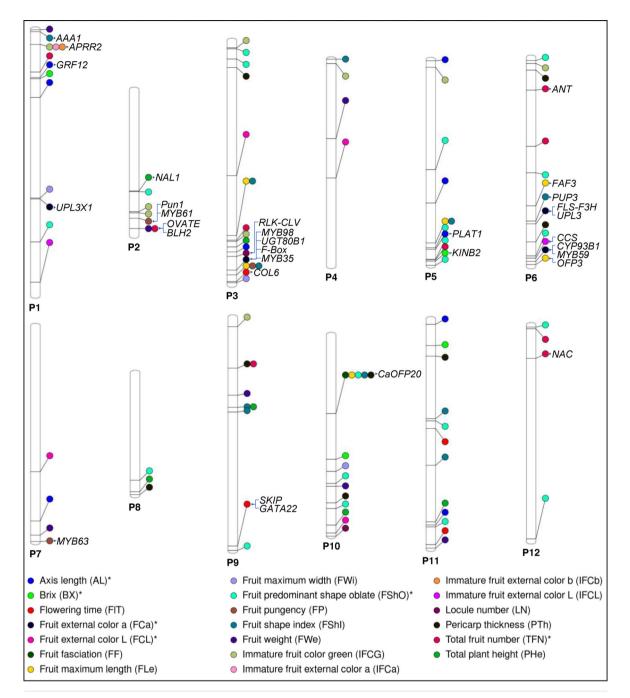


Figure 5. Physical position of the 114 highest-confidence QTLs and their best candidate genes mapped on the 12 pepper chromosomes.

The 31 candidate genes discussed in the manuscript are indicated next to the QTL positions. Candidate gene names are detailed in Table S9, along with the name and position of their corresponding QTLs. Asterisks (*) indicate which traits were never analyzed by GWAS before. AAA1, homolog of KATANIN P60 ATPASE-CONTAINING SUBUNIT A1; ANT, homolog of AINTEGUMENTA; APRR2, homolog of ARABIDOPSIS PSEUDO RESPONSE REGULATOR2; BLH2, homolog of BEL1-LIKE HOMEODOMAIN PROTEIN 2; CaOFP20, C. annuum OVATE FAMILY PROTEIN 20; CCS, CAPSANTHIN-CAPSORUBIN SYNTHASE; COL6, homolog of CONSTANS-LIKE 6; CYP93B1, homologs of a LICODIONE SYNTHASE (CYP93B1); FAF3, homolog of FANTASTIC FOUR 3; F-Box, homolog of the F-box/ kelch-repeat protein At3g18720; FLS-F3H, homolog of Flavonol synthase and flavanone 3-hydroxylase; GATA22, homolog of GATA TRANSCRIPTION FACTOR 22; GRF12, homolog of GROWTH-REGULATING FACTOR 12; KINB2, homolog of SNF1-RELATED PROTEIN KINASE REGULATORY SUBUNIT BETA-2; MYB35, homolog of MYB35 transcription factor; MYB61, homolog of MYB69 transcription factor; MYB61, homolog of VAB86 transcription factor; MYB63, homolog of NARB0W LEAF 1; OFP3, homolog of OVATE FAMILY PROTEIN 3; OVATE, homolog of OVATE; PLAT1, homolog of PLAT DOMAIN PROTEIN 1; Pun1, Pun1 gene; PUP3, homolog of PURINE PERMEASE 3; RLK-CLV, homolog of leucine-rich repeat receptor-like protein kinase of the CLAVATA family At5g63303; SKIP, homolog of SNW/SKI-INTERACTING PROTEIN; UGT80B1, homolog of a STEROL 3-BETA-GLUCOSYLTRANSFERASE (UGT80B1); UPL3, homolog of E3 UBIQUITIN-PROTEIN LIGASE 3; UPL3X1, homolog of UPL3 LIN-like isoform X1.

pepper germplasm across 10 038 accessions from collections conserved in 10 major genebanks using 26 566 SNPs (Tripodi et al., 2021).

To fully exploit this unprecedented diversity survey, we constructed a core collection of 423 pepper accessions of rich genotypic and phenotypic variability. Representatives from the secondary and tertiary genepools were included in the core collection, inter alia for an exploratory screening of non-annuum species for specific traits such as responses to biotic and abiotic stresses. The core collection shows a species-wise genetic structure, and the stratification among its 350 C. annuum accessions used for GWAS is partly correlated to their geographical origin (Figure 2c,d), which is consistent with the previous results on different sets of C. annuum accessions from Nicolaï et al. (2013), Lee et al. (2016), Taranto et al. (2016) and Tripodi et al. (2021). Therefore, the inherent structure of the core collection needs to be addressed in future genotypephenotype associations studies, as it was addressed here. This core collection paves the way for a broad range of studies. For instance, our suite of genomic trait association scans revealed candidate genes for relevant traits for pepper agriculture.

