Homeologous recombination in *Solanum lycopersicoides* introgression lines of cultivated tomato.

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ABSTRACT

A library of ‘introgression lines’ containing *Solanum lycopersicoides* chromosome segments in the genetic background of cultivated tomato (*Lycopersicon esculentum*) was used to study factors affecting homeologous recombination. Recombination rates were estimated in progeny of 42 heterozygous introgressions and whole chromosome substitution lines, together representing 11 of the 12 tomato chromosomes. Recombination within homeologous segments was reduced to as little as 0-10% of expected frequencies. Relative recombination rates were positively correlated with the length of introgressed segments on the tomato map. The highest recombination (up to 40-50% of normal) was observed in long introgressions or substitution lines. Double introgression lines containing two homeologous segments on opposite chromosome arms were synthesized to increase their combined length. Recombination was higher in the double than in the single segment lines, despite a preference for crossovers in the region of homology between segments. A greater increase in homeologous recombination was obtained by crossing the *S. lycopersicoides* introgression lines to *L. pennellii* – a phylogenetically intermediate species – or to *L. esculentum* lines containing single *L. pennellii* segments on the same chromosome. Recombination rates were highest in regions of overlap between *S. lycopersicoides* and *L. pennellii* segments. The potential application of these results to breeding with introgression lines are discussed.
INTRODUCTION

As plant breeders broaden their search for novel traits and allelic diversity, it is frequently necessary to search in the more distant wild relatives of crop plants. Introgression of genes from exotics can be restricted by pre- and post-zygotic barriers that prevent or impede gene transfer. In cases where interspecific hybrids are viable but sterile, the sterility may result from genic and/or chromosomal effects (STEBBINS 1958). Sterility of the latter type occurs when genomes are so diverged that homeologous chromosomes of different species fail to recombine, leading to abnormal assortment at meiosis. Meiotic recombination can also be suppressed in backcross generations, leading to linkage drag, i.e. the unintended transfer of large blocks of DNA surrounding a gene of interest.

The cultivated tomato, Lycopersicon esculentum (syn. Solanum lycopersicum), can be experimentally hybridized with each of the 9-13 species in the tomato clade (i.e. genus Lycopersicon or Solanum section Lycopersicon). The resulting F1 hybrids are relatively fertile, and chromosomes undergo normal meiotic pairing and recombination processes. Comparative genetic maps constructed from interspecific tomato populations show a high degree of colinearity between genomes of all tomato species. However, recombination is typically reduced after backcrossing to cultivated tomato (RICK 1969; RICK 1971), and somewhat lower in male than in female gametes (DEVICENTE and TANKSLEY 1991; VAN OIJEN et al. 1994).

Hybrids between cultivated tomato and the related nightshades Solanum lycopersicoides or Solanum sitiens (syn. S. rickii) are highly sterile, due at least in part to reduced chromosome pairing and recombination (RICK 1951; DEVERNA et al. 1990). A comparative linkage map of the S. lycopersicoides / S. sitiens genome shows it is mostly colinear with the genomes of species in the tomato clade, but with one chromosome arm (10L) involved in a paracentric inversion.
Recombination in the F1 $L. \text{esculentum} \times S. \text{lycopersicoides}$ hybrid is reduced genome-wide by about $\sim 27\%$ relative to other tomato maps (Chetelat et al. 2000). While the inversion accounts for part of this reduction – recombination is completely blocked in genotypes heterozygous for the inverted region – the genome-wide effects must be due in part to excessive sequence divergence between the parental species. The $L. \text{esculentum}$ and $S. \text{lycopersicoides}$ genomes are readily distinguished by genomic in situ hybridization, and differ in the copy number and/or locations of certain repetitive DNA elements (Ji et al. 2004). Differences were also found between them with respect to chromosome size and timing of condensation (Menzel 1962; Rick et al. 1986). Allotetraploid hybrids show preferential pairing of homologous chromosomes, with complete bivalent formation, and consequently greater fertility than $2x$ hybrids (Menzel 1964), suggesting the genomes of $S. \text{lycopersicoides} / S. \text{sitiens}$ are homeologous (partially homologous) with that of cultivated tomato.

Heterozygous substitution lines, in which one tomato chromosome is substituted with one $S. \text{lycopersicoides}$ homeologous chromosome, recombine at less than 50% of the rate observed in the F1 interspecific hybrid, indicating strong background effects (Ji and Chetelat 2003). Recombination is even lower (0-2% of normal) in corresponding monosomic addition lines, in which the $S. \text{lycopersicoides}$ chromosome is present as an extra (i.e. $2n+1$), suggesting that exchange between homeologous chromosomes is antagonized by homologous associations.

We recently reported synthesis of a set of $S. \text{lycopersicoides}$ introgression lines in the background of cultivated tomato (Canady et al. 2005). Each line contains one to several donor segments. Approximately 96% of the nightshade genome is captured in 56 such lines. In the present study, homeologous segments from different regions of the genome were compared with respect to their tendency to recombine with the tomato chromosomes. Several properties of
introgressions that might affect the rate of homeologous recombination, such as segment length and position within the chromosome, were examined. Double introgression lines of various types were synthesized to evaluate their potential for increasing recombination frequency in a region of interest. These experiments tested the importance of two principle variables affecting homeologous recombination in interspecific hybrids: the preference for homologous exchange within the same chromosome, and the degree of sequence divergence between introgressed segments and the recipient genome.

