Diversity and linkage disequilibrium analysis within a selected set of cultivated tomatoes

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Abstract Within the Dutch genomics initiative the “Centre for Biosystems Genomics” (CBSG) a major research effort is directed at the identification and unraveling of processes and mechanisms affecting fruit quality in tomato. The basis of this fruit quality program was a diverse set of 94 cultivated tomato cultivars, representing a wide spectrum of phenotypes for quality related traits. This paper describes a diversity study performed on these cultivars, using information of 882 AFLP markers, of which 304 markers had a known map position. The AFLP markers were scored as much as possible in a co-dominant fashion. We investigated genome distribution and coverage for the mapped markers and conclude that it proved difficult to arrive at a dense and uniformly distributed coverage of the genome with markers. Mapped markers and unmapped markers were used to investigate population structure. A clear substructure was observed which seemed to coincide with a grouping based on fruit size. Finally, we studied amount and decay of linkage disequilibrium (LD) along the chromosomes. LD was observed over considerable (genetic) distances. We discuss the feasibility of marker-trait association studies and conclude that the amount of genetic variation in our set of cultivars is limited, but that there exists scope for association studies.

Introduction

Five years ago, the Dutch plant genomics initiative the “Centre for Biosystems Genomics” (CBSG) was set up to stimulate plant genomics research in general and to investigate more specifically for potato and tomato the phenotypic and genetic variation in consumer quality, environmental quality, technological aspects and societal aspects (e.g., public acceptance of genomics methodologies). To provide a benchmark for research advances in those agricultural crops, Arabidopsis was included as a model crop for which state of the art resources were available.

A considerable part of the CBSG resources was concentrated on the improvement of tomato fruit quality. As part of that research, diversity and fruit quality was assessed for a panel of 94 greenhouse cultivated tomatoes, containing both older as well as current tomato cultivars. Breeding companies that are partners within CSBG were asked to provide a sample of their germplasm taking care to include contrasts on relevant quality aspects. Tomato quality was investigated in terms of metabolic profiling and in terms of sensory and consumer appreciation studies on ripe tomato fruits. This paper focuses on genetic diversity, as sampled by molecular markers. The results of the quality assessments and marker—trait association studies for quality will be presented elsewhere.

Sufficient genetic variation, or diversity, in target traits is a condition for progress in plant breeding. Genetic diversity
can be investigated in many ways, for example, through morphological and biochemical traits, pedigree analysis or by molecular markers (Franco et al. 2001; Zhao et al. 2005a). Among these methods, morphological traits are intuitive and practical, but as they are subject to environmental influences and selection pressure during domestication and breeding, the interpretation of the results of diversity studies based on such traits can be difficult. Pedigree records are often incomplete or have limited availability. The advent of DNA markers stimulated diversity studies in plants, as DNA markers allow extensive sampling at the DNA level. Within tomato (Solanum lycopersicum), many studies have been performed on genetic diversity and variety identification, using molecular markers such as RFLP (Miller and Tanksley 1990), RAPD (Williams and St. Clair 1993; Villand et al. 1998; Noli et al. 1999; Carelli et al. 2006), AFLP (Park et al. 2004) and SSR (Bredemeijer et al. 2002).

For the diversity study in this paper, we used AFLP markers (Vos et al. 1995). AFLP genotyping has the advantage of sampling a large number of informative fragments with random genomic origin (Mueller and LaReesa Wolfenbarger 1999). Although other choices of marker types would have been possible, AFLP markers have some desirable properties in the light of application in statistical diversity analyses (see Meudt and Clarke 2007; Bonin et al. 2007 for a more elaborate discussion on this subject). We investigated the distribution of AFLP markers across the genome, and assessed genotypic frequencies for the marker loci. Bayesian and nearest neighbor clustering methods were used for identifying possible population structure. Finally, as a precursor to later association studies, we studied the magnitude of linkage disequilibrium (LD) and the decay of LD as a function of genetic (map) distance.

