

Genetic Architecture of Chile Pepper (*Capsicum* Spp.) QTLome Revealed Using Meta-QTL Analysis

Dennis N. Lozada (✉ dlozada@nmsu.edu)

Department of Plant and Environmental Sciences, New Mexico State University, Las Cruces, NM 88003

Lanie Whelpley

New Mexico State University

Andrea Acuña-Galindo

University of Arkansas at Fayetteville

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Abstract

Chile peppers (*Capsicum* spp.) are among the most important vegetable crops in the world due to their health-related, economic, and industrial uses. In recent years, quantitative trait loci (QTL) mapping approaches have been widely implemented to identify genomic regions affecting variation for different traits for marker-assisted selection (MAS) in peppers. Meta-QTL analysis for different traits in *Capsicum* remains lacking, and therefore it would be necessary to re-evaluate identified QTL for a more precise MAS for genetic improvement. We report the first known meta-QTL analysis for diverse traits in the chile pepper QTLome. A literature survey using 29 published linkage mapping studies identified 766 individual QTL from five different trait classes. A total of 311 QTL were projected into a consensus map. Meta-analysis identified 30 meta-QTL regions distributed across the 12 chromosomes of *Capsicum*. *MQTL5.1* and *MQTL5.2* related to *Phytophthora capsici* fruit and root rot resistance were delimited to < 1.0 cM confidence intervals in chromosome P5. Candidate gene analysis for the flanking sequences for the P5 meta-QTL revealed biological functions related to DNA repair and transcription regulation. Moreover, epigenetic mechanisms such as histone and RNA methylation and demethylation were predicted, indicating the potential role of epigenetics for *P. capsici* resistance. Allele specific SNP markers for the meta-QTL will be developed and validated using different breeding populations of *Capsicum* for MAS of *P. capsici* resistant lines. Altogether, results from meta-QTL analysis for chile pepper QTLome rendered further insights into the genetic architecture of different traits for this valuable horticultural crop.

Introduction

The wealth of genomic information available for different crop species and diverse traits has expanded significantly in recent years due to the decreasing costs of high-throughput genotyping and the development of novel and powerful tools to dissect quantitative trait loci (QTL), which are genomic regions affecting variation for quantitative traits¹. A 'QTLome' refers to the collection of QTL and their allelic variation shown to affect any quantitative trait for a given trait and species^{2,3}. The first step to interpret information from QTLome is meta-QTL analysis^{2,4}. QTL meta-analysis integrates results from multiple QTL studies and could provide more insights into the genetic architecture of traits associated with different genomic regions⁵. The QTL identified from a group of different QTL using meta-analysis at a 95% confidence interval are called meta-QTL⁶. In meta-QTL analysis, various models can be implemented for identifying the consensus QTL from different studies thereby validating results and refining positions of QTL in the consensus map^{4,7}. Using information from meta-analysis could therefore allow for a more precise marker-assisted breeding for the genetic improvement of traits across different crops.

Chile peppers, belonging to the genus *Capsicum* and family Solanaceae, are one of the most important horticultural crops in the world due to their culinary uses, health benefits, and economic impact. Nutrients such as vitamins A, C, and folate are present in varying degrees in peppers⁸. Chile peppers have been used to help combat chronic pain, and it was suggested that capsaicin, the compound found predominantly in pungent lines, can induce depletion in nerve sensory terminals, and is commonly found in pain relieving creams⁹. In the state of New Mexico in the United States, chile peppers are important cash crops for farmers¹⁰. A present disadvantage of the cultivation of *Capsicum* species, however, is an inadequate supply due to low yield from farmers. Low yield can be attributed to many factors, including pests and diseases, and undesired agronomic traits such as subpar fruit size or uneven biomass distribution¹¹⁻¹³. In the past, marker-assisted selection through genetic mapping has been implemented in chile peppers to facilitate genetic improvement for different traits including yield and resistance to major diseases.

While many QTL for diverse traits have been identified for chile peppers, there is no known report of meta-study across diverse traits in *Capsicum*. Most of the meta-QTL studies by far focused on diverse sets of traits across the Solanaceous relatives of chile peppers such as potatoes (*Solanum tuberosum*)¹⁴ and tomatoes (*S. lycopersicum*)¹⁵; and in major field crops such as rice (*Oryza sativa*)^{6,16}, wheat (*Triticum aestivum*)^{17,18}, barley (*Hordeum vulgare*)¹⁹, maize (*Zea mays*)^{20,21}, cotton (*Gossypium* spp.)²², and soybeans (*Glycine max*)²³. In chile peppers, the only known report for meta-QTL to date was a study by Mallard et al.²⁴, who identified meta-genomic regions for *P. capsici* resistance, designated as *MetaPc5.1*, *MetaPc5.2*, and *MetaPc5.3*, in chromosome P5. Meta-analysis indicated that *MetaPc1* confers resistance against eight isolates, whereas *MetaPc5.2* and *MetaPc5.3* exhibit resistance for three isolates of *P. capsici*.

