$_{\scriptscriptstyle 1}$ Life-history trade-offs and the genetic basis of fitness in Arabidopsis

thaliana

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₂ Abstract

Local adaptation may entail trade-offs in resource allocation to survival and reproduction, but also an increase in overall availability of resources ("condition") through improved capacity for coping with local environmental factors. We examined the relative contribution of trade-offs and increased condition to adaptive evolution in a recombinant-inbred-line population of Arabidopsis thaliana planted at the native sites of the parental ecotypes in Italy and Sweden in two years. We estimated genetic correlations among fitness components based on genotypic means, and explored their causes with QTL mapping. Local ecotypes produced more seeds per fruit 18 than did non-local ecotypes, reflected in stronger adaptive differentiation than was previously shown based 19 on survival and fruit number only. Genetic correlations between survival and overall fecundity, and between number of fruits and number of seeds per fruit, were positive, and there was little evidence of a trade-off 21 between seed size and number. Quantitative trait loci for these traits tended to map to the same regions of the genome, and showed positive pleiotropic effects. The results indicate that adaptive differentiation between 23 the two focal populations largely reflects the evolution of increased ability to acquire resources in the local environment, rather than shifts in the relative allocation to different life-history traits. Differentiation both in phenology and tolerance to cold are likely to contribute to the advantage of the local genotype at the two sites.

₇ Introduction

Adaptation to the local environment is reflected in an increase in mean population fitness in response to local selection pressures (Williams 1966a; Kawecki & Ebert 2004). This may be associated with a shift in relative allocation of resources to optimise the balance between components of fitness, such as between reproduction and survival, or offspring size and number (Williams 1966a; b; Smith & Fretwell 1974; Schluter et al. 1991). Adaptation may also be associated with the evolution of traits increasing the amount of resources available to individuals (their resource status or "condition") in the local environment, which could supersede the effects of variation in relative allocation to different functions (Van Noordwijk & Jong 1986; Schluter et al. 1991). Condition is likely to be a complex phenotype affected by many traits, including those influencing the ability to tolerate local environmental stresses, and those influencing the match between the timing of life-history transitions and seasonal changes in resource availability. Variation in condition versus allocation have distinct implications for local adaptation. With variation in allocation, trade-offs place a constraint on adaptive evolution, and will be reflected as negative correlations between components of fitness. By contrast,

variation in condition should allow for increased allocation to multiple components of fitness and thus positive correlations between fitness components. A full understanding of local adaptation and life-history evolution therefore requires knowledge of the genetic basis of overall fitness and the genetic basis of correlations between different components of fitness.

If fitness components are correlated, this raises two questions about their genetic architecture. First, to what
extent are fitness components affected by loci with pleiotropic effects on multiple components? Second, if loci
do show pleiotropy, are the directions of allelic effects on affected traits consistent across all loci and with the
signs of correlations among components of fitness? If there are genetic trade-offs among fitness components,
we expect to observe antagonistic pleiotropic effects of individual quantitative trait loci (QTL), whereby
an allele is associated with an increase in one component of fitness, but a decrease in one or more other
components (Hazel 1943; Falconer & Mackay 1996). On the other hand, if variation in condition is large we
expect positive pleiotropy, whereby alleles at QTL affecting resource status are associated with changes in two
or more fitness components in the same direction, causing phenotypes to be positively correlated (Houle 1991).
In reality, both processes are likely to be acting, and it is the relative strength of trade-offs and variation in
condition that will determine whether negative or positive correlations are observed.

In plants, three trade-off relationships are likely to be especially relevant for overall fitness. Firstly, resources invested in reproduction are not available for growth and defence, causing a trade-off between fecundity and survival. A trade-off between reproduction and subsequent survival has been documented in many iteroparous species (Williams 1966b; Edward & Chapman 2011). However, a trade-off between reproduction and survival can also be expected in semelparous organisms if traits increasing the chance of juvenile survival reduces resources available to reproduction. Thus, we expect a negative correlation between fecundity and survival. Second, total seed production is a function of both the number of fruits produced and the number of seeds per fruit, and there may be a trade-off between these two components of fecundity. For practical reasons, studies of local adaptation in plants typically focus on either fruit production or estimates of total seed production as a measure of fecundity (e.g. Latta 2009; Hall et al. 2010; Fournier-Level et al. 2011; Ågren et al. 2013). To quantify components of fecundity, it is necessary to estimate both number of fruits and number of seeds per fruit, and substantial additional effort is required to collect and process data on two components compared to just one (e.g. Maddox & Antonovics 1983; Verhoeven et al. 2004; Hall & Willis 2006; Leinonen et al. 2011; Ågren & Schemske 2012). If investment in fruit production is negatively correlated with investment in seed production per fruit, relying on estimates of only one of these components of fecundity will overestimate

variation in total fecundity. If investment in seed and fruit production are positively correlated, the opposite would be true.

Third, theory predicts a trade-off between investment in individual offspring and the total number of offspring 72 (Lack 1954; Smith & Fretwell 1974; Lloyd 1987). In plants this would be expressed as a negative correlation 73 between seed size and number (Harper et al. 1970; Leishman et al. 2000), and selection for larger seeds may thus constrain the evolution of increased fecundity. Negative correlations between seed size and total seed number have been documented across species (Šerá & Šerý 2004) and within crop species (Sadras 2007). Meanwhile studies within natural plant populations have found positive, negative, and negligible correlations between seed size and number (Silvertown 1989; Venable 1992). In Arabidopsis thaliana, trade-offs between seed size and number of seeds per fruit have been observed for recombinant lines grown in controlled environments (Alonso-Blanco et al. 1999; Gnan et al. 2014), and both linkage mapping and mutant screens have found pleiotropic or closely linked QTL with antagonistic effects on seed size and number of seeds per 81 fruit (Alonso-Blanco et al. 1999; Van Daele et al. 2012; Gnan et al. 2014). However, the extent to which these relationships translate into a negative correlation between seed size and overall fecundity is not known and should depend on the direction and magnitude of the correlation between the two components of fecundity: number of seeds per fruit and number of fruits.

Previous work on the genetics of correlations among fitness components and seed size in A. thaliana has largely been conducted in controlled conditions, and it is unclear whether results reflect the situation in natural environments (Alonso-Blanco et al. 1999; Van Daele et al. 2012; Gnan et al. 2014). On the one hand, trade-offs may be more likely to be expressed in less benign natural environments, where resource availability is lower and stress in the form of frost, drought, and antagonistic biotic interaction more common. On the other hand, in genetically variable populations, variation in ability to meet the challenges of harsh environmental conditions may still result in sufficient variation in plant condition to mask variation in allocation strategy. For example, studies of natural populations of species other than A. thaliana have found that the the heritability of seed size, the direction of the correlation between seed size and number, and the impact of variation in seed size on fitness depends critically on the environment in which they are grown (reviewed by Silvertown 1989). To understand properly the relationship between trade-offs, variation in condition and local adaptation, we need to compare genotypes in the environments from which they originate.

In this study, we investigate the contribution of individual components of fitness to estimates of local adaptation, and the genetic basis of correlations among components of fitness. We use a population of recombinant

inbred lines (RILs) derived from a cross between two locally-adapted ecotypes of A thaliana from close to 100 the southern (Italy) and northern (Sweden) margins of the native range in Europe. Reciprocal transplants 101 have shown that the two source populations display strong adaptive differentiation expressed through higher 102 survival and fruit production of the local ecotype (Ågren & Schemske 2012; Ågren et al. 2013), and there is 103 some evidence that the local ecotype also produces more seeds per fruit compared to the non-local ecotype 104 (Ågren & Schemske 2012). QTL mapping in the RIL population identified a total of 15 QTL for an estimate 105 of overall fitness (number of fruits per seedling planted) at the sites of the two source populations (Ågren et 106 al. 2013). However, this estimate of overall fitness did not include possible variation in seed production per 107 fruit, and it is therefore not clear how inclusion of this fitness component would affect estimates of selection 108 against the non-local ecotype, correlations between fecundity and survival, or the genetic basis of fecundity 109 and overall fitness.

Here, we quantify seed output per fruit and mean seed size per fruit of the parental ecotypes and of >300 111 RILs planted at the sites of the source populations in two years. We combine these data with previously 112 published data on survival and fruit production to ask: (1) Does the local ecotype produce more seeds per 113 fruit than does the non-local ecotype, which would result in an even larger estimate of selection against the 114 non-local ecotype than an estimate previously reported based on survival and fruit production only? (2) Are 115 there correlations between fecundity and survival, between components of fecundity (number of fruits and 116 number of seeds per fruit), and between offspring number and size, and are these negative or positive? (3) Are 117 there pleiotropic effects of QTL for number of seeds per fruit and seed mass on other components of fitness, 118 and are these effects positive or negative? 119

$_{\scriptscriptstyle 120}$ Materials and methods

Data collection

We estimated seed traits for recombinant inbred lines (RIL) and parental accessions in reciprocal transplant experiments conducted at the native sites of the source populations in two years (2010-2011 and 2011-2012; henceforth 2010 and 2011). These experiments have previously been described by Ågren & Schemske (2012) and Ågren et al. (2013), who quantified survival to reproduction, number of fruits per reproductive plant, and number of fruits per seedling planted for the parental ecotypes and 398 RILs. We expanded these data by quantifying the number of seeds per fruit (henceforth "seeds/fruit") and mean seed mass per fruit. In each

site × year combination we sampled a single mature fruit from between 1 and 12 plants per RIL and between 128 23 and 100 parental plants; sample sizes varied because lines varied in how well they survived in different sites 129 and years. For each fruit, we counted the number of seeds and determined total seed mass to the nearest 0.01 130 mg on an AT261 balance (Mettler Toledo, Columbus, United States). We calculated mean seed mass as the 131 mass of all seeds in a fruit, divided by the total number of seeds. We estimated genetic values for each line in 132 each site × year combination as the mean across all individuals of the same RIL or parental ecotype. Not all 133 RILs produced at least one fruit that could be harvested, but we could estimate genetic values for seeds/fruit 134 and seed mass in 395 (Italy 2010), 398 (Italy 2011), 395 (Sweden 2010), and 394 (Sweden in 2011) of the total 135 of 398 RILs. As such, there is unlikely to be substantial bias due to the RILs not included. 136

We combined data on seeds/fruit with previously published data to obtain estimates of overall fecundity and 137 overall fitness that include information on seed number. In previous analyses of data from these experiments 138 (Ågren & Schemske 2012; Ågren et al. 2013, 2017), fecundity was defined as number of fruits per reproductive 139 plant ("fruits/RP") and overall fitness as number of fruits per seedling planted ("fruits/seedling"). Here, we 140 estimated the overall fecundity of reproductive plants (henceforth: "seeds/RP") by multiplying individual fruits/RP by line-mean seeds/fruit. We chose to estimate fecundity this way because we did not have data on 142 seeds/fruit for all individuals for which data on fruits/RP were available. Moreover, it was impractical to sample more than one fruit per plant, precluding any estimate of within-plant variation in seeds/fruit. We 144 quantified total fitness as the number of seeds per seedling planted ("seeds/seedling"; zero for plants that did 145 not survive to reproduce). 146

We estimated broad-sense heritability (H²) as the proportion of total phenotypic variation among individuals
that is explained by RIL genotype in each site-year combination. We used a mixed-effect ANOVA estimated
using the package lme4 (Bates et al. 2015), with block as a fixed effect and RIL genotype as a random effect.
To assess the uncertainty around these estimates we performed parametric bootstrapping on model parameters
using the function bootMer, and estimated 95% confidence intervals as the 2.5% and 97.5% quantiles of 1000
bootstrap draws. We carried out data handling and statistical analyses in RStudio 1.1.442 using R 3.6.3
(RStudio Team 2016; R Core Team 2018)).

Fitness differences between parental lines

To assess the influence of different fitness components on estimates of adaptive differentiation, we quantified selection against the non-local ecotype by calculating selection coefficients based on individual components of

fitness, and on estimates of overall fitness. We calculated selection coefficients $s = 1 - w_{min}/w_{max}$, where w_{min} is the fitness of the less fit ecotype and w_{max} that of the fitter ecotype. For cases where the non-local ecotype had higher fitness than the local ecotype, we multiplied the selection coefficient by -1. We calculated selection coefficients based on two estimates of overall fitness: fruits/seedling and seeds/seedling, reflecting fitness estimates excluding and including information on seeds/fruit. We also calculated selection coefficients based on survival, and on two components of fecundity (fruits/RP and seeds/fruit).

We estimated confidence intervals for selection coefficients by non-parametric bootstrapping. We drew 1000 bootstrap re-samples by sampling data with replacement from within experimental blocks (N = 30 blocks in 164 each site x year combination). We calculated selection coefficients for each bootstrap sample and estimated 95\% confidence intervals for each coefficient as the 2.5\% and 97.5\% quantiles of these values. We tested the 166 null hypothesis that there is no adaptive differentiation using two-tailed p-values, calculated as twice the 167 proportion of bootstrap values overlapping zero. It is more difficult to determine whether selection coefficients 168 for the two measures of overall fitness, fruits/seedling and seeds/seedling, differ from one another because 169 both estimates include common data on fruit number, and as such are not independent. Rather than perform 170 a formal test, we simply asked whether the selection coefficient based on seeds/seedling was beyond the 95% 171 confidence interval of that based on fruits/seedling. We compared differences between parental lines in mean 172 seed mass in each site × year combination using Wilcox-signed-rank tests. 173

74 Correlations between traits

For each site × year combination we quantified the genetic correlations between pairs of traits by calculating Pearson correlation coefficients, r, between RIL means. A genetic correlation is the correlation between 176 genetic values of genotypes, which are obtained by averaging over individuals within each genotype (Falconer 177 & MacKay 1989). We estimated 95% confidence intervals around point estimates of genetic correlations 178 by drawing 1000 non-parametric bootstrap samples from vectors of RIL means, recalculating correlation 179 coefficients, and taking the 2.5% and 97.5% quantiles of the distribution of correlation coefficients across 180 these resamples. We examined relationships between three pairs of traits: (1) survival and overall fecundity 181 (seeds/RP), (2) the two components of fecundity (fruits/RP and seeds/fruit), and (3) offspring size (mean seed mass) and overall fecundity (seeds/RP). We compared seed mass with seeds/RP rather than seeds/fruit 183 or fruits/RP because the former is a more complete estimate of fecundity, but to allow comparison with other studies we also report the correlations between seed mass and seeds/fruit. 185

186 QTL mapping

We mapped QTL for fitness and its components using the R/qtl package in R (Broman et al. 2003; Broman & 187 Šen 2009) using additional visualisation tools from the package arghqtl (Ellis 2018). We mapped QTL for seed 188 mass, seeds/fruit, seeds/RP, and seeds/seedling. Mapping results for survival, fruits/RP and fruits/seedling 189 were previously reported by Ågren et al. (2013) based on 398 RILs. However, the number and positions of QTL 190 detected can be affected by the number of RILs included because different subsets of lines contain different 191 recombination events. To allow comparisons of QTL positions and examination of evidence of pleiotropic 192 QTL effects, we therefore repeated QTL mapping for survival, fruits/RP and fruits/seedling including only 193 those RILs with information on seed size and number in each site × year combination as described in "Data 194 collection". 195

We performed mapping based on RIL mean data for each site × year combination separately. We used 196 Haley–Knott regression using genotype probabilities of the genetic markers and pseudomarkers in gaps >2 197 cM (Haley & Knott 1992). We performed a two-QTL scan of the genome with 10,000 permutations of the phenotypic data to determine 5% LOD-significance thresholds for inclusion of QTL and epistatic interactions 199 (Doerge & Churchill 1996; Broman & Šen 2009). Based on these thresholds, we used R/qtl's automated stepwise model selection procedure to identify significant additive QTL and epistatic interactions. We applied 201 a quantile normal transformation to phenotypes before model selection. Finally, we fitted a multiple-QTL 202 model to untransformed data to calculate, for each locus, the proportion of the total phenotypic variance 203 among RILs explained (PVE), and the effect size (in units of the trait) of a substitution of the Swedish 204 homozygous genotype. 205

To investigate whether QTL showed pleiotropic effects on multiple traits, we examined whether QTL for 206 different traits map (co-localise) to the same region. There are currently no clear guidelines on how to formally delineate QTL in linkage-mapping studies, so we rely on a set of heuristic rules used in previous studies (Ågren 208 et al. 2013, 2017; Dittmar et al. 2014; Oakley et al. 2014; Postma & Ågren 2018). We considered any pair of 209 QTL to co-localise and represent the same QTL if the 95% Bayesian credible intervals for these estimates 210 overlapped. Based on these criteria, we identified 'pleiotropic' regions associated with multiple traits if they 211 contained co-localising QTL for two or more traits that were directly observed (fruits/RP, seeds/fruit, survival 212 and seed mass). We excluded "poorly-defined" QTL whose credible intervals for QTL position were greater 213 than one quarter of the length of the shortest chromosome (15.2cM) from assessments of co-localisation, because such QTL provide little information about position.

$_{^{216}}$ Results

Number of seeds per fruit influences estimates of selection

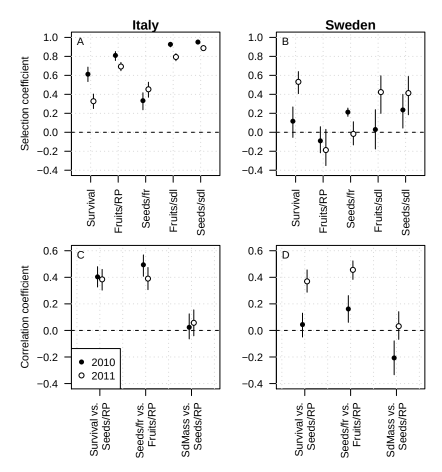


Figure 1: Selection against the non-local ecotype and genetic correlations among components of fitness, and between mean seed mass and fecundity in experiments established in 2010 and 2011 at sites in Italy and Sweden. Selection against the non-local ecotype was estimated based on three components of fitness: survival to reproduction, number of fruits per reproductive plant (Fruits/RP), and number of seeds per fruit (Seeds/fr), and based on two estimates of total fitness: number of fruits per seedling planted (Fruits/sdl), and number of seeds per seedling planted (Seeds/sdl) at the Italian and Swedish sites (A, B). Error bars show 95% bootstrap confidence intervals. Positive selection coefficients indicate selection favouring the local ecotype, negative values selection favouring the non-local ecotype. Genetic correlations between components of fitness and between mean seed mass (SdMass) and number of seeds per reproductive plant (Seeds/RP) were quantified as the correlations among RIL means at the Italian and Swedish sites (C,D).

Selection through seeds/fruit favoured the local genotype in both years in Italy, and in one year in Sweden (figure 1A and 1B). In Italy, the local ecotype produced significantly more seeds/fruit in both 2010 (means:

36.1 vs 24.0) and 2011 (means: 35.6 vs 19.5). In Sweden, the local ecotype produced significantly more

seeds/fruit than the non-local ecotype in 2010 (means: 36.6 vs 28.8) but not in 2011 (means: 23.0 vs 23.4). 221 Selection through seeds/fruit in Sweden in 2010 is a novel source of selection as none was previously detected 222 for survival or fruits/RP in Sweden this year (figure 1B; Ågren & Schemske 2012). The only significant source 223 of selection in Sweden 2011 was via survival (figure 1B; Ågren et al. 2013). 224 Differences in seeds/fruit affected estimates of differences in overall fitness. When information on seeds/fruit was 225 included in estimates of overall fitness (i.e. fitness quantified as seeds/seedling rather than as fruits/seedling), 226 estimates of selection favouring the local ecotype in Italy increased from 0.93 to 0.95 in 2010 and from 0.79 to 227 0.89 in 2011 compared to when fecundity was based on fruit production only (figure 1A). In Sweden, including 228 information on seeds/fruit increased the estimated selection against the non-local ecotype in 2010 from 0.03 to 0.24 and reduced it from 0.42 to 0.41 in 2011 (figure 1B). 230

$_{\scriptscriptstyle{231}}$ QTL for seeds/fruit contribute to differences in fecundity

We found QTL for seeds/fruit across all five chromosomes that together explained between 24.3% and 29.0% 232 of the variance in mean seeds/fruit among RILs in each site-year combination (figure 2, tables S1 and S2). For 233 all QTL detected in Italy, the non-local Swedish allele was associated with fewer seeds per fruit. In Sweden, 234 the local allele was associated with an increase in seeds/fruit in all four QTL detected in 2010, and in three 235 out of five QTL in 2011. For the other two QTL the local allele was associated with a decrease in seeds/fruit 236 (figure 2). Swedish alleles at QTL for seeds/fruit were thus associated with reduced seed output per fruit in Italy, whereas the direction of effects varied in Sweden. In addition, we detected two pairs of loci showing 238 significant epistatic interactions for seeds/fruit in Sweden (figure S1, table S2). In 2010, RILs with at least one Swedish genotype at 1@22.7 or 5@70.6 produced more seeds/fruit compared to plants with the Italian 240 genotype at both loci. In 2011, RILs with Swedish genotypes at both 1@61.1 and 3@21.0 produced the fewest 241 seeds/fruit of any genotype combination, and the most seeds/fruit was recorded for RILs that were Swedish 242 at 1@61.1 and Italian at 3@21.0 (figure S1).

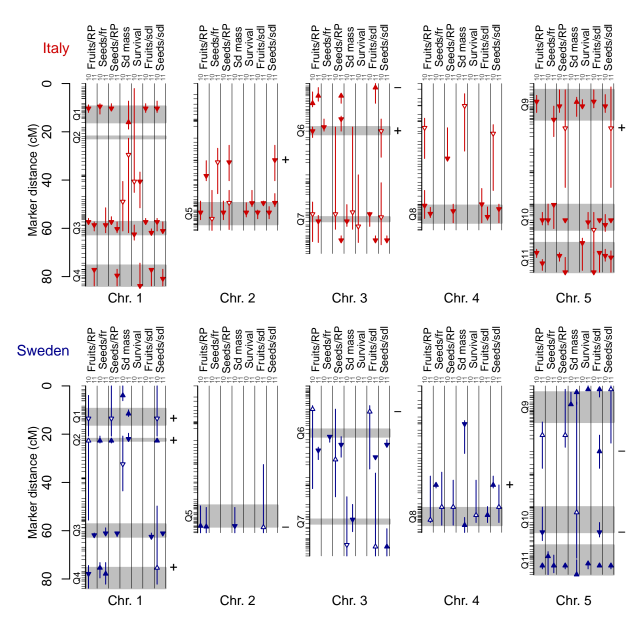


Figure 2: QTL for fitness components, seed mass and estimates of total fitness. Lanes show QTL for number of fruits per reproductive plant (Fruits/RP), number of seeds per fruit (Seeds/fr), number of seeds per reproductive plant (Seeds/RP), seed mass (Sd mass), survival, number of fruits per seedling planted (Fruits/sdl), and number of seeds per seedling planted (Seeds/sdl). Arrows indicate most-likely QTL position and direction of effect of the Swedish genotype on the phenotype (upward: increased; downward: decreased) in Italy (red, upper panels) and Sweden (blue, lower panels) in the 2010 (10) and 2011 (11) experiments, respectively. Vertical bars show the 95% Bayesian credible intervals for QTL position. Open arrows show QTL with credible intervals wider than 15.2cM. Grey boxes indicate regions Q1 to Q11 harbouring QTL with pleiotropic effects on two or more of those traits that were directly observed (Fruits/RP, Seeds/fr, survival and seed mass). Tracks to the left of chromosomes show marker positions in cM. "+" and "-" symbols indicate QTL detected for seeds/seedling not detected for fruits/seedling in the same year and vice versa.

Including information on seeds/fruit in estimates of overall fitness affected the number of fitness QTL detected

(figure 2, tables S3 and S4). Most QTL for seeds/seedling corresponded to a nearby QTL for fruits/seedling, 245 and are likely to reflect the same QTL. However, in Italy, when fitness is quantified as seeds/seedling we detected four additional QTL that were not detected in the same year when fitness is defined as fruits/seedling 247 (indicated with "+" in figure 2), but did not detect one QTL that was previously found for fruits/seedling 248 (indicated with "-" in figure 2). In Sweden, none of the four QTL detected for number of seeds per seedling 249 planted in 2010 were detected when fitness is quantified as fruits/seedling, whereas four QTL detected for fruits/seedling in 2010 or 2011 were not detected for seeds/seedling in the same year. When information 251 on seed number is incorporated into estimates of overall fitness, there was thus a net gain in the number of 252 fitness QTL detected, although this was offset by the loss of some QTL detected when fitness is estimated 253 based on survival and fruit production only. 254

Limited differentiation in seed mass

Italian parent produced 18% larger seeds than did the Swedish parent at the Italian site (W = 1449.5, p =257 0.003) and 3% larger seeds at the Swedish site (W = 5130, p = 0.023). No significant difference in seed mass between the two parental ecotypes was recorded in the second year at either the Italian site (W = 753.5, p =259 0.306) or the Swedish site (W = 720.5, p = 0.519). Both ecotypes produced larger seeds at the site in Sweden 260 compared to that in Italy. In Italy, we identified two QTL for seed mass in 2010 and five in 2011 explaining 8.6% and 15.1% of the 262 phenotypic variation among RILs respectively (figure 2, table S5). The Swedish allele was associated with a decrease in seed mass for both QTL detected in 2010, and for three of the five QTL detected in 2011. In 264 Sweden, we detected five QTL for seed mass in 2010 and eight in 2011, explaining 26.0% and 29.0% of the 265 phenotypic variation among RILs respectively. The local Swedish alleles were associated with an increase 266 in seed mass at two of the five QTL detected in 2010, and at five of the eight QTL detected in 2011. We 267 detected a significant epistatic interaction for seed mass between one pair of loci in Sweden in 2011 (figure S3, table S2). Plants with the Swedish genotype at 3@55.4cM had smaller seeds, but only in the presence of the 269 Italian genotype at 5@78.2cM.

Differences in seed mass between the two ecotypes depended on both site and year (figure S2). In 2010, the

271 Positive correlations dominate among fitness components

Components of fitness were mostly positively correlated with one another. Genetic correlations between 272 fruits/RP and seeds/fruit, as well as between survival and seeds/RP were positive in both years in Italy and 273 in Sweden in 2011 ($r \ge 0.37$; $p \le 0.0001$; $df \ge 392$; figures 1C, 1D and S4). In Sweden in 2010, the positive 274 correlation between fruits/RP and seeds/fruit was weaker but still significant (r = 0.16, p = 0.001, df = 393), 275 while survival and seeds/RP were not significantly correlated (r = 0.04, p = 0.381, df = 393). Seed mass showed either negligible or negative correlations with (seeds/RP) and the fecundity component 277 seeds/fruit (figures 1C, 1D and S4). In Sweden in 2010, seed mass was negatively correlated with seeds/RP $(r = -0.21, p \le 0.0001, df = 393)$, whereas no significant correlation was detected between seed mass and 279 overall fecundity in Sweden in 2011 (r = 0.03, p = 0.529, df = 392), nor in Italy in either year $(|r| \le 0.06;$ 280 $p \ge 0.253$; $df \ge 393$). Correlations between seed mass and seeds/fruit were very similar to those between seed mass and seeds/RP (table S6), with a significant negative correlation in Sweden in 2010 (r = -0.19, 282 p = 0.0002, df = 393), but no significant correlations in Sweden in 2011 or Italy in either year ($|r| \le 0.06$; $p \ge 0.06$), $p \ge 0.06$ 0.266; $df \ge 392$). 284

285 QTL show positive pleiotropy

QTL for components of fitness that could be resolved to within 15.2 cM tended to map to one of eleven distinct regions of the linkage map (Q1-Q11, indicated in grey in figure 2; tables S1, S7 and S8). Seven regions in Italy 287 and three in Sweden harboured QTL for both survival and seeds/RP. Likewise, six regions in Italy and five in 288 Sweden harboured QTL for both seeds/fruit and fruits/RP. In all but two of these cases, local alleles at the respective sites were associated with an increase in both fitness components. In Sweden in 2011, Q3 showed a 290 significant epistatic interaction with Q6 for both seeds/fruit and fruits/RP (figures S1 and S5) as well as for 291 both overall fecundity and overall fitness (figures S6 and S7; see also Ågren et al. 2013). Plants with the 292 Swedish genotype at Q3 had higher fecundity and fitness than plants with the Italian genotype at this locus, but only in the presence of the Italian genotype at Q6, whereas plants with the Swedish genotype at both loci 294 had the lowest fitness of any of the genotypic combinations. Taken together, these observations indicate that regions containing pleiotropic QTL tend to affect pairs of fitness components in the same direction. 296

We found one notable exception to the preponderance of positive pleiotropic effects. In Sweden in 2010, the
Swedish allele at Q4 was associated with an increase in seeds/fruit, but a decrease in fruits/RP (figure 2).

This antagonistic effect represents a trade-off between the two components of fecundity.

Three QTL for seed mass in Italy and seven in Sweden were also found in pleiotropic regions (Q1, Q7, Q9 in Italy; Q1, Q2, Q5, Q7, Q8, Q9 and Q11 in Sweden; figure 2, table S5). At one of these regions in Italy (Q7) and three regions in Sweden (Q8, Q9, Q11) where QTL for seeds/RP were detected, local alleles were associated with an increase in both seed mass and fecundity. At two regions each in Italy (Q1, Q9) and Sweden (Q1, Q2), alleles were associated with effects in opposite directions on seed mass and seeds/RP. For a subset of these regions, overlap was also observed between positions of QTL for seed mass and seeds/fruit: local alleles were associated with an increase in both seed mass and seeds/fruit in two regions in Sweden (Q8, Q11), and with effects in opposite directions in two regions in Italy (Q1, Q9) and one region in Sweden (Q2; figure 2).

Discussion

The present study demonstrates that genetic differences influencing the number of seeds per fruit can make 310 an important contribution to adaptive differentiation and the genetic basis of fitness variation among natural populations of Arabidopsis thaliana. In a reciprocal transplant between an Italian population located close to 312 the southern margin of the European native range and a Swedish population located close to the northern range margin, the local ecotype produced more seeds per fruit than did the non-local ecotype, and including 314 information about seed number per fruit thus increased the estimated magnitude of the fitness advantage of the local ecotype compared to estimates based on differences in fruit production and survival alone. Genetic correlations between fecundity and survival, and between components of fecundity were generally positive, 317 and there was little evidence of a trade-off between fecundity and seed size. The genetic correlations were 318 reflected in widespread pleiotropic effects of QTL for fecundity and survival, with allelic effects typically in 319 the same directions. Below we discuss the results in relation to processes affecting adaptive differentiation and pleiotropic interactions among traits. 321

Adaptive differentiation for seed number per fruit

The contribution of variation in seeds/fruit to adaptive differentiation varied between sites. In both years in Italy, the local ecotype produced more seeds/fruit than did the non-local ecotype, and had a greater overall fitness advantage when fitness was estimated including information about seeds/fruit (figure 1A). However,

because selection through seeds/fruit was not as strong as selection through fruits/RP or survival (figure 326 1A), and because seeds/fruit was positively correlated with fruits/RP (figure 1C), much of the adaptive 327 differentiation measured as seeds/seedling in Italy would have been captured by measuring fruits/seedling 328 only. This is consistent with observations from similar experiments using Arabidopsis lyrata (Leinonen et 329 al. 2011) and Hordeum spontaneum (Verhoeven et al. 2004) that found that variation in fruit production 330 makes a greater contribution to adaptive differentiation than does seeds/fruit, but contrasts with those in 331 Mimulus quttatus where the opposite pattern was observed (Hall & Willis 2006). At the Italian site, the 332 marked difference in flowering time of the two ecotypes coupled with strong selection for early flowering can 333 explain much of the large local advantage in terms of fruit production (Ågren et al. 2017). Compared to the 334 non-local Swedish ecotype, the local ecotype began to flower about 1.5 months earlier and produced 5.3 and 335 3.2 times more fruits before the onset of summer in the two years of study (Ågren et al. 2013, 2017). Despite strong selection through seeds/fruit in Italy, differences in fruit production makes a much larger contribution 337 to local adaptation at this site.

In Sweden, by contrast, differences in seeds/fruit made a significant contribution to local adaptation, but this 339 contribution varied among years. In 2010, seeds/fruit was the only fitness component for which the parental 340 ecotypes differed (figure 1B) and showed high heritability among the RILs (figure S8). This was reflected in a significant overall advantage to the local ecotype that year (figure 1B), as well as the detection of four QTL 342 for overall fitness that were detected only when this component was included in the estimate of overall fitness (figure 2). In that experiment, plants experienced a mild winter and heavy damage from rodents (Ågren 344 & Schemske 2012). However, in the 2011 experiment, local adaptation was expressed only as a difference in survival, and no selection through seeds/fruit was detected. By comparison, no significant differences in 346 fruits/RP between the two ecotypes were detected in either year, nor in four additional years studied by 347 Ågren & Schemske (2012). This suggests that selection through fruits/RP in Sweden is weak, and consistently 348 weak across years, which at least partly can be attributed to the modest difference in flowering time between 349 the two ecotypes at this site (3 and 9 days in the years of study; Ågren et al. 2017). These findings highlight that the relative importance of individual fitness components for local adaptation can fluctuate from year to 351 year, and also the value of experiments conducted over multiple seasons.

Including information on seed production into estimates of overall fitness allowed us to detect several QTL
that were not detected when fitness is estimated based only on fruit production and survival (figure 2). In four
of the seven cases where a QTL was detected for seeds/seedling but not fruits/seedling the QTL co-localised
with both a QTL for fruits/RP and a QTL for seeds/fruit in the same year. In one case it co-localised only

with a QTL for fruits/RP and in two cases only with a QTL for seeds/fruit. This indicates that that QTL for 357 seeds/seedling act through both seeds/fruit and fruits/RP, and that the gain in power to detect additional QTL is due to a refinement of the phenotype that allows QTL close to the threshold of significance to be 359 detected. In three of the four cases where QTL for overall fitness co-localised with QTL for both fruits/RP and QTL for seeds/fruit, the direction of effect was the same at the two component QTL, which should 361 have facilitated the detection of the QTL for overall fitness. The exception to this pattern is the QTL for seeds/seedling at the end of chromosome 1 detected in Sweden in 2010 (75.4cm; figure 2), where the Swedish 363 allele was associated with higher overall fitness and an increase in seeds/fruit, but a decrease in fruits/RP. This discrepancy in the direction of allelic effects of the component QTL indicates that this QTL for fitness 365 was detected due to the strong effects via seeds/fruit. Despite this example of a genetic trade-off between 366 components of fecundity, the results suggest that many QTL for seeds/seedling affect fruit and seed production in the same direction. 368

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Surprisingly, some QTL for fitness estimated as fruits/seedling were not detected when fitness was estimated as seeds/seedling (indicated by '-' in figure 2). One explanation for this could be that QTL have weakly negative pleiotropic effects on one or more combinations of seeds/fruit, fruits/RP and survival. However, with the exception of Q4 in Sweden, wherever pleiotropy for seeds/fruit and fruits/RP was observed, the effects were positive (figure 2), so this seems unlikely. Alternatively, our estimates of seeds/fruit might be less precise than for fruits/RP, which would inflate the residual variance of our estimates of seeds/seedling. Although sample sizes for seeds/fruit and seed mass were much smaller than for survival and fruit production, this was not reflected in reduced heritability of seed traits (figure S8), so this explanation also appears unlikely. A third explanation is that there are many loci affecting fitness, many of which have effects close to the threshold of statistical significance. This would be consistent with classical population-genetic theory, which posits that mutations with small effects on fitness dominate as populations get closer towards a local fitness optimum (Fisher 1930; Kimura 1983; Orr 1998). Such subtle effects would be sensitive to the precise way in which fitness is defined, as well as to fluctuations in environmental noise. Because linkage mapping is designed to detect relatively few loci of large effect (Beavis 1998; Xu 2003), this would cause some stochasticity in the loci detected and their map positions indicated by the stepwise-regression approach used in QTL mapping (Harrell 2001; Broman & Šen 2009). Consistent with this, the QTL that differed in models for fruits/seedling and seeds/seedling tended to have broader credible intervals around the estimates of their location (figure 2) and show the weakest allelic effects (tables S3, S4). The apparent disappearance of fitness QTL when information on seed number is included could thus reflect a highly polygenic nature of fitness.

Positive pleiotropic effects on multiple fitness components

Both genetic correlations in the RIL population and the QTL mapping showed evidence of positive pleiotropy between different components of fitness. Correlations between components of fecundity, and between fecundity 390 and survival were positive, except in Sweden in 2010 when no significant correlation was observed between 391 fecundity and survival (figure 1D). Moreover, QTL for these components of fitness tended to map to the same 392 regions of the genome, and allelic effects were in the same direction in all cases, except for the fruits/RP 393 and seeds/fruit QTL at the end of chromosome 1 in Sweden (figure 2). These regions did not correspond 394 to those of two QTL with pleiotropic effects on fruit number and survival identified in a panel of natural 395 accessions tested in outdoor common-garden experiments, although pleiotropic region Q1 overlapped with a 396 SNP associated with fruit production in that study (Fournier-Level et al. 2011). Taken together, the overall positive genetic correlations among fitness components were thus reflected in positive pleiotropic effects of the 398 underlying genetic loci in the present cross.

In addition, we found that the epistatic interaction detected in Sweden 2011 between Q3 and Q6 for seeds/fruit, fruits/RP and both measures of fitness was "positively" pleiotropic, that is, the rank order of two-locusgenotype fitness estimates were the same across phenotypes (figures S1, S6, S7). Counter to what one would expect under local adaptation, plants with Swedish genotypes at both loci had lower fitness than plants with recombinant genotypes. This epistatic interaction should contribute to the transgressive variation observed in fitness at the Swedish site (Ågren et al. 2013), and suggests that not only additive effects but also epistatic interactions among local alleles may be maladaptive. Possible causes of such maladaptive effects include higher genetic load in the Swedish population, lagging adaptation, and temporal variation in selection (Ågren et al. 2013).

The preponderance of positive genetic correlations and positive pleiotropy indicates that variation in overall condition, where the fittest genotypes have more resources overall to invest across fitness components, 410 predominates over variation in relative allocation to different components of fitness in the RIL population 411 (Van Noordwijk & Jong 1986; Houle 1991). The greater condition, and thus higher fitness, of local genotypes 412 likely reflects adaptive differentiation via many traits. For example, the Swedish ecotype has higher freezing 413 tolerance (Oakley et al. 2014) and a greater ability to optimize photosynthesis at cold, but non-freezing 414 temperatures (Cohu et al. 2013; Adams et al. 2014; Oakley et al. 2017). This should provide a fitness 415 advantage at the Swedish site where plants are exposed to cold conditions for an extended period, but may be associated with a fitness cost in the milder climate at the Italian site (Oakley et al. 2014). Consistent with 417

this hypothesis, four of the seven QTL for freezing tolerance identified by Oakley *et al.* (2014) are located in pleiotropic regions identified here (Q4, Q5, Q9, Q11). QTL affecting these traits can thus be expected to have pleiotropic effects on overall fitness and its components.

QTL for phenological variation may also play a key role in stress tolerance and resource acquisition that 421 would contribute to variation in condition. Previous work on the same populations has demonstrated strong 422 selection to tune the timing of germination (Akiyama & Ågren 2014; Postma & Ågren 2016; Zacchello et al. 423 2020) and flowering time to match the local climate (Ågren et al. 2017). The timing of life history traits can have cascading effects on later life-history stages (Lindström 1999; Beckerman et al. 2002; Donohue 2014; 425 Postma & Ågren 2016; Hepworth et al. 2020; Martinez-Berdeja et al. 2020), meaning that direct effects on early traits can cause pleiotropic effects on multiple fitness components. Consistent with this, all pleiotropic 427 regions bar Q1 and Q2 identified here also harbour well-resolved QTL associated with flowering time at 428 one or both sites (Ågren et al. 2017), and loci such as FT (Q3), FLC (Q9) and VIN3 (Q11) with effects 429 on flowering time documented in field experiments (e.g. Caicedo et al. 2004; Korves et al. 2007; Taylor et 430 al. 2019; Hepworth et al. 2020). Furthermore, Q10 overlaps with the primary QTL explaining variation 431 in seed dormancy, which contains the candidate gene DOG1 (Postma & Agren 2016). QTL affecting both 432 physiological and phenological traits can thus be expected to influence condition and thereby have pleiotropic 433 effects on multiple components of fitness. 434

The role of variation in condition in local adaptation is a broadly important topic in evolutionary biology. In 435 a meta-analysis of local adaptation across 74 studies of plants, animals, fungi and protists, Hereford (2009) 436 was not able to test explicitly for correlations between selection through components of fitness, but did 437 demonstrate that the advantage of local populations was greater when estimated based on overall fitness 438 than when estimated based on survival or fecundity alone. Although not a formal test, this pattern would be 439 expected if selection acts to increase overall condition, causing components of fitness to be positively correlated within populations. These observations indicate that adaptation frequently entails increased condition in the 441 local environment, and that while local adaptation is reflected as trade-offs in performance across environments, it may often also be associated with positive genetic correlations among fitness components within a given 443 environment in situations where there is genetic variation for overall fitness.

Limited evidence for a trade-off between offspring size and number

There was substantial overlap between the positions of seed mass QTL detected in the present study and those of QTL that have previously been identified as affecting seed size. On one hand, Ren et al. (2019) found that the genetic architecture of seed mass in a panel of natural accession was characterised by many loci of very 448 small effect spread across the genome, and the single large-effect QTL detected does not correspond to any 449 locus found in this study. On the other hand, using a RIL population derived from a cross between the Ler450 and Cvi accessions, Alonso-Blanco et al. (1999) found QTL for seed mass in positions close to those of Q1, Q2 451 and Q8. Furthermore, Gnan et al. (2014) found QTL for seed size that co-localise with Q8, Q9, Q10 and Q11 452 in an intercross population derived from 16 natural accessions. Finally, using a panel of knock-out mutant 453 lines, Van Daele et al. (2012) identified thirteen candidate genes affecting seed mass, seven of which are found within the pleiotropic regions affecting seed mass identified here (DWARF11 in Q1, GW2 in Q1, CKX1 in Q2, 455 ANT in Q8, GASA4 in Q9, CKX3 in Q11, and ARF2 in Q11). These genes are mostly transcription factors involved in cytokinin metabolism (Orozco-Arroyo et al. 2015). The overlap between these studies indicates 457 that alleles influencing seed mass are segregating in A. thaliana populations beyond the cross examined here. 458 Even if QTL affecting seed mass are common, three observations indicate that selection on seed mass does not constrain the evolution of increased fecundity (seeds/RP). Firstly, differences in seed mass between the 460 parents were small to modest. The greatest difference we observed was in Italy in 2010, where the local 461 genotype had 18% larger seeds than the Swedish genotype (figure S2). For comparison, Alonso-Blanco et 462 al. (1999) found an 81% difference in seed mass between the Ler and Cvi accessions. However, we cannot 463 exclude the possibility that variation in seed mass does contribute to differences in fitness via the earliest life-history stages, as has been shown under laboratory conditions for seedling survival (Krannitz et al. 1991) 465 and early growth rate (El-Lithy et al. 2004) in A. thaliana. The present experiment was started with seedlings, precluding the detection of effects of seed size on germination rate and seedling establishment, and reducing 467 the chance of observing effects of these life-history stages on overall fitness. Secondly, there was little evidence of a consistent genetically based trade-off between seed size and fecundity 469 in the RIL population. Although seed mass and seeds/RP were negatively genetically correlated in Sweden in 470 2010, no significant genetic correlation was detected in the other three site × year combinations (figures 1C, 471 1D and S4). This was also true for correlations between seed mass and seeds/fruit. Previous studies have 472 documented a negative correlation between seed size and seeds/fruit in mapping populations grown under controlled conditions and derived from crosses between other sets of A. thaliana accessions (Alonso-Blanco

et al. 1999; Gnan et al. 2014). The negative correlation found in Sweden in 2010 shows that there are 475 circumstances under which a genetic trade-off between seed size and number, albeit weak, can be expressed in 476 the cross examined here. As previously noted, the plants in that experiment experienced an especially mild 477 winter and the difference in fitness between the two parental genotypes was unusually small, so the expression of a trade-off may reflect reduced variation in condition compared to the other three site x year combinations. 479 The results suggest that correlations between seed size and measures of fecundity may vary among crosses, and highlight the importance of field studies in understanding how the sign and magnitude of correlations 481 between fitness components affect local adaptation. 482 Thirdly, if the correlation between fecundity and seed size constrains the evolution of increased fecundity, we would expect to see trade-offs reflected in the directions of allelic effects at pleiotropic QTL affecting both 484 seed mass and fecundity. In fact, although QTL for seed mass often co-localised with QTL for fecundity and seeds/fruit, pleiotropic effects on these two traits were as likely to be positive as antagonistic (figure 486 2), which is consistent with the weak overall genetic correlation between the two traits. In contrast, only 487 one of the ten seed mass QTL detected in the mapping population studied by Alonso-Blanco et al. (1999), and one of the seed mass QTL documented by Gnan et al. (2014) showed evidence of a pleiotropic effect 489 on seeds/fruit, and in both cases effects were antagonistic. Moreover, when single-gene-knock-out mutants 490 showed pleiotropic effects on seed size and seeds/fruit, the effects were always antagonistic (Van Daele et al. 491 2012). Further crosses to generate additional recombination are required to determine conclusively whether the co-localising QTL for seed size and number documented in the present study reflect pleiotropic loci, or 493 distinct non-pleiotropic loci that are too close to each other to be resolved in this RIL population. In summary, the lack of consistent differences between parents, weak correlations between seed size and fecundity, and 495 the limited evidence for widespread antagonistic pleiotropy for QTL affecting seed size and number do not

498 Conclusions

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In conclusion, this study has examined how variation in number of seeds per fruit and other components of fitness contribute to overall adaptive differentiation in *A. thaliana*. Our results show that there is adaptive variation in seed production independent of variation in fruit number, and that the advantage to local genotypes can be underestimated if this is ignored. Moreover, we demonstrate consistent positive pleiotropy among components of fitness reflected in both genetic correlations among phenotypes and effects of underlying

support a role for variation in seed mass in local adaptation between the two parental populations.

QTL, and very little evidence of a trade-off between offspring size and number. These findings indicate that
the process of population divergence has been due in large part to the fixation of alleles that increase overall
vigour or condition at each site.

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Data availability

Data, R scripts, and the R markdown document used to create this manuscript will be uploaded to a suitable public server on publication. In the meantime, they are available at https://github.com/ellisztamas/fecundity_components.

519 Competing interests

The authors declare no conflict of interest.

Authors' contributions

TJE performed analyses and wrote the manuscript. JÅ conceived the study, and JÅ, FMP and CGO co-ordinated data collection and critically revised the manuscript.

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553 Supplementary material

⁶⁵⁴ Supplementary tables

Table S1: QTL for number of seeds per fruit identifed in individual experiments. QTL positions are shown with their 95% Bayesian credible intervals. Effect sizes are given as additive effects and as a percentage of variance among RIL means. Effect sizes are quantified as the least-square mean difference in seeds/fruit between genotypes homozygous for the Swedish and Italian allele, respectively; negative values indicate that the Swedish genotype is associated with fewer seeds/fruit.

Name	Site	Year	Chr.	Position (cM)	LOD	Effect size	% var. explained
1@9.5	Italy	2010	1	9.5 (8.4-12.6)	9.3	-1.431	8.7
2@56.0	Italy	2010	2	56.0 (44.3-60.9)	3.9	-0.906	3.5
3@18.0	Italy	2010	3	18.0 (14.7-19.7)	5.9	-1.130	5.4
5@56.6	Italy	2010	5	56.6 (49.9-58.6)	6.9	-1.211	6.3
1@58.6	Italy	2011	1	58.6 (51.5-61.8)	4.7	-1.071	4.0
2@32.6	Italy	2011	2	32.6 (26.8-45.6)	6.3	-1.246	5.3
5@15.0	Italy	2011	5	15.0 (9.4-22.0)	4.0	-1.001	3.4
5@50.2	Italy	2011	5	50.2 (49.9-60.1)	13.8	-1.924	12.3
1@22.7	Sweden	2010	1	22.7 (20.8-23.7)	4.9	0.815	4.4
1@75.4	Sweden	2010	1	75.4 (73.2-79.6)	7.3	1.334	6.6
4@41.0	Sweden	2010	4	41.0 (41.0-42.3)	9.2	1.508	8.4
5@70.6	Sweden	2010	5	70.6 (68.7-78.2)	7.1	1.093	6.4
1@61.1	Sweden	2011	1	61.1 (58.8-61.1)	9.9	-0.819	8.7
1@77.9	Sweden	2011	1	77.9 (73.2-82.1)	2.5	0.909	2.1
3@21.0	Sweden	2011	3	21.0 (21.0-23.3)	12.8	-1.332	11.5
4@50.2	Sweden	2011	4	50.2 (35.8-57.7)	6.9	1.380	6.0
5@74.5	Sweden	2011	5	74.5 (70.1-77.4)	9.3	1.597	8.2

Table S2: Pairs of significant epistatic QTL detected across traits.

Trait	Site	Year	QTL	LOD	% var. explained
Fruits/RP	Italy	2010	5@7.5 x 5@70.1	0.55	0.35
Fruits/RP	Italy	2011	$1@58.6 \times 5@74.5$	6.42	3.33
Fruits/RP	Sweden	2011	$1@61.8 \times 3@26.7$	8.43	6.93
Seeds/fruit	Sweden	2010	$1@22.7 \pm 5@70.6$	2.38	2.08
Seeds/fruit	Sweden	2011	1@61.1 x 3@21.0	7.41	6.45
Seeds/RP	Italy	2010	$5@9.4 \times 5@71.4$	1.19	0.85
Seeds/RP	Sweden	2011	1@61.1 x 3@24.4	10.11	8.59
Seeds/seedling	Italy	2010	$5@9.4 \times 5@71.4$	0.77	0.48
Seeds/seedling	Sweden	2011	1@61.1 x 3@24.4	9.07	6.49
Seed mass	Sweden	2011	3@55.4 x 5@78.2	3.54	3.00

Table S3: QTL for number of fruits per seedling identified in individual experiments. QTL positions are shown with their 95% Bayesian credible intervals. Effect sizes are given as additive effects, and as a percentage of variance among RIL means. Effect sizes are quantified as the least-square mean difference in fruits/seedling between genotypes homozygous for the Swedish and Italian allele, respectively; negative values indicate that the Swedish genotype is associated with fewer fruits/seedling.

Name	Site	Year	Chr.	Position (cM)	LOD	Effect size	% var. explained
1@10.3	Italy	2010	1	10.3 (8.4-12.0)	9.9	-0.774	6.2
1@57.3	Italy	2010	1	57.3 (55.9-57.3)	11.1	-0.823	7.0
2@53.4	Italy	2010	2	53.4 (52.5-56.0)	7.6	-0.664	4.7
3@54.1	Italy	2010	3	54.1 (53.3-59.2)	7.2	-0.646	4.4
4@50.0	Italy	2010	4	50.0 (43.5-57.7)	4.4	-0.541	2.6
5@7.5	Italy	2010	5	7.5 (2.6-10.0)	6.9	-0.677	4.2
5@60.6	Italy	2010	5	60.6 (53.4-70.1)	2.9	-0.465	1.7
5@78.2	Italy	2010	5	78.2 (70.1-78.2)	4.5	-0.583	2.8
1@61.8	Italy	2011	1	61.8 (60.2-61.8)	13.6	-0.761	6.8
1@77.2	Italy	2011	1	77.2 (77.0-83.8)	4.4	-0.421	2.1
2@49.5	Italy	2011	2	49.5 (48.7-50.6)	14.3	-0.675	7.2
3@1.7	Italy	2011	3	1.7 (0.0- 8.2)	3.6	0.332	1.7
3@64.9	Italy	2011	3	64.9 (63.1-65.9)	16.3	-0.727	8.3
4@55.2	Italy	2011	4	55.2 (51.1-56.6)	10.6	-0.593	5.2
5@56.6	Italy	2011	5	56.6 (56.6-57.7)	8.3	-0.622	4.0
5@70.1	Italy	2011	5	70.1 (70.1-78.2)	5.6	-0.514	2.7
3@10.7	Sweden	2010	3	10.7 (8.2-34.4)	2.5	0.296	2.9
1@62.5	Sweden	2011	1	62.5 (61.0-62.5)	13.1	-0.646	9.4
2@58.6	Sweden	2011	2	58.6 (32.6-60.9)	3.0	0.608	2.0
3@29.5	Sweden	2011	3	29.5 (29.5-30.5)	12.1	-0.725	8.6
3@66.6	Sweden	2011	3	66.6 (36.1-70.5)	2.2	0.522	1.5
4@53.5	Sweden	2011	4	53.5 (50.0-56.6)	7.9	1.046	5.5
5@1.4	Sweden	2011	5	1.4 (1.4- 4.6)	4.7	0.807	3.2
5@27.1	Sweden	2011	5	27.1 (20.4-34.1)	2.7	0.616	1.8
5@60.6	Sweden	2011	5	60.6 (56.6-62.6)	3.0	-0.740	2.0

Name	Site	Year	Chr.	Position (cM)	LOD	Effect size	% var. explained
5@74.5	Sweden	2011	5	74.5 (74.5-74.5)	18.6	1.892	13.8

Table S4: QTL for number of seeds per seedling identified in individual experiments. QTL positions are shown with their 95% Bayesian credible intervals. Effect sizes are given as additive effects and as a percentage of variance among RIL means. Effect sizes are quantified as the least-square mean difference in seeds/seedling between genotypes homozygous for the Swedish and Italian allele, respectively; negative values indicate that the Swedish genotype is associated with fewer seeds/seedling.

Name	Site	Year	Chr.	Position (cM)	LOD	Effect size	% var. explained
1@10.3	Italy	2010	1	10.3 (7.3-12.0)	12.5	-33.121	8.3
1@57.3	Italy	2010	1	57.3 (55.9-61.1)	9.4	-28.474	6.2
2@53.4	Italy	2010	2	53.4 (51.0-56.0)	7.2	-24.485	4.7
3@19.7	Italy	2010	3	19.7 (14.7-30.5)	1.2	-10.411	0.8
3@55.4	Italy	2010	3	55.4 (41.1-66.8)	3.0	-16.202	1.9
4@20.7	Italy	2010	4	20.7 (17.1-44.3)	2.8	-15.534	1.8
5@9.4	Italy	2010	5	9.4 (4.6-12.0)	5.9	-20.793	3.8
5@58.6	Italy	2010	5	58.6 (54.3-61.3)	3.5	-20.923	2.2
5@71.4	Italy	2010	5	71.4 (68.7-73.6)	2.9	-15.678	1.9
1@61.1	Italy	2011	1	61.1 (57.3-61.8)	12.5	-28.327	6.7
1@80.9	Italy	2011	1	80.9 (77.0-83.8)	3.8	-15.287	1.9
2@31.7	Italy	2011	2	31.7 (25.6-40.3)	3.1	-13.873	1.6
2@49.4	Italy	2011	2	49.4 (45.6-51.0)	7.5	-21.523	3.9
3@64.9	Italy	2011	3	64.9 (63.1-65.9)	7.7	-19.639	4.0
4@52.0	Italy	2011	4	52.0 (50.0-57.7)	4.4	-15.495	2.2
5@18.5	Italy	2011	5	18.5 (1.4-43.0)	2.7	-11.992	1.3
5@56.6	Italy	2011	5	56.6 (56.6-57.7)	11.7	-29.901	6.2
5@72.1	Italy	2011	5	72.1 (69.4-78.2)	4.7	-18.419	2.4
1@13.5	Sweden	2010	1	13.5 (0.0-20.8)	2.5	-29.376	2.4
1@22.7	Sweden	2010	1	22.7 (21.9-22.7)	5.7	45.258	5.7
1@75.4	Sweden	2010	1	75.4 (49.8-82.1)	3.1	24.579	3.1
4@41.0	Sweden	2010	4	41.0 (37.4-42.3)	6.0	34.597	6.1
1@61.1	Sweden	2011	1	61.1 (61.0-61.8)	12.1	-20.792	8.8
3@24.4	Sweden	2011	3	24.4 (22.3-25.4)	15.5	-42.447	11.5
3@66.6	Sweden	2011	3	66.6 (59.2-70.5)	2.9	25.576	2.0
4@50.2	Sweden	2011	4	50.2 (41.0-56.6)	9.0	46.843	6.4

Name	Site	Year	Chr.	Position (cM)	LOD	Effect size	% var. explained
5@1.4	Sweden	2011	5	1.4 (0.0-23.7)	5.3	34.857	3.7
5@74.5	Sweden	2011	5	74.5 (74.5-75.8)	18.7	67.112	14.2

Table S5: QTL for seed mass identifed in individual experiments. QTL positions are shown with their 95% Bayesian credible intervals. Effect sizes are given as additive effects and as a percentage of variance among RIL means. Effect sizes are quantified as the least-square mean difference in seed mass between genotypes homozygous for the Swedish and Italian allele, respectively; negative values indicate that the Swedish genotype is associated with lower seed mass.

Name	Site	Year	Chr.	Position (cM)	LOD	Effect size	% var. explained
1@48.9	Italy	2010	1	48.9 (40.6-61.1)	3.3	-0.570	3.5
3@57.1	Italy	2010	3	57.1 (50.2-59.2)	4.7	-0.688	5.2
1@16.1	Italy	2011	1	16.1 (7.3-18.9)	3.7	0.768	3.7
1@29.5	Italy	2011	1	29.5 (22.7-61.8)	3.7	-0.765	3.7
3@53.3	Italy	2011	3	53.3 (18.0-60.4)	4.0	-0.655	4.0
4@9.3	Italy	2011	4	9.3 (4.2-28.0)	3.5	-0.612	3.5
5@7.7	Italy	2011	5	7.7 (5.7-13.6)	3.6	0.619	3.6
1@3.9	Sweden	2010	1	3.9 (0.0- 6.1)	8.9	0.749	8.1
1@32.4	Sweden	2010	1	32.4 (20.8-43.5)	3.4	-0.454	3.0
2@58.0	Sweden	2010	2	58.0 (50.4-60.9)	3.1	-0.413	2.7
3@65.9	Sweden	2010	3	65.9 (45.9-70.5)	4.8	-0.514	4.3
5@7.7	Sweden	2010	5	7.7 (2.6- 7.7)	11.4	0.821	10.6
1@11.5	Sweden	2011	1	11.5 (9.9-12.6)	5.4	0.841	4.6
1@21.9	Sweden	2011	1	21.9 (19.6-22.7)	5.1	-0.820	4.4
3@55.4	Sweden	2011	3	55.4 (49.1-60.4)	5.1	-0.360	4.3
4@15.8	Sweden	2011	4	15.8 (14.4-28.0)	3.6	-0.534	3.0
4@57.7	Sweden	2011	4	57.7 (54.5-60.7)	3.2	0.516	2.7
5@2.6	Sweden	2011	5	2.6 (1.4- 7.7)	10.3	0.930	9.1
5@52.4	Sweden	2011	5	52.4 (6.7-71.4)	2.8	0.496	2.4
5@78.2	Sweden	2011	5	78.2 (72.1-78.2)	4.7	0.282	4.0

Table S6: Pearson correlations between seed mass and number of seeds per fruit.

Site	Year	r	Deg. freedom	p-value
Italy	2010	0.02	393	0.655
Italy	2011	0.02	396	0.761
Sweden	2010	-0.19	393	0.000
Sweden	2011	-0.06	392	0.266

Table S7: QTL for survival to reproduction identified in individual experiments. QTL positions are shown with their 95% Bayesian credible intervals. Effect sizes are given as additive effects, as a percentage of variance among RIL means, and as a percentage of the difference between parental means. Effect sizes are quantified as the least-square mean difference in survival between genotypes homozygous for the Swedish and Italian allele, respectively; negative values indicate that the Swedish genotype is associated with lower survival.

Name	Site	Year	Chr.	Position (cM)	LOD	Effect size	% var. explained
1@40.6	Italy	2010	1	40.6 (2.1-45.1)	2.9	-0.024	2.1
1@62.5	Italy	2010	1	62.5 (58.6-64.9)	5.5	-0.033	3.9
2@53.4	Italy	2010	2	53.4 (49.4-55.3)	12.6	-0.046	9.3
3@59.2	Italy	2010	3	59.2 (37.9-65.9)	4.6	-0.027	3.2
5@9.4	Italy	2010	5	9.4 (6.7-10.0)	10.1	-0.041	7.3
5@66.1	Italy	2010	5	66.1 (64.1-77.4)	9.1	-0.039	6.6
1@40.6	Italy	2011	1	40.6 (36.7-51.5)	6.7	-0.027	5.8
1@83.8	Italy	2011	1	83.8 (74.4-83.8)	4.8	-0.023	4.1
2@49.5	Italy	2011	2	49.5 (44.3-51.0)	10.0	-0.033	9.0
5@57.5	Italy	2011	5	57.5 (53.4-60.1)	9.2	-0.032	8.1
4@53.5	Sweden	2011	4	53.5 (39.1-56.6)	6.2	0.037	5.1
5@1.4	Sweden	2011	5	1.4 (0.0- 1.4)	10.8	0.049	9.2
5@73.6	Sweden	2011	5	73.6 (73.6-75.8)	17.1	0.061	15.0

Table S8: QTL for number of fruits per reproductive plant identifed in individual experiments. QTL positions are shown with their 95% Bayesian credible intervals. Effect sizes are given as additive effects and as a percentage of variance among RIL means. Effect sizes are quantified as the least-square mean difference in fruits/RP between genotypes homozygous for the Swedish and Italian allele, respectively; negative values indicate that the Swedish genotype is associated with fewer fruits/RP.

Name	Site	Year	Chr.	Position (cM)	LOD	Effect size	% var. explained
1@10.3	Italy	2010	1	10.3 (8.8-12.0)	13.2	-1.576	9.0
1@57.3	Italy	2010	1	57.3 (55.9-57.3)	9.4	-1.312	6.2
2@53.4	Italy	2010	2	53.4 (48.7-56.4)	2.8	-0.700	1.8
3@8.2	Italy	2010	3	8.2 (3.5- 9.6)	4.7	1.179	3.0
3@19.7	Italy	2010	3	19.7 (18.0-22.3)	2.2	-0.846	1.4
3@54.1	Italy	2010	3	54.1 (49.1-65.5)	4.3	-0.888	2.7
4@18.3	Italy	2010	4	18.3 (14.4-31.0)	2.6	-0.697	1.7
4@50.6	Italy	2010	4	50.6 (44.3-55.9)	4.3	-0.967	2.8
5@7.5	Italy	2010	5	7.5 (4.9-12.0)	4.9	-0.914	3.2
5@70.1	Italy	2010	5	70.1 (68.7-72.1)	7.3	-1.060	4.8
1@58.6	Italy	2011	1	58.6 (57.3-61.1)	15.1	-0.916	8.2
1@77.2	Italy	2011	1	77.2 (77.0-83.8)	4.8	-0.597	2.5
2@38.2	Italy	2011	2	38.2 (32.0-40.3)	8.1	-0.689	4.2
3@5.1	Italy	2011	3	5.1 (1.7- 7.3)	5.4	0.562	2.8
3@57.1	Italy	2011	3	57.1 (54.7-65.9)	13.3	-0.903	7.2
4@53.9	Italy	2011	4	53.9 (51.3-55.5)	12.2	-0.878	6.6
5@56.6	Italy	2011	5	56.6 (53.4-58.6)	4.9	-0.616	2.5
5@74.5	Italy	2011	5	74.5 (72.9-77.4)	12.5	-0.676	6.7
1@13.5	Sweden	2010	1	13.5 (3.9-20.8)	3.4	-3.414	3.6
1@22.7	Sweden	2010	1	22.7 (21.9-55.6)	3.7	3.559	3.8
1@77.9	Sweden	2010	1	77.9 (74.4-83.8)	3.3	-2.487	3.4
2@58.0	Sweden	2010	2	58.0 (49.7-60.9)	2.1	1.956	2.2
3@9.6	Sweden	2010	3	9.6 (8.2-42.6)	2.4	2.115	2.5
1@61.8	Sweden	2011	1	61.8 (61.0-62.5)	13.4	-1.934	11.3
2@58.3	Sweden	2011	2	58.3 (50.4-60.9)	4.2	2.065	3.4
3@26.7	Sweden	2011	3	26.7 (25.4-30.5)	10.7	-1.754	8.9

Name	Site	Year	Chr.	Position (cM)	LOD	Effect size	% var. explained
4@55.5	Sweden	2011	4	55.5 (37.4-56.6)	4.1	2.073	3.3
5@20.4	Sweden	2011	5	20.4 (15.0-34.1)	4.7	2.186	3.8
5@60.6	Sweden	2011	5	60.6 (50.2-64.1)	3.3	-2.166	2.7
5@74.5	Sweden	2011	5	74.5 (74.5-75.8)	11.0	4.043	9.2

Table S9: QTL for number of seeds per reproductive plant identifed in individual experiments. QTL positions are shown with their 95% Bayesian credible intervals. Effect sizes are given as additive effects and as a percentage of variance among RIL means. Effect sizes are quantified as the least-square mean difference in seeds/RP between genotypes homozygous for the Swedish and Italian allele, respectively; negative values indicate that the Swedish genotype is associated with fewer seeds/RP.

Name	Site	Year	Chr.	Position (cM)	LOD	Effect size	% var. explained
1@10.3	Italy	2010	1	10.3 (8.4-10.3)	12.2	-58.194	9.3
1@57.3	Italy	2010	1	57.3 (55.2-57.3)	7.8	-45.710	5.7
2@53.4	Italy	2010	2	53.4 (48.7-56.4)	3.2	-28.643	2.3
3@54.1	Italy	2010	3	54.1 (18.0-60.4)	4.2	-32.925	3.1
4@31.0	Italy	2010	4	31.0 (18.3-32.1)	5.6	-39.584	4.1
5@9.4	Italy	2010	5	9.4 (2.6-12.0)	4.5	-28.700	3.3
5@71.4	Italy	2010	5	71.4 (70.1-73.6)	7.9	-39.604	5.8
1@60.2	Italy	2011	1	60.2 (57.3-61.8)	9.3	-31.861	5.1
1@79.6	Italy	2011	1	79.6 (77.0-83.8)	3.5	-19.123	1.9
2@32.6	Italy	2011	2	32.6 (25.6-40.3)	2.9	-17.713	1.5
2@49.4	Italy	2011	2	49.4 (44.3-60.5)	3.5	-19.166	1.8
3@5.1	Italy	2011	3	5.1 (4.5- 8.2)	3.3	23.424	1.8
3@14.7	Italy	2011	3	14.7 (13.5-19.7)	2.0	-18.386	1.0
3@64.9	Italy	2011	3	64.9 (63.1-65.9)	8.0	-26.330	4.4
4@52.8	Italy	2011	4	52.8 (50.0-57.7)	5.0	-21.542	2.7
5@18.5	Italy	2011	5	18.5 (2.6-43.0)	2.8	-16.171	1.5
5@56.6	Italy	2011	5	56.6 (56.6-57.7)	11.3	-35.522	6.3
5@78.2	Italy	2011	5	78.2 (72.1-78.2)	4.8	-22.641	2.6
1@13.5	Sweden	2010	1	13.5 (0.0-20.8)	3.7	-151.237	4.0
1@22.7	Sweden	2010	1	22.7 (21.9-23.7)	5.2	178.597	5.6
3@30.5	Sweden	2010	3	30.5 (9.6-45.9)	3.1	103.634	3.3
1@61.1	Sweden	2011	1	61.1 (61.0-62.5)	12.7	-48.232	11.0
3@24.4	Sweden	2011	3	24.4 (21.0-29.5)	13.1	-74.780	11.3
4@50.2	Sweden	2011	4	50.2 (39.1-57.7)	4.1	80.002	3.4
5@20.4	Sweden	2011	5	20.4 (1.4-25.4)	3.9	76.310	3.2

Name	Site	Year	Chr.	Position (cM)	LOD	Effect size	% var. explained
5@74.5	Sweden	2011	5	74.5 (74.5-75.8)	10.0	122.523	8.5

Supplementary figures

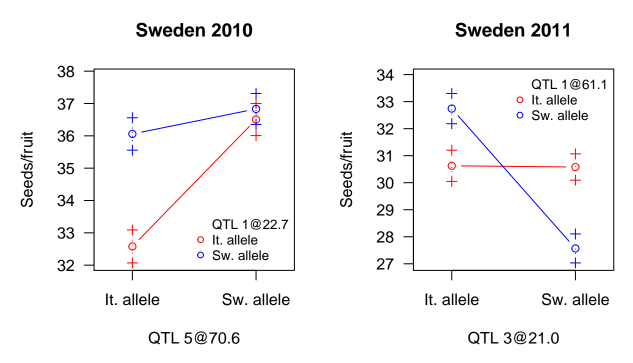


Figure S1: Interaction plots for epistatic interactions detected for number of seeds per fruit, showing mean and standard errors for each genotype-by-genotype combination.

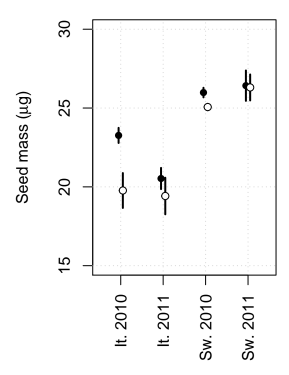


Figure S2: Mean seed mass of the Italian (closed symbols) and Swedish (open symbols) parental ecotypes at the field sites in Italy (It.) and Sweden (Sw.) in 2010 and 2011. Bars indicate standard error.

Sweden 2011

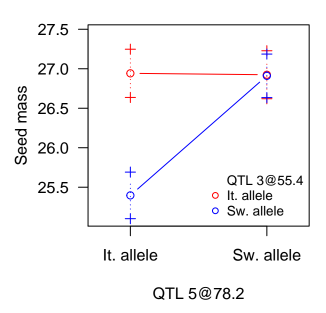


Figure S3: Interaction plots for epistatic interactions detected for seed mass, showing mean and standard errors for each genotype-by-genotype combination.

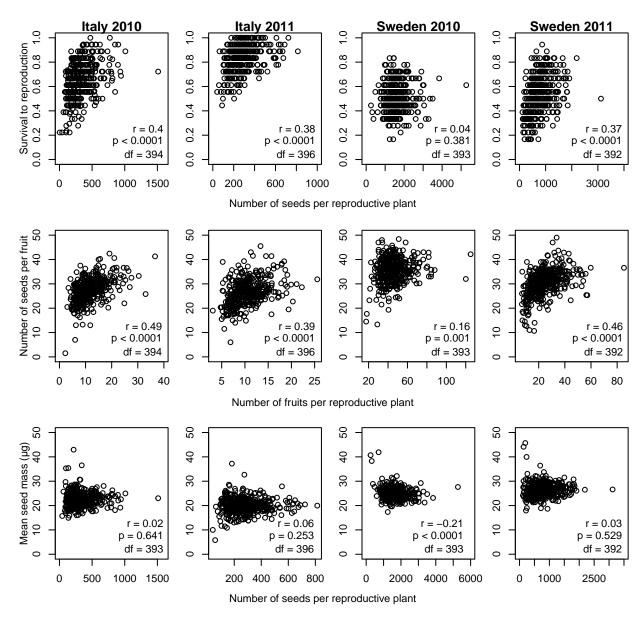


Figure S4: Relationships between RIL-means for survival and fecundity (top), components of fecundity (middle), and between fecundity and seed mass (bottom). Pearson correlation coefficients (r), associated p-values and degrees of freedom (df) are indicated.

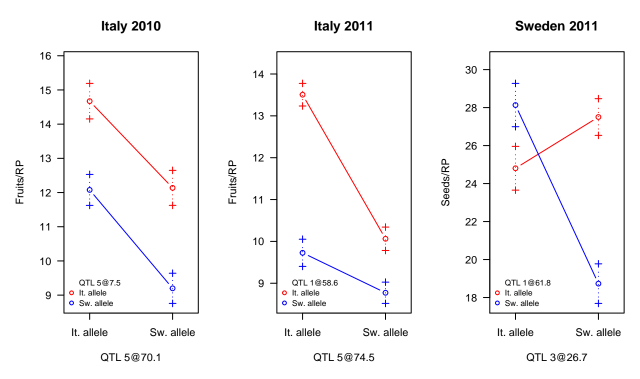


Figure S5: Interaction plots for epistatic interactions detected for number of fruits per reproductive plant (fruits/RP), showing mean and standard errors for each genotype-by-genotype combination.

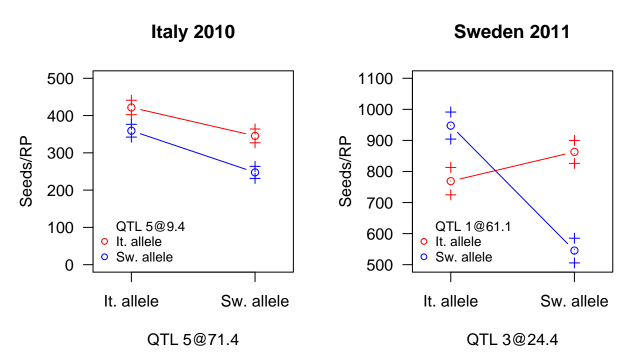


Figure S6: Interaction plots for epistatic interactions detected for number of seeds per reproductive plant, showing mean and standard errors for each genotype-by-genotype combination.

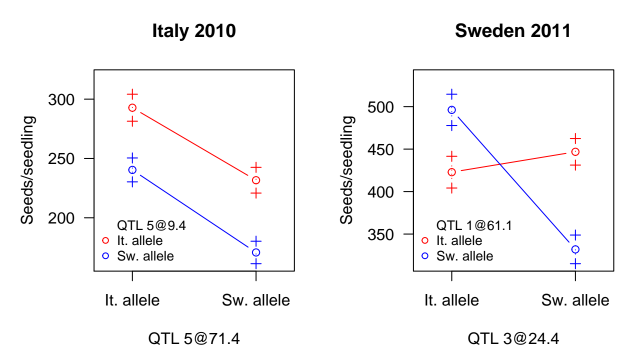


Figure S7: Interaction plots for epistatic interactions detected for number of seeds per seedling, showing mean and standard errors for each genotype-by-genotype combination.

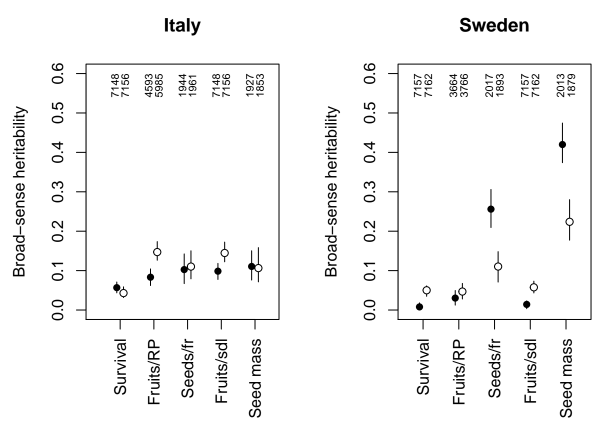


Figure S8: Broad-sense heritability estimates for survival to reproduction, number of fruits per reproductive plants (fruits/RP), number of seeds per fruit (Seeds/fr), number of fruits per seedling (Fruits/sdl), and seed mass. Open and closed symbols show values in 2010 and 2011 respectively. Error bars show 95% parametric bootstrap confidence intervals. Numbers indicate the number of individuals each estimate is based on.