Materials and methods

Germplasm

Five internationally operating vegetable breeding companies (Enza, De Ruiter Seeds, Rijk Zwaan, Seminis Vegetable Seeds and Syngenta Seeds), all participating in the research consortium CBSG (see Van Berloo et al. 2007 for more details), were asked to provide tomato cultivars from their collections of current and historic cultivated tomato germplasm. These companies together collected a set of ~150 cultivars. Based on previously gathered sensory and morphological data, a set of 94 cultivars was selected from this collection. This set can be considered representative, with regard to phenotypic diversity, for the available germplasm used in commercial European greenhouse tomato cultivation. The phenotypic diversity of the sampled germplasm was compared to other studies in tomato for two traits: Brix (soluble solid content, expressed as a percentage) and average fruit weight. Thus, we first compared the observed ranges in our germplasm to the observations of Chaib et al. (2006), who investigated tomato fruit quality aspects in a set of 144 RILs derived from an inter-specific cross and in approximately 100 derived backcross lines. In addition, we looked at the observations of Helyes et al. (2003), who described phenotypic variation in an experiment involving 18 tomato cultivars.

Seeds of the tomato germplasm discussed in this paper are available on request. Please contact the CBSG consortium secretariat (http://www.cbsg.nl for contact info).

Morphological and fruit observations

The chosen cultivars were grown in replicated trials in the greenhouse, and fruits were harvested and subjected to a number of observations and analyses, which are discussed in van Berloo et al. (2007), Ursem et al. (2008) and other papers in preparation. Twelve morphological and anatomical plant characteristics were scored: total plant length at the end of the cultivation period; total weight of vegetative tissues (leaves, stem) at the end of the cultivation period; accumulated weight of all harvested fruits during cultivation; average number of flowers per inflorescence; average stem inter-node length; fresh weight and dry weight of the vegetative system (dry weight measured after storage in an oven at 80°C for 48 h); total leaf area (which was measured at the end of the cultivation period using a dedicated apparatus with semi-automatic pass-through similar to the device described by Murata and Hayashi 1967); shape of the fruits (expressed as a length/width ratio); total number of harvested fruits during cultivation; and average fruit weight and fruit firmness (which was measured as the amount of deformation of the fruit when submitted to a force of 3 N). These observations can be used to assess the phenotypic diversity among the set of cultivars, and compare this type of diversity with the other types of diversity like that based on molecular markers, the latter being the main theme of this paper. In addition to the above traits, Brix (soluble solid content of the tomato fruit) was measured using a refractometer (GMK-701R; Nie-Co Products, Aalsmeer, NL).

AFLP genotyping

Fifty AFLP primer combinations were run on DNA samples of the 94 tomato cultivars. This large number of primer combinations was used because it is known that the amount of marker polymorphism in cultivated tomato is generally low (Miller and Tanksley 1990; Archak et al. 2002). However, when polymorphism rate is low the AFLP technology is often still capable of uncovering genetic differences
The integrated map we used has not been published, but (Rouppe van der Voort et al. 1995). EcoRI-Mse and PstI-Mse primer combinations were used. Selective nucleotide adaptors were selected, based on previous experience in tomato, in such a way that the expected number of polymorphic loci was optimal. Genotyping resulted in the scoring of 882 polymorphic AFLP fragments. For 304 AFLP fragments the map position could be established from an integrated inter-specific linkage map.

AFLP scoring is standardly done by registration of presence and absence of bands, i.e., in a dominant fashion. We developed a co-dominant scoring methodology which is based on the application of mixture models to the quantitative gel intensity values for the fragments. This method extends the mixture model methods described by Piepho and Koch (2000) and Jansen et al. (2001) for biparental populations to germplasm panels of arbitrary composition. The essence of our mixture model is that we estimated the fractions of homozygotes and heterozygotes from the distribution of band intensity data itself, where the earlier methods for biparental populations fixed those fractions in relation to the type of population. For example, for an F2 population, the fractions will be 0.25 – 0.50 – 0.25, respectively, for homozygote band present, heterozygote band present, and homozygote band absent.