**MATERIALS AND METHODS**

**Plant material:** A group of 42 *S. lycopersicoides* introgression lines and 2 substitution lines were used in this study (Table 1). The parental genotypes and breeding strategy used for selecting the lines were described previously (CANDY et al. 2005; Ji and CHETELAT 2003). Briefly, both types of prebreds were derived from *S. lycopersicoides* accession LA2951 by backcrossing to *L. esculentum* cv. VF36. The lines used in the current study cover most of the genome. Several genomic regions are excluded from this analysis, including regions of chromosomes 2, 3 and 4 for which no introgressed segments were recovered, and chromosomes 5, 9 and 11 for which heterozygous lines were not available.

Double introgression lines were constructed by crossing selected lines with single segments. Heterozygosity for both segments was confirmed with RFLP markers. Some *S. lycopersicoides* introgression lines were also crossed to *L. pennellii* (syn. *S. pennellii*) accession LA0716, a self-compatible and completely homozygous strain of this wild species. Others were crossed to stocks containing single introgressed segments from *L. pennellii* in the background of *L. esculentum* cv. M82, developed by ESHED and ZAMIR (1995) and LIU and ZAMIR (1999).
Seeds of all plant materials used in this study were obtained from the C.M. Rick Tomato Genetics Resource Center at UC-Davis.

Recombination frequency was measured by genotyping F2 (and in a few instances, BC) progeny of heterozygous introgression or substitution lines. F2 seeds were obtained by allowing heterozygotes to self-pollinate, backcross seeds by crossing heterozygotes to ‘VF36’, the background genotype for these lines. Seeds of segregating F2 and BC populations were treated for 30 minutes with 50% household bleach (equivalent to 2.75% sodium hypochlorite), rinsed under running tap water for several minutes, and plated on germination paper in plastic boxes, and incubated at 25ºC under a 12 hr photoperiod. Seedlings with fully expanded cotyledons were transplanted to artificial soil mix in the greenhouse, and were genotyped at the 3-4 true leaf stage.

**Marker analysis:** DNA marker analysis was used to measure recombination within introgressed segments. For each introgressed segment, a minimum of two markers were used, one from each end. Markers, primarily RFLPs, were chosen to mark the ends of each segment, according to their locations on the high density molecular linkage map of tomato (TANKSLEY et al. 1992). For substitutions and longer introgressed segments, more than two markers were used in order to detect multiple crossovers. DNA extractions and RFLP analysis were performed as previously described (CANADY et al. 2005).

**Statistical analysis:** In cases where marker data was available from two or more independent progeny tests of the same heterozygous introgression line, a chi-square test for independence was used to decide if data from the separate tests could be pooled. The maximum likelihood method was used to estimate recombination rates using either LINKAGE-1 (SUITER et al. 1983) or Mapmaker v3.0 (LANDER et al. 1987). The threshold parameters used for detecting
linkage were a chi-square test with $P<0.05$ for LINKAGE-1 or a LOD $\geq 3.0$ and recombination fraction $\leq 30\%$ for Mapmaker. Genetic distances in centimorgans (cM) were calculated from recombination fraction estimates using the Kosambi mapping function. A Student’s $t$ test was used to make pair-wise comparisons of mean recombination values in peri- vs. paracentric, and terminal vs. interstitial *S. lycopersicoides* segments.

Expected values for recombination rates were obtained from the genetic distances in the corresponding marker intervals on the RFLP map of tomato. Based on an interspecific cross of *L. esculentum* × *L. pennellii*, the reference map is not, strictly speaking, a control for purely homologous recombination rates. However, due to the lack of marker polymorphism within the cultivated genepool, the interspecific map provides the best available estimates of recombination rates useful for comparisons across different mapping populations. Furthermore, *L. esculentum* × *L. pennellii* hybrids are fertile, and their chromosomes pair normally during meiosis (KUSH and RICK 1963), indicating they are functionally homologous at this level. Recombination frequencies in the double introgressions were compared to the single segment controls using chi-square contingency tests on the numbers of parental vs. recombinant progeny. To compare $F_2$ with BC populations, the chi-square test was carried out on the numbers of parental vs. recombinant gametes rather than individual plants.

**RESULTS**

**Recombination in single introgression lines:** Estimates of homeologous recombination frequencies were obtained from $F_2$ and BC progeny of 42 different introgression and substitution lines, representing 11 out of 12 *S. lycopersicoides* chromosomes. The total genetic length of the marker intervals for which these lines were heterozygous (i.e. in which recombination could be
measured) constituted approx. 68% of the map units in the tomato genome. Most of the lines were nearly isogenic, i.e. contained a single introgressed segment, while a small number contained one to several extra segments on other chromosomes (Canady et al. 2005). Certain introgressions on chromosomes 3, 9 and 12 were homozygous for S. lycopersicoides markers at one end of the introgressed region or at the other end of the chromosome (Figure 1).

Recombination between the L. esculentum chromosome and the introgressed S. lycopersicoides segments was greatly reduced, in most cases to only 0-10% of the expected values (Table 1, Figure 2). This reduction was genome-wide, occurring on all tested chromosomes. Double crossover genotypes were extremely rare, indicating a strong crossover interference in most cases (Table 1). Crossovers were approximately randomly distributed within the introgressed segments (Figure 1). Intervals between more distant markers (on the tomato RFLP map) generally had more recombination events, as expected. However, for introgressed segments in which more than one marker interval was tested, there seemed no preference for crossovers in distal vs. proximal regions (Figure 1). Despite the overall lower level of homeologous recombination, crossover events were recorded in most regions of the genome, with a few exceptions, such as chromosome 10 and the short arm of chromosome 6 (Figure 1).

