478

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Quantitative trait locus mapping associated with earliness and fruit weight in tomato

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Introduction

Earliness is a characteristic that has grown in importance over the course of the history of plant breeding. This trait has been managed with great success due to its intermediate-high heritability (Banerjee, 1989; Kemble and Gardner, 1992), which has led to plant breeders obtaining earlier crops with high efficiency (Foolad, 2007).

During the domestication process of tomato (Solanum lycopersicum L.), the selection of large fruit has brought genetic changes in certain components of earliness. For example, genotypes with large fruit often present a longer flowering period than the wild ancestor species S. pimpinellifolium L. (Grandillo and Tanksley, 1996). Several correlations between flowering time and other traits have been reported by classical and molecular studies (Sumugat et al., 2010; Grandillo and Tanksley, 1996; Doganlar et al., 2002; Georgelis et al., 2004; Monamodi et al., 2013). Such studies showed that the number of leaves produced before the floral transition plays an important role during the time required by tomato plants to develop their first flower (Samach and Lotan, 2007). This effect has been demonstrated in mutants such as FALSIFLORA (fa) and SINGLE FLOWER TRUSS (sft) where a mutation in one of their floral pathway integrators caused an increase in the flowering period due to an increased number of leaves formed in the initial simpodial segment (Molinero-Rosales et al., 1999; Molinero-Rosales et al., 2004).

The co-localization of QTLs associated with days to flowering of the first truss and fruit size on chromosomes 1, 4 and 6 has revealed a number of causes of the positive association identified in several types of map-

ABSTRACT: The flowering time is regarded as an important factor that affects yield in various crops. In order to understand how the molecular basis controlling main components of earliness in tomato (Solanum lycopersicum L.), and to deduce whether the correlation between fruit weight, days to flowering and seed weight, is caused by pleiotropic effects or genetic linkage, a QTLs analysis was carried out using an F2 interspecific population derived from the cross of S. lycopersicum and S. pimpinellifolium. The analysis revealed that most of the components related to earliness were independent due to the absence of phenotypic correlation and lack of co-localization of their QTLs. QTLs affecting the flowering time showed considerable variation over time in values of explained phenotypic variation and average effects, which suggested dominance becomes more evident over time. The path analysis showed that traits such as days to flowering, seed weight, and length of the first leaf had a significant effect on the expression of fruit weight, confirming that their correlations were due to linkage. This result was also confirmed in two genomic regions located on chromosomes 1 and 4, where despite showing high colocalization of QTLs associated to days to flowering, seed weight and fruit weight, the presence and absence of epistasis in $dfft1.1 \times dftt4.1$ and $fw1.1 \times fw4.1$, suggested that the linkage was the main cause of the co-localization.

Keywords: Solanum lycopersicum, QTLs, epistasis, days to flowering, path analysis

ping populations (e.g., F_2 , inbred backcross lines, near isogenic lines and recombinant inbred lines) (Grandillo and Tanksley, 1996; Doganlar et al., 2002; Monforte and Tanksley, 2000; Yates et al., 2004; Sumugat et al., 2010). However, it is not clear if the cause is due to pleiotropic effects or to gene linkage.

Recently in tomato, several studies have reported the influence of epistasis on fruit shape, locule number, seed size and fruit weight during the process of domestication (Lippman and Tanksley, 2001; van der Knaap et al., 2002; Khan et al., 2012). In few studies have the epistatic interactions between the components of earliness and fruit weight been examined, and it is equally unclear if tomato earliness is controlled by a single genetic mechanism or something more complex. Therefore, this work focused on the identification of QTLs affecting the main components of earliness and fruit weight. Furthermore, we were interested in identifying the epistatic interactions between these QTLs. Such information was useful for determining if the positive associations of fruit weight with the period of flowering and seed weight were due to pleiotropic effects or to gene linkage.