For quantification and investigation of genetic diversity, we used both the co-dominant genotypic classification on the basis of AFLP band intensities explained above, and log transformed numerical band intensity values. The reason for using both types of marker derived information is mainly one of convenience in later calculations whose results will be presented elsewhere. For the present paper, the choice of log band intensity or discrete (band) allele number did not influence results or interpretations related to diversity.

Linkage map

Since the marker data were scored on a set of cultivars, and not in a segregating population, we could not create a genetic linkage map from the data at hand. Therefore, we used an existing high density linkage map created from integrating three inter-specific linkage maps. Identity of newly scored AFLP fragments and previously mapped fragments followed from equality of their mobility numbers (Rouppe Van der Voort et al. 1997; Vuylsteke et al. 1999). The integrated map we used has not been published, but can be considered an extension of the previously published integrated map by Haanstra et al. (1999).

Genome coverage, marker distribution and test for Hardy–Weinberg equilibrium

Only a minority of the scored AFLP markers could be assigned map positions: 304 out of 882. Through our estimate of the decay of LD we could estimate the fraction of the genome sampled by the available set of mapped markers. The allocation of markers to genome positions was analyzed by looking at the distribution of distances between neighboring markers. We assessed the general distribution of markers over the chromosomes in the form of box plots of inter marker distances as well as the uniformity of markers along the chromosome through Chi-square tests that compared the observed numbers of markers in 10 cM chromosome intervals with expected numbers assuming a uniform distribution. Estimated allele frequencies were used to calculate expected genotype frequencies, which were compared to observed genotype frequencies in tests for Hardy–Weinberg (HW) equilibrium (Falconer and Mackay 1996, Chap. 1).

Population structure analyses

Possible structure in the population was investigated in various ways. First, to the co-dominant genotypes we applied the Bayesian clustering method implemented in the STRUCTURE 2.1 package (Pritchard et al. 2000). This method is based on the creation of groups that are internally characterized by HW equilibrium and absence of LD, while between groups LD exists. A subset of the markers was used for this analysis. This subset covered the genome with evenly spaced markers at approximately 10 cM inter marker distance, thereby reducing the bias following from sampling closely linked markers. STRUCTURE settings were left at their default values and 10,000 cycles were applied for burn-in, followed by 100,000 cycles for the actual analysis, assuming admixture of populations. Secondly, we evaluated population structure by inspection of a neighbor joining dendrogram built from genotypic distances, which were calculated as Euclidean distances using the log band intensities. Genotypic distances were calculated by the GENSTAT statistical package (Genstat 2005). The MEGA 3 package (Kumar et al. 2004) was used for Neighbor Joining cluster analysis (Saitou and Nei 1987). A complementary analysis studying between group separation was done for within and between group distances, taking the a priori sets of cherry, beef, and round tomatoes as groups.

Relationships between genotypes were further studied with network methodology following Ursem et al. (2008). The log band intensities of the markers selected for use in the STRUCTURE analysis were now used to calculate Euclidean distances between genotypes. These Euclidean distances
were transformed to similarities which served to join genotypes in a network representation created with the Pajek package (Bagatelj and Mrvar 2003), whenever the observed similarity exceeded the 99 percentile of the null distribution for the similarities. This null distribution was obtained from permutation of the log band intensities with respect to the genotypes. The network was inspected for connectivity. Clustering coefficients were calculated to quantify the density of the network around individual genotypes (Watts and Strogatz 1998). A clustering coefficient expresses the number of observed links between neighboring genotypes in relation to the total number of possible links between neighbors. Neighboring genotypes are pairs of genotypes that are both connected to a third genotype. Neighboring genotypes need not themselves be connected again, i.e., they might be linked to a third genotype without being directly linked themselves.

For comparison with the marker information, a second neighbor joining analysis was performed on distances obtained from morphological plant characteristics. Euclidean distances were derived from the set of 12 numerically scored morphological observations introduced in a previous paragraph.