Given that meta-QTL studies remain lacking in chile peppers, it would be necessary to implement meta-analysis to determine significant genomic regions involved in variation for important traits. The current study aims to perform analysis of meta-QTL to identify regions associated with diverse traits for targeted genetic improvement in chiles. Specifically, our objectives were to (1) develop a consensus genetic map in chile peppers using SNP markers; (2) identify meta-QTL related with different traits such as heat levels (pungency), yield, adaptation, and resistance to diseases in peppers; (3) determine SNP markers linked to meta-QTL; and (4) identify candidate genes present in the meta-QTL regions. We report the first known meta-QTL study of the chile pepper QTLome that could provide insights into the genetic architecture of different traits for this important horticultural crop.

Results

Classification of QTL for diverse traits in *Capsicum* spp. Overall, 766 individual QTL from 130 unique traits and five different trait classes across the 12 chromosomes of chile pepper were identified. These results were based from 29 linkage mapping studies published within a 10-year period (2010–2020) (Table 1; Fig. 1). Among these studies, majority (90%) of the reported QTL were identified using either a recombinant inbred line (RIL) or an F₂ biparental mapping population. The number of individuals used for QTL mapping ranged between 63 and 440. Marker types used in identifying QTL included amplified fragment length polymorphism (AFLP), simple sequence repeats (SSR), and single nucleotide polymorphism (SNP) markers, among others. Across the various linkage-mapping studies surveyed, among the most commonly used parents in creating a biparental mapping population included ‘CM-334’, a *P. capsici* resistant line for identification of QTL linked to chile pepper blight resistance; ‘Bhut Jolokia’, a ‘superhot’ chile pepper for the discovery of QTL associated with capsaicinoid content; and ‘Yolo Wonder’, a bell-pepper type, for linkage mapping of stem and fruit-related traits and disease resistance QTL.

Table 1

Summary of QTL mapping studies used for meta-analysis of diverse traits for the *Capsicum* QTLome. ^a *BC*- Backcross; *DH* Double haploid; *RIL*- Recombinant inbred line. ^b *AFLP*- Amplified fragment length polymorphism; *CAPS*- Cleaved amplified polymorphic sequence; *HRM*- High resolution melting; *InDel*- Insertion/Deletion; *ISSR*- Inter simple sequence repeat; *RAPD*- Random amplified polymorphic DNA; *SCAR*- Sequence characterized amplified region; *SLAF*- Specific locus amplified fragment sequence; *SNP*- Single nucleotide polymorphism; *SPP*- Single position polymorphism; *SSR*- Simple sequence repeats. ^c F_1 progenies of MP crossed with GNM and SM; BC_1 crossed with GNM; For SP, F_1 and BC_1 crossed with GNM.

Study No.	Reference	Parents used in crossing	No. of individuals	Mapping population ^a	Trait(s)	Marker type ^b
1	Alimi et al. ⁵⁵	Yolo Wonder x CM334	149	RIL	Fruit and stem-related traits	SSR
2	Arjun et al. ⁵⁶	<i>C. annuum</i> FL 201 x <i>C. galapagoense</i> TC 07245	210	F_2	Fruit Length	SSR
3	Chunthawodtiporn et al. ⁵⁷	<i>C. annuum</i> Maor x CM334	120	RIL	Horticultural traits	SNP
4	Du at al. ⁵⁸	BVRC 25 x BVRC 1	440	F_2	Bacterial wilt resistance	SNP, InDel
5	Dwivedi et al. ⁵⁹	California Wonder x LCA235	74	RIL	Plant height and yield related traits	SSR, SCAR, RAPD
6	Dwivedi et al. ⁶⁰	California Wonder x LCA235	74	RIL	Oleoresin content	SSR, SCAR, RAPD
7	Eggink et al. ⁶¹	MT x PEN45 (MP); SM x PEN45 (SP); GNM ^c	250	BC	Agronomic, biochemical, physiological traits	AFLP, SNP
8	Han et al. ⁴¹	Perennial x Dempsey (PD); TF68 x Habanero (TH)	56 (PD), 85 (TH)	RIL	Capsaicinoid content	SNP
9	Kim et al. ⁶²	<i>C. baccatum</i> var. Pendulum x Golden-aji	126	F_2	Anthraxnose resistance	SSR, AFLP, SRAP
10	Lee et al. ⁶³	NB1 x Bhut Jolokia	175	F_2	Capsaicinoid content	HRM, SSR, CAPS, Gene-based
11	Lee et al. ⁶⁴	CV4 x Jeju	300	F_2	Chili veinal mottle virus	SNP
12	Lee et al. ⁶⁵	Perennial x Dempsey	120	RIL	Fruit-related traits	SNP
13	Li et al. ⁶⁶	BJ0747 x XJ0630	195	F_2	Cucumber mosaic virus resistance	SLAF-SNP