Multi-environment GWAS delivers robust QTLs

Various studies in pepper have shown that traits related to morpho-agronomic performance and phytochemical composition can be impacted by $G \times E$ interactions (Alimi et al., 2013; Cabral et al., 2017; Tripodi et al., 2019). To assess the influence of the environment and of G × E interactions, envirotyping of our core collection was performed with trials from five geographically and climatically contrasted locations, and for two consecutive years in one of them (Table S4). By addressing the environment and $G \times E$ interaction effects, we refined the estimations of the part of phenotypic variance explained by the genotype and of the heritability of traits in various environments (Malosetti et al., 2013). The use of multiple environments also differentiated environment-specific QTLs from more robust QTLs. The ANOVA models explaining the phenotypic variance fitted the data satisfactorily, and heritability values of the traits were high despite the $G \times E$ interaction effects (Figure 3), suggesting that the QTLs detected could be used efficiently in pepper breeding through markerassisted selection, particularly for fruit morphology (FWe, FLe, FWi, FShI, and PTh), color (FCL, FCa, and FCb) and pungency (FP) traits. Eight robust QTLs were located in the vicinity of the already reported pepper genes Pun1, CCS, OFP20, and OVATE. In addition, we found that two robust QTLs may correspond to already reported QTLs from GWAS (Table S1). QTL FW-P10.1 on chromosome P10 (interval 175 298 168-175 898 168 bp) was in the vicinity of a QTL determining fruit weight at 173 645 289 bp (Lozada et al., 2022). QTL PHe-P2.1 on chromosome P2 (interval

Multi-environment GWAS in a pepper core collection 11

116 761 596–117 561 596 bp) was located between two QTLs at 111 478 856 bp and 124 942 237 bp (Lozada et al., 2022). This underlines that QTLs detected by GWAS depend not only on the environment, but also on other factors, such as the phenotyping methodology, the SNP coverage over the whole genome, and the nature of the phenotyped core collection influencing the proportion of rare alleles and the extent of LD decay.

Altogether, phenotyping a representation of the global C. annuum diversity for a large number of traits in multiple environments allowed an unprecedented in-depth insight into the genetic architecture of the major agronomical traits of cultivated pepper. The observed correlations between two main antagonistic clusters of traits suggest how these traits, especially fruit morphology and fruit flavor, have been predominant in the breeding history of pepper which led to the development of divergent types. The colocalization on a few chromosomal regions of a priori unrelated traits (Table S8; Figure 5) could stem from pleiotropic genes or from several linked QTLs, which should be determined by further fine mapping. As an example, the intervals of the FShI-P3.4 and FP-P3.3 QTLs, whose most strongly associated markers are 17 kb apart, overlapped by 783 kb (Figure S8a,b). It is therefore not possible to conclude whether these two QTLs are determined by a single pleiotropic gene or by two physically linked genes. We observed that the GG haplotype, combining the alleles for narrow and elongated fruits (high FShI) and for low pungency (low FP), was absent from the set of 350 C. annuum (Figure S8c,d). The presence of the other three haplotypes indicates recombination between these two markers and possibly between the two associated causal genes, while the absence of the fourth haplotype may arise from a negative effect on another trait, and therefore its counterselection. However, FShI and FP are not completely dependent, as the core collection includes a few accessions with long, thin and sweet fruits, indicating the influence of other QTLs on these traits. The possible physiological antagonism of the clusters of traits should be taken into consideration for breeding.

Candidate genes were identified for 53 of the highestconfidence QTLs

We defined a set of candidate genes within the most promising QTL regions (Table S9; Figure 5), proven or suspected to be involved in molecular pathways determining the trait, or governing the same or similar traits in pepper or other species (Pflieger et al., 2001). QTL intervals were defined according to chromosome-averaged LD decay assessments, although it is known that there are some local LD decay variations along chromosomes, which mostly show suppression of recombination in pericentromeric regions and high recombination rates in the distal chromosome regions (Fuentes et al., 2022). Thus, some QTL intervals

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could be slightly misestimated. Here, four reported wellknown candidate genes (*Pun1, CCS, OFP20,* and *OVATE*) are somewhat outside the calculated QTL interval (Table S9). Additionally, the lack of markers in some regions reduces the chances to find associations near a known gene. In some cases, this may also be due to the difference in type of studied population: alleles with large effects that are segregating in a biparental population may be rare in the germplasm, hence difficultly identified by GWAS (Korte & Farlow, 2013). To conclude, the candidate genes may be valuable targets for marker-assisted selection.

Candidate genes for fruit flavor

Among the eight candidate genes for fruit pungency and Brix, we highlighted four strong candidates for three QTLs. The well-known Pun1 gene, corresponding to the locus C (Deshpande, 1935), determines pungency qualitatively through the presence or absence of capsaicinoids (Blum et al., 2002; Stewart et al., 2005) and is linked to our strongest marker-pungency association. This putative acyltransferase is indeed less than 50 kb apart from the QTL FP-P2.1 on chromosome P2. Furthermore, various members of the MYB transcription factor family, and particularly genes of the R2R3-MYB subfamily, were demonstrated to play key regulatory roles in capsaicinoid biosynthetic pathway (Arce-Rodríguez & Ochoa-Alejo, 2017; Han et al., 2019; Ju et al., 2020; Sun et al., 2019, 2020; Wang, Liu, et al., 2020). In our study, homologs of R2R3-MYB family genes MYB61 and MYB63, underlying QTLs FP-P2.1 and FP-P7.3, respectively, could therefore control pungency.

