

Prediction of hybrid biomass in *Arabidopsis thaliana* by selected parental SNP and metabolic markers

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Abstract A recombinant inbred line (RIL) population, derived from two *Arabidopsis thaliana* accessions, and the corresponding testcrosses with these two original accessions were used for the development and validation of machine learning models to predict the biomass of hybrids. Genetic and metabolic information of the RILs served as predictors. Feature selection reduced the number of variables (genetic and metabolic markers) in the models by more than 80% without impairing the predictive power. Thus, potential biomarkers have been revealed. Metabolites were shown to bear information on inherited macroscopic phenotypes. This proof of concept could be interesting for breeders. The example population exhibits substantial mid-parent biomass heterosis. The results of feature selection could therefore be used to shed light on the origin of heterosis. In this respect, mainly dominance effects were detected.

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Introduction

The main objective of the work presented in this article is to develop methods which serve to improve the prediction of hybrid properties based on their potential parents, a fundamental aspect in many breeding programs. Today breeders often use genetic information to identify specific lines whose progeny are likely to manifest positive traits (McCouch 2004). The aim is to accelerate the otherwise laborious process of quality assessment and selection.

We focus on the development and validation of machine learning methods designed to improve the prediction of traits of new crosses using molecular data from different sources. Molecular data often are described by a huge number of features, the importance of which for the traits under investigation is generally not known. We present a procedure that combines variables/feature selection with regression and dimensionality reduction. The selected variables serve as potential biomarkers allowing the prognosis of progeny properties.

Several methods have been developed to predict hybrid performance in maize using genetic markers (Maenhout et al. 2009; Reif et al. 2003; Schrag et al. 2007; Schrag et al. 2009a, b; Vuylsteke et al. 2000) or gene expression analysis (Frisch et al. 2009). A combination of genetic markers with morphological characters, isozymes, and proteins was employed for the same purpose in oilseed rape (Yu et al. 2005). We present a proof of concept of a new complementary approach that involves the utilization of metabolite profiles as predictors in addition to SNP markers and the introduction of a new feature selection procedure.

The detection of important markers is closely related to the understanding of the interactions between them and the resulting implications for the progeny traits. The

advancement of this understanding is a further objective of the present study.

The introduction of hybrids was a successful development in plant breeding, especially with respect to yield (Birchler et al. 2003). This is due to the effect called heterosis, which describes the superiority of heterozygous hybrids in comparison to their homozygous parents (Shull 1948). Three hypotheses have been put forward in early studies to explain this phenomenon: the dominance (Bruce 1910; Davenport 1908), overdominance (Crow 1948; Hull 1945), and epistasis hypothesis (Powers 1944; Williams 1959). However, in spite of enormous efforts, the molecular basis of this phenomenon remains largely obscure.

For the validation of our procedure we chose a recombinant inbred line (RIL) population derived from *Arabidopsis* accessions Col-0 and C24 (Törjék et al. 2006) and the heterozygous testcrosses of its lines with both parents. This population meets the following requirements of a validation population: it has been genotyped using SNP markers (Törjék et al. 2006), manifests significant biomass heterosis (Meyer et al. 2009), shows sufficient variance of the trait in the crosses and consists of a sufficient number of genotypes. In addition a large amount of established biochemical knowledge on the population is available (Lisec et al. 2008, 2009; Meyer et al. 2007). Especially, quantitative trait loci (QTL) for biomass per se, biomass heterosis, and metabolites are known. Furthermore, we performed for the present study a QTL analysis for testcross biomass per se. The simple design of the testcross population with one parent being kept constant facilitates both, prediction and interpretation. A further advantage of using *Arabidopsis thaliana* is the existence of comprehensive databases such as AraCyc (www.arabidopsis.org/biocyc/), which contain information about predicted and experimentally determined pathways, reactions, compounds, genes, and enzymes.