Study No.	Reference	Parents used in crossing	No. of individuals	Mapping population ^a	Trait(s)	Marker type ^b
14	Liu et al. ³⁵	YCM334 x Tean	128	RIL	<i>P.capsici</i> resistance	SNP
15	Lu et al. ⁶⁷	YCM334 x Taean	126	RIL	Agronomic and morphological traits; <i>P. capsici</i> resistance	SNP
16	Mahasuk et al. ⁶⁸	Bangchang x PBC932 (BP), PBC80 x CA1316 (PC)	126 (BP), 146 (PC)	F ₂	Anthraco-nose resistance	SNP
17	Mallard et al. ²⁴	H3 x Vania (HV), Perennial x Yolo Wonder (PY), YW x CM334 (F5YC)	101 (HV), 114 (PY), 297 (RIL)	DH (PY, HV), RIL (F5YC)	<i>P.capsici</i> resistance	AFLP, CAPS, RFLP
18	Naegele et al. ⁶⁹	CM334 x EJ	63	RIL	<i>P.capsici</i> resistance	SNP
19	Park et al. ⁴²	Habanero x Jolokia (HJ), SNU11-001 x Jolokia (SJ)	87 (HJ), 124 (SJ)	F ₂	Capsaicinoid content	SNP
20	Park et al. ⁷⁰	M5 x AG13-3	96	F ₂	AGI activity	SNP
21	Rehrig et al. ⁷¹	CM334 x EJ	66	RIL	<i>P.capsici</i> resistance	SNP
22	Siddique et al. ³⁴	CM334 x ECW30R	188	RIL	<i>P.capsici</i> resistance	SNP
23	Sun et al. ⁷²	<i>C. annuum</i> 77013 x <i>C. chinense</i> PBC932	186	BC	Anthraco-nose resistance	SSR, InDel, CAPS
24	Tan et al. ⁷³	<i>C. annuum</i> BA3 x <i>C. frutescens</i> YNXML	154	F ₂	Flowering time	SSR, InDel
25	Wei et al. ⁷⁴	<i>Capsicum annuum</i> 007EA x <i>Capsicum frutescens</i> P1512	120	F ₂	Horticultural traits	SNP
26	Yao et al. ⁷⁵	BJ0747-1-3-1-1 x XJ0630-2-1-2-1-1	334	F ₂	Cucumber mosaic virus resistance	SSR, ISSR
27	Yarnes et al. ⁴³	2814-6 x NuMex R Naky	105	RIL	Plant architecture, phenology, fruit quality	SPP
28	Zhang et al. ⁷⁶	PM702 x FS871	146	RIL	First flower node	SLAF/SNP

Study No.	Reference	Parents used in crossing	No. of individuals	Mapping population ^a	Trait(s)	Marker type ^b
29	Zhu et al. ⁴⁶	<i>C. chinense</i> 740 x <i>C. annuum</i> CA1	150	F ₂	Flower number, flowering time	SLAF

Meta-QTL analysis and candidate gene mining. The *Capsicum* consensus map generated from three previously published linkage maps consisted of 14,952 total markers distributed across 12 chromosomes, with an average of 1,246 markers per chromosome (Table 2). Chromosomes P3 (2,231 SNP markers), P7 (1,996), and P6 (1,711) had the greatest number of loci, whereas P8 (431), P4 (707), and P10 (722) had the least. Average length of the consensus map ranged between 94.97 (P6) and 175.09 cM (P8). From the preliminary pool of 766 QTL, 418 remained after further processing and quality control (**Supplementary File 1**). The consensus map projected a total of 304 QTL, with P1 having the greatest number of projected QTL (61), followed by P5 (45), and P4 (29). Chromosomes P9, P7, and P8 had the least number of QTL projected in the consensus map, with seven, eight, and 11, respectively.

Analysis further identified 30 meta-QTL distributed in the 12 chromosomes of chile pepper, with a mean position of 58.83 cM across all chromosomes (Fig. 2; Table 3). Four meta-QTL were identified for chromosomes P1 and P5, whereas three meta-QTL were detected for P3, P11, and P12. Only a single meta-QTL was identified for chromosome P7. Mean genetic position of individual meta-QTL ranged between 12.19 (*MQTL2.1*) and 156.87 cM (*MQTL5.4*). Two meta-QTL in chromosome P5, *MQTL5.1* and *MQTL5.2*, comprised of clusters of QTL for *P. capsici* fruit and root rot resistance had confidence intervals of < 1.0 cM. *MQTL1.3* had the greatest number of individual QTL (33), followed by *MQTL5.2* (24), and *MQTL4.2* (19). Conversely, *MQTL5.4*, *MQTL6.1*, *MQTL8.2*, and *MQTL12.2* had only two individual QTL comprising the meta-QTL. QTL related with resistance to *P. capsici* was found in 11 out of the 30 meta-QTL detected (36.6%), whereas loci associated with anthracnose and fruit rot resistance was found in nine (30.0%) and seven (23.3%) of the identified meta-QTL, respectively. Biochemical and physiological traits such as AGI activity and transverse fruit perimeter QTL belong to six different meta-QTL. Akaike information criterion (AIC) values for determining the best QTL models in identifying meta-QTL ranged between 82.13 (P7) and 555.66 (P1), with majority of these models with the lowest AIC value being model 5. Two other chromosomes, P7 and P12, had QTL models 3 and 4 as the best model in identifying meta-QTL, respectively.

Table 2
 Length and number of markers for each chromosome
 of the constructed Capsicum spp. consensus map.