In general, lines with ‘longer’ segments (i.e. greater map units on the RFLP map) recombined more frequently than those with shorter segments, as might be expected (Table 1). Significantly, this trend was true even when recombination frequencies were normalized for expected genetic length based on the RFLP map. In fact, the highest ‘relative’ recombination frequencies were observed in the substitution lines, which were heterozygous for an intact S. lycopersicoides chromosome. A positive correlation was observed between the expected genetic
lengths of introgressed segments and the observed relative rates of homeologous recombination in the introgressed region (Figure 2). This correlation was also observed within individual chromosomes. For instance, on chromosomes 7 and 8, the highest recombination frequencies were observed in the whole chromosome substitutions. Introgression lines with relatively long homeologous segments (>50% of the length of the chromosome) were intermediate, and those with short segments recombined at the lowest rates (Table 1, Figure 2).

A few exceptions to these trends were noted. Line LA4234, which contained a fairly short *S. lycopersicoides* introgression (31.9 cM on the tomato map) on chromosome 1, recombined at a rate higher than average (34.2% of expected) (Table 1). In contrast, no recombination was detected in line LA4242, which contained a longer introgressed segment (over 50cM) on chromosome 3. As expected, no recombination was observed in LA4276, which contained a segment spanning the whole long arm of chromosome 10; recombination in this region is prevented by a paracentric inversion in *S. lycopersicoides* relative to *L. esculentum*, and thus is irrelevant to the relationship between length and recombination frequency.

Average recombination estimates were calculated for introgressions according to the relative positions of *S. lycopersicoides* and *L. esculentum* segments within the chromosomes (Table 2). A chromosome segment was considered ‘terminal’ if it included one end of the chromosome, or ‘interstitial’ if it did not. Alien segments that spanned the centromere (‘pericentric’) were distinguished from those that were limited to one arm (‘paracentric’). On average, terminal segments tended to recombine at a higher rate than interstitial ones, however the difference was not significant (*t* = 1.42, *P*<0.1). Although pericentric segments showed a higher recombination rate than paracentric ones, the former were also longer on average. Thus any difference in recombination frequency between these two categories is confounded by
segment length in addition to position effects. As mentioned previously, the substituted chromosomes recombined at higher rates than any of the introgressed segments.

Recombination in ‘target / driver’ introgression lines: The positive correlation we observed between recombination frequency and the length of *S. lycopersicoides* segments suggested a possible strategy to increase the probability of recovering recombinants. By combining two introgressed segments from different regions of the same chromosome, the total length of homeology could be increased (Figure 3A). We refer to such a double introgression as a ‘target / driver’ genotype, wherein the ‘target’ segment would contain a gene of interest (e.g. a disease resistance locus), around which recombinants are desired. The ‘driver’ segment would contain a different *S. lycopersicoides* introgression, preferably on the opposite chromosome arm. In the doubly heterozygous F1 hybrid, these two segments would initially be oriented in repulsion phase. Recombinants would be selected in F2 progeny by marker analysis. In theory, two overlapping terminal segments combined in this fashion would be similar or equivalent to a substitution line, except for their linkage phase. To test this hypothesis, we made 4 pairs of double introgression lines combining *S. lycopersicoides* segments on chromosomes 1, 2, and 7 (Figure 4). These represented several different configurations of the two segments: two interstitial segments on the same arm (chromosome 1), one interstitial and one terminal on the same arm (chromosome 2), and two terminal segments on opposite arms, with either a small or a large region of homology between them (chromosome 7). The controls for the target / driver genotypes were the corresponding single segment introgression lines.

The chromosome 1 and 2 target / driver combinations showed some evidence of increased homeologous recombination relative to the single segment controls (Figure 4A, 4B). For each chromosome, one of the two *S. lycopersicoides* segments (1A and 2A, respectively)
recombined at higher rates than the controls, but only in the BC populations, not the F2’s. For segment 1A, an increase of over 6× was observed ($\chi^2 = 4.37, P<0.05$), and for segment 2A, recombination increased from 0 to 2.8cM ($\chi^2 = 7.84, P<0.01$). Interestingly, it was the more proximal segment on both chromosomes that showed the increase, while the distal segments showed little or no change. Both double introgression lines for chromosome 7 showed elevated recombination frequencies for the segments on the long arm in each pair, 7B and 7C (Figure 4C). Recombination within the 7B segment increased by approx. 3× in the target / driver genotypes relative to the single segment control ($\chi^2 = 5.6, P<0.025$ for the F2 and $\chi^2 = 5.4, P<0.025$ for the BC population). Recombination rate in the 7C segment increased from 0 in the control, to up to 1.3cM in the F2 combination stock ($\chi^2 = 4.2, P<0.05$). The frequency of recombination within the longer segment 7B was approx. 10× higher than that observed in the shorter segment 7C. This is consistent with our observation of lower recombination in short than in long introgressed segments (Figure 2).

A more pronounced increase in recombination was observed in the interval between the introgressed segments, i.e. in the intercalary stretch of homology (Figure 4). Recombination frequencies in these ‘gaps’ could be estimated because each target / driver combination (e.g. 1A+1B) was heterozygous for markers flanking the homozygous interval. In contrast, the lines with a single homeologous segment provided no information on recombination in these regions, because they were heterozygous for markers on one side only. We therefore used genetic distances between the same markers on the reference map of tomato (from F2 L. esculentum × L. pennellii) as ‘controls’. In each case, recombination rate in these homozygous regions between the paired segments was increased relative to the reference map. For example, on chromosome 1, the length of the TG71 -- TG83 interval increased from 19.6cM (from the reference map) to
approx. 50cM in the double segment lines. In half of the mapping populations, the increase in recombination within these gaps was so great that linkage between the flanking markers could not be detected. These suggest that reduced recombination within regions of homeology (i.e. S. lycopersicoides segments) is compensated by an increase in recombination within adjacent – in this case intercalary – regions of homology.

