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QTL analysis of early stage heterosis for biomass in Arabidopsis

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Abstract The main objective of this study was to identify genomic regions involved in biomass heterosis using QTL, generation means, and mode-of-inheritance classification analyses. In a modified North Carolina Design III we backcrossed 429 recombinant inbred line and 140 introgression line populations to the two parental accessions, C24 and Col-0, whose F_1 hybrid exhibited 44% heterosis for biomass. Mid-parent heterosis in the RILs ranged from –31 to 99% for dry weight and from –58 to 143% for leaf area. We detected ten genomic positions involved in biomass heterosis at an early developmental stage, individually explaining between 2.4 and 15.7% of the phenotypic

variation. While overdominant gene action was prevalent in heterotic QTL, our results suggest that a combination of dominance, overdominance and epistasis is involved in biomass heterosis in this Arabidopsis cross.

Introduction

The term ‘heterosis’ describes the superiority of F_1 hybrids manifested as increased size, fertility, or yield compared with the corresponding parental homozygous lines (Shull 1948). The phenomenon of heterosis has been exploited extensively in plant breeding (Birchler et al. 2003), however, the underlying genetic and molecular mechanisms are still largely unknown. Early studies put forward three genetic models as explanation for the extreme hybrid phenotype. The dominance hypothesis (Bruce 1910; Davenport 1908) attributes heterosis to the joint action of favorable

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dominant alleles at multiple loci from both parents combined in the hybrid. The overdominance hypothesis (Crow 1948; Hull 1945) postulates the existence of loci at which the heterozygous state is superior to either homozygote, i.e. a positive interaction between alleles at a heterozygous locus. Pseudo-overdominance (Crow 1952), in contrast, refers to the situation of tightly linked genes with favorable dominant alleles linked in repulsion. The epistasis hypothesis (Powers 1944; Williams 1959) explains heterosis by the interaction of favorable alleles at different loci contributed by the two parents, which themselves may show additive, dominant, or overdominant action.

One of the most promising approaches to unravel the genetic basis for heterosis at the molecular level emerged through the availability of molecular markers. They provided the means to perform quantitative trait locus (QTL) analyses, a powerful approach to map and subsequently identify genes involved in complex traits. In recent studies evidence was provided for the occurrence of dominance (Abdelkhalik et al. 2005; Xiao et al. 1995) or overdominance (Li et al. 2001; Semel et al. 2006) at multiple loci, and several reports highlighted the importance of epistasis (Kusterer et al. 2007a; Luo et al. 2001; Mei et al. 2005; Melchinger et al. 2007a; Monforte and Tanksley 2000; Yu et al. 2005). These diverse results indicate that heterosis may be caused by combinations of these mutually non-exclusive mechanisms.

Heterosis has been shown to be a widespread occurring phenomenon in *Arabidopsis thaliana* and has been described for such varying traits as stem length (Griffing and Langridge 1963), photosynthetic efficiency (Sharma et al. 1979), seedling viability (Mitchell-Olds 1995), seed number (Alonso-Blanco et al. 1999), seed size (Stokes et al. 2007), phosphate efficiency (Narang and Altmann 2001), cold tolerance (Korn et al. 2008; Rohde et al. 2004), rosette diameter and plant height (Barth et al. 2003; Syed and Chen 2004), and biomass (Meyer et al. 2004). The occurrence of heterosis in the model plant *Arabidopsis* opens the possibility to combine QTL analyses and the rapid identification of the causal DNA variation with transcript and metabolite profiling to discover physiological and molecular processes involved in heterosis. This process should establish a precedence that may guide the analysis of heterosis in crops.

We study heterosis in *A. thaliana* with the aim to identify genomic sequences that contribute to heterosis. The first step toward this end was a forward genetics approach of defining genomic regions influencing heterosis for biomass with the use of QTL mapping experiments in segregating populations developed from two divergent accessions, C24 and Col-0, which in previous studies exhibited transgressive segregation and heterosis for biomass (Meyer et al. 2004). The main objective of this study

was to identify QTL for biomass heterosis and to characterize the genetic mode of action of heterotic QTL using the approach proposed by Melchinger et al. (2007b).

Materials and methods

Plant materials

Analyses were performed on homozygous mapping populations derived from the two *A. thaliana* accessions C24 and Col-0 (P_1 and P_2). The recombinant inbred line (RIL) population was derived from reciprocal crosses C24 \times Col-0 (201 lines) and Col-0 \times C24 (228 lines) as described in Törjék et al. (2006). Two large reciprocal sets of introgression lines (ILs) were created through backcrossing and selfing using marker assisted selection (Törjék et al. 2008). In both IL sets (78 ILs with Col-0 background and 62 ILs with C24 background) the donor segments cover almost the entire genome with an average substitution size of 18.3 cM.

To allow analyses of heterotic effects, testcrosses (TC) with parents Col-0 and C24 were produced for 429 RILs (422 RIL-TC_{C24} and 416 RIL-TC_{Col}) and 42 ILs (20 IL-TC_{C24} and 22 IL-TC_{Col}).