A homolog of the gene encoding the *SNF1-RELATED PROTEIN KINASE REGULATORY SUBUNIT BETA-2 (KINB2)* underlies *BX-P5.2* and may control Brix. Indeed, the Snf1related protein kinase (SnRK) complex plays a central role in the regulation of carbohydrate metabolism to deal with sugar limitation during stress conditions in Arabidopsis (Guérinier et al., 2013), and the inactivation of the SnRK complex subunit SIP1 leads to reduced levels of sugars (Anderson & Kohorn, 2001).

Candidate genes for fruit size and shape

Regarding the 28 candidate genes probably affecting fruit size and shape, we highlighted eight strong candidates for 11 QTLs. Ovate Family Proteins (OFP) are transcriptional repressors controlling multiple aspects of plant growth and development in multiple species including pepper (Wang et al., 2016). *OVATE* was the first gene identified as responsible for a tomato fruit shape QTL (Liu et al., 2002), and *CaOvate*'s down-regulation in pepper is likely a promoter of fruit elongation (Tsaballa et al., 2011). The *OVATE* homolog positioned at 155 Mb on chromosome P2, is closely linked to the fruit weight QTL *fw2.1* (Zygier et al., 2005; Table S1) and close to the QTL *FWe-P2.4*, detected

with the strongest marker-FWe association at position 160 Mb. Moreover, *C. annuum OVATE FAMILY PROTEIN* 20 (*CaOFP20*), known to control fruit elongation through cell expansion and replication (Borovsky et al., 2022), is 1.2 Mb distant from our strongest associations for FLe, FShI, PTh, and FF (*FLe-P10.1, FShI-P10.1, PTh-P10.1,* and *FF-P10.1*) and one of our strongest associations for FShO (-*FShO-P10.1*), which are all colocalized on chromosome P10. In peach, a chromosomal inversion downstream of an OFP gene is responsible for the flat fruit shape, comparable to the oblate pepper phenotype (Zhou et al., 2021). A homolog of the gene encoding the transcription repressor Ovate Family Protein 3 (*OFP3*) underlying the QTL *FLe-P6.4* could also determine fruit maximum length (FLe).

Transcription factors from the BEL1-LIKE HOMEODO-MAIN (BLH) family were found to interact with OFPs in Arabidopsis, cotton, rice, and banana (Hackbusch et al., 2005; Li et al., 2011; Liu & Douglas, 2015; Pagnussat et al., 2007; Schmitz et al., 2015; Wang et al., 2016; Zhang et al., 2016). Moreover, Bernard et al. (2021) found that a BLH protein was likely associated with walnut kernel weight. A homolog of *BEL1-LIKE HOMEODOMAIN PROTEIN 2* (*BLH2*) underlying the QTL *FWe-P2.4*, detected with the strongest marker-FWe association, may thus affect fruit weight.

In tomato, *CELL SIZE REGULATOR (CSR)* encodes a FAF-like protein controlling fruit weight (Mu et al., 2017). In pepper, a FAF protein interacting with protein WUSCHEL might regulate the size of shoot meristems as in Arabidopsis (Wahl et al., 2010), as well as fruit weight (Nimmakayala et al., 2016). A homolog of FANTASTIC FOUR 3 (FAF3) underlying the QTL *FLe-P6.2* may thus determine FLe.

Two members of the purine permeases (PUPs) transporter family independently induce a remarkable increase in rice grain size and length, probably by cytokinin transport and allocation into vascular bundle cells (Qi & Xiong, 2013; Xiao et al., 2019). The KATANIN microtubulesevering enzymes are AAA ATPases also involved in the progression of cell division, or cytokinesis, and cell division plane orientation (Luptovčiak et al., 2017). Wang et al. (2021) showed that a cucumber homolog of *KATANIN P60 ATPASE-CONTAINING SUBUNIT A1 (CsKTN1)* is involved in fruit elongation. Therefore, homologs of *PURINE PERMEASE 3 (PUP3)* and *KATANIN P60 ATPASE-CONTAINING SUBUNIT A1 (AAA1)* may govern fruit shape index, as they underlie QTLs *FShI-P6.3* and *FShI-P1.1*, respectively.

In tomato, the *lc* locus encoding WUSCHEL and the *fas* locus encoding the receptor-like kinase CLAVATA3 (CLV3) synergistically control the locule number (Chu et al., 2019; Muños et al., 2011). The cucumber homolog of *CLV3* also impacts carpel number (Che et al., 2020) and *CLAVATA1* (*CLV1*) may impact fruit size in *C. chinense* (Nimmakayala et al., 2021). A homolog of the probable leucine-rich repeat receptor-like protein kinase of the

CLAVATA family *At5g63930*, underlying the QTL *LN-P3.1*, wit may thus determine locule number. lear