This effect appears to be due in part to selection towards some recombinants genotypes. Significant segregation distortion was observed in many of the target / driver genotypes (Table 3). In every case, the recombinant classes showed an excess of genotypes that had lost one segment through recombination (i.e. +-+), and a deficiency of genotypes that had gained a segment (i.e. A-B in coupling phase). The most pronounced distortion of this type was on chromosome 1, where the BC population produced 47 +-+ vs. 0 A-B recombinants. On chromosome 7, the ratio of +-+ to A-B recombinants was as high as 4:1.

Recombination between S. lycopersicoides and L. pennellii: The reduced recombination we observed between tomato chromosomes and the homeologous S. lycopersicoides segments likely result from excessive sequence divergence between the two genomes. We therefore hypothesized that S. lycopersicoides introgressed segments might recombine more readily with DNA from a species of closer sequence similarity. To test this concept, we crossed an introgression line containing a S. lycopersicoides segment on chromosome 2, with L. pennellii, a wild species that is phylogenetically intermediate between tomato and the nightshade (Figure 5). In the genetic background of L. esculentum, this introgressed region recombined at only 4.7% of the expected value. However, in F2 progeny of the introgression line × L. pennellii hybrid, the total genetic length of the S. lycopersicoides segment was increased to 21cM, an approx. 10× increase, but still less than the homologous
recombination rate inferred from the F₂ L. esculentum × L. pennellii map. This result suggests that S. lycopersicoides chromosome segments recombine more readily with L. pennellii than with L. esculentum DNA. Interestingly, recombination along the rest of the chromosome, i.e. between L. esculentum and L. pennellii DNA, was slightly increased compared to the reference map, particularly in the intervals flanking the S. lycopersicoides segment. This provides further evidence that reduced recombination within a region of homeology is accompanied by increased crossing-over in the adjacent homologous regions.

However, the increased recombination between the S. lycopersicoides segment and the L. pennellii chromosome could result from genetic background effects. For example, there might be genes affecting pairing or recombination on other L. pennellii chromosomes (i.e. other than chromosome 2, in this case). Alternatively, there could be a genome-wide enhancement of recombination due to the interspecific nature of the hybrids with L. pennellii. Finally, from a breeding standpoint, crossing prebreds already in a cultivated tomato background to pure L. pennellii would be an inefficient method to increase recombination since extensive backcrossing and selection would be needed to eliminate the genome of the latter species.

To address these issues, we took advantage of a similar set of introgression lines containing L. pennellii segments in the background of cultivated tomato. We crossed lines containing a segment from S. lycopersicoides to those containing one from L. pennellii on the same chromosome (Figure 3B). We refer to the L. pennellii introgressions in this context as ‘bridging introgressions’ to reflect their phylogenetic connection to both the nightshade and cultivated tomato. If the rate of homeologous recombination is primarily limited by the degree of sequence homology, then the bridging genotypes should increase recombination relative to the single segment controls.
To test this hypothesis, we measured recombination within a segment from *S. lycopersicoides* chromosome 7 (LA4259), either alone or in combination with three different *L. pennellii* segments on the same chromosome (Figure 6). To eliminate the effect of differences in the genetic background between the two sets of introgressions – ‘M82’ was used for the *L. pennellii* and ‘VF36’ for the *S. lycopersicoides* derivatives – each single introgression was crossed to the parent of the other set so that all recombination measurements were made in a constant genetic background, equivalent to VF36 × M82. Recombination rates were normalized to expected frequencies from the reference map of tomato in order to compare relative recombination rates across different marker intervals.

Alone, the *S. lycopersicoides* segment in LA4259 recombined at only 2.4% of the expected rate. Recombination within the single segment *L. pennellii* line was higher -- 10 to 15% of normal -- consistent with the closer homology between *L. pennellii* and cultivated tomato. In the LA4259 × IL7-1 hybrid, recombination within the homeologous *S. lycopersicoides* segment increased by over 6×, to 18% of normal, but only in the region of overlap with *L. pennellii* DNA; no recombination events were recovered in the non-overlapping region. In the LA4259 × IL7-3 cross, recombination was increased to 51% of normal in the overlapping region between *L. pennellii* and *S. lycopersicoides* segments; again, little or no change in the recombination rate was detected in the flanking regions. The last combination tested, LA4259 × IL7-4-1, consisted of two non-overlapping segments on opposite arms of chromosome 7. Recombination was increased in both the *S. lycopersicoides* and the *L. pennellii* segments – to 27% of normal in the former, 39% in the latter – relative to their single segment controls. These results suggest that in addition to the role of sequence homology, the bridging
introgression lines may elevate homeologous recombination by a mechanism similar to that of the target/driver genotypes.

**DISCUSSION**

Genetic improvement of crop plants has depended to a large extent on disease resistances and other novel traits originating in wild relatives. Various types of prebred stocks have been used for this purpose in different crop plants. Sets of introgression lines representing whole genomes of related wild species, though time consuming to synthesize, provide permanent resources for mapping projects, and superior starting material for breeding programs (ZAMIR 2001). However, a potential disadvantage is that recombination between alien (homeologous) segments and the recipient genome can be reduced.