LD analyses

LD between markers was assessed by calculation of squared correlation coefficients \(R^2\); Zhao et al. 2005b) between marker intensity patterns, using the GGT 2.0 software package (Van Berloo 2008, 1999). LD statistics were calculated per chromosome for all marker pairs and subsequently, aggregated over all 12 chromosomes, the decay of LD with map distance was evaluated. As a threshold for significance, a 5% false discovery rate was chosen, following the two-step procedure described by Benjamini et al. (2005).

The strength of LD between marker pairs across the full genome was plotted in a heatmap, using the GGT 2.0 package. Separate heatmaps were created for the beef and round tomatoes on the one hand and the cherry tomatoes on the other hand. In addition to the heatmaps, correlations between markers on different chromosomes were calculated.

**Table 1** Range of phenotypic observations for Brix and fruit weight of the cultivars described in this study compared with three diverse research populations from literature

|                          | Brix (%) | Fruit weight (g) |
|--------------------------|----------|------------------|
|                          | Minimum  | Mean  | Maximum | Minimum | Mean  | Maximum |
| Helyes et al.: 18 cultivars | 5.20     | –     | 8.70     | –       | –     | –       |
| Chaib et al.: 144 RILs    | 4.38     | 7.13  | 9.38     | 10.8    | 27.0  | 71.3    |
| Chaib et al.: 100 BC3S1   | 5.72     | 7.12  | 8.40     | 35.4    | 72.5  | 117.7   |
| CBSG: 94 cultivars        | 3.83     | 5.17  | 9.27     | 6.8     | 58.2  | 161.4   |
of markers over the chromosomes was not uniform ($P < 0.001$) and that for chromosomes 4, 6, 9 and 11 the distribution of markers within the chromosome was not uniform either. Some deviations from HW equilibrium were observed for markers on all chromosomes, but for some chromosomes (4, 5 and 8) HW equilibrium was almost completely absent. The non-uniform dispersion of markers along the chromosomes becomes apparent when we look at the distribution of markers over the chromosomes in Fig. 1. To investigate the distribution of inter-marker distances per chromosome, we constructed box-plots that visualize the distribution of distances between consecutive markers (Fig. 2). Figure 2 and Table 2 show considerable variability for coverage and distribution of markers. Several regions are well or very well covered while other chromosomes are only sparsely covered. Chromosomes 4, 6 and 9 have an average distance between consecutive markers of less than 1 cM, and 75% of all marker distances between neighboring markers are less than 3 cM. On the other hand for chromosomes 2, 3 and 7 more than half of the distances between consecutive markers are larger than 3 cM.

### Population structure analyses

The analysis with the Structure 2.1 software package indicated the presence of population structure, because the model likelihood increased considerably when more than one population was assumed. However, estimating the number of populations was difficult as the likelihood for three to six populations was very similar and only increased when seven or more populations were assumed (data not shown). A closer look at the membership probabilities for the genotypes showed us that, when four or more subpopulations were assumed, all genotypes became “distributed” over at least two populations. We therefore chose the simplest solution that still assigned a large part of the genotypes to a single population, which was at three (sub-) populations. A second Structure run using all 882 codominant markers was performed (data not shown) and confirmed the earlier results. Figure 3 gives subpopulation memberships in the form of a bar plot for the three cluster solution. To initiate the analysis, the tomato types (beef, cherry, and round) were used as prior population identifiers. These types are indicated by the numbers below the bars. The plot clearly demonstrates that one of the identified populations coincides almost completely with the cherry tomato type (cluster 1 and group C in the plot). Although most cultivars belonging to group B (beef tomatoes) have a