Candidate genes for fruit color

Among the 16 candidate genes for immature fruit color, we highlighted two strong candidate genes for four QTLs. Homologs of transcription factor ARABIDOPSIS PSEUDO RESPONSE REGULATOR2 (APRR2) determine mature or immature fruit color in numerous cucurbits and solanaceous crops including melon (Oren et al., 2019), watermelon (Oren et al., 2019), cucumber (Liu et al., 2016; Tang et al., 2018), zucchini (Montero-Pau et al., 2017), wax gourd (Ma et al., 2021), tomato (Pan et al., 2013), eggplant (Arrones et al., 2022; Fang et al., 2023), as well as stem color in squash (Zhu et al., 2022). The pepper homolog of APRR2 on chromosome P1 corresponds to the well-known mature fruit color locus c1 (Kormos & Kormos, 1960), which was also shown to regulate chlorophyll content and immature fruit color (Jeong et al., 2020; Lee, Kim, et al., 2020; Pan et al., 2013) via regulating thylakoid numbers and chloroplast size (Lee, Kim, et al., 2020). Nonsense mutations resulting in premature stop codons were reported for this gene in all these crops, except zucchini (Arrones et al., 2022; Fang et al., 2023; Jeong et al., 2020; Lee, Kim, et al., 2020; Liu et al., 2016; Ma et al., 2021; Oren et al., 2019; Pan et al., 2013; Song et al., 2022; Tang et al., 2018; Zhu et al., 2022). Therefore, the APRR2 homolog nonsense mutation identified in our findings is a strong candidate for the colocalizing QTLs IFCG-P1.2, IFCa-P1.1, and IFCb-P1.1, corresponding to the strongest marker associations with three immature external fruit color traits.

The CAPSANTHIN-CAPSORUBIN SYNTHASE gene (CCS), involved in the synthesis of red carotenoid pigments in the mature fruit, corresponds to the y locus determining red versus yellow ripe fruit color in pepper (Lefebvre et al., 1998; Li et al., 2013; Popovsky & Paran, 2000). This gene is positioned less than 300 kb apart from our strongest marker association for immature fruit external color L (*IFCL-P6.1*), suggesting that it might also determine the intensity of the immature fruit color.

Within the 16 candidate genes pertaining to ripe fruit color, we highlighted 10 strong candidates for four QTLs, potentially related to pepper pigments. Flavonol synthase and flavanone 3-hydroxylase are involved in flavonoid synthesis (Preuß et al., 2009; Winkel-Shirley, 2001), while a sterol glucosyltransferase (*UGT80B1*) is required for flavonoid accumulation in the Arabidopsis seed coat (DeBolt et al., 2009). Their respective pepper homologs are thus plausible candidates for QTLs *FCa-P6.1* and *FCa-P3.1*, the latter being detected by the strongest marker-FCa association. Moreover, two homologs of a *LICODIONE SYNTHASE* (*CYP93B1*) underlie *FCa-P6.2* and may govern fruit color, as Mizuno et al. (2016) suggested that the encoded enzyme was related to the synthesis of flavones and associated

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with dark or light brown coloration in tan-colored injured leaves of sorghum.

Transcription factors of the MYB family regulate the expression of structural genes of the anthocyanin biosynthetic pathway in plants (Yan et al., 2021) and more specifically in C. annuum (Stommel et al., 2009), leading to changes in fruit coloration through anthocyanin accumulation (Borovsky et al., 2004). Moreover, CaMYB306 expression was higher in red fruit and its homolog's activity in tomato increases carotenoid accumulation and decreases anthocyanin and chlorophyll content (Ma et al., 2022). The two QTLs FCa-P3.1 (detected by the strongest marker-FCa association) and FCa-P6.2 are underlain by homologs of the three genes MYB35, MYB59 and MYB98 which may determine FCa. Additionally, the ubiquitin E3 ligase CON-STITUTIVE PHOTOMORPHOGENIC1 (MdCOP1) modulates the degradation of a MYB transcription factor in order to regulate light-induced anthocyanin biosynthesis in apple and probably in other species, as well as the red coloration of apple fruit (Li et al., 2012). Homologs of E3 UBIQUITIN-PROTEIN LIGASE 3 (UPL3) and a putative UPL3 LIN-like isoform X1 respectively underlie QTLs FCa-P6.1 and FCa-P1.2, and may thus determine FCa.

Lastly, FLAVIN-BINDING, KELCH REPEAT, F-BOX 1 (SIFKF1) may impact lycopene concentrations in tomato (Shibuya et al., 2021), suggesting that the pepper homolog of the F-box/kelch-repeat protein At3g18720 underlying FCa-P3.1 could impact FCa.

Candidate genes for plant vigor and earliness

Among the seven plant vigor candidate genes, we highlighted three strong candidates for three QTLs. The *PLAT DOMAIN PROTEIN 1 (PLAT1*), a potential downstream target of the abscisic acid signaling pathway, is involved in plant growth in Arabidopsis (Hyun et al., 2014), while members of the growth-regulating factor (GRF) family are involved in rice stem elongation (Cao et al., 2017; van der Knaap et al., 2000). Homologs of *PLAT1* and *GRF12* may therefore govern axis length as they underlie QTLs *AL-P5.4* and *AL-P1.2*, respectively. Moreover, the total plant height QTL *PHe-P2.1* is underlain by a homolog of *NAR-ROW LEAF 1 (NAL1)* which encodes a putative trypsin-like serine and cysteine protease affecting plant height in rice (Qi et al., 2008; Subudhi et al., 2020).