In tomato, recombination within segments derived from *L. pennellii* and *L. hirsutum* is typically about 15-30% of normal levels (ALPERT and TANKSLEY 1996; MONFORTE and TANKSLEY 2000; VAN WORDRAGEN *et al.* 1996). The recombination rates we observed for *L. pennellii* chromosome 7 introgression lines also fell into this range. Previous studies using morphological markers had shown that recombination between *L. esculentum* and *L. pennellii* chromosomes drops off during successive backcross generations (RICK 1969, 1971). Thus one of the advantages of introgression lines — their more uniform genetic background — is a potential obstacle when searching for recombinants. This problem is exacerbated in genomic regions subject to low recombination rates. The pericentromeric regions of each tomato chromosome have lower than average recombination rates per unit of physical distance (SHERMAN and STACK 1995; TANKSLEY *et al.* 1992). As a result, genes located near the centromeres, such as the nematode resistance gene *Mi* (KALOSHIAN *et al.* 1998), can be difficult to isolate by positional cloning. In contrast, genes located in recombination hotspots are more amenable to map-based
cloning. For example, recombination in the region of the soluble solids QTL $\text{Brix9-2-5}$ was so high that the effects of the $L. \text{pennellii}$ allele could be mapped to a single amino acid (Fridman et al. 2004). From a breeding standpoint, linkage drag can make it difficult to combine tightly linked resistance genes from different sources (i.e. originally in trans configuration) into a single inbred parent (i.e. in cis orientation).

**Recombination in the introgression lines is limited by sequence divergence:** In the present study, we observed a genome-wide reduction in recombination frequencies within introgressed $S. \text{lycopersicoides}$ segments, often to as low as 0-10% of normal levels. These values are generally lower than previously reported for similar $L. \text{pennellii}$ or $L. \text{hirsutum}$ derivatives, consistent with molecular systematic studies which indicate that $S. \text{lycopersicoides}$ is more distantly related to cultivated tomato (Peralta and Spooner 2001; Spooner et al. 2005). Evidence from other model systems, including bacteria (Shen and Huang 1986), yeast (Datta et al. 1996) and Arabidopsis (Li et al. 2006), among others, has clearly demonstrated that recombination is strongly dependent on the degree of sequence identity, and can be inhibited by as little as a single nucleotide mismatch. In a sample of coding and noncoding sequences, the nucleotide divergence between $L. \text{esculentum}$ and $L. \text{pennellii}$ varied from 0 – 6.3% (Nesbitt and Tanksley 2002). Our analysis of published waxy gene sequences (from Peralta and Spooner 2001) found approx. 6% divergence between $L. \text{esculentum}$ and $S. \text{lycopersicoides}$, compared to 2% between $L. \text{pennellii}$ and $L. \text{esculentum}$ (data not shown). These data provide further evidence that recombination between tomato chromosomes and orthologous segments introgressed from wild relatives is strongly influenced by their degree of sequence homology.

The above considerations do not take into account potential disruptions in chromosomal synteny that might differentiate these species and could potentially suppress recombination
independent of nucleotide divergence. In hexaploid wheat, for example, the homeologous A, B and D genomes differ by gene duplications and deletions (AKHUNOV et al. 2003); the B genome showed the greatest loss of synteny, which may explain why this genome undergoes less pairing with homeologous chromosomes of the A and D genomes (in the absence of Phl) than A and D homeologues with each other. Larger scale rearrangements, such as chromosomal inversions and translocations, would be expected to strongly suppress recombination. Comparative linkage maps have so far revealed few rearrangements among species within the Lycopersicon clade. One exception is a paracentric inversion involving part of the short arm of chromosome 7 in *L. pennellii* relative to *L. esculentum* (VAN DER KNAAP et al. 2004). This inverted region is located within the introgressed *L. pennellii* segment in the IL 7-4-1 line, which was included in the present study. This line recombined at approx. 15% of normal, similar to the other *L. pennellii* derivatives. However, our reference for the expected recombination frequencies was the high density RFLP map of tomato (from TANKSLEY et al. 1992), which was based on F2 *L. esculentum × L. pennellii*. Thus, any effect of the chromosome 7 inversion would be factored into the reference map, so that the relatively lower recombination observed for the same marker interval in IL 7-4-1 should be due to other factors, such as sequence divergence or genetic background.

The position of introgressed segments within chromosomes appears to play a relatively minor role. Average recombination rates were slightly higher in terminal and pericentric segments than in interstitial and paracentric segments, respectively. However these trends were not statistically significant. A more pronounced difference was observed between homeologous segments, irrespective of their position within the chromosome, and the substitution lines, which contained intact *S. lycopersicoides* chromosomes. The latter had much higher recombination rates than observed in introgressed segments of the same wild species chromosomes. These
results are consistent with previous research on wheat in which substitution lines containing whole *Triticum monococcum* chromosomes recombined at higher rates than the segmental lines, but less than half of the homologous recombination frequency (LUO et al. 2000). Thus, in both wheat and tomato, recombination rates are determined by the level of sequence divergence within a region of homeology, and whether it is contained within an intact alien chromosome or a segmental introgression.

**Recombination within homeologous vs. homologous regions:** We detected a positive correlation between the rate of homeologous recombination, expressed as a percentage of the expected value, and the length of the introgressed segments on the genetic map of tomato. This correlation could be due to a ‘preference’ for recombination within homologous over homeologous regions of the chromosome. Our data suggest a process whereby chromosomes are scanned for homology, and crossovers allocated based on the degree of similarity. Such a process could involve the DNA mismatch repair system, which restricts recombination between homeologous sequences in other model systems, such as *E. coli* (ZAHRT and MALOY 1997), yeast (DATTA et al. 1996) and Arabidopsis (LI et al. 2006).