### Table 2: Details of marker distribution among the 12 tomato chromosomes, coverage of the genetic map obtained with the scored markers, and tests for Hardy–Weinberg equilibrium (see text for details)

| Chromosome | Number of markers | Length (cM) | $P$-value $\chi^2$ test for uniform distribution | Percentage of chromosome with sufficient coverage | Percentage of bands in Hardy–Weinberg equilibrium |
|------------|------------------|-------------|-----------------------------------------------|-----------------------------------------------|-----------------------------------------------|
| 1          | 14               | 119         | 0.21                                          | 89.4                                          | 31                                            |
| 2          | 5                | 94          | 0.88                                          | 88.9                                          | 40                                            |
| 3          | 3                | 126         | 0.32                                          | 53.5                                          | 100                                           |
| 4          | 35               | 90          | <0.01                                         | 80.6                                          | 6                                             |
| 5          | 28               | 86          | 0.12                                          | 65.9                                          | 18                                            |
| 6          | 44               | 100         | <0.01                                         | 58.8                                          | 95                                            |
| 7          | 6                | 128.5       | 0.71                                          | 67.2                                          | 67                                            |
| 8          | 7                | 88.2        | 0.47                                          | 87.7                                          | 14                                            |
| 9          | 123              | 94          | <0.01                                         | 89.1                                          | 93                                            |
| 10         | 6                | 75          | 0.18                                          | 68.8                                          | 33                                            |
| 11         | 19               | 86          | <0.01                                         | 98.7                                          | 63                                            |
| 12         | 14               | 90          | 0.29                                          | 84.6                                          | 50                                            |
| Overall    | 304              | 1,176.7     | 77.8                                          | 65                                            |                                               |
high probability to belong to the population defined by cluster 2, and many cultivars belonging to group R (round tomatoes) have a high probability of belonging to the population defined by cluster 3, tomato type and subpopulation do not completely coincide for beef and round tomato types. Further evidence for this can be found in the membership proportions, as given in the structure output for each of the three a-priori defined types, presented in Table 3. Each of the three a priori tomato types coincides to a certain degree with one of the a posteriori clusters. This is most obvious for the cherry group, for which the a-posteriori membership probabilities were on average 0.87 for cluster 1. The beef and round tomato cultivars also coincide with proper clusters, but with more exceptions.

Table 3  Membership proportions of three tomato type subgroups in each of the three inferred clusters resulting from Structure analysis

|          | Cluster 1 | Cluster 2 | Cluster 3 |
|----------|-----------|-----------|-----------|
| Cherry   | 0.87      | 0.07      | 0.06      |
| Beef     | 0.01      | 0.69      | 0.31      |
| Round    | 0.02      | 0.36      | 0.62      |

The cluster numbers refer to the clusters given in Fig. 3.

A neighbor joining dendrogram (Fig. 4) derived from Euclidean distances on (log transformed) band intensities also indicated the presence of groups in the set of genotypes. Especially the cherry tomatoes stand out as a separate group. Beef and round tomatoes are not clearly separated and showed smaller distances among their cultivars than the cherry cultivars. To check the stability of the dendrogram we used bootstrapped cluster analysis in MEGA 3, using for convenience a dominant scoring of AFLP markers. We found that the main separation of subclusters was very stable (bootstrap values of 99% for the main subdividing nodes). We did not observe indications for clustering of genotypes related to the breeding origin of the cultivars, as most genotypes that originated from a particular company ended up scattered in a seemingly random fashion throughout the dendrogram (data not shown).

Figure 5 gives the genetic distances within and between the groups (cherry vs beef + round) defined by the tomato types. The cherry group showed larger within-group genetic distances than the beef + round group, but the between group distances were clearly larger than the within group distances.

The network constructed on the basis of band intensities in Fig. 6 emphasized once again the difference between the cherry tomatoes and the beef and round tomatoes. Figure 6 shows that the connectivity in the cherry cluster was extremely low with an average cluster coefficient of 0.24, while the connectivity in the beef and round cluster was
very high with a clustering coefficient of 0.70. Both clusters were uniquely connected through the genotype pair R131 and C132. Interestingly, the single outlying genotype R068 was also connected to R131. These three genotypes happened to have in common an elongated Roma-like fruit shape. Other unique genotypes were R038 and B098. (Unique genotypes have cluster coefficient zero). Looking at the distributions of clustering coefficients for the genotypes stemming from a particular breeding company (Table 4), it was observed that the median clustering coefficient for company 4 was the lowest, and the coefficient for company 5 was the highest, with the companies 1, 2 and 3 having more or less the same intermediate value. This means that the amount of genotypic variation selected by company 4 was highest, that by company 5 lowest. Still, differences were very small.