Chr.	Mean length (cM)	No. of SNP markers
P1	134.90	986
P2	113.95	1,442
P3	130.42	2,231
P4	116.43	707
P5	147.70	1,165
P6	94.97	1,711
P7	127.70	1,996
P8	175.09	431
P9	97.36	878
P10	112.47	722
P11	137.91	1,529
P12	120.63	1,154
Total	1,509.52	14,952
Average	125.79	1,246

Table 3

Meta-QTL identified for diverse traits in chile peppers (*Capsicum* spp.). a Akaike information criterion value for the best QTL model. b Meta-QTL identified in at least two independent linkage mapping studies. c Confidence interval.

Chr.	No. of QTL	No. of projected QTL	No. of Meta-QTL	QTL model	AIC value ^a	Meta-QTL ^b	No. of QTL in the meta-QTL	Mean genetic position (cM)	CI ^c (cM)
P1	73	61	4	5	555.66	<i>MQTL1.1</i>	15	16.75	2.59
						<i>MQTL1.2</i>	6	40.95	5.27
						<i>MQTL1.3</i>	36	67.62	2.11
						<i>MQTL1.4</i>	4	129.82	4.25
P2	64	27	2	5	424.59	<i>MQTL2.1</i>	13	12.19	1.6
						<i>MQTL2.2</i>	14	71.44	2.9
P3	43	27	3	5	307.38	<i>MQTL3.1</i>	6	22.33	4.05
						<i>MQTL3.2</i>	6	94.16	5.14
						<i>MQTL3.3</i>	15	146.12	1.65
P4	35	29	2	5	295.78	<i>MQTL4.1</i>	10	20.52	3.79
						<i>MQTL4.2</i>	19	84.92	2.02
P5	51	45	4	5	316.37	<i>MQTL5.1</i>	13	23.03	0.79
						<i>MQTL5.2</i>	25	29.16	0.55
						<i>MQTL5.3</i>	7	87.33	1.07
						<i>MQTL5.4</i>	2	156.87	12.42
P6	27	20	2	5	217.58	<i>MQTL6.1</i>	2	14.7	10.49
						<i>MQTL6.2</i>	19	54.73	1.65
P7	11	8	1	3	82.13	<i>MQTL7.1</i>	8	38.99	2.53
P8	15	14	2	5	124.72	<i>MQTL8.1</i>	9	22.13	5.61
						<i>MQTL8.2</i>	2	62.24	4.01
P9	17	10	2	5	124.95	<i>MQTL9.1</i>	4	43.8	7.87
						<i>MQTL9.2</i>	3	74.89	9.43
P10	35	26	2	5	231.24	<i>MQTL10.1</i>	13	22.39	1.86
						<i>MQTL10.2</i>	13	83.87	2.29
P11	24	23	3	5	214.53	<i>MQTL11.1</i>	5	19.3	5.72
						<i>MQTL11.2</i>	11	54.56	4.01
						<i>MQTL11.3</i>	7	84.22	3.0
P12	24	21	3	4	242.70	<i>MQTL12.1</i>	13	23.51	2.82

Chr.	No. of QTL	No. of projected QTL	No. of Meta-QTL	QTL model	AIC value ^a	Meta-QTL ^b	No. of QTL in the meta-QTL	Mean genetic position (cM)	CI ^c (cM)
						<i>MQTL12.2</i>	2	55.69	10.97
						<i>MQTL12.3</i>	6	106.76	14.90
Total	419	311	30				308		

A total of 346 candidate genes with known biological functions were identified using sequence information for the flanking markers for *MQTL5.1* (*Scaffold 3155.426970* and *Marker1044376*) and *MQTL5.2* (*Cb-HRM288* and *Marker635294*) in chromosome P5 (**Supplementary Table S1**). These ranged between five and 318 candidate genes for *Scaffold 3155.426970* (*MQTL5.1*) and *Marker635294* (*MQTL5.2*). A wide range of biological functions for the candidate genes, including those related with DNA replication, repair, transcription regulation, phosphorylation, and glycosylation, among others, for *MQTL5.1* and *MQTL5.2* has been identified. Moreover, several functions related with epigenetic mechanisms such as RNA, DNA, tRNA, and histone methylation have been predicted. Among the candidate genes identified for *MQTL5.2*, *PHT67052* and *PHT65976*, have functions related to transcription regulation, histone H3-K36 methylation, and histone lysine methylation; and regulation of gene expression, respectively, in *C. annuum*. *Solyc11g033270.2.1* (*MQTL5.2*) has functions associated with regulation of defense response to bacterial and fungal infections, as well as activation of protein kinase activity, and stress-activated protein kinase signaling cascade in *S. lycopersicum*. Gene *Solyc11g013370.2.1* has roles in defense response to fungus, leaf morphogenesis, seed dormancy, and positive regulation of cell division in tomatoes. Genes *OIS99998* and *OIS96773* have roles in methylation in wild tobacco (*Nicotiana attenuata*), whereas *PGSC0003DMT400072321* has functions related to double-strand break repair via homologous recombination; DNA replication, recombination, and repair, and *PGSC0003DMT400003724* has functions in tRNA N2-guanine methylation, tRNA processing, and methylation in *S. tuberosum*.