Regarding the eight flowering time candidate genes, we highlighted three strong candidates for two QTLs. *CON-STANS* and *CONSTANS*-like genes are crucial mediators of the photoperiodic induction of flowering in numerous plants including Arabidopsis, aspen trees, rice, Ipomoea, chrysanthemum and tomato (Böhlenius et al., 2006; Fu et al., 2015; Liu et al., 2001; Suárez-López et al., 2001; Yang et al., 2020; Yano et al., 2000). Their function may thus be conserved across angiosperms, making the homolog of Zinc finger protein *CONSTANS-LIKE* 6 (*COL6*) a strong

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The Plant Journal, (2023), doi: 10.1111/tpj.16425

candidate for the QTL *FIT-P3.3*. Moreover, *SNW/SKI-INTERACTING PROTEIN* (*SKIP*) affects both circadian clock and flowering time in Arabidopsis through alternative splicing (Cao et al., 2015; Cui et al., 2017; Wang et al., 2012), and two paralogous genes encoding GATA-type transcription factors repress flowering time, probably by repressing gibberellin and auxin signaling (Hudson et al., 2011; Richter et al., 2010, 2013). Homologs of *SKIP* and of putative *GATA22* underlie *FIT-P9.2* and could thus affect FIT.

Candidate genes for plant productivity

Among the 14 candidate genes for total fruit number, we highlighted two strong candidates for two QTLs. The QTL *TFN-P6.1*, corresponding to the strongest marker association with TFN, is underlain by a homolog of *AINTEGU-MENNTA (ANT)*. In Arabidopsis, this gene and its homolog *AINTEGUMENTA-LIKE 6 (AIL6)* have well-established importance in flower organs initiation and development, and affect flower number (Elliott et al., 1996; Krizek, 2009; Krizek et al., 2020; Long & Barton, 2000). A possible NAC transcription factor (*CA03g28100*) underlying *TFN-P12.2* may also affect TFN, given that different transcription factors from the NAC gene family influence fruit number in tomato (Lira et al., 2017; Ma et al., 2018; Wang, Zheng, et al., 2020).

EXPERIMENTAL PROCEDURES

Plant material

A core collection of Capsicum spp. was constructed using the GBS dataset described in Tripodi et al. (2021), corresponding to 10 038 accessions originated from 10 genebanks (Ozalp et al., 2020; Salinier et al., 2022). A Euclidean distance matrix was computed using the function dist from the stats R package (R Core Team, 2021). A core collection of 449 accessions was created with the coreCollection (v0.9.5)package (https://cran.r-project.org/package= coreCollection), using the Accession-to-Nearest-Entry "A-NE" criterion as introduced by Jansen and van Hintum (2007), the "split" adjusted group method, the "randomDescent" algorithm, and default values for the remaining parameters, with a preselected set of 26 wild and 84 cultivated accessions used as controls by the G2P-SOL project partners. For phytosanitary reasons, 26 of the 449 accessions could not be sent by the genebanks, reducing the G2P-SOL core collection to 423 Capsicum spp. accessions. From this core collection, 350 C. annuum accessions with less than 5% heterozygosity and less than 30% of missing data were retained for GWAS purposes. Accessions with a higher heterozygosity level were probably mixed genotypes since they were often scored as phenotypically heterogeneous in more than one trial.

Genotyping

DNA was extracted from leaf tissue samples of a single plant of each accession of the core collection only, and a second genotyping experiment was performed following the same *Mspl/Pstl* GBS protocol as reported in Tripodi et al. (2021) for the primary genotyping. Reads were aligned with *C. annuum* cv. CM334 reference genome version 1.6 (Kim et al., 2017), using the Burrows-Wheeler

Aligner (BWA) program (v0.7.17-r1188) and the 'mem' command with default parameters (Li, 2013). BAM files were processed and used for the SNP calling using GATK (v4.1.4.1; DePristo et al., 2011). SNPs were filtered using VariantFiltration, with QualBy-Depth, FisherStrand, RMSMappingQuality, MappingQualityRank-SumTest, and ReadPosRankSumTest arguments (QD < 2.0 \parallel FS > 60.0 \parallel MQ < 40.0 \parallel MQRankSum < -12.5 \parallel ReadPosRankSum < -8.0). The markers that were polymorphic within the 350 *C. annuum* accessions used for GWAS were combined from both genotyping experiments. A final dataset of 10 195 markers was obtained after filtering with Bcftools (v1.15; Li, 2011) to retain biallelic SNPs with a MAF greater than 0.01, a mean read depth greater than 20% of the accessions. Finally, accessions showing a heterozygosity higher than 2% were removed.

Field trials

Six trials (independent experiments from different years and/or locations) were organized. Under a Mediterranean climate, the National Research Institute for Agriculture, Food and Environment (INRAE) conducted two trials in France in 2019 and 2020, the Council for Agricultural Research and Economics (CREA) conducted one trial in Italy in 2019, the Agricultural Research Organization (ARO) conducted one trial in Israel in 2020, and Bati Akdeniz Agricultural Research Institute (BATEM) conducted one trial in Türkiye over 2020 and 2021. The World Vegetable Center (WorldVeg) conducted one trial in Taiwan over 2020 and 2021, under a subtropical climate. Depending on the trial, the plants were transplanted 5–7 weeks after sowing, in a randomized complete block design with two to three blocks, and two to four plants per accession per block. Metadata for each trial and each trait are listed in Tables S4 and S5, respectively.