The Euclidean distances, based on numerical trait values for 12 morphological and harvest related traits, were used to create another neighbor joining dendrogram (Fig. 7). This dendrogram shows a large cluster and a smaller cluster. The smaller cluster clearly stands out and is dominated by cherry cultivars, but no exclusive separation of cherry and other tomatoes can be observed.

Our analyses of both types of genetic distances indicate the presence of at least two subgroups: a group containing all cherry type cultivars and a large group containing beef and round cultivars. This second group could still be subjected to further subdivision, but no obvious partitioning
related to tomato type can be found, which confirms the results obtained using the STRUCTURE package.

**Linkage disequilibrium analysis**

As separate analyses of LD decay for the cherry and beef-round tomatoes yielded very similar results, only the decay for the cherry group is presented (Fig. 8). Extensive LD can be observed and reasonably strong LD can be found for markers up to 20 cM apart, indicating good possibilities to explore LD based marker-trait associations in this material. We used a slightly conservative estimate of the average extent of LD of 15 cM to determine the genome coverage presented in Fig. 1.

Figure 9 shows separate LD heat maps for the beef-round (Fig. 9a) and cherry (Fig. 9b) tomato types. Both heat maps show that strong LD is limited to certain hot-spots, and is mostly concentrated along the diagonal of these plots, as was expected. Strong LD between markers on different chromosomes is rare, although it does occur at some spots. In the heat map for the cherry group, considerable LD can be observed between markers on chromosomes 2 and 4, 4 and 8, 4 and 11, 1 and 9 and 4 and 9. The heat map for the beef-round tomato types shows inter-chromosome LD for markers on chromosomes 4 and 10, 4 and 11, 1 and 9 and 4 and 9.

**Discussion**

In this paper we have shown that AFLP markers are useful tools for genetic analysis, and that they allow us to get an estimate of the genetic diversity. The number of markers obtained using AFLP methodology is of course highly dependent on the genotyping and protocol conditions, but an average of 30–40 polymorphic markers per primer combination is often feasible (e.g. Haanstra et al. 1999; Isidore et al. 2003; Keygene, unpublished data). In our case on average only 18 polymorphic markers per primer combination could be scored co-dominantly, which is an indication that the amount of genotypic variability present in the tested set of cultivars is limited.
Fig. 7 Neighbor joining tree for the 94 tomato cultivars. Genotypic distances are calculated as Euclidean distances based on 12 morphological and harvest related traits (see text for details). The different shapes indicate the three different tomato types (filled square beef tomato, filled circle round tomato, filled diamond cherry tomato).

Fig. 8 Linkage disequilibrium $R^2$, versus map distance in cM, for the cherry cultivars. The red line indicates a smoothing spline. The 0.05 false discovery rate threshold is $R^2 = 0.37$. 
Fig. 9 Heatmap display of LD as $R^2$ for all marker-pairs. 

**a** Heatmap of LD between markers for the beef and round tomato types. 

**b** Heatmap of LD between markers for the cherry tomato type. In both figures on the x and y axis the 304 mapped markers are arranged in map order. LD between pairs of markers is plotted in the bottom diagonal part of the figure. Along the diagonal the chromosomes and marker positions are indicated. The color intensity reflects the amount of LD.
We used normalized gel intensity data to analyze individual marker behavior in more detail, and identified and scored a substantial number of markers co-dominantly. Especially in a situation like the one studied in this paper, where analyses are performed on plant material that is to a large extent (expected to be) heterozygous, the ability to discriminate between presence of a marker band in single or double quantity can be beneficial to increase power in diversity and association analyses.