Materials and Methods

Mapping population

To identify the different types of gene action (additive, dominance and overdominance) expressed by the QTLs identified in this study, an F_2 mapping population was obtained from the interspecific cross between two F_5 inbred lines belonging to *S. lycopersicum* collection LOR82 (L) and *S. pimpinellifolium* collection 11904 (P). The F_1 hybrid was generated, and then it was selfed to produce the F_2 population. LOR82 is a local cultivar from the state of Puebla, Mexico. This line produces large fruits with characteristics similar to the Saladette type tomato and it is locally known as "Chino Criollo". *S. pimpinellifolium* is a wild relative of tomato and has a small rounded fruit. A number of 172 F_2 plants were evaluated along with the parents and the F_1 generation, under a randomized complete block design with four replications. The evaluation of the experiment was conducted under green-

house conditions during the 2012 spring-summer growing season in Texcoco (19°28'4.26" N, 98°53'42.18" W, 2250 m.a.s.l.), state of Mexico, Mexico.

Phenotypic characterization

Individual F2 plants were evaluated for eleven quantitative traits. Fruit weight (FW), in grams, was scored as the average weight of five fruits per plant; number of flowers per truss (NFT) was measured by averaging the number of flowers from the third and fourth trusses; days to emergence (DE), in days, were obtained as the amount of time each plant took to emerge from planting date; days to flowering of the first (DFFT), second (DFST) and third truss (DFTT), were determined as the number of days elapsed from the date of sowing until the opening of the first flower in the first, second and third truss. Days to ripening (DR) were obtained by counting the days elapsed from the date of sowing to production of the first ripe fruit. Seed weight (SW), in grams, was estimated as the average weight of 100 seeds harvested of each F2 plant. This score was used to estimate the weight of 1,000 seeds. Finally, the traits seedling height (SH), width and length of the first leaf (WFL and LFL, respectively) were measured at 25 days after planting; plant height was measured as the length of the stem while leaf length and leaf width as the horizontal and vertical length of the first true leaf, respectively.

Genotypic analysis

DNA extraction was performed according to the protocol established by Bernatzky and Tanksley (1986). A total of 61 molecular markers located on twelve chromosomes were scored for construction of the linkage map. These SSR markers were obtained from Tomato Mapping Resource Database (http://164.107.85.47:8003) and Sol Genomics Network (http://solgenomics.net/) based on their map positions. Each PCR reaction containing a final volume of 25 μ L was prepared with the following final concentrations: PCR Buffer (1X), MgCl₂ (1.5 mM), dNTP's (0.1 mM by nucleotide), F and R Primer (0.4 μ M), Taq Polymerase (0.1 u μ L⁻¹) and DNA (1.6 ng μ L⁻¹). Finally, the PCR products were run on acrylamide gels as described by Creste et al. (2001).

Data analysis

Means, normality test (Anderson-Darling's test), tests for homogeneity of variance (Bartlett's test) and Pearson's correlation, were calculated for each trait using R software version 3.0.1 (R Core Team, 2013). Significant differences between the generation means were estimated using one-way ANOVA followed by a Tukey test. Path analysis was carried out to estimate the direct and indirect effects of each earliness component on fruit weight according to the procedures described by McGiffen et al. (1994). This analysis is employed to elucidate how other variables mediate the relationship formed by two traits (McGiffen et al., 1994).