We previously developed a procedure to predict mid-parent heterosis from a combination of SNP markers and metabolite profiles of the homozygous population (Gärtner et al. 2009). This multivariate procedure combined regression, dimensionality reduction, and feature selection methods. In the work presented here, we predict directly the biomass of the hybrids, from data obtained from the parents. The predicted trait—in contrast to the mid-parent heterosis—and its predictors are entirely derived from different genotypes allowing for the validation of the method.

Methods and materials

The recombinant inbred line population and testcrosses

The homozygous RIL population was created from a reciprocal cross between the *Arabidopsis thaliana* accessions

C24 and Col-0. F₂ plants were propagated by single seed descent to the F₈ generation. A set of 110 SNP markers served for the genotyping of the RIL population (Törjék et al. 2003; Törjék et al. 2006). In the present study, we included 359 RILs and 718 testcrosses with both parents, for which both genetic and metabolic data were available.

Plant cultivation

All plants were grown together under controlled conditions in 1:1 mixture of GS 90 soil and vermiculite (Gebrüder Patzer, Sinntal-Jossa, Germany), under long-day regime (16 h fluorescent light at 20°C and 60% relative humidity/ 8 h darkness at 18°C and 75% humidity (Lisec et al. 2008)). Six plants of the same line were grown in one pot.

Shoot dry weight

The shoot dry weight was measured 15 days after sowing (DAS). Mean shoot dry weight in milligram per plant was estimated by using the linear mixed model $G + E:E \cdot G + E \cdot GC + E \cdot GC \cdot T$ where E is experiment, G is genotype, GC is growth chamber, and T is tray (Meyer et al. 2007; Piepho et al. 2003).

Metabolite data

The metabolite profiles for each line were measured by gas chromatography–mass spectrometry (GC–MS). The samples for this measurement were harvested simultaneously with those for the dry weight measurement at 15 DAS. The details of measurement and GC–MS analysis are described by Lisec et al. (2006). The metabolite profiles contain 181 different metabolites. Since the lines were measured on different days the effect of detector sensitivity were corrected by dividing the intensity of each metabolite by the median of all intensities of that metabolite per measurement day. The normalization was done as described by Lisec et al. (2008).

Search for gene metabolite connections

We used the AraCyc 4.5 database to connect metabolites, SNP markers, and genes as described by Lisec et al. (2009) and identified genes directly involved in the conversion of the respective metabolite. For such a gene, the closest SNP marker is regarded to be associated with that metabolite. Alternatively the SNP markers closest to the metabolic QTL found by Lisec et al. (2008) were included.

QTL analyses

The QTL analyses for the biomass of the testcrosses followed the approach described by Meyer et al. (2009).

Composite interval mapping (CIM) was performed using the software package PLABQTL (Utz and Melchinger 1996). Cofactors were automatically selected by forward stepwise regression. Empirical logarithm of the odds (LOD) thresholds were determined by 1,000 permutations (Churchill and Doerge 1994). The genetic map used in all QTL analyses is based on the map presented by Törjék et al. (2006), with additional SNP markers (Meyer et al. 2009).

Machine learning procedure

The objective of the study presented here is to learn to predict the biomass of the progeny from molecular data of the ancestors. The machine learning procedure to achieve this purpose is divided into two parts: (i) variable or feature selection and (ii) regression. In the first step, the molecular quantities that are best suited to predict the trait are identified in order to reduce the number of variables without compromising the predictive power of the data set—defined here as the correlation between estimated and measured trait. In the second step, regression models are estimated using only the selected variables. Here, the actual prediction is performed.

The variables selection was subjected to a robustness evaluation. The combination of feature selection and regression was subjected to cross validation and permutation tests.

Variable selection methods

The variables selection method used in the present study is a modification of the approach described by Gärtner et al. (2009). In both approaches the variables are first ordered according to the same measure of importance. In the second step the actual selection takes place.