However, our results might also be influenced by purely stochastic processes. Chromosomes with relatively long introgressions contain shorter regions of homology, thus a higher probability of crossovers occurring in the homeologous segments. Another factor that might contribute to the observed correlation is gametic selection. A failure to form at least one crossover, either in a region of homology or homeology, would result in unpaired chromosomes during the first meiotic division. Pairing failure, in turn, would lead to unbalanced gametes, which, in the case of deficiencies, are not viable during gametogenesis of tomato, and in the case of duplications, would be less competitive during pollination. The result would be to increase
the overall frequency of recombinant progeny relative to the rate of crossing-over. We previously quantified the rate of pairing failure, as indicated by univalent formation, in heterozygous substitution lines, and found that it had a relatively small effect on recombination estimates (Ji and Chetelat 2003).

Segregation distortion, a common feature of interspecific crosses and their derivatives, can potentially bias recombination estimates in some circumstances (Liu 1998). In practice this has not prevented construction of high resolution genetic maps in tomato, most of which -- due to limited polymorphism in the cultivated genepool -- are based on interspecific crosses. We previously described the inheritance of S. lycopersicoides introgressions in progeny of heterozygotes (Canady et al. 2005). For many regions, we observed a deficiency of plants homozygous for the S. lycopersicoides segments, indicating selection against alleles of the wild species during gametogenesis, pollination, and/or zygote development. In this context, recombinant progeny should have a selective advantage since they contain shorter S. lycopersicoides segments, with potentially fewer genes subject to selection. This might contribute to the higher recombination rates observed in long vs. short introgressions. However, the cases of non-Mendelian transmission we observed in the introgression lines could be explained by a small number of segregation distorter loci, no more than one or two per chromosome (Canady et al. 2005). Also, short segments were just as likely as long ones to be subject to these effects. We conclude that the effects of selection might bias some recombination estimates, but probably cannot account for the genome-wide recombination reduction, or the correlation with segment length that we observed.

**Recombination in double vs. single introgressed segments:** Our observations of higher recombination rates in lines with intact S. lycopersicoides chromosomes or long
introgressed segments led us to construct double introgression lines containing two segments on opposite chromosome arms. These target / driver genotypes were designed to increase the total length of homeology, and thereby reduce the opportunity for homologous recombination. Surprisingly, the compound stocks resulted in only modest increases in recombination within the homeologous segments, much less than values obtained for the corresponding substitution lines. At the same time, recombination in the region of homology between paired \textit{S. lycopersicoides} segments was greatly enhanced.

One factor that may explain these observations is linkage phase: the target / driver segments were oriented in repulsion phase, in contrast to the substitutions which contain a single \textit{S. lycopersicoides} chromosome (i.e. all markers in coupling phase). For paired segments in repulsion, the parental chromosomes each contain a single alien introgression, and therefore fewer \textit{S. lycopersicoides} genes that could be under negative selection than gametes that acquire all or part of both homeologous segments as a result of a crossover within either. On the other hand, recombination in the homologous interval between segments would result in some gametes containing no \textit{S. lycopersicoides} genetic material, which should have few if any detrimental effects (i.e. reduced linkage drag). This interpretation is consistent with our observations of a bias towards recombinant genotypes that involve a loss of one or both alien segments, relative to those that gained a segment. Target / driver genotypes oriented initially in coupling phase should be less subject to this bias, since a crossover anywhere within or between either segment would produce gametes with less alien genetic material. This prediction could be tested by comparing recombination in coupling vs. repulsion phase stocks.

Another factor that may explain the lower recombination in target / driver introgressions than in substitution lines is the presence in the former of an intercalary region of homology
between the pair of homeologous segments. This ‘gap’ provides the opportunity for crossovers between perfectly homologous sequences, which would compete against crossovers within the homeologous segments. Assuming recombination occurs preferentially in regions of homology, then crossover interference would limit the number of recombination events elsewhere on the chromosome, including within either of the introgressed segments. This leads to the prediction that homeologous recombination frequency would be highest if the paired segments cover all or most of the chromosome (i.e. with the shortest possible gap). This is what we observed with the two sets of chromosome 7 target / driver lines.

**L. pennellii introgressions can increase recombination in a target region:** Our data on recombination in *S. lycopersicoides* introgression and substitution lines generally support the hypothesis that homeologous recombination can be enhanced by reducing the opportunity for homologous interaction elsewhere on the same chromosome. Another strategy we explored was to increase the level of sequence homology vis-à-vis the *S. lycopersicoides* segment by introducing an overlapping introgression from *L. pennellii*. Recent molecular systematic studies suggest that *L. pennellii* is a basal taxon in the Lycopersicon clade, phylogenetically intermediate between *L. esculentum* and *S. lycopersicoides* (PERALTA and SPOONER 2001; SPOONER et al. 2005). In agreement with the recent molecular phylogenies, the earliest taxonomic descriptions of *L. pennellii* by CORRELL (1958) highlighted its distinctive anther morphology, which displays characteristics of both Solanum (lack of a sterile tip) and Lycopersicon (longitudinal dehiscence). It may also be significant that *L. pennellii* is the only species in the tomato clade that can be experimentally hybridized with both *L. esculentum* and *S. lycopersicoides* (RICK 1979). Further evidence of these genetic relationships is that *L. pennellii* can serve as a ‘bridge’
to overcome the unilateral incompatibility of *S. lycopersicoides* and facilitate introgression of traits into cultivated tomato (Chetelat and DeVerna 1991; Rick et al. 1988).