The linkage map was constructed using the Map-Disto software version 1.7.7 under a LOD of 3, and the Kosambi mapping function (Kosambi, 1944) was employed to convert the recombination frequencies into map units expressed in centimorgans (cM). The location of linkage groups was based on the genetic map published by Robbins et al. (2011). QTL analysis and phenotypic variance were performed using the mapping software Windows QTL Cartographer 2.5. Putative QTLs were detected employing composite interval analysis (Jansen and Stam, 1994; Zeng, 1994) where the significance level to claim a QTL was determined using 1,000 permutations. For genomic scan a window size of 10 cM and walk speed of 1 cM were used. The dominance degree of alleles for a QTL was estimated based on the ratio |d/a|, where d = (LP) - 1/2(LL + PP), a = (LL) - 1/2(LL + PP), LP =frequency of heterozygotes individuals, LL = frequency of S. lycopersicum homozygote individuals and PP = frequency of S. pimpinellifolium homozygote individuals, thus when |d/a| < 0.2 the trait was considered additive, $0.2 \le |d/a| < 0.8$ partially dominant, $0.8 \le |d/a| < 1.2$ completely dominant and $|d/a| \ge 1.2$ overdominant (Stuber et al., 1987). The average effects of the L and P alleles for each QTL were estimated according to: q[a + d(q-p)] for L and -q[a + d(-qp)] for P, where p and q = gene frequency of the L and P alleles, respectively (Falconer and Mackay, 1996).

Finally, epistatic interactions were identified by the following factorial design with two factors: $y = \mu + M_i + Mj + (MM)_{ij} + e_{ijk'}$ where $M_i = effect$ of marker i, $M_j = effect$ of marker j, $(MM)_{ij} = interaction$ between markers i and j and $e_{ijk} = residual error$. Thus, where $(MM)_{ij}$ was significant (p < 0.05), epistasis was claimed. Broad-sense heritability was calculated using the equation $H^2 = \sigma_g^2 / (\sigma_g^2 + \sigma_e^2)$ where σ_g^2 is the genetic variance and σ_e^2 is the environmental variance estimated with the ANOVA error term.

Results

Phenotypic distributions and correlations between quantitative traits

Several significant differences (p < 0.05) between parents (L and P), and generations F_1 and F_2 were observed in almost all the traits evaluated. Specifically, the parents differed statistically (p < 0.05) in nine of the eleven traits analyzed (Table 1). Furthermore, all variables showed a normal distribution, indicating that these traits were governed by complex inheritance mechanisms (Figure 1). However, traits such as FW, DE and DFFT showed a normal distribution when these were Hernández-Bautista et al.

Traitª	S. lycopersicum	S. pimpinellifolium	F ₁		Durshush	MODe		
	Mean	Mean	Mean	Mean \pm SE	Min	max	- P-value	INI2D _c
FW	128.1 a ^d	0.7 b	6.6 b	8.4 ± 0.44 b	1.0	40.0	* * *	11.4
NFT	5.7 c	25.7 a	9.5 cb	11.0 ± 0.23 b	5.0	23.0	* * *	4.7
DE	7.8 a	8.3 a	7.4 a	7.8 ± 0.10 a	5.0	15.0	ns	2.1
DFFT	77.1 a	60.6 b	65.7 b	66.1 ± 0.43 b	54.0	86.0	* * *	9.2
DFST	96.0 a	69.9 b	76.2 b	77.9 ± 0.35 b	67.0	91.0	* * *	9.4
DFTT	106.7 a	76.1 b	84.4 b	84.7 ± 0.46 b	72.0	103.0	* * *	9.4
DR	140.6 a	124.2 b	119.4 b	121.7 ± 0.40b	108.0	142.0	* * *	8.1
WS	4.0 a	1.3 c	2.7 b	2.5 ± 0.04 b	0.4	3.7	* * *	0.8
SH	4.5 a	2.7 b	4.1 a	2.8 ± 0.04 b	1.5	4.3	* * *	0.8
WFL	0.79 a	0.72 ba	0.87 ba	0.66 ± 0.01 b	0.20	0.90	* * *	0.18
LFL	3.5 a	2.0 bc	3.0 ba	2.5 ± 0.03 c	0.6	3.3	* * *	0.6

Table 1 – Mean phenotypic values, standard deviations for parents, F₁ and F₂ population.