The variables are ordered according to their variables importance in the projection (VIP) (Chong and Jun 2005; Pérez-Enciso and Tenenhaus 2003). The VIP method is based on the partial least squares (PLS) approach (Eriksson et al. 2001; Wold 1975). PLS looks for linear combinations of the original predictor variables that maximize the covariance with a dependent variable also called response. These combinations, called PLS components, are orthogonal, in our application. Thus, by taking only a small number h of components PLS can be used for dimensionality reduction. There are different ways to determine h , as explained below.

The weight of the j th original variable in the linear combination resulting in the i th PLS component is denoted by w_{ij} . The VIP of the j th variable depends basically on the sum of the squared w_{ij} ($i = 1, \dots, h$) multiplied by the correlation of the i th PLS component with the response.

In the approach by Gärtner et al. (2009) the VIP of each original variable is calculated on the basis of the complete data set. The number of PLS components h in the corresponding PLS model is determined by maximizing the squared correlation between the true dry weight and the dry weight predicted in cross validation. Afterward, subsets of variables are considered, the size of which varies between 1 and the total number of variables. The k th subset comprises the variables with the k th highest VIPs. For several subsets PLS regression models are tested using cross validation. This cross validation is performed in the training set only and is repeated for each training set. Thus, two different subsets are determined: a set with maximal predictive power and another set, the predictive power of which is not significantly lower than that of the maximal set. In order to estimate the significance of the deviation from the maximal value, confidence intervals are calculated by jackknife procedures. The minimal set is the set of selected variables that will be used in the subsequent prediction procedure.

The modified procedure proposed here also calculates the VIP of each original variable, but the determination of the number h of PLS components in the corresponding PLS model is achieved by applying F tests. The modified procedure starts with the maximum VIP variable. For the next variable in the VIP order an F test is performed, which decides if the new variable yields additional information about the response, i.e., the null hypothesis of the test states that the regression coefficient of the new variable is zero. The F statistic we used is defined by $F = \text{RSS}(\beta_k) - \text{RSS}(\beta_{k+1}) / (\text{RSS}(\beta_{k+1}) / (n - k - 2))$, RSS denoting the residual sum of squares for the models expressed by the coefficient vectors β_k and β_{k+1} , and k and $k + 1$ representing the number of variables already selected. If the P value for the new variable is lower than 0.05 the variable is included in the subset, if not, the next variable in the VIP order is tested as described above.

Regression methods

The choice of the regression model depends on the properties of predictor data. Since the two procedures of variables selection differ, we used two different regression models.

The variable selection method by Gärtner et al. (2009) does not consider orthogonality directly. Therefore, a dimensionality reduction method rendering the predictor matrix orthogonal was required. Gärtner et al. (2009) used PLS regression.

For the data sets generated by the modified approach we applied, in addition to the PLS regression, ordinary least squares (OLS) models, which maximize the correlation between a combination of the predictor variables and the

response. The advantage of this method lies in its unbiased estimation of the model. The disadvantage is that correlation and co-linearity of the predictors result in a large variance of the estimation. However, since the selection of variables is biased to orthogonal variables, the application of OLS models is appropriate.

Evaluation of the robustness of the feature selection

We tested the robustness of the feature selection against possible loss of information by the reduction of the number of lines by applying bootstrap-like resamplings. In the first test, 1077 ($= 3 \times 359$) samples were drawn with replacement from the set of all (359) RILs. This specific number of samples was chosen because the expected proportion of lines drawn at least once was then approximately 95%. Thus, 18 lines were expected not to be included in the resampled sets. For the second test, 359 samples were drawn from the set of all RILs, thus in average around 35% of the lines were left out. The resampling was replicated 100 times in both cases. Variable selection was performed on the generated data sets as described in the previous sections.

We also evaluated the effect of small perturbations. For this purpose one observation was removed from the original data set 20 times. The question whether some of the selected markers could be replaced by others if there are only small changes in the data set was approached that way.