Phenotyping

A total of 23 agronomic traits mainly based on pepper descriptors described by the International Plant Genetic Resources Institute (IPGRI) were assessed (Table 1; Table S5). Fifteen of the 18 quantitative traits had a continuous measurement (axis length, AL; Brix, BX; fruit external color, FCL, FCa, and FCb; fruit maximum length, FLe; fruit shape index, FShI; fruit weight, FWe; fruit maximum width, FWi; immature fruit external color, IFCL, IFCa, and IFCb; total plant height, PHe; pericarp thickness, PTh; total fruit weight, TFWe) and three had discrete values (flowering time, FIT; locule number, LN; total fruit number, TFN). Three of the five qualitative traits were ordinal traits visually assessed following categorical values in a range (fruit fasciation, FF; fruit load, FLo; external immature fruit color green, IFCG), as opposed to the two binary traits assessed as presence/absence (fruit pungency, FP; fruit predominant shape oblate, FShO).

Data analysis

Statistical analyses using R software were performed with version 4.1.2 (R Core Team, 2021).

Population structure

To investigate the population structure, the SNP panel (without Indels) was pruned based on LD using plink2 (Purcell & Chang, 2019) with the command --indep-pairwise 50 10 0.2. PCA was performed using the SNPRelate R package (v3.13; Zheng et al., 2012) with default parameters. In addition, the population structure was analyzed by ADMIXTURE (v1.3.0; Alexander et al., 2009) with the cluster number K ranging from 1 to 15, using a CV value of 15 (--cv = 15). Each K was run with 20 replicates and the outputs were analyzed using pong (v1.5; Behr et al., 2016) and plotted with pophelper (v2.3.1; Francis, 2017). CV error was used to determine a suitable value for K. Individuals were assigned to one of the K populations when their membership coefficient in that group was \geq 0.70.

LD decay study

LD was estimated as pairwise squared allele-frequency correlation (r^2) between SNP markers, correcting for relatedness using a kinship matrix by the R package LDcorSV (v1.3.3; Mangin et al., 2012). The kinship matrix was estimated as covariance between accessions. The overall LD decay estimated per chromosome was likely not significantly affected by non-recombinant heterochromatic regions, since gene-rich euchromatic regions were preferentially sequenced because of the use of methylation-sensitive *Mspl/Pst* restriction enzymes for the GBS protocol. The r^2 values were binned according to the physical distances (Mb) between SNP pairs, and the 95th percentile of each bin was fitted by a polynomial curve. The extent of LD decay was estimated as the distance (Mb) where LD decayed below the arbitrarily chosen baseline of 0.2.

Phenotypic data processing

The proportions of explained variance were calculated following a fixed-effects linear model: $Y_{ijk} = \mu + G_i + E_j + G_i \times E_j + \varepsilon_{ijk}$, where Y_{ijk} represents the phenotypic value of the plant *k* from the genotype *i* (G_i) in the environment *j* (E_j), $G_i \times E_j$ the interaction between genotype and environment, and ε_{ijk} the residual error. The model was fitted using the function Im from the stats R package (R Core Team, 2021). The proportions of explained variance were calculated as the ratio of the sum of squares of the genetic, environmental, or G \times E effect considered to the total sum of squares.

The broad-sense heritability was computed following the approach proposed by Cullis et al. (2006) implemented in the inti R package (v0.5.5; Lozano-Isla, 2022). It uses genotypic BLUPs calculation and is particularly suitable for unbalanced datasets (Piepho & Möhring, 2007). First, a random-effects linear model was fitted: $Y_{ijk} = \mu + G_i + E_j + G_i \times E_j + e_{ijk}$, where Y_{ijk} represents the phenotype value of the plant *k* from the genotype *i* (G_i) in the environment *j* (E_j), $G_i \times E_j$ the interaction between genotype and environment and e_{ijk} the residual error. Then, heritability was calculated as $h^2 = 1 - \overline{v}_{\Delta}^{\text{BLUP}} / (2 \times \sigma_g^2)$, where $\overline{v}_{\Delta}^{\text{BLUP}}$ corresponds to the mean variance of a difference of two BLUPs for the genotypic effect and σ_g^2 to the genotypic variance. Environment *j* (E_j) represents the trial *j* in the models used for computing the proportion of explained variance and the heritability, except for external immature fruit color parameters measured at BATEM only (for IFCL, IFCa and IFCb, E_i represents the block *j*).