^aFW = fruit weight (g); NFT = Number of flowers per truss; DE = Days to emergence; DFFT = Days to flowering in first truss; DFST = Days to flowering in second truss; DFTT = Days to flowering in third truss; DR = Days to ripening; SW = Weight of 1000 seed (g); SH = Seedling height (mm); WFL = Width of the first leaf (mm); LFL = Length of the first leaf (mm); ^bP-value = Threshold set to < 0.05, ns = non-significant, *** significant at p < 0.001; ^cMSD = Minimum Significant Difference; ^dTukey test; genotypes sharing same letter are statistically equal under (p < 0.05).

adjusted with \log_{10} or square root transformations. The phenomenon of transgressive segregation was observed in most of the variables (NFT, DE, DFFT, DFST, DFTT, DR, SW, SH, WFL and LFL) (Table 1), where interestingly some plants expressed a shorter flowering period to that of *S. pimpinellifolium* (Table 1).

Several significant positive and negative correlations (p < 0.002) between traits were observed in the F₂ population (Figure 2). NFT showed a significant negative correlation with FW (-0.34), DFFT (-0.24), DFST (-0.29), DFTT (-0.40), SW (-0.35) and SH (-0.26). On the other hand, DE only had a negative correlation (p < 0.001)with WFL and LFL, whose values were less than 0.35. In addition, we found statistically significant correlations between DFFT and five other variables (FW, DFST, DFTT, SW and SH). Likewise, flowering traits such as DFST and DFTT variables showed significant correlations with FW, SW and SH, whose association values were below 0.5. Seed weight (SW) presented six correlations with other traits, some of which resulted from the association with FW (0.43), NFT (-0.35), DFST (0.30), DFTT (0.34) and LFL (0.34). Finally, LFL also showed significant correlation with SH (0.34) and WFL (0.47).

Direct and indirect effects of the components of earliness on fruit weight

A path analysis was conducted to identify the direct and indirect effects of the components of earliness affecting fruit size. Due to the high correlation between DFFT, DFST and DFTT, we only used DFTT as the flowering variable affecting fruit weight. The analysis showed to DFTT, SW and LFL as the main components of earliness affecting to FW (p < 0.05), where their path coefficients for the direct effects were 0.31, 0.31 and 0.24, respectively (Figure 3).

Although we observed an association between FW with NFT and SH in the correlation analysis (p < 0.002), these characteristics showed no effect on FW in the path analysis.

QTL analysis

The map spanned 815.71 cM of the tomato genome with an average marker spacing of 13.37 cM. A total of 27 QTLs were identified in eight of the eleven traits evaluated. Out of the 27 QTLs identified, only two (*de8.1* and *dr7.1*) showed an opposite effect (Table 2 and Figure 4) and *dftt1.1* had the highest value (25 %) in the individual explained variance (\mathbb{R}^2). The trait that presented the highest value of total explained variance (TEV) was SW with 43 % while DE had the lowest value of 2 %. Additionally, seed weight was the trait that presented the highest number of QTLs identified by trait, where a set of seven QTLs governing this quantitative trait was detected on chromosomes 1 (*sw1.1* and *sw1.2*), 2 (*sw2.1*), 3 (*sw3.1* and *sw3.2*), 4 (*sw4.1*) and 7 (*sw7.1*).

Epistatic interactions

As the square root transformation reduced skewness in FW, we performed the analysis using transformed data for all traits; however, non-transformed data have been used in the epistasis schemes to improve the observation of the interactions. Based on the results of the twoway analysis of variance, four significant interactions between QTLs (p < 0.05) were identified for FW and DFTT variables (Figure 5). For FW, three significant interactions were detected: $fw1.1 \times fw3.1$, $fw3.1 \times fw1.2$ and $fw1.2 \times$ fw4.1. As regards the fw3.1 \times fw1.1 interaction, the two loci under a homozygous condition (LL) expressed a disproportionate increase in the expected phenotypic value. However, this increase was greater when fw1.1 was selected as the independent variable, since the phenotypic variation between homozygous genotypes (LLLL and PPLL) (Figure 5A) was greater (8.65 g) than that of the reciprocal scheme (Figure 5B). This result indicated that fw1.1 had a greater effect on increasing fruit weight when fw3.1 S. lycopersicum alleles had been previously fixed in the individuals. Figure 5C and Figure 5D show that the interaction $fw3.1 \times fw1.2$ had a similar effect on the two genotypes LLLL and PPLL compared to the effect gen-

481



Figure 1 – Frequency distribution histograms of non-normalized data for earliness-related traits and fruit weight in the interspecific F_2 population. P = S. pimpinellifolium mean, L = S. lycopersicum mean, $F_1 = F_1$ mean.