Cross validation

The cross validation was performed according to the leave-one-out (LOO) procedure: the predictor matrix $X_{n \times p}$ (with n number of samples, i.e., RILs; and p number of variables, i.e., SNP or metabolites) and the response, i.e., the dry weight vector, are divided into subsets. All but one subset are used to train a model including feature selection and regression. The model is then applied to the remaining subset in order to predict response Y of the test set. The pseudo code displayed in Supplementary Fig. 1 illustrates this procedure.

Permutation tests

Permutation tests were performed to determine the statistical significance of the estimation of the response (i.e., the dry weight) from the predictor data sets. The null hypothesis assumes that there is no relationship between the considered set of markers and the testcross dry weight. Therefore, the dry weight vectors were permuted 1,000 times. The complete machine learning procedure as described above, including the variables selection, was

applied to each of these permuted dry weight vectors, while the marker set remained unchanged. For each permutation the correlation between the permuted vector and its prediction was calculated. These correlations were compared to the predictive power of the procedure, when applied to the real data. The significance of the procedure is measured by a P value, which is defined as the number of random correlations higher than the predictive power divided by the number of permutations: (number of $R_{\text{perm},i} > R_{\text{true}}$ /number of permutations). The procedure of permutation test is represented by the pseudo code in Supplementary Fig. 2.

Results

The parental dry weight as predictor for hybrid biomass

The biomass ratio of the biggest to the smallest RIL is 1.8, the corresponding ratios of C24 and Col-0 testcrosses are 2.6 and 3.3, respectively. The mean dry weight values are 1.08, 1.59, and 1.55 mg plant⁻¹ for RILs, C24, and Col-0 testcrosses, respectively. The power of the prediction of the testcross biomass from parental biomass was evaluated separately for the C24 and the Col-0 testcross population. Since only the RIL parent has a variable dry weight in this experimental set-up, the Pearson correlation of the RIL dry weight with the biomass of both types of testcrosses is considered as a measure for predictive power. For the C24 testcrosses that correlation is very low (0.21) but still statistically significant (P value = 6×10^{-5}). The corresponding correlation with Col-0 testcrosses biomass is even lower (0.08 with a P value of 0.11).

Prediction of dry weight of testcrosses by parental molecular data from different sources of the parents

The following four data sets were used as predictors: metabolite profiles containing relative levels of 181 metabolites, 110 SNP markers, the combination of SNP markers and metabolite profiles, the combination of SNP markers, metabolites, and the RIL dry weight. All mentioned predictor variables are measurements on RIL parents only. In the following, we refer to these sets as METAB, SNP, METAB-SNP, and METAB-SNP-DW. The response to be predicted was the dry weight of the C24 testcrosses and the dry weight of the Col-0 testcrosses.

Before applying the feature selection, the predictive power of the predictor complete sets was determined using the OLS and the PLS regression method. A cross validation was performed as described in the “Methods” part. The best result was obtained for the C24 testcross population using the SNP data set ($R = 0.48$). For all other data sets

much lower values were obtained, especially in the case the OLS regression is applied (Table 1).

Using our modified feature selection method we sought to minimize the size of biomarker sets without significantly lowering the predictive power in cross validation. The predictive power of each data set/combination is given in Table 1. For the METAB-SNP set the OLS prediction with feature selection is improved in comparison to or equally good as the PLS results for the complete data set. Figure 1 illustrates these results and their application using the example of the combination of parental SNP markers, metabolites, and dry weight as predictors for biomass in C24 testcrosses. We have shown that the plants predicted to be the 10% biggest C24 testcrosses applying the OLS regression method had indeed a significantly higher mean value (1.79 mg) than the entire C24 testcross population (1.59 mg). This was done by a one sample *t* test. The *P* value was lower than 10^{-16} . This indicates a significant biomass difference of the population mean as compared to the value 1.79 mg.