A possible explanation for the uneven distribution of markers along the chromosomes may lie in the origin of the integrated genetic map that was at our disposal. The research that formed the basis for the construction of that map was directed at a number of specific tomato genomic regions, which contained genes of interest, like resistance loci (e.g., Van Berloo and Lindhout 2001). Therefore the region on chromosome 6, bearing resistance against tomato yellow leaf curling virus and the region on chromosome 9, harboring tomato mosaic virus resistance, have become more saturated with markers than other regions. From Table 2 we observe that especially chromosome 4 and to a lesser extent chromosomes 5 and 8 show strong deviation from HW equilibrium. Chromosome 4 also shows considerable LD with markers on other chromosomes. Recent breeding efforts focusing on (regions of) chromosome 4 could be responsible for these observations.

Even though we selected, among the available cultivated material for greenhouse cultivation, a set of varieties that was quite diverse at the phenotypic level, we discovered that the diversity at marker level was limited. The dendrogram based on phenotypic observations (Fig. 7) shows larger diversity than the dendrogram based on molecular marker data (Fig. 4). This observation is in line with other studies in cultivated tomato (e.g., Archak et al. 2002; Park et al. 2004; Garcia-Martinez et al. 2006) and could be a consequence of a strong selection pressure that was applied to the current cultivated germplasm. Also the fact that several regions of the genome were poorly covered by markers does not necessarily reflect a shortage of markers, but a lack of polymorphic markers, since many markers that are polymorphic when sampled in wider genetic backgrounds, like a population derived from an inter-specific cross, did not show polymorphism among our set of 94 cultivated varieties.

Based on fruit morphology we could clearly define three groups within our set of cultivars. A large fruited group (beef), a group with regular sized (round) table tomatoes and a group with much smaller fruited (cherry) tomatoes. The network analysis confirmed the classification in groups, but also revealed that this classification may be extended with a classification based on fruit shape, as cultivars with different fruit sizes but with a similar elongated shape remained closely connected in this diagram. At the genomic (AFLP) level the distinction between the cherry tomatoes and the round and beef tomatoes remained clear. Minor differences between the round and beef tomatoes were observed but these differences were far less striking than the dominating cherry versus beef + round division. Most likely this is caused by the shared breeding history of these tomato types, which is much longer for round and beef tomato cultivars (and parental breeding lines), while the cherry tomatoes have only recently become more popular and subjected to breeding efforts.

The magnitude of LD and the length of genome intervals with strong LD appear to be larger than that observed in studies dealing with other plant species like sugar beet (Kraft et al. 2000), Arabidopsis (Nordborg et al. 2002) and barley (Kraakman et al. 2004). A reason for the strong LD could be the self pollinated history of the tomato species, combined with a very limited variability, due to strong and continued selection by plant breeders and re-usage of donor lines from a limited gene pool. Although new donor segments have been introduced from more variable related species into the tomato germplasm, mainly to add resistance genes, the time for recombination and breaking down of LD within these donor segments has been rather short, and the process to achieve linkage equilibrium is still ongoing. Furthermore, new donor genome segments are continuously being introduced by breeders, thereby introducing new LD into the cultivated germplasm.

LD that was observed between markers that are located on different chromosomes could be caused by errors in the map positions of some of these markers, or by simultaneous selection of certain regions, that remain together because of advantageous epistatic effects, but further research would be needed to confirm this hypothesis.

The LD analyses presented in this paper indicate strong LD among markers, and LD extending over relatively large regions. These results may be biased by chromosome fragments that have been introgressed from wild germplasm in the past decades, i.e., fragments containing resistance genes. However, the high amount of LD that is found indicates that there is sufficient prospect for association mapping. Even regions that are only sparsely covered by markers may still show sufficient LD to be interesting for marker-trait association studies. The downside of the strong LD that is found over a considerable genetic distance will be that any results of association studies will only be roughly indicative for the genomic position of effects that are observed.

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