	NFT	DE	DFFT	DFST	DFTT	DR	SW	SH	WFL	LFL
FW	-0.34***	0.01 ns	0.28***	0.33***	0.42***	–0.02 ns	0.43***	0.24**	–0.05 ns	0.17*
	NFT	0.06 ns	-0.24**	-0.29***	-0.40***	0.04 ns	-0.35***	-0.26***	0.02 ns	-0.11 ns
		DE	0.11 ns	0.07 ns	0.04***	0.02 ns	–0.09 ns	-0.19*	-0.34***	-0.30***
			DFFT	0.50***	0.64***	0.16*	0.27***	0.26***	-0.15*	-0.19**
				DFST	0.79***	0.16*	0.30***	0.35***	-0.18*	–0.15 ns
					DFTT	0.16*	0.34***	0.41***	-0.22***	-0.08 ns
						DR	0.01 ns	–0.03 ns	0.02 ns	-0.11 ns
							SW	0.28***	0.03 ns	0.04*
								SH	0.08 ns	0.34***
									WFL	0.47***
										LFL

Figure 2 – Pearson's coefficient of correlations between trait pairs in the F_2 population. FW = fruit weight; NFT = Number of flowers per truss; DE = Days to emergence; DFFT = Days to flowering in first truss; DFST = Days to flowering in second truss; DFTT = Days to flowering in third truss; DR = Days to ripening; SW = Weight of 1000 seed; SH = Seedling height; WFL = Width of the first leaf; LFL = Length of the first leaf; ns = non-significant; *, **, *** = significant at p < 0.05, 0.01, 0.001, respectively.

Table 2 – Significant QTLs associated with earliness traits and fruit weight detected on F_2 population.

	-	-				-	2				
Traita	QTL	Nearest Marker	Chr⁵	Position (cM)	LOD score	R ² (%) ^c	α^d_L	α^{d}_{P}	d∕a e	TEV ^f	H ^{2 g}
FW	fw1.1	TOM202	1	0.01	3.01	9.16	1.16	-1.37	0.01		
	fw1.2	SSR42	1	96.95	4.10	8.39	1.25	-1.24	0.47		
	fw3.1	SSR111	3	0.01	4.38	11.05	1.26	-1.38	0.02		
	fw4.1	SSR306	4	60.12	2.54	5.76	1.12	-0.98	0.35	29.31	0.57
NFT	nft1.1	TOM202	1	2.01	2.52	4.01	-0.45	0.54	0.81		
	nft3.1	SSR320	3	57.00	2.33	5.57	-0.49	0.49	0.76		
	nft4.1	TOM184	4	137.93	3.11	5.30	-0.50	0.47	0.30		
	nft5.1	SSR115	5	14.30	2.95	5.14	-0.74	0.64	0.05	21.44	0.43
DE	de8.1	SSR327	8	37.80	2.01	2.00	-0.23	0.25	0.45	2.01	0.09
DFFT	dfft1.1	TOM202	1	0.01	4.56	5.34	1.02	-1.21	1.21		
	dfft4.1	SSR306	4	75.54	3.55	8.25	1.27	-1.12	0.09		
	dfft7.1	SSR45	7	116.50	3.12	7.75	0.99	-1.02	0.66	17.71	0.41
DFST	dfst1.1	TOM202	1	0.01	8.32	11.22	1.65	-1.95	0.78		
	dfst4.1	SSR306	4	77.54	3.48	7.69	1.17	-1.03	0.11		
	dfst7.1	SSR45	7	114.63	2.86	7.02	0.95	-0.98	0.66	23.38	0.50
DFTT	dftt1.1	TOM202	1	0.01	15.76	25.20	2.25	-2.66	0.61		
	dftt4.1	SSR306	4	83.54	3.50	5.60	1.07	-0.94	0.12		
	dftt7.1	SSR45	7	112.70	2.35	5.74	0.90	-0.93	0.64	34.43	0.65
DR	dr7.1	SSR286	7	10.01	2.96	3.94	-0.70	0.78	1.07		
	dr10.1	SSR248	10	32.00	2.97	7.63	1.12	-1.09	0.79	11.69	0.27
SW	sw1.1	TOM202	1	1.01	2.04	4.16	0.08	-0.09	0.08		
	sw1.2	SSR42	1	101.95	2.56	6.85	0.09	-0.09	0.38		
	sw2.1	TOM188	2	0.01	3.29	8.63	0.12	-0.10	0.34		
	sw3.1	SSR111	3	0.01	6.31	17.27	0.13	-0.15	0.30		
	sw3.2	SSR320	3	57.00	2.18	9.27	0.09	-0.09	0.46		
	sw4.1	SSR306	4	75.54	3.03	8.27	0.10	-0.09	0.35		
	sw7.1	SSR45	7	113.49	3.60	6.56	0.03	-0.04	0.43	42.77	0.87