For these reasons, we hypothesized that the *L. pennellii* might recombine readily with both the *S. lycopersicoides* and *L. esculentum* chromosomes. In support of this concept, we found that recombination within a *S. lycopersicoides* segment on chromosome 2 increased nearly 10× in the hybrid with *L. pennellii*. However, in this interspecific cross, the effect of sequence homology within the tested chromosome is confounded by the influence of overall genetic background. These background effects are not insignificant; for example, the chromosomes of *L. pennellii* recombine readily with those of *L. esculentum* in the F₁ interspecific hybrid, yet once bred into cultivated tomato, recombination between them is greatly reduced (Rick 1969; 1971).

To eliminate the influence of genetic background, we constructed double introgression lines containing *S. lycopersicoides* and *L. pennellii* segments on the same pair of chromosomes (bridging introgressions). We observed that recombination within the *S. lycopersicoides* segment was substantially elevated within the region of overlap with the *L. pennellii* segment. Outside the overlap region, recombination was relatively unaffected. These data provide further evidence that the level of sequence homology within the introgressed segment exerts a strong influence on recombination rate, consistent with data from other systems. However, factors unrelated to DNA sequence homology of introgressed segments may also play a role. For example, chromosomes of the parental species differ cytologically in their pericentromeric heterochromatin (Khush and Rick 1963; Menzel 1962), and DNA packaged as heterochromatin is known to be less recombinogenic than euchromatin (Sherman and Stack 1995). Thus differences between the two introgressed segments that affect chromatin packaging or the location of heterochromatin/euchromatin boundaries could influence recombination.
Surprisingly, recombination was also enhanced in a double introgression line that had the *L. pennellii* and *S. lycopersicoides* segments on opposite arms of the same chromosome (i.e. no overlap). This result presumably reflects the same processes – either during meiotic recombination, or selection during gametophytic or sporophytic phases – that were responsible for similar results from the target / driver genotypes.

**Conclusions and outlook:** Several potential practical applications emerge from our studies with the *S. lycopersicoides* derivatives. First, long introgressed segments or substituted chromosomes are a richer source of recombinants than short ones. Therefore, in order to reduce linkage drag associated with wild species introgressions, the most expeditious – and somewhat counterintuitive – method would be to start with very large original segments containing the gene of interest, select for single crossovers close to the target locus, and only then generate secondary recombinants on the other side. Sears (1977) made similar recommendations for wheat, and pointed out that segments with crossovers on opposite sides could be allowed to recombine, resulting in the shortest possible introgressions. Second, recombination within a homeologous segment can be increased by constructing a double introgression with greater homology. In the present experiments, segments from *L. pennellii* recombined readily with both *S. lycopersicoides* and *L. esculentum*, however we have not tested similar stocks from other wild relatives. Thus, the source of the bridging segments might be manipulated in order to maximize recombination frequency. Thirdly, the *S. lycopersicoides* segments could be used to suppress recombination on one arm of a chromosome in order to increase crossover frequency within a target introgression (e.g. from more closely related species) on the other arm.

A significant disadvantage of these chromosome engineering strategies is the extra time involved in constructing compound stocks then eliminating residual genetic material in later
generations. Situations that might justify the extra effort include genes located in regions of suppressed recombination (e.g. near centromeres), or map-based cloning projects where the ordering of genes, not variety development, is the primary goal. In other cases, a ‘brute force’ screening for rare recombinants using high throughput marker technologies might be more efficient.

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Table 1. Frequency of parental and recombinant genotypes in progeny of *S. lycopersicoides* introgression lines.

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<th>Position&lt;sup&gt;b&lt;/sup&gt;</th>
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<th>SCOs</th>
<th>DCOs</th>
<th>Obs.</th>
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<td>F2</td>
<td>29</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>13.8</td>
<td>0</td>
</tr>
<tr>
<td>12</td>
<td>LA4283</td>
<td>TG111 – CT156</td>
<td>I/Pa</td>
<td>F2</td>
<td>30</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>25.0</td>
<td>0</td>
</tr>
<tr>
<td>12</td>
<td>LA4282</td>
<td>TG180 – TG111</td>
<td>T/Pe</td>
<td>F2</td>
<td>60</td>
<td>15</td>
<td>1 (0)</td>
<td>12.1</td>
<td>47.1</td>
<td>25.69</td>
</tr>
</tbody>
</table>

*a* Represent end markers on each introgressed segment.

*b* I = interstitial, T = terminal, Pa = paracentric, Pe = pericentric, S = substituted chromosome

*c* BC♀ = backcross, heterozygote used as female parent

*d* NR = nonrecombinant, SCO = single crossover genotypes, DCO = double crossover genotypes (in parentheses, the # of DCOs that must have occurred in the same gamete).

*e* Expected recombination rate for the same marker interval from the reference map of tomato (*Tanksley et al*. 1992).
Table 2. Average recombination frequencies within *S. lycopersicoides* introgressed segments according to their positions within chromosomes relative to telomeres and centromeres.