Discussion

The identification and analysis of notable QTL regions associated with different traits has been one of the cornerstones of modern molecular breeding for plant genetic improvement. In *Capsicum* spp., previous studies focused on identifying QTL linked to diverse traits including but not limited to resistance to major diseases such as chile pepper blight caused by the oomycete *P. capsici*, yield and yield components, capsaicin (heat) content, and agronomic traits, among others (Table 1). Given this wealth of information from previous QTL studies, it would be necessary to re-evaluate results from linkage mapping using a meta-analysis approach to refine genomic regions associated with important traits resulting in a more efficient implementation of MAS in chile pepper breeding programs. QTL meta-analysis for different traits in *Capsicum* remains lacking, where a major focus in the past has been the identification of meta-QTL for resistance to *P. capsici*²⁴. This status quo of meta-studies for chile peppers has thus driven us to explore meta-QTL for the *Capsicum* QTLome for diverse traits, i.e. not only for those QTL related with disease resistance, but also for those loci linked with other important yield and agronomic characters in chiles. Here, we report the first known meta-analysis of the chile pepper QTLome rendering deeper insights into the genetic architecture of diverse sets of traits for this valuable crop.

We employed a relatively stringent method in declaring a QTL cluster as a meta-QTL: (1) each meta-QTL should be composed of at least two different QTL; and (2) these QTL should come from at least two independent studies.

Accordingly, from an initial set of 39 meta-QTL, only 30 were regarded to be 'true' meta-QTL across the 12 chromosomes of chile peppers, with confidence intervals between 0.55 cM (*MQTL5.2*) and 14.90 cM (*MQTL12.3*). These criteria were therefore relevant for a more accurate representation of the meta-QTL identified for the chile pepper QTLome. In other crop species, varying numbers of meta-QTL have been identified. Only 11 meta-QTL were detected for seedling stage salinity tolerance in rice⁶, whereas 60 meta-QTL were identified for *Fusarium* head blight resistance in wheat²⁵. In another meta-study of QTL in pea plants, 27 meta-QTL were resolved for seed protein content and yield-related traits²⁶. Such differences could be a consequence of the genome size, reliability of the consensus map used for meta-analysis, number of QTL regions identified, as well as the intrinsic properties of the reported QTL, such as phenotypic variation explained and LOD scores. As precision in QTL positions are dependent on population size and trait variation explained⁷, re-calculating positions based on the type of mapping populations used for analyses could facilitate a better representation of the genetic positions for each of the QTL evaluated.

One of the objectives of a meta-QTL study is to delimit the region of a QTL using information from multiple linkage-mapping studies. Chromosome P5 represents a major chromosome for *P. capsici* resistance in chile peppers, with large-effect QTL reported in previous studies²⁷⁻²⁹. In the current study, we reported two meta-QTL regions in chromosome P5, namely *MQTL5.1*, and *MQTL5.2* delimited to < 1.0 cM confidence interval, i.e. 0.79 cM and 0.55 cM, respectively, comprised of QTL mapped for *P. capsici* fruit and root rot resistance. These corresponded to the genomic regions having the most refined genetic distance among all the meta-QTL identified in the present work. Similarly, in peanut (*Arachis hypogea*), a recent meta-analysis of QTL for late leaf spot resistance delimited a region to 0.38 cM and 0.70 cM³⁰, whereas in wheat, genomic regions associated with *Fusarium* head blight resistance and root-related traits were narrowed to 0.82 cM²⁵ and 0.50 cM intervals³¹, respectively. *Capsicum* spp. *MQTL5.1* and *MQTL5.2* consisted largely of major effect QTL, with percent variation explained ranging between 10 and 52.7% (*MQTL5.1*) and 8.9 and 67.7% (*MQTL5.2*) identified from five independent QTL mapping studies, with *N* ranging between 63 and 297 individuals. Notably, these constituent QTL also represent those with the highest phenotypic trait variation explained in the *Capsicum* QTLome evaluated; this could be a reason for a more refined meta-QTL region for disease resistance. Furthermore, *MQTL5.2* consisted of 24 QTL, which was next to *MQTL1.3* identified to having the highest number of individual QTL. Among the criteria for choosing a meta-QTL for selection are (1) a small confidence interval, (2) a high number individual QTL comprising the meta-QTL, and (3) a high trait variation explained of initial QTL⁷. Considering these factors, *MQTL5.1* and *MQTL5.2* could serve as potential targets for marker-assisted breeding and selection for improved *P. capsici* resistance in chile peppers. The identification of meta-QTL linked DNA-based markers will help prioritize different QTL for introgression through MAS in plant breeding programs^{3,31}. In this regard, information from the flanking sequences for *MQTL5.1* and *MQTL5.2* identified in chromosome P5 will be utilized for the development of Kompetitive allele specific (KASP®)³² SNP assays for marker-assisted breeding. These KASP assays will be further validated using a recombinant inbred line population previously developed at New Mexico State University³³, and on a diverse population of New Mexican chile peppers to screen for resistance to different races of *P. capsici*.