^eFW = fruit weight (g); NFT = Number of flowers per truss; DE = Days to emergence; DFFT = Days to flowering in first truss; DFST = Days to flowering in third truss; DR = Days to ripening; SW = Weight of 1000 seed (g); ^bChromosome on which QTL was detected; ^cPercentage of phenotypic variation explained by the QTL; ^dAverage allelic effect provided by *S. lycopersicum* and *S. pimpinellifolium*, respectively; ^eDegree of dominance expressed by the QTL; ^fTotal explained variance by all QTL in each trait; ^gBroad-sense heritability.

Hernández-Bautista et al.

				Indirect	effects				
Direct effects		NFT	DR	DFTT	DR	SW	SH	WFL	LFL
	NFT	-0.10 ns	0.00	-0.12	0.00	-0.11	0.02	0.00	-0.03
	DE	-0.01	0.07 ns	0.01	0.00	-0.03	0.01	0.03	-0.07
	DFTT	0.04	0.00	0.31**	-0.01	0.10	-0.03	0.02	-0.02
	DR	0.00	0.00	0.05	-0.04 ns	0.00	0.00	0.00	-0.03
	SW	0.03	-0.01	0.10	0.00	0.31***	-0.02	0.00	0.01
	SH	0.03	-0.01	0.13	0.00	0.09	-0.06 ns	-0.01	0.08
	WFL	0.00	-0.02	-0.07	0.00	0.01	0.00	-0.07 ns	0.12
	LFL	0.01	-0.02	-0.02	0.00	0.01	-0.02	-0.04	-0.24**
	Correlation	-0.34***	0.01 ns	0.42	-0.02 ns	0.43***	0.24**	–0.05 ns	0.17*

Figure 3 – Direct and indirect effects of the earliness components over fruit weight identified by the path analysis. NFT = Number of flowers per truss; DE = Days to emergence; DFTT = Days to flowering in third truss; DR = Days to ripening; SW = Weight of 1000 seed; SH = Seedling height; WFL = Width of the first leaf; LFL = Length of the first leaf. The direct effects are indicated in bold type. ns = non-significant; * = significant at p < 0.05.