Plant cultivation

Plants were grown in 1:1 mixture of GS 90 soil and vermiculite (Gebrüder Patzer, Sinntal-Jossa, Germany) in 96-well-trays under a long-day regime (16 h fluorescent light ($120 \mu\text{mol m}^{-2} \text{s}^{-1}$) at 20°C and 60% relative humidity/8 h dark at 18°C and 75% relative humidity). Six plants of the same line were grown per well. To avoid position effects, trays were rotated around the growth chamber every 2 days. In the first experiment, all RILs and RIL-TCs were cultivated together with parents C24 and Col-0 (P_1 and P_2) and their reciprocal F_1 -hybrids C24 \times Col-0 and Col-0 \times C24 (F_{1-a} and F_{1-b}) in a split-plot design (Liseč et al. 2008). At least three replicates per line were grown.

For the second experiment, ILs were selected to cover the QTL regions determined in the RIL experiment. Plants were grown in two blocks and six subplots per block. Each subplot contained 42 ILs, 42 IL-testcrosses (IL-TCs) to the recurrent parent, and the controls twice (P_1 , P_2 , F_{1-a} , F_{1-b}). The position within the subplot was random. In addition, 'unselected' ILs without IL-TCs were grown in the same experiment. In this case, each subplot consisted of 56 ILs and 36 controls.

Molecular markers and linkage map

The RIL and IL mapping populations were genotyped with a set of 110 framework SNP markers (Törjék et al. 2003).

A linkage map was created from the RIL genotypes as described in detail in Törjék et al. (2006). Additional SNP markers were added to the framework map using Joinmap version 3.0 (Van Ooijen and Voorrips 2001) (Supplementary Fig. 1).

Determination of leaf area and shoot dry biomass

Images of plants were taken 6, 8 and 10 days after sowing (DAS) and whole leaf area was determined using the imaging system and software described in Walter et al. (2007). Due to different light source selection, parameter settings in the automated part of the analysis were adapted accordingly. Editing tools allowing for manual post-processing like removal of false positives (e.g. due to light reflection on vermiculite) were adapted to plant arrangement and their user-friendliness improved.

Shoot dry biomass was determined 15 DAS. The harvested aerial parts of the plants were placed in a vacuum oven at 80°C for 48 h. Dry biomass was measured using an analysis balance. Mean shoot dry biomass in mg plant^{-1} and mean leaf area in $\text{mm}^2 \text{plant}^{-1}$ were estimated using a linear mixed model as described in Meyer et al. (2007).

Calculation of biomass heterosis

F_1 mid-parent-heterosis (MPH) was calculated from the estimated mean values of parents and reciprocal F_1 -hybrids using the equation $\text{MPH} = 100(\overline{F}_1 - \overline{P})/\overline{P}$, where $\overline{F}_1 = (F_{1-a} + F_{1-b})/2$ and $\overline{P} = (P_1 + P_2)/2$ (Falconer and Mackay 1996). The corresponding testcross relative mid-parent-heterosis of RIL-TCs was calculated as $r\text{MPH}_{P_1} = 100(\text{TC}_{P_1,i} - 0.5(\text{RIL}_i + \overline{P}_1))/0.5(\text{RIL}_i + \overline{P}_1)$ for crosses to C24 and as $r\text{MPH}_{P_2} = 100(\text{TC}_{P_2,i} - 0.5(\text{RIL}_i + \overline{P}_2))/0.5(\text{RIL}_i + \overline{P}_2)$ for crosses to Col-0. For statistical analyses absolute testcross mid-parent-heterosis values were calculated from RIL-TCs with C24 as $a\text{MPH}_{P_1} = \text{TC}_{P_1,i} - 0.5(\text{RIL}_i + \overline{P}_1)$ and with Col-0 as $a\text{MPH}_{P_2} = \text{TC}_{P_2,i} - 0.5(\text{RIL}_i + \overline{P}_2)$ (Lamkey and Edwards 1999). To determine the contribution of different genetic effects to MPH of the quantitative trait under study, the linear contrasts for augmented additive effect (additive effect confounded with epistasis, $Z_1 = \text{TC}_{P_1,i} + \text{TC}_{P_2,i}$) and augmented dominance effect (dominance effect confounded with epistasis, $Z_2 = \text{TC}_{P_1,i} - \text{TC}_{P_2,i}$) as defined by Melchinger et al. (2007b) were calculated.

Estimation of heritability

Broad-sense heritability (h^2) for dry weight and leaf area in the RILs was calculated as $\text{Var}_G/(\text{Var}_G + \text{Var}_E)$, where Var_G = genotypic variance and Var_E = environmental

variance estimated using the method of restricted maximum likelihood (REML) with genotype as random factor.

Correlation analyses

Procedures FCORRELATION and PCORRELATION in GenStat Release 10.2 were used for correlation analyses.