<table>
<thead>
<tr>
<th>Position Relative to</th>
<th>No. Lines (n)&lt;sup&gt;c&lt;/sup&gt;</th>
<th>Exp. Recomb. (cM)&lt;sup&gt;d&lt;/sup&gt;</th>
<th>Obs. Recombination (cM) (% of expected)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Telomere&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Interstitial</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Paracentric</td>
<td>8</td>
<td>22.9</td>
<td>0.48 (1.6)</td>
</tr>
<tr>
<td>Pericentric</td>
<td>5</td>
<td>50.4</td>
<td>2.6 (4.1)</td>
</tr>
<tr>
<td>Either</td>
<td>13</td>
<td>33.5</td>
<td>1.3 (2.6)</td>
</tr>
<tr>
<td>Terminal</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Paracentric</td>
<td>15</td>
<td>27.0</td>
<td>1.3 (4.6)</td>
</tr>
<tr>
<td>Pericentric</td>
<td>10</td>
<td>43.7</td>
<td>4.0 (8.4)</td>
</tr>
<tr>
<td>Either</td>
<td>25</td>
<td>33.7</td>
<td>2.4 (6.1)</td>
</tr>
<tr>
<td>Substitution</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pericentric</td>
<td>4</td>
<td>93.4</td>
<td>31.8 (34.1)</td>
</tr>
<tr>
<td>All</td>
<td>--</td>
<td>42</td>
<td>4.9 (7.7)</td>
</tr>
</tbody>
</table>

<sup>a</sup>Terminal = includes one end of the chromosome, Interstitial = does not include a chromosome end; Substitution = an intact alien chromosome.

<sup>b</sup>Pericentric = includes the centromeric region, Paracentric = does not include the centromere.

<sup>c</sup>Number of lines in each category.

<sup>d</sup>Expected recombination rates are estimated as the length of introgressed segments on the reference map (Tanksley et al. 1992).
Table 3. Segregation in progeny of double introgression lines containing two *S. lycopersicoides* segments on the same chromosome pair, oriented in repulsion phase, and corresponding single segment control lines.

<table>
<thead>
<tr>
<th>Chrom.</th>
<th>Cross (♀ × ♂)a</th>
<th>Line</th>
<th>A</th>
<th>+ B</th>
<th>A B</th>
<th>+ B</th>
<th>++</th>
<th>A</th>
<th>+ B</th>
<th>A B</th>
<th>A B</th>
<th>Exp.</th>
<th>1:2:1</th>
<th>χ²d</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Control F₂ (1A/+) × self LA4296</td>
<td>20</td>
<td>--</td>
<td>--</td>
<td>16</td>
<td>53</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>89</td>
<td>1:2:1</td>
<td>3.60ns</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>Control F₂ (1B/+) × self LA4233</td>
<td>--</td>
<td>--</td>
<td>1</td>
<td>6</td>
<td>--</td>
<td>45</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>52</td>
<td>1:2:1</td>
<td>28.7***</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>F₂ (1A ++ 1B) × self LA4233</td>
<td>3</td>
<td>11</td>
<td>3</td>
<td>1</td>
<td>10</td>
<td>5</td>
<td>2</td>
<td>8</td>
<td>0</td>
<td>43</td>
<td>1:2:1</td>
<td>1.47ns</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>BC (1A ++ 1B) × ++</td>
<td>--</td>
<td>0</td>
<td>--</td>
<td>47</td>
<td>39</td>
<td>42</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>128</td>
<td>1:1</td>
<td>0.049ns</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Control F₂ (2A/+) × self LA4237</td>
<td>13</td>
<td>--</td>
<td>--</td>
<td>32</td>
<td>49</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>94</td>
<td>1:2:1</td>
<td>7.85***</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Control F₂ (2B/+) × self LA4239</td>
<td>--</td>
<td>--</td>
<td>16</td>
<td>44</td>
<td>--</td>
<td>52</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>112</td>
<td>1:2:1</td>
<td>14.6***</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>F₂ (2A ++ 2B) × self LA4239</td>
<td>2</td>
<td>25</td>
<td>9</td>
<td>1</td>
<td>15</td>
<td>13</td>
<td>5</td>
<td>11</td>
<td>0</td>
<td>81</td>
<td>1:2:1</td>
<td>6.52*</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>BC (2A ++ 2B) × ++</td>
<td>--</td>
<td>17</td>
<td>--</td>
<td>23</td>
<td>20</td>
<td>36</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>96</td>
<td>1:1</td>
<td>4.02*</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>BC ++ × (2A ++ 2B)</td>
<td>--</td>
<td>24</td>
<td>--</td>
<td>33</td>
<td>51</td>
<td>67</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>175</td>
<td>1:1</td>
<td>1.91ns</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cross</td>
<td>Population</td>
<td>A</td>
<td>B</td>
<td>C</td>
<td>D</td>
<td>E</td>
<td>F</td>
<td>Sample Size</td>
<td>Ratio</td>
<td>Significance</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>----</td>
<td>----------------------------------</td>
<td>------------</td>
<td>----</td>
<td>----</td>
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<td>----</td>
<td>-------------</td>
<td>-------</td>
<td>--------------</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>Control F₂ (7A/+) × self</td>
<td>LA4261</td>
<td>4</td>
<td>--</td>
<td>--</td>
<td>43</td>
<td>17</td>
<td>--</td>
<td>64</td>
<td>1:2:1</td>
<td>61.6***</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>Control F₂ (7B/+) × self</td>
<td>LA4259</td>
<td>--</td>
<td>--</td>
<td>20</td>
<td>24</td>
<td>--</td>
<td>52</td>
<td>100</td>
<td>1:2:1</td>
<td>1.00ns</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>Control F₂ (7C/+) × self</td>
<td>LA4258</td>
<td>--</td>
<td>--</td>
<td>37</td>
<td>31</td>
<td>--</td>
<td>41</td>
<td>109</td>
<td>1:2:1</td>
<td>7.35***</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>F₂ (7A ++ 7B) × self</td>
<td></td>
<td>0</td>
<td>8</td>
<td>18</td>
<td>3</td>
<td>4</td>
<td>15</td>
<td>53</td>
<td>1:2:1</td>