Figure 4 – Heatmap of QTLs identified for earliness-related traits and fruit weight. Only chromosomes with significant QTLs are displayed. SW = Weight of 1000 seed; NFT = Number of flowers per truss; FW = fruit weight; DR = Days to ripening; DFTT = Days to flowering in third truss; DFST = Days to flowering in second truss; DFFT = Days to flowering in first truss; DE = Days to emergence.

erated for $fw3.1 \times fw1.1$. However, the effect shown by fw1.2 was higher considering a background already fixed for LL alleles at fw3.1. In $fw1.2 \times fw4.1$, the greatest effect was caused by fw4.1, indicating that the fw4.1 *S. lycopersicum* alleles were subsequently selected over those of fw1.2 (Figure 5E and Figure 5F).

Finally, for DFTT only one partially dominant \times additive interaction was observed between *dftt1.1* \times *dftt4.1*. In this interaction, a disproportionate increase is observed when both loci are homozygous for LL alleles, which evidenced the phenomenon of epistasis (Figure 5G and Figure 5H).

Discussion

Different genetic pathways controlling earliness

During the study, a considerable phenotypic variation was found between parents and the F_2 population. On the other hand, the absence of correlation between DE and DR indicated that the first emerging plants were

not the first to produce the first ripe fruit. This same trend in reproductive development was observed during the period from DFTT to DR, where the plants which showed an earlier anthesis did not form their first fruit as quickly. Similar results were obtained by Burdick (1954), who evaluated eight tomato inbred-lines and 28 F₁ genotypes obtained under a diallel cross and, found that some F, hybrids were later than S. pimpinellifolium during the period between days to emergence and days to flowering to first truss, but were earlier in the last stage (days to first truss-days to maturity). In contrast, the close relationship between DFFT and WS caused by the co-localization of their QTLs indicated that plants derived from seeds with more weight expressed a later floral transition than those emerging from seeds with less weight. Similar associations were also obtained between DFST and DFTT with WS, which suggested that seed weight showed a positive effect on the development time of sympodial units formed after the floral transiHernández-Bautista et al.



Figure 5 – Interaction plots for fruit weight (FW) (A - F) and days to flowering to third truss (DFTT) (G and H). The reciprocal plots are shown on the right column.

tion due to gene linkage. In this regard, similar effects caused by the seed size and weight in tomato seedlings were reported by Khan et al. (2012), who concluded that effects on seedling growth were due to the genetic variation in the amount of food reserve (starch) and maternal effects. In addition to seed weight, various factors such as the number of leaves produced before the first inflo-

rescence, length of floral development, light intensity and temperature have been reported to affect the floral transition (Dieleman and Heuvelink, 1992; Samach and Lotan, 2007; Sumugat et al., 2010). This indicates that earliness in tomato estimated through the various trusses is affected by both genetic and non-genetic factors, which makes this trait even more complex to study.

To date, several mapping studies have been conducted in characteristics related to earliness of tomato (Grandillo and Tanksley, 1996; Doganlar et al., 2002; Cagas et al., 2007; Sumugat et al., 2010). However, there is no path analysis for earliness which can identify the progress of the effects of QTLs controlling earliness in each component. In this study, analyzing various components of earliness, a total of 27 QTLs were identified, where most of these QTLs were identified in regions where previously, several studies had reported QTLs associated to earliness (Grandillo and Tanksley 1996; Doganlar et al., 2002; Cagas et al., 2007; Jimenez-Gomez et al., 2007; Sumugat et al., 2010). In general, most QTLs identified were unique to each stage of earliness, except for dftt1.1, dftt4.1 and dftt7.1, which were consistent in the three stages of flowering evaluated (DFFT, DFST, DFTT). These results were similar to those reported by Grandillo and Tanksley (1996) and Doganlar et al. (2002), who observed no co-localization between QTLs for days to flowering and days to fruit ripening. Finally, the lack of correlation and localization between QTLs associated with DE and DR, as well as with flowering variables (DFFT, DFST and DFTT) suggests that earliness is controlled by different genetic mechanisms.