Quantitative traits were screened for the presence of outliers within each trial using a median absolute deviation (MAD)-based Bonferroni-Holm test (Bernal-Vasquez et al., 2016) with $\alpha = 0.01$, applied on the MAD-standardized residuals of a linear model correcting for genotype, block and genotype-block interaction effects (Table S5). The model was fitted using the function Im from the stats R package (R Core Team, 2021). Locule number was treated as an ordinal trait. When the degrees of freedom were insufficient due to a lower number of measurements, the correction for interaction effect was not applied. When the block information was not available for a particular trial, only the correction for genotype effect was applied. Simple descriptive statistics were computed for accession means of each trait after outlier removal, with the stat.desc function from the pastecs R package (v1.3.21; Grosjean et al., 2018). Between-trial Pearson correlations were

Multi-environment GWAS in a pepper core collection 15

computed for each trait on the accession means per environment, to detect less correlated trials. The Box-Cox family of power transformations was then applied to the most correlated trials together to optimize normality of residuals for each trait, after fitting a model including all the main effects for genotype, trial and block (and trial-block interaction when the number of measurements was sufficient) (Table S5). Prior to Box-Cox transformation, traits containing negative values were transformed into only-positive ones adding an appropriate constant. The less correlated trials were transformed separately. This procedure was performed using the functions powerTransform and bcPower from the R package car (v3.0.13; Fox & Weisberg, 2019). Adjusted means per accession to be used as phenotypic input for GWAS were estimated using a BLUP model with the 'Ime4' R package (Bates et al., 2015). These BLUPs were calculated both for single environments (correcting for block and genotype-block interaction effects when the number of measurements was sufficient) and across highly correlated environments (correcting for block nested in the trial effects and for genotype-block interaction effects when the number of measurements was sufficient). The lesser correlated trials were analyzed through single environment GWAS only, not to bias the multi-environment GWAS conducted on the higher correlated trials. Accessions with extreme adjusted means that were more than three MADs apart from the median were then removed, as they could bias the estimation of marker effects (Table S5).

Ordinal traits and locule number were visually screened for outliers, and between-trial Spearman correlations were computed on the accession means per environment, in order to detect less correlated trials. The Box-Cox family of power transformations was applied and adjusted means per accession were calculated as for the quantitative traits. Single-environment BLUPs were not computed for the less correlated trials. The lower correlations for these ordinal and more subjective traits may have arisen from divergent appreciations of the different trial assessors. Accessions with extreme adjusted means were removed by visual inspection, using the same lower and upper bounds for the singleenvironment BLUPs and the multi-environment BLUPs (Table S5).

The fruit pungency (FP) and fruit predominant shape oblate (FShO) binary traits were coded as 0 versus 1. Between-trial Spearman correlations were computed to detect less correlated trials. Adjusted means per accession were then calculated as for the other traits.

Genome-wide association study

GWAS were conducted on both single-environment and multienvironment BLUPs, using the R package "Genomic Association and Prediction Integrated Tool" (GAPIT Version 3; Wang & Zhang, 2021). The mean dosage value per marker across accessions was used for imputing the remaining missing values (5% of the genotypic matrix). Three GWAS models were used: MLM (Yu et al., 2006), MLMM (Segura et al., 2012), and BLINK (Huang et al., 2019). The first four PCs of a PCA of marker genotypes were added to all the GWAS models as covariate variables, to reduce the number of false positives due to population stratification. In addition, MLM and MLMM included a random term to correct for relatedness according to a kinship matrix based on the VanRaden formula, calculated by GAPIT3. To control familywise error rate, a significance threshold of 5.31 was calculated using the Bonferroni procedure (i.e., dividing the desired significance of 0.05 by the number of tested markers 10 195). Circular Manhattan plots were generated using the R package CMplot (v4.2; Yin et al., 2021).

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The Plant Journal, (2023), doi: 10.1111/tpj.16425

QTLs and candidate genes

QTL intervals were calculated in order to group together highly linked QTNs (i.e., the SNP and Indel markers significantly associated to a trait) identified using the three different GWAS models (MLM, MLMM, and BLINK) on up to six environments or across all environments. Intervals were defined as the first and last QTN position \pm the extent of specific chromosomal LD decay. Each QTL name was defined as the associated trait code, followed by the chromosome number and the QTL's order among all QTLs for this trait along the chromosome. Some of the QTLs were categorized as more promising, when they were robust (i.e., significant in at least two environments), or detected with at least two GAPIT models, or with a -log10(P-value) higher than five above the threshold (i.e., higher than 10.31). Since quantitative external immature fruit color traits were measured in one trial only, their QTLs with a -log10(P-value) higher than 2 above the threshold (i.e., higher than 7.31) were also included. Annotations for genes within the most promising QTL bounds were retrieved from C. annuum cv. CM334 reference genome version 1.55 (Kim et al., 2014) and version 1.6 (Kim et al., 2017). Their function in plants was then investigated in the literature, and genes involved in traits linked to the pepper associated trait were therefore proposed as putative causal genes.