The power of meta-QTL analysis lies in determining genomic regions that are most frequently involved in phenotypic variation and in delimiting the QTL intervals, therefore enabling candidate gene identification for positional cloning³¹. Also, meta-QTL are potentially genomic regions that are highly rich in genes²⁵ thereby facilitating pyramiding or stacking of important loci. Putative blight resistant protein homologues and leucine rich repeat (LRR) receptor-like serine/threonine protein kinases have been previously identified as candidate genes for *P. capsici* resistance in chile peppers^{34,35}. In the current study, candidate gene analysis using sequences for markers flanking *MQTL5.1* and *MQTL 5.2* in chromosome P5 identified genes with diverse biological functions related to

disease resistance, including DNA repair, DNA strand renaturation, ion transport, and several epigenetic mechanisms such as DNA, RNA, and histone methylation/demethylation, indicating the possible function of epigenetics in controlling gene expression for disease resistance in chile peppers. Epigenetics and its relationship with conferring disease resistance has been well recognized in other crops such as *Arabidopsis*³⁶, rice³⁷, and maize³⁸. The denser cytosine methylation profile of the *Capsicum* genome relative to that of the tomato and potato genomes³⁹ could indicate the relevance of epigenetics for the expression of different genes in peppers. Accordingly, identifying epialleles near the meta-QTL regions in chromosome P5 could be important in breeding towards improving resistance to *P. capsici* in chile peppers. Nevertheless, while the candidate genes identified here represent promising targets for future breeding, it is not known whether they are the true functional regulators of the detected meta-QTL, as many other genes could be present within the meta-QTL regions⁵. It would therefore be relevant to perform functional validation of the effects of these candidate genes using different chile pepper germplasm. Overall, meta-QTL analysis confirmed the relevance of chromosome P5 as a major genomic region harboring QTL and different candidate genes for *P. capsici* resistance in *Capsicum*.

Chile peppers are unique among the members of family Solanaceae due to their ability to produce capsaicinoids which render distinct flavors and heat profiles. Previously, gene mapping, allele sequence data, and expression profile analyses collectively identified the pungency gene *Pun1* in chromosome P2 responsible for the biosynthesis of capsaicinoids in chile peppers⁴⁰. More recently, the quantitative nature of capsaicinoid levels in chile peppers have also been demonstrated through linkage mapping which identified heat level-related QTL on chromosomes P1, P6, and P10⁴¹⁻⁴³. Several meta-QTL (e.g. *MQTL2.1*, *MQTL2.2*) identified in the present study have pungency-related QTL co-localized with QTL for agronomic and disease resistance traits such as fruit wall (pericarp) thickness, biomass, number of leaves on primary axis, and anthracnose resistance, among others. This demonstrated potential pleiotropy and/or effects of close linkage between the underlying QTL⁴⁴. Such colocalization of QTL related with diverse sets of traits for the identified meta-QTL across different chromosomes of chile pepper indicates the possibility of multi-trait improvement using genomic information from multiple linkage mapping studies.

In the current study, meta-QTL analysis was used to dissect the genetic architecture of diverse traits in *Capsicum*. Genomic regions for disease resistance to *P. capsici* in chile peppers were refined, and the role of chromosome P5 as a major genomic region harboring disease resistance QTL has been confirmed. Two meta-QTL, *MQTL5.1* and *MQTL5.2*, in chromosome P5 have been delimited to < 1.0 cM intervals. Analysis of candidate genes for these meta-QTL revealed biological functions related to DNA repair, response to bacterial and fungal infection, and DNA, RNA, and histone methylation, which demonstrate the potential role of epigenetics on resistance to *P. capsici*. The colocalization of several unrelated QTL on similar chromosomal regions demonstrates potential pleiotropic effects and the effect of linkage due to location. SNP assays will be developed for these meta-QTL and will be used for MAS for resistance to pepper blight. This study by far is the largest reported meta-analysis of different traits and the first known study of the *Capsicum* QTLome. The information presented here could serve as a valuable resource for the genomic improvement of diverse sets of traits in chile peppers.

Materials And Methods

Collection and characterization of QTL for different traits in chile peppers. A comprehensive bibliographic review of 29 QTL mapping studies published between 2010 and 2020 (Table 1) was first conducted to generate a QTL database for *Capsicum*. The evaluated traits in chile peppers were divided into five major categories, namely (1) adaptation; (2) agronomic, quality, and yield; (3) disease resistance; (4) heat profile (pungency); and (5) biochemical

and physiological traits. Each QTL was characterized according to the number of lines used for QTL mapping, type of mapping population (e.g. F₂, RIL, BC, DH), QTL name, trait, chromosome and linkage group designations, LOD, phenotypic variation explained (R^2), and chromosome positions (in cM). QTL with LOD scores of < 2.0 and with R^2 values not reported in the original study were excluded in further analysis. Genomewide association studies were not included in the literature review.

Projection to a consensus map and analysis of meta-QTL. Three different genetic maps from four cultivated species of *Capsicum* were used to develop a consensus map for the analysis of meta-QTL in chile peppers. These included an interspecific SNP genetic map derived from the hybridization between *C. annuum* and *C. frutescens*⁴⁵; a SLAF-based SNP array resulting from genotyping an F₂ population of a cross between *C. chinense* and *C. annuum*⁴⁶; and an intraspecific SNP linkage map derived from the hybridization between two *C. baccatum* varieties⁴⁷. Consensus maps for each of the 12 chromosomes of chile pepper were created using the 'LPmerge' package⁴⁸ in R⁴⁹. This function implements a linear programming (LP) algorithm to effectively reduce the mean absolute error in combining different genetic or linkage maps.