QTL analyses

Composite interval mapping (CIM) in RILs was performed using the software package PLABQTL (Utz and Melchinger 1996) with an extension for calculation of the Bayesian information criterion (BIC) according to the method of Burnham and Anderson (2004) to accommodate selection of cofactors and comparison of the models with and without digenic epistatic interactions (Kusterer et al. 2007b). Cofactors were automatically selected by forward stepwise regression. Significant LOD thresholds were determined by 5000 permutations. Data were also submitted to QTLNetwork-2.0 (Yang et al. 2008) to validate main-effect QTL and to perform a 2D-genome scan. QTLNetwork-2.0 implements a mixed-model based composite interval mapping method and allows simultaneous mapping of main-effect and interaction QTL. The genetic map used in all QTL analyses is based on the map presented by Törjék et al. (2006), with additional SNP markers (Supplementary Fig. 1). The augmented degree of dominance ($d_i^*/|a_i^*|$) of significant per se or heterotic QTL was determined by estimating Z_1 and Z_2 at the respective positions (Kusterer et al. 2007b). QTL were classified according to their augmented dominance ratio as additive (A; $|d^*/a^*| < 0.2$), partially dominant (PD; $0.2 \leq |d^*/a^*| < 0.8$), dominant (D; $0.8 \leq |d^*/a^*| < 1.2$), and overdominant (OD; $|d^*/a^*| \geq 1.2$) (Stuber et al. 1987). In contrast to Kusterer et al. (2007b), our experimental design did not include testcrosses of the RILs to the F_1 , therefore epistatic effects (linear contrast Z_3) could not be estimated.

Heterosis data obtained in the IL population were analyzed using ANOVA in GenStat Release 10.2 with the appropriate contrasts, i.e. comparing the IL-TC with the mean of IL and recurrent parent.

Generation means analysis

An overall generation means analysis (GMA) including parental lines, F_1 -hybrids, RILs and RIL-TCs was performed as described in Kusterer et al. (2007b). All parameters in the models were defined according to the F_2 -metric (Cockerham 1954; Yang 2004). Neither cytoplasmic nor maternal effects were significant ($P < 0.01$) in the exploratory analysis and were excluded from further analyses. Model 1 contained the overall mean μ , the overall

additive effect [a] and the overall dominance effect [d]. In addition to these parameters, model 2 also included epistatic effects between unlinked pairs of loci [aa]. Adequacy of each model was tested with a Chi-square (χ^2) test (Mather and Jinks 1982). The coefficient of determination (R^2) was calculated to estimate the proportion of the variation among generation means accounted for by each model.

In the second experiment analyzing introgression lines, genetic effects for individual ILs were estimated by GMA as described in Melchinger et al. (2007a), but using only ILs and their testcrosses to the recurrent parent.

Qualitative mode-of-inheritance classification of QTL

To complement the generation means analysis of ILs, we performed QTL mapping and a classification of the mode-of-inheritance according to Semel et al. (2006) as described in Kusterer et al. (2007a). Each IL and IL-TC was compared by a t-test with the recurrent parent P_i as well as to each other. If either of them was significantly different from parent P_i , the corresponding IL was considered as harboring a QTL. Because the number of replications for the IL and IL-TC was identical but the number of replications of the parental lines was higher, the comparison of IL and IL-TC with the parent was tested at a significance level of $P < 0.01$, whereas the comparison of IL with IL-TC was tested at $P < 0.05$, adopting the procedure of Semel et al. (2006).

Results

Description of traits

We analyzed performance data [dry weight at 15 DAS (DW15) and leaf area at 6, 8, 10 DAS (LA06, LA08, LA10)], as well as data sets calculated from adjusted mean values of the performance data in RILs and RIL-TCs (relative mid-parent heterosis rMPH, absolute mid-parent heterosis aMPH, linear contrast Z_1 and linear contrast Z_2). Heritabilities (h^2) for the performance traits were calculated as proportion of total variance allocated to genotype (Supplementary Table 1). Values were moderate for DW15 (42%) and moderately high for leaf area (58, 59, 53%). Means and variances for all traits averaged over the RILs are given in Supplementary Tables 1 and 2. The distributions of the performance traits show a distinct shift toward higher values in the RIL-TCs compared to RILs for both leaf area and dry weight (Fig. 1). This is reflected in the mid-parent-heterosis values (Table 1). Significant ($P < 0.001$) differences in the performance (DW15, LA06, LA08, LA10) were detected between the two testcross