Outlook

We constructed the G2P-SOL core collection, a unique and valuable resource for the pepper community. Its great genotypic and phenotypic variability, representative of the worldwide diversity, will accelerate the comprehensive understanding of the genetic basis of multiple traits. The core collection will be available on request, along with the genotypic and phenotypic data collected as part of the G2P-SOL project. As research resources, we also provided comprehensive up-to-date summaries of previously detected QTLs for agronomic traits in pepper and previously published GWAS in C. annuum. We conducted multi-environment GWAS on a great variety of plant and fruit traits, and GWAS for biotic stress resistance to multiple pathogens, heat stress tolerance and metabolic compounds is underway. We identified 97 new or already known genes potentially underlying 53 of the 119 highestconfidence QTLs. Further validation of specific genes will be needed to confirm their incidence on the phenotype. Nevertheless, these results demonstrate the usefulness and reliability of our core collection, phenotypic dataset and methodology. Traditionally, candidate genes identified for traits with simple inheritance can be used as diagnostic markers or as part of trait-specific markerassisted selection. In addition, the genomic and phenotypic matrices provided by this study should improve multivariate genomic prediction at the whole-genome level, and should speed up the identification of candidate accessions that are phenotypically closest to a predefined genetic ideotype, adapted to a targeted environment or to several environments (Yang et al., 2022). The gene pool formed by these candidate lines, whose genetic backgrounds may be different, would greatly influence the simultaneous prebreeding of multiple quantitative traits, and contribute to the development of genetically improved climate-resilient pepper crops.

AUTHOR CONTRIBUTIONS

VL, IP, PT, LB, and GG conceived the study. LM, VL, JS, CG, PT, HFB, RO, IP, YB, RS, and DB performed the agronomic trials and phenotyping. NS and MTR-W coordinated the genotyping-by-sequencing. LM, VL, LB, GT, MB, RF, and PT realized the bioinformatic and statistical analyses. LM, VL, LB, IP, PT, GT, and RV wrote the manuscript. All authors approved the manuscript.

ACKNOWLEDGMENTS

Authors gratefully acknowledge ARO, BATEM, CREA, INRAE, WorldVeg, IPK (Andreas Boerner), MVCRI (Maritsa Vegetable Crops Research Institute, Gancho Pasev), UPV (Universitat Politècnica de València, Maria-Jose Diez), CGN (Centre for Genetic Resources, Willem van Dooijeweert) for providing plant seeds to build the core collection. In this matter, the World Vegetable Center obtained additional support from their long-term strategic donors Taiwan, UK aid from the UK government, United States Agency for International Development (USAID), Australian Centre for International Agricultural Research (ACIAR), Germany, Thailand, Philippines, Korea and Japan. The G2P-SOL pepper core collection is now maintained and distributed by the INRAE GAFL's genebank, CRB-Lég. We are grateful to the members of the CRB-Lég and ReDD teams of INRAE GAFL unit and of the INRAE Mediterranean Fruit and Vegetables Experimental Facility (A2M, https://doi.org/10.17180/nxpc-kv48) for their technical assistance for the trials. This work was funded by the G2P-SOL project (Linking genetic resources, genomes, and phenotypes of Solanaceous crops) financed by the European Union's Horizon 2020 Research and Innovation Program under the grant agreement number 677379, and by the PRO-GRACE project (Promoting a Plant Genetic Resource Community for Europe) financed by the Horizon Europe Program under the grant agreement number 101094738. LM received a 3-year fellowship from the project G2P-SOL and from INRAE Plant Biology and Breeding Division.

CONFLICT OF INTEREST

The authors declare no conflicts of interest.

DATA AVAILABILITY STATEMENT

The raw phenotypic and VCF genotypic datasets generated for this study are accessible at https://doi.org/10.57745/ S7JVEM. Seeds from the G2P-SOL core collection will be available upon reasonable request at INRAE GAFL CRB-Lég, considering the sharing of costs for their production and their shipping (e-mail veronique.lefebvre@inrae.fr).

SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article.

Figure S1. Genotypic dataset characteristics.

Figure S2. Average linkage disequilibrium (LD) decay plots for the 12 chromosomes.

Figure S3. A cluster number K of 11 derived from an ADMIXTURE population structure analysis of the 350 accessions used for GWAS.

Figure S4. Principal component analysis (PCA) using accession means of all 23 traits for each of the six independent trials.

Figure S5. Visualization of between-trials correlation matrices for the 23 traits.

Figure S6. Between-traits pairwise-complete Pearson correlation heatmap for each environment.

Figure S7. Physical position of all 423 quantitative trait loci (QTLs) detected for each of the 23 traits, mapped on the 12 pepper chromosomes.

Figure S8. *FShI-P3.4* and *FP-P3.3* fruit shape index (FShI) and fruit pungency (FP) overlapping QTLs on chromosome P3 lack a haplo-type combining the alleles for narrow and elongated fruit with low pungency.

Table S1. Summary of *Capsicum* spp. QTLs in the literature from1998 to 2022.

Table S2. Summary of genome-wide association studies on *Capsicum annuum* from 2014 to 2023.

 Table S3. List of Capsicum spp. accessions included in the G2P-SOL core collection.

Table S4. Metadata of the six phenotypic trials.

Table S5. Characteristics of the 23 traits phenotyped in up to six trials.

Table S6. Number of marker-phenotype associations found according to the trial and to the genome-wide association study model.

 Table S7. Full list of the 423 quantitative trait loci detected for the 23 traits.

 Table S8. List of highest-confidence quantitative trait loci colocalized for several traits.

 Table S9. Full list of candidate genes for the highest-confidence

 QTLs and their annotations.

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The Plant Journal published by Society for Experimental Biology and John Wiley & Sons Ltd., *The Plant Journal*, (2023), doi: 10.1111/tpj.16425

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