For QTL projection to the consensus map, the confidence interval (CI) for each QTL were calculated according to Darvasi and Soller⁵⁰ and Guo et al.⁵¹ based on the type of population used for linkage mapping using the following formula: Backcross (BC) and F₂: $CI = 530/NR^2$; Double haploid (DH): $CI = 287/NR^2$; and RIL: $CI = 163/NR^2$, where N is the number of lines used in QTL mapping and R^2 is the phenotypic variation explained by the QTL. QTL were projected to the *Capsicum* consensus map using the QTL projection function in Biomercator v.4.2.3⁵².

Meta-analysis was performed using Biomercator v.4.2.3 which implements a maximum likelihood algorithm developed by Goffinet and Gerber⁴. In this method, an N number of QTL linked to the same trait or set of related traits detected in independent experiments and located in the same genomic regions is determined to be consistent with five different QTL models, namely, 1, 2, 3, 4, and 5- N QTL models. An Akaike information criterion was used to determine the best model in identifying a meta-QTL, or "real" QTL which best represent the original QTL²⁵. Additionally, only those genomic regions where QTL from at least two different genetic mapping studies co-localized to form a meta-QTL were regarded as a meta-QTL⁵. Therefore, no meta-QTL identified in this study consist of only a single QTL (i.e. a singleton). Designations for each identified meta-QTL were based on the corresponding chromosome number and their position relative to the short arm of the chromosome (e.g. "MQTL1.1", "MQTL2.3"). Pepper chromosomes with the identified meta-QTL and their flanking markers were redrawn using the 'LinkageMapView' package⁵³ in R.

Candidate gene identification for the meta-QTL. Identification of candidate genes was conducted using the sequences of the flanking markers for the identified meta-QTL that has a < 1.0 cM confidence interval. Flanking sequences were BLASTn searched in EnsemblPlants (<https://plants.ensembl.org/index.html>)⁵⁴ against the genome of *C. annuum* and annotated genes and their biological functions were listed. Additionally, orthologous genes from Solanaceous plant species, including tomato (*S. lycopersicum*), potato (*S. tuberosum*), and wild tobacco (*N. attenuata*) were identified.

Declarations

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Author Contributions

D.N.L. conceived the research and wrote the first draft of the manuscript, L.W. and D.N.L. performed the meta-QTL and candidate gene analysis, A.A. edited the manuscript. All authors reviewed the manuscript.

Conflict of interest: The authors declare no competing interests.