Statistical significance of the procedure

The statistical significance of the OLS regression results including our new variable selection procedure was evaluated by permutation tests as described in the “Methods” section. The highest *P* value was 0.085 for the metabolite data set applied to C24 testcrosses. For all other potential biomarker sets the *P* value was smaller than 0.001 in both testcross set-ups.

For the METAB set permutation tests with 100 permutations were performed using PLS and the variable selection method of Gärtner et al. (2009). *P* values of 0.04 and

Table 1 Predictive power in cross validation of the molecular data from different sources and their combination

Data set	C24		Col	
	Complete	VS	Complete	VS
OLS regression				
METAB	0.11	0.05	0.11	0.14
SNP	0.36	0.41	0.28	0.38
METAB-SNP	0.16	0.42	0.14	0.32
METAB-SNP-DW	0.16	0.41	0.14	0.31
PLS regression				
METAB	0.12	0.06	0.23	0.21
SNP	0.48	0.40	0.35	0.37
METAB-SNP	0.39	0.40	0.28	0.32
METAB-SNP-DW	0.39	0.40	0.29	0.27

The columns 2–5 contain the predictive power for both effects (C24 and Col-0) and for the use of the variables selection (VS) and the use of the complete sets. The table contains the results for both OLS and PLS regression

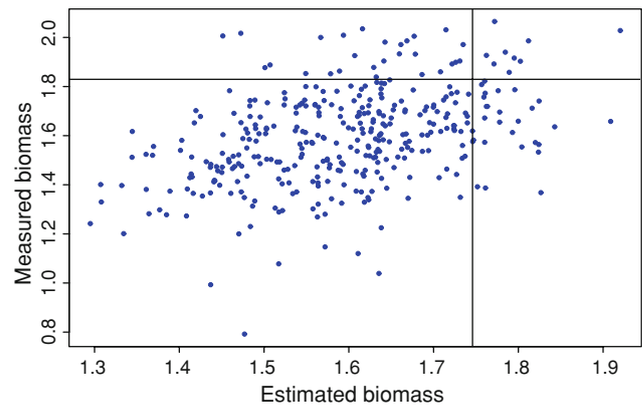


Fig. 1 Plot of the dry weight observed in the C24 testcrosses against the dry weight predicted by the METAB-SNP-DW set after variables selection. The vertical line indicates the 90% quantile of the predicted dry weight values. Thus, the testcrosses corresponding to the data points to the right of this line have been predicted to be the biggest 10% of the crosses. The horizontal line indicates the 90% quantile of the true biomass values

0.02 were calculated for the Col-0 and C24 effect, respectively.

The detected markers

The set of variables selected from the METAB-SNP set contained six metabolites and six SNP markers for the C24 testcrosses' biomass and 12 metabolites and three SNP markers for the Col-0 testcrosses' biomass. The overlap between the two testcross set-ups comprises one metabolite and one SNP marker. All three SNP markers found for the Col-0 effect, and five of six SNP markers found for the C24 effect in the METAB-SNP set were also selected in the SNP set. Lists of the selected markers are arranged in the Supplementary Tables 1–3.

Robustness of the selected marker sets

To evaluate the effect of small changes one observation was removed randomly 20 times from the METAB-SNP set. The whole procedure was then applied to the reduced data set and the selected features for each repetition were stored. Eight of the 12 selected C24 testcross markers were identified in all 20 subsets. Two further markers were used more than 10 times. A further 31 markers were detected at least once. The corresponding numbers for the Col-0 effect are three, eight, and 39 (Supplementary Table 3). The predictive power for the reduced sets ranged from 0.535 to 0.564 for C24 and from 0.451 to 0.483 for Col-0.

The results of robustness evaluation of the METAB-SNP variable selection by bootstrap-like resamplings in the case of the threefold resampling is presented in

Supplementary 5. For the C24 effect eight markers were found at least 90 times. Six of these markers belong to the 12 markers detected in the real data set (cf. Supplementary Table 3). This includes the only marker that was detected 100 times (see Supplementary Table 5). For the Col-0 effect four markers were detected at least 90 times including two of the selected markers (cf. Supplementary Table 3). One SNP marker from chromosome 4 was found 100 times. This marker also belongs to the set of markers detected within the real data set.