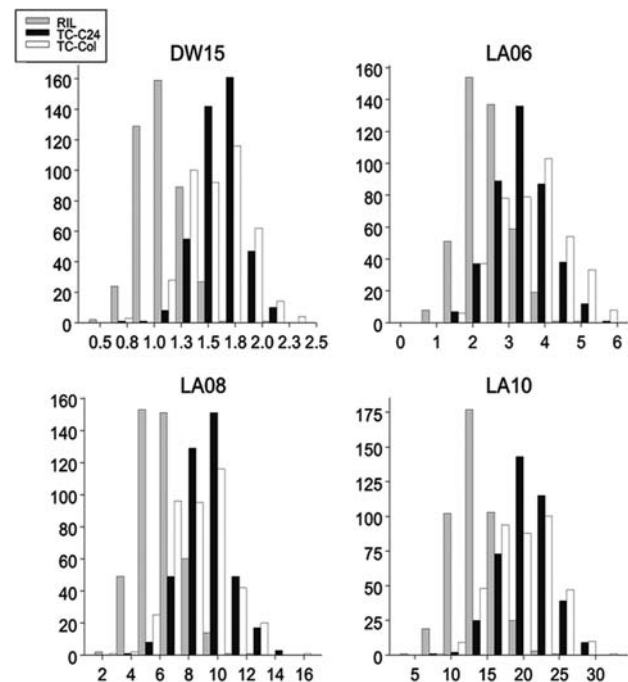


Fig. 1 Distribution of dry weight and leaf area in the RIL and RIL-TC populations. Shown are the distributions of the means (x axis) for dry weight at 15 DAS (mg plant^{-1}) and leaf area at 6, 8, 10 DAS ($\text{mm}^2 \text{plant}^{-1}$). The y axis indicates the number of lines

Table 1 Biometrical analyses of phenotypic data for the six traits in F_1 and RILs

	Trait	MPH (%) F_1	Mean rMPH (%) RILs	rMPH (%) Range in RILs
MPH-C24	DW15	43.71	36.60 ± 18.39	−31.13 to 88.32
	LA06	28.11	29.68 ± 32.10	−45.38 to 126.50
	LA08	24.43	28.42 ± 25.14	−50.48 to 105.60
	LA10	26.15	25.94 ± 22.71	−49.96 to 92.15
MPH-Col	DW15	43.07	34.45 ± 24.99	−36.33 to 99.10
	LA06	32.45	28.59 ± 34.95	−57.93 to 133.03
	LA08	22.02	28.20 ± 30.67	−52.54 to 142.94
	LA10	18.55	26.44 ± 28.01	−46.35 to 92.87

Mid-parent-heterosis (MPH) for the original cross (F_1 , with $F_{1-\text{Col} \times \text{C24}}$ in MPH-C24 and $F_{1-\text{C24} \times \text{Col}}$ in MPH-Col), mean test-cross relative MPH (rMPH) calculated from RILs and RIL-TCs and averaged over the whole population (RIL) and the range of rMPH detected using RILs and RIL-TCs (range)

populations but not the reciprocal RIL populations using Kolmogorov-Smirnov two-sample tests (Conover 1971).

Correlation analyses

Pearson correlation coefficients between dry weight and leaf area within RILs were positive and moderately high (Table 2). The respective means across both RIL-TCs

Table 2 Correlation coefficients between RIL per se traits (below diagonal) and between RIL per se and the mean across RIL-TCs (equivalent to Z_1) (diagonal)

	DW15	LA06	LA08	LA10
DW15	0.17***			
LA06	0.72***	0.11*		
LA08	0.75***	0.93***	0.10*	
LA10	0.73***	0.91***	0.96***	0.15**

***, **, * Significant at $P < 0.001$, $P < 0.01$, $P < 0.05$, respectively

Table 3 Mean absolute correlation coefficients between heterosis ($aMPH_{C24}$, $aMPH_{Col}$) and linear transformations (Z_1 , Z_2), and marker heterozygosity

Trait	$aMPH_{C24}$	$aMPH_{Col}$	Z_1	Z_2
DW15	0.342***	0.204***	0.063 ^{ns}	0.436***
LA06	0.170**	0.122*	0.037 ^{ns}	0.258***
LA08	0.195***	0.107 ^{ns}	0.022 ^{ns}	0.271***
LA10	0.194***	0.119*	0.022 ^{ns}	0.283***

^{ns} Not significant

***, **, * Significant at $P < 0.001$, $P < 0.01$, $P < 0.05$, respectively

(equivalent to Z_1) showed significant but weak positive correlations with the RIL per se performance (Table 2). The linear contrast Z_1 shows no correlation to marker heterozygosity, whereas the linear contrast Z_2 and $aMPH$ show moderate and weak correlations, respectively, with marker heterozygosity (Table 3).

Analyses of RILs and RIL-TCs

A global generation means analysis for dry weight and leaf area was performed on parental lines, F_1 -hybrids, RILs and RIL-TCs (Table 4). Model 1 (not including additive \times additive effects) accounted for 96–98%, and model 2 (including [aa] for 99% of the variation among generation means for all traits). The χ^2 values for the goodness-of-fit of both models were significant. Both models yielded similar estimates of [a] and [d]. Dominance effects were significant ($P < 0.01$) and had a positive sign for all traits. Estimates of [aa] under model 2 were significant ($P < 0.01$) and positive for all traits.