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Figures

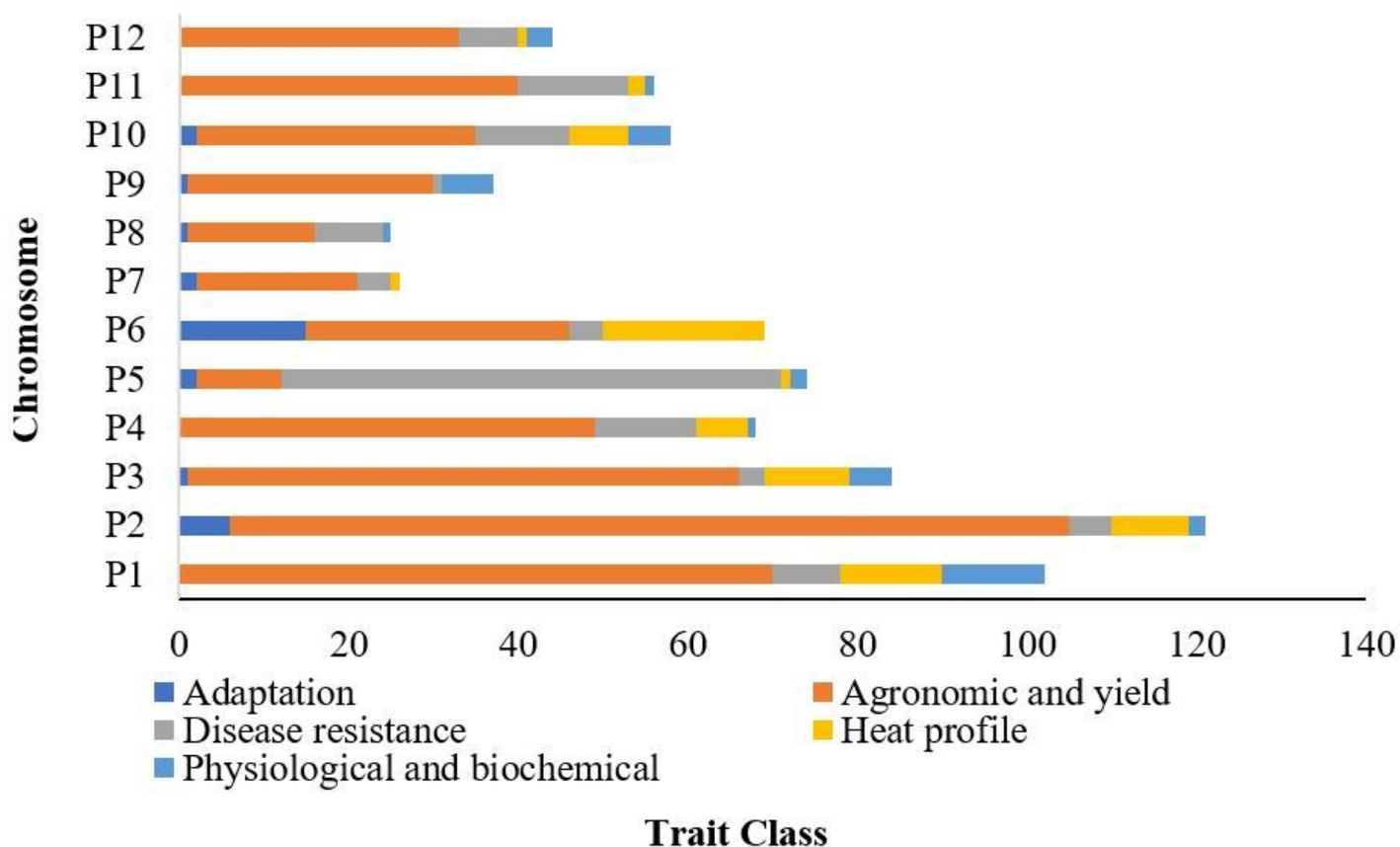


Figure 1

Distribution of 766 QTL surveyed in 29 linkage mapping studies across five different trait classes in chile peppers (*Capsicum* spp.).

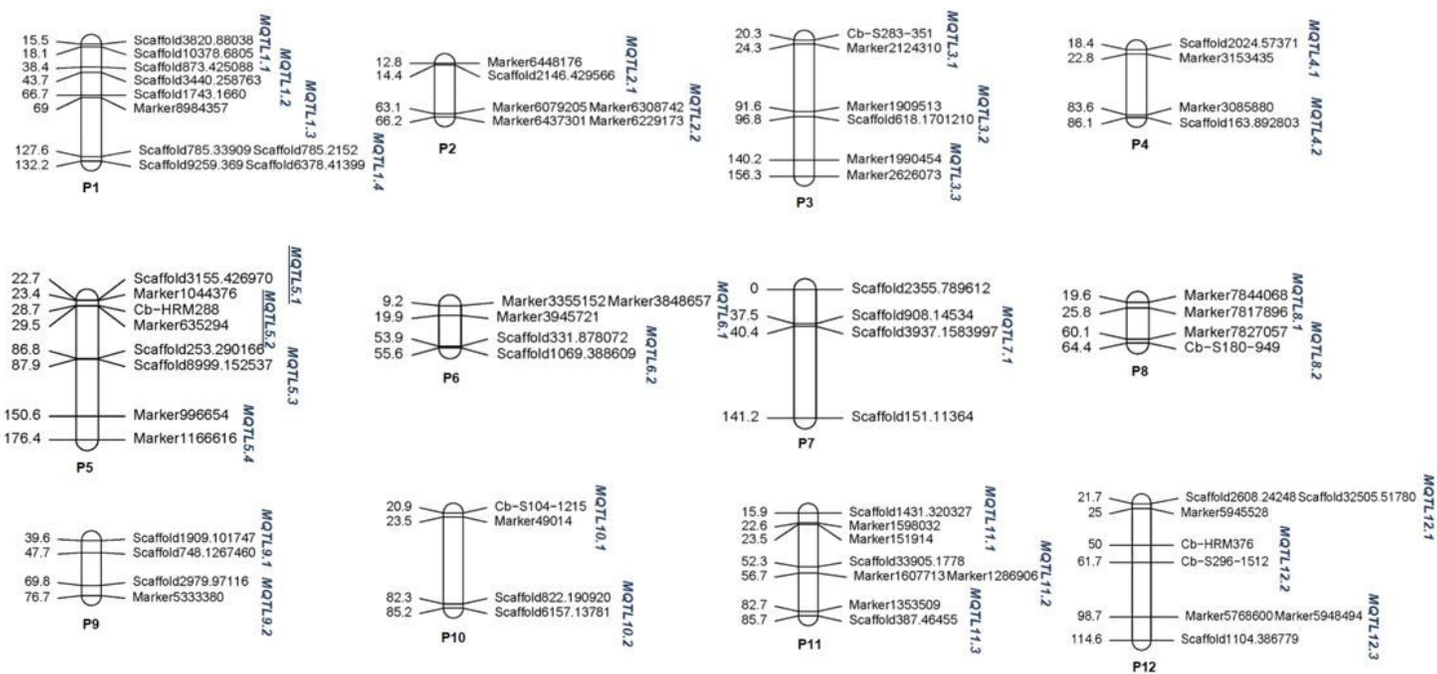


Figure 2

Meta-QTL and their flanking markers identified using 29 linkage mapping studies for diverse sets of traits for *Capsicum* spp. QTLome. Meta-QTL with underline are those with confidence intervals delimited to < 1.0 cM.

Supplementary Files

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