With the second, more stringent resampling strategy (see “Methods and materials” part), one marker for the C24 effect and two markers for the Col-0 effect were found at least 90 times. One metabolite marker of unknown chemical identity was selected for both effects. For the Col-0 effect the SNP marker MASC04123 located on chromosome 4 was selected in addition. Both markers were also detected within the real data set.

Methods comparison

We compared our results to those obtained applying the method proposed by Gärtner et al. (2009). For this purpose the variables selected by their method were subjected to a cross validation, i.e., the variables selection is not included in the cross validation. The procedure is similar to those explained by the pseudocode for the permutation loop. Therefore, the markers selected by the new feature selection were subjected to the same kind of cross validation. Here, only the results for PLS regression were compared (Table 2). In most cases the predictive power is similar for both methods.

However, for the C24 effect metabolites have a higher predictive power, if the procedure by Gärtner et al. (2009) is applied, whereas the application of the new approach on SNP markers yields better or equally good results. In most cases fewer variables are needed when the new approach is employed (Table 2). There is large overlap between the markers detected with both methods. Notably, all of the

Table 2 Predictive power of molecular data from different sources, obtained with the procedure as described by Gärtner et al. (2009) and the new method

	C24		Col-0	
	Gärtner	New	Gärtner	New
METAB	0.33 (56)	0.26 (9)	0.37 (18)	0.38 (13)
SNP	0.52 (17)	0.54 (9)	0.41 (9)	0.45 (8)
METAB-SNP	0.53 (56)	0.54 (12)	0.44 (30)	0.45 (15)
METAB-SNP-DW	0.54 (55)	0.56 (14)	0.44 (30)	0.46 (15)

The feature selection was performed outside the cross validation loop. The numbers in brackets signify the number of selected variables

markers robust against small changes are found with both methods.

Selected metabolites and SNP markers connected to them

We found in the AraCyc 4.5 database all genes connected directly to the metabolites of known chemical identity selected from the METAB-SNP set. In the second step we found for each such gene the SNP closest to it on the chromosome. Using an *F* test we determined, whether integrating the set of these SNPs in the reduced METAB-SNP model (excluding the metabolites of known chemical identity) significantly raises the predictive power.

For the Col-0 testcrosses 22 SNPs were linked to six metabolites from the METAB-SNP set. The *F* test resulted in a *P* value of 0.23, the inclusion of the six metabolites in a *P* value of 0.02. For the C24 effect 12 SNPs were determined as belonging to four metabolites. The corresponding *P* values are 0.65 and 0.04.

Alternatively, the SNPs closest to the QTL found for those metabolites by Liseć et al. (2008) were used, allowing us to include also SNPs for metabolites of unknown chemical identity. For the Col-0 effect four SNPs were linked to the 11 metabolites from the METAB-SNP selection. There was no significant gain in information by the inclusion of the SNPs (*P* value 0.18) in contrast to the inclusion of the metabolites (see above). For the C24 effect we found eight SNPs belonging to the six metabolites. Again the *F* test yielded no significant *P* value (0.57) for the inclusion of the set of SNPs, while a *P* value of 3×10^{-4} was obtained for the inclusions of the metabolites.

Comparison with per se, biomass heterosis, and testcross biomass QTL

The SNP markers selected as important features were compared to the SNP markers closely linked to detected biomass QTL (Meyer et al. 2009). The results are summarized in Table 3. We found that four of the eight SNP

Table 3 Comparison between QTL and SNP markers detected from the SNP set

Effect	C24		Col-0	
	# of QTL	# of co-location	# of QTL	# of co-location
per se	7	2	7	4
Biomass heterosis	6	3	1	1
Testcross biomass	6	5	2	2

For both C24 and Col-0, the number of QTL and the number of co-locations of the QTL with the 9 and 8 SNPs, respectively, are given

markers selected from the SNP set for the prediction of dry weight in Col-0 testcrosses, are co-locating with one of the seven per se biomass QTL. The only Col-0 QTL for biomass heterosis is co-located with one of the markers selected from the SNP set.