In order to identify genomic regions responsible for the occurrence of heterosis in the Col-0/C24 cross, absolute mid-parent-heterosis ($aMPH$) and the linear contrasts Z_1 and Z_2 for shoot biomass and leaf area were used as traits in QTL analyses (composite interval mapping) in PLABQTL and QTLNetwork-2.0. A complete list and description of QTL detected for biomass heterosis and leaf growth is given in Supplementary Table 3. For biomass heterosis at 15 DAS, six QTL for $aMPH_{C24}$, one QTL for $aMPH_{Col}$ and six QTL for Z_2 were found, jointly accounting for 20.6 ± 3.5 , 18.0 ± 3.4 and $30.5 \pm 3.8\%$ of the phenotypic variation, respectively. Seven QTL for DW15 per se explain $20.2 \pm 3.5\%$, and four QTL for Z_1 account for $18.9 \pm 3.5\%$ of the phenotypic variation. Individual QTL effects range from 1.95 to 18.03%. We could also detect per se and heterotic QTL for leaf area, with joint R^2 ranging from 2.4 to 17.6% (Supplementary Table 3). Augmented dominance ratios for significant QTL were

Table 4 Generation means analysis using RILs and RIL-TCs

Parameters	DW15	LA06	LA08	LA10
Model 1				
μ	$1.56 \pm 0.02^{**}$	$3.61 \pm 0.07^{**}$	$8.99 \pm 0.19^{**}$	$20.25 \pm 0.45^{**}$
[a]	-0.01 ± 0.04	-0.15 ± 0.14	-0.33 ± 0.38	-0.42 ± 0.87
[d]	$0.90 \pm 0.06^{**}$	$2.21 \pm 0.19^{**}$	$5.78 \pm 0.53^{**}$	$13.06 \pm 1.23^*$
R^2 (%)	98.2	96.3	96.0	95.9
$\chi^2_{(5)}^a$	103.0****	76.6****	127.0****	144.9****
Model 2				
μ	$1.52 \pm 0.02^{**}$	$3.45 \pm 0.05^{**}$	$8.48 \pm 0.15^{**}$	$19.09 \pm 0.26^{**}$
[a]	-0.01 ± 0.02	-0.16 ± 0.06	-0.32 ± 0.17	-0.40 ± 0.31
[d]	$0.82 \pm 0.03^{**}$	$1.96 \pm 0.21^{**}$	$4.91 \pm 0.32^{**}$	$11.12 \pm 0.55^{**}$
[aa]	$0.20 \pm 0.05^*$	$0.67 \pm 0.15^*$	$2.19 \pm 0.49^*$	$4.95 \pm 0.85^{**}$
R^2 (%)	99.6	99.4	99.3	99.6
$\chi^2_{(4)}^a$	20.7***	13.3*	21.6***	15.2*

Generation means were calculated without cytoplasmic effect c, which was not significant in the exploratory analysis. Parameters included are overall mean μ , overall additive effect [a], overall dominance effect [d], overall additive \times additive epistasis [aa] (model 2 only)

****, ***, **, * Significant at $P < 0.0001$, $P < 0.001$, $P < 0.01$, $P < 0.05$, respectively

^a χ^2 statistic with degrees of freedom in parentheses

Table 5 Augmented degree of dominance of significant QTL estimated from linear contrasts Z_1 and Z_2

Interval	DW						LA06					
	per se	aMPH _{C24}	aMPH _{Col}	Z_1	Z_2	Gene action	per se	aMPH _{C24}	aMPH _{Col}	Z_1	Z_2	Gene action
1/4–28	0.15	–0.15			–0.20	OD	0.16					OD
1/82–102	–0.13	–0.22		–0.30		A	–0.13			–0.23		A
2/42–60		–0.20			–0.20	OD						
3/0–12	0.26					PD	0.16					D
3/32–38					–0.14	OD						
3/50–72	–0.21	–0.29		–0.18	–0.26	PD		–0.18			–0.23	D
3/82–86												
4/0–13	–0.22	0.13	0.44	0.27	–0.30	PD	–0.15		0.37	0.25	–0.23	PD
4/30–72	–0.20			–0.17		PD	–0.27					PD
5/32–42											–0.16	OD
5/70–94	0.19	–0.20			–0.20	OD	0.16					PD

Interval	LA08						LA10					
	per se	aMPH _{C24}	aMPH _{Col}	Z_1	Z_2	Gene action	per se	aMPH _{C24}	aMPH _{Col}	Z_1	Z_2	Gene action
1/4–28	0.17					OD	0.15					OD
1/82–102		–0.18		–0.23		A	–0.17	–0.19		–0.26		A
2/42–60												
3/0–12	0.24					D	0.22					D
3/32–38												
3/50–72		–0.18			–0.26	D		–0.16			–0.23	D
3/82–86							–0.19					OD
4/0–13			0.37	0.25	–0.23	PD			0.36	0.26	–0.23	PD
4/30–72	–0.26					PD	–0.19			–0.15		PD
5/32–42					–0.16	OD					–0.18	OD
5/70–94	0.19					OD						