Genetic variation in truss production over time is affected by dominance

In spite of the strong correlation, same number and co-localization of the QTLs identified for DFFT, DFST and DFTT, differences in the phenotypic variation of their OTLs were observed, as well as changes in the average effect of their alleles. The QTL located on chromosome 1 (dftt1.1) showed an increase in its phenotypic variance in each sympodial segment, while dftt4.1 and dftt7.1 exhibited a decrease per truss. These changes in allelic effects were attributed to the increase (in *dftt1.1*) and decrease (in *dftt4.1* and *dftt7.1*) of genotypic range established by the two homozygous genotypes at each stage, where the dominance of S. pimpinellifolium alleles was the main factor influencing this variation of the genotypic range. These findings support the idea proposed by Burdick (1954), who found that the dominance of S. pimpinellifolium alleles is more evident over time. On the other hand, the major effect and dominance expressed by dftt1.1 supports the hypothesis suggested by Buckler et al. (2009), who proposed that the flowering time in cross-pollinated species is controlled by genes with small additive effects, while in autogamous species, such as tomato, it is affected by genes with large effects and dominance action.

The QTL named here as dftt1.1 has been the most frequently mapped in most mapping populations, indicating that due to its major effect on the floral transition, it can be detected under different environments. This observation supports the general conclusion arrived at by Tanksley (1993), who concluded that QTLs showing major effects have a null genotype \times environment interaction. However, our study is the first reporting a major effect of dftt1.1 on flowering time (> 2 days), which could be identified with such magnitude due to the contrast shown by progenitors conforming the mapping population. Furthermore, we observed that despite the large physical distance between the marker TOM202 and PHYB1 gene, the *dftt1.1* effect could be caused by PHYB1, due to a low recombination reported in this region (Gonzalo and van der Knaap, 2008); however, additional NILs populations are required to demonstrate this idea.

Gene linkage: leading cause of association fruit size-flowering period and fruit size-seed size

During the domestication process of tomato, selection of plants with larger fruit has indirectly caused changes in the flowering period, number of flowers per truss and seed weight. Genetic linkage and pleiotropic effects have been proposed as the main causes of the indirect effects on some quality traits (Paterson et al., 1991). However, in a number of chromosomal regions located in the final part of chromosome 1, it has not been possible to discern whether the effect shown for several characteristics such as fruit weight and external color of the fruit is caused by pleiotropic effects or gene linkage (Monforte and Tanksley, 2000).

In this research we identified three chromosomal regions on chromosomes 1, 3 and 4, which presented a co-localization of QTLs for days to flowering, seed weight and fruit weight (Figure 4). In the two regions located on chromosomes 1 and 4, it was observed that $dftt1.1 \times dftt4.1$ exhibited epistasis, while epistasis could not be observed for $sw1.1 \times sw4.1$ and $fw1.1 \times fw4.1$. Similar results of a non-constant epistasis between QTLs co-locating in two same regions was also observed between the chromosome 1 and 3, where the QTLs controlling fruit weight ($fw1.1 \times fw3.1$ and fw1.2 and fw3.1) expressed a highly significant epistasis, a characteristic not shown in both cases for $sw1.1 \times sw3.1$ and $sw1.2 \times$ sw3.1. These results suggest that co-localization of QTLs in these chromosomal regions is caused by gene linkage rather than pleiotropic effects, given that if the phenomenon of pleiotropy was the main reason, a constant epistasis between epistatic QTLs controlling various traits would be constant throughout the traits. Similar results, though between the content of soluble solids and weight of fruits, have been observed in F₂ populations of sub NILs derived from the cross between S. lycopersicum with S. hirsutum and S. peruvianum, where Yates et al. (2004) found that high co-localization of QTLs controlling fruit weight and soluble solids content in the region of chromosome 4 was caused by strong genetic linkage. Additionally Monforte and Tanksley (2000) concluded that high co-localization of QTLs controlling different fruit variables in the terminal part of chromosome 1 was not caused by pleiotropic effects but by gene linkage. Therefore, the linkage found in the initial region of chromosome 1 may be the main cause of positive associations found between the flowering period-fruit weight and seed weight-fruit weight as reported in this study.

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