For the C24 testcrosses three out of nine SNP markers selected are also in the support intervals of per se biomass QTL. For the six C24 QTL for biomass heterosis we found three co-located SNP markers with the variables selection.

In addition, a QTL search for the testcross biomass was performed in the present study with the same methods as used by Meyer et al. (2009), cf. Chap. “Methods and materials”. We found six QTL for the C24 testcrosses and two for the Col-0 testcrosses (Supplementary Table 4). The markers co-locating with QTL are indicated by arrows in Fig. 2. One SNP marker on chromosome 1 is found for both effects. The signs of the impact of the markers obtained from the feature selection on the hybrid biomass indicate in the most cases a biomass increase when the corresponding position is heterozygous. There are two exceptions for the C24 effect, but the corresponding markers are less important than the others. One marker on chromosome 1 for both effects shows a decrease in the hybrid biomass, if the RIL parent had a C24 allele at this position.

The SNP markers located in the support intervals of the testcross biomass QTL were used to predict the corresponding dry weight in cross validation. The predictive power of these predictors was determined with 0.48 and 0.37 for C24 and Col-0, respectively. Since the same response trait (i.e., testcross dry weight) was used these results could be in principle compared to the predictive power of the markers obtained by feature selection. However, the cross validation did not include the QTL search. Therefore, the SNPs detected by the feature selection were

subjected to the same kind of cross validation (i.e., the feature selection is not included in the cross validation). Their predictive power is then 0.54 and 0.45 for the C24 and Col-0 effect, respectively.

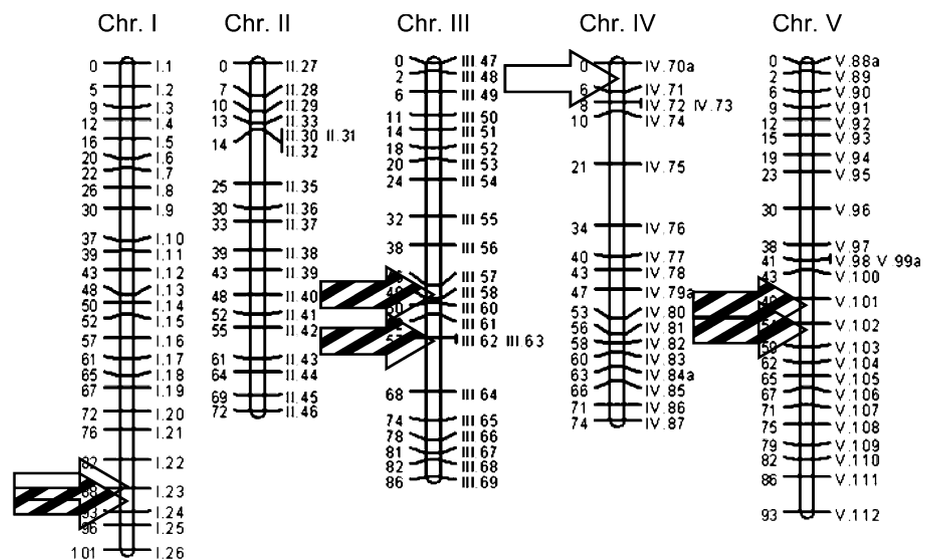
Discussion

We developed a new feature selection method that represents a complementary approach to previous works in the field (e.g., Frisch et al. 2009; Maenhout et al. 2009; Schrag et al. 2007, 2009a, b; Yu et al. 2005). For the proof of concept presented here we used a model population of *Arabidopsis*, therefore a direct comparison with the more agricultural applications of these authors is difficult. However, the predictive power and the reduction of the number of markers achieved by our procedure indicate its potential usefulness for breeding programs. Additionally, metabolites are introduced as useful markers.