Only significant effects ($P < 0.01$) are shown. A negative value signifies an increasing effect of the Col-0 allele. The dominance ratio d^*/a^* for all significant QTL was calculated using effects estimated in Z_1 (a^*) and Z_2 (d^*). QTL were classified as additive (A; $|d^*/a^*| < 0.2$), partially dominant (PD; $0.2 \leq |d^*/a^*| < 0.8$), dominant (D; $0.8 \leq |d^*/a^*| < 1.2$), and overdominant (OD; $|d^*/a^*| \geq 1.2$)

estimated from the linear contrasts Z_1 and Z_2 (Table 5). For shoot biomass four of six QTL detected in Z_2 show overdominant effects and two QTL show partial dominance. The QTL for aMPH_{C24} at position 1/90, where there is no corresponding QTL for Z_2 , has additive effect. Gene actions defined for leaf area are consistent with the biomass data. We tested for presence of digenic epistatic effects between all pairs of marker loci using QTLNetwork-2.0, and could find two significant additive \times additive interactions between unlinked pairs of markers on chromosomes 1 and 5 for DW15 and LA08, and on chromosomes 1 and 4 for DW15, LA08 and LA10, respectively (Supplementary Table 4).

Verification of detected QTL in ILs and IL-TCs

The QTL detected in the RIL population were verified using lines of the two corresponding, reciprocal IL populations (Törjék et al. 2008). In a first step, heterosis data

were analyzed as linear contrasts in ANOVA. All six biomass heterosis QTL at positions 1/12, 1/92, 2/46–56, 3/56–62, 4/4, 5/74 could be verified (Supplementary Table 5). Additional biomass heterosis QTL could be identified in the intervals 1/65–68, 1/77–83, 2/74, 3/74–79, 4/60–78 and 5/14–44. Most (80.5%) IL/IL-TC combinations showed significant heterosis (Fig. 2), revealing a highly complex situation. The IL data were also analyzed using the generation means approach (Melchinger et al. 2007a). Significant dominance effects were detected for previously identified heterotic QTL at positions 1/90–92, 2/46, 2/56, 3/56–62 and 4/4–6 (Supplementary Table 5). Significant additive effects were detected for previously identified biomass QTL (Liseć et al. 2008) at positions 3/60 and 5/86. The mode-of-inheritance classification according to Semel et al. (2006) identified a total of 12 heterotic (overdominant) QTL, across both reciprocal IL subpopulations, most with a positive overdominance effect (Supplementary Table 6).

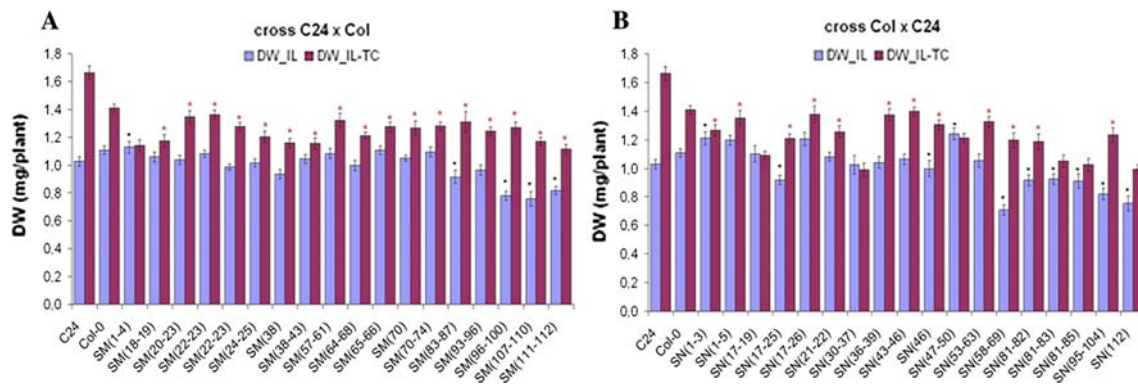


Fig. 2 Biomass in ILs and ILs-TC from the C24xCol (a) and ColxC24 (b) IL populations. Mean values and standard deviations of 24 replicates are shown. ILs are denoted with the substituted marker intervals, with SM indicating C24 background and SN Col-0

background. ILs with biomass significantly ($P < 0.05$) different from the recurrent parent are indicated by a black asterisk; IL-TC displaying significant ($P < 0.05$) mid-parent-heterosis are marked with a red asterisk above the bars

Discussion

The distributions of the performance traits show clear differences between RILs and their testcrosses for dry weight and leaf area, identifying them as heterotic traits. Testcross relative mid-parent-heterosis (rMPH) for all traits behaves like a segregating trait in the RILs, displaying a broad range of values and thus making it possible to use it as a trait in QTL analyses (Stuber et al. 1992). The two testcross populations differ significantly for all performance traits. The differences are also reflected by the fact that only 1 (of 10) heterotic QTL region, at the top of chromosome 4, contributes to both $aMPH_{C24}$ and $aMPH_{Col}$ for dry weight. This can be explained by different heterozygous regions involved in the realization of heterosis between the two populations. Xiao et al. (1995) described a similar situation in rice, where they detected most of the QTL (72%) in only one of the two testcrosses. The authors concluded that dominance complementation is the major genetic basis of heterosis in rice. In a recent maize study, one testcross was significantly higher for all eight traits analyzed, confirming the prevalence of alleles with increasing effect in one of the parents (Frascaroli et al. 2007). The situation is not as clear-cut in the analyzed Arabidopsis populations, with both parents contributing ‘increasing alleles’ as illustrated by the observed transgressive segregation. In our study, augmented dominance ratios of the detected QTL and mode of inheritance classification provide evidence that both dominance and overdominance are involved in biomass heterosis in Arabidopsis. We identified more QTL for performance per se than for Z_1 . This is consistent with the low correlation detected between the two measures. Z_1 represents the augmented additive effect, i.e. the additive effect for the respective QTL minus half the sum of dominance \times additive epistatic interactions (Melchinger et al. 2007b). The discrepancy between QTL detected for performance per se

and for Z_1 points toward the existence of dominance \times additive epistasis, which reduces the power of Z_1 to detect QTL.

Treating mid-parent-heterosis as a quantitative trait, we could identify heterotic QTL for biomass and leaf area. As expected from the high phenotypic correlation between dry weight and leaf area we identified similar performance and heterotic QTL for these traits, not only concerning position, but also effect, increasing allele and gene action, with LA10 being most similar to DW15. There is also good coincidence between QTL detected for $aMPH$ and Z_2 , in agreement with the correlation coefficients. In these cases, the QTL representing the augmented dominance effect (dominance effect for respective QTL minus half the sum of additive \times additive epistatic interactions (Melchinger et al. 2007b)) for the trait under consideration also indicates a heterotic effect. QTL detected for transformation Z_2 show at least partial dominance; most (4 of 6) have an overdominant gene action. The additive effect estimated for the $aMPH$ QTL for DW15 at position 1/90 (no overlap with Z_2) can be attributed to the fact that $aMPH$ is confounded with additive effect (Melchinger et al. 2007b). The same genomic region was found to significantly influence heterosis in the IL/IL-TC study, with overdominant gene action identified in the generation means and mode of inheritance analyses. Positive dominance effects can be counterbalanced by positive additive \times additive effects, thus reducing the power of Z_2 to detect heterotic QTL (Melchinger et al. 2007b). The existence of positive additive \times additive effects is suggested by the overall generation means analysis in the RILs and substantiated by the mixed model QTL analysis. In GMA the model including epistasis detected significant dominance and additive \times additive effects for all traits. The inclusion of epistasis in the model considerably improved the fit, explaining around 99% of the variation among generation means for all traits (substantially increased R^2 ,

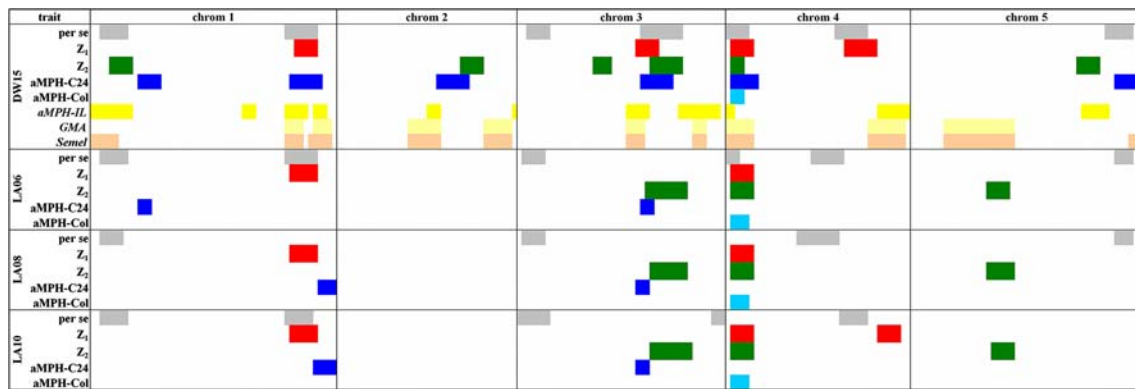


Fig. 3 Schematic representation of growth related per se and heterotic QTL. QTL positions are indicated by *boxes* corresponding to the intervals covered by the respective confidence intervals. *DW15* Shoot dry weight at 15 DAS, Z_1 augmented additive effect,

Z_2 augmented dominance effect, $aMPH_{C24}$, $aMPH_{Col}$ absolute mid-parent-heterosis in cross with C24 and Col-0, respectively, *LAN* leaf area at n DAS

while the χ^2 test remained significant). Furthermore, Kusterer et al. (2007b) using a triple test cross approach on the same RIL population at a later growth stage, did detect additive \times additive epistatic effects. Overdominance and epistasis have been reported to play an important role in heterosis in a number of organisms for various traits (reviewed in Lippman and Zamir (2007)). In *Brassica rapa*, 15 of 23 QTL detected for biomass and associated traits showed overdominance, and 444 digenic epistatic interactions were found (Dong et al. 2007). Analyzing heterosis for yield and its component traits in *Brassica napus*, Radoev et al. (2008) concluded that epistasis together with all levels of dominance form the genetic basis of heterosis in this crop plant. QTL analyses of heterosis in maize revealed an association of QTL with additive to dominance effects with traits with low heterosis, whereas QTL for traits with high heterosis mostly showed dominance to overdominance effects (Frascaroli et al. 2007). While our experimental design did not allow us to estimate directly the involvement of digenic epistatic effects on heterosis via the transformation Z_3 proposed by Melchinger et al. (2007b), we found evidence for additive \times additive epistasis between two pairs of loci. The interaction between loci on chromosomes 1 and 5, with parental allele combinations increasing biomass, does not include main-effect QTL. In contrast, the two interacting loci on chromosomes 1 and 4 have been identified previously as main-effect QTL. Furthermore, both have been detected as aMPH QTL and a recombinant allele combination increased biomass, consistent with a role in heterosis (increased performance of a double heterozygote over both parental homozygotes).

QTL for different traits and transformations co-locate in three main genomic regions on chromosomes 1, 3, and 4. In a parallel study (Liseć et al. 2009) focusing on metabolites we identified hot spots for heterotic metabolite QTL in similar positions. The cluster containing the strong heterotic

biomass QTL identified at position 4/4 corresponds to the ‘hot spot’ described for metabolic and biomass QTL in the same Arabidopsis population (Liseć et al. 2008) and co-locates with a cluster of 23 metabolic heterotic QTL described in the metabolic study (Liseć et al. 2009). Interestingly, this ‘hot spot’ is also involved in additive \times additive epistasis (cf. Supplementary Table 3). This could indicate the presence of a major growth/metabolism regulator in this region, although we cannot rule out tightly linked QTL, as demonstrated by Kroymann and Mitchell-Olds (2005) who detected two interacting QTL within 210 kb. The ‘hot spot’ on top of chromosome 4 currently spans 155 kb and contains 66 genes, including the *FRIGIDA* gene (*FRI*, At4g00650), a well-known major determinant of natural variation in Arabidopsis for flowering time (Johanson et al. 2000). However, the location of the QTL peak and preliminary analyses of subILs covering this region suggest a minor role for *FRI* in the manifestation of heterosis.

The parallel use of RIL and IL populations and several analysis methods proved very beneficial, as both populations and the application of the various methods complemented each other. This did not only allow us to validate QTL found in one population, but also to detect additional QTL in the other. The main difference between RILs and ILs is the absence of variation for ‘background’ epistasis in the ILs, thereby increasing the power to detect QTL (Keurentjes et al. 2007; Reif et al. 2009). However, as epistasis has been shown to play an important role in heterosis, either both types of populations (Keurentjes et al. 2007), or specially selected double or multiple introgression ILs (Reif et al. 2009) are needed to detect and quantify digenic, or higher order interactions.

In our study QTL for growth traits co-locate at only two positions (on chromosome 3) with growth QTL (Z_1) identified previously in the same populations in a different environment at a later developmental stage (Kusterer et al. 2007b).

The specifically detected QTL indicate that different factors are of relevance under different conditions. In contrast, eight of the ten heterotic QTL (Z_2 and Z_3) reported by Kusterer et al. (2007b) and Melchinger et al. (2007a) coincide with heterotic QTL identified in this study (cf. Fig. 3), including the main candidate regions at the top of chromosome 4 and bottom of chromosome 3. In these regions, the earlier study identified QTL for the linear contrast Z_3 , estimating additive \times additive epistatic effects, further underlining the importance of epistasis for heterosis. The complex trait ‘heterosis’ is expected to be controlled by many genes, the combination and interaction of which may depend on the organism and trait under study (Korn et al. 2008; Li et al. 2008). In each cross, only a subset of specific combinations may be relevant for heterosis, as illustrated by the seemingly contradictory findings concerning the genetic basis of heterosis reported in different studies (reviewed in Lippman and Zamir (2007)). The joint detection of heterotic QTL in our two studies suggests the existence of genomic regions of more general importance for the manifestation of heterosis. These regions constitute primary targets for further fine-mapping with the ultimate goal to identify and characterize the underlying genes. The identification of heterotic QTL in *Arabidopsis* and the superior tools for the identification of natural genetic determinants of trait variation (Peters et al. 2003; Weigel and Nordborg 2005), including the rapid advances in genome re-sequencing in this species (Ossowski et al. 2008) opens the opportunity to identify the DNA variation responsible for or contributing to heterosis.

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