We employed three types of potential predictors: macroscopic phenotypes, genetic markers, and metabolites. The small amount of variance of the hybrid biomass explained by the parents' biomass shows the insufficiency of phenotypic markers as good predictors. The application of machine-learning procedures to molecular data is therefore a relevant alternative for the prediction of hybrid performance in this population.

We could show that the prediction of the trait under investigation is clearly increased by the use of SNP markers and metabolite profiles in comparison to the use of parent dry weight only. Permutation tests showed with one exception, that the metabolites significantly predict the testcross biomass. We conclude therefore that these substances are potential biomarkers for hybrid performance.

Fig. 2 Location of the SNP markers used in the present study. Markers detected as biomarkers in both feature selection as well as QTL search are indicated by arrows. Banded and unbanded arrows indicate C24 and Col-0 biomarkers, respectively. The arrows on chromosome one point to the same SNP



With our procedure we were able to reduce the number of variables employed for at least one of the two testcross classes from 291 SNP markers and metabolites to 12 features. The markers found by feature selection prove to be robust against small changes in the data set. Some markers are exchangeable without compromising the predictive power. Overall, only a small proportion of the available markers are used, e.g., for the C24 effect 31 out of 291. A much smaller number (eight and four for the C24 and Col-0 effect, respectively) of markers are robust against the loss of about 5% of the lines and only two markers are robust against greater changes as shown by the results of the bootstrap-like resampling.

The selection of small sets of markers is important for three reasons. First, the prediction of the trait could be improved for most of the predictor sets (see Table 1). Second, the selection of a small set of markers reduces the cost of measurement. Finally, a targeted measurement of metabolite concentrations will result in a higher accuracy. The selection of few important metabolites enables such targeted measurements. This in turn is likely to improve the predictive power of the procedure.

The modification of our original method (Gärtner et al. 2009) lowers the number of variables necessary to make predictions with a nearly equal predictive power. As described above, this is advantageous for the direct application in breeding. However, when we are interested not only in the prediction of an observed effect but in an explanation of it in molecular terms, it is reasonable to take into account also markers that improve the prediction only slightly. Therefore, the markers identified by the method of Gärtner et al. (2009) should be considered in such investigations, e.g., modelling approaches. Furthermore, the modified method has the tendency to fail in case of predictors with considerable measurement errors. This is shown by the worse results for the metabolites in the case of the C24 testcrosses, where a significant prediction was computed with our original method.

The metabolites of known chemical identity found to be important in the METAB-SNP set, could be related to SNP markers using information from the AraCyc 4.5 database. In contrast to the metabolites, this set of SNP markers does not add to the predictive power of the set of SNP markers found to be important. The use of SNP markers derived from metabolic QTL lead to the same conclusion. The metabolite concentrations can not be explained sufficiently by a linear combination of the genes known to be related to these metabolites.

The SNP markers found by feature selection overlap substantially with the QTL determined by Meyer et al. (2009). This is also true for the testcross biomass QTL determined in the present study. Mainly dominance effects are found. The only QTL found for both C24 and Col-0

shows additive effects. The predictive power of the SNP marker set detected by the variables selection is clearly higher than that of the combination of QTL. This shows that our method can find new interesting regions on the chromosomes, represented by the SNP markers. In our approach interactions between SNP markers are not considered and therefore, epistasis could not be detected, directly. However, the role of the metabolites as presented above nevertheless clearly indicates the presence of epistasis.

To improve our approach, we plan to include gene interaction in our model. Here, the knowledge of important metabolites and their connection to genes will be helpful. The use of more complex populations, i.e., derived from more than two accessions, would be a further possibility to test our method. In combination these two applications will increase the area of potential applications of our procedure for plant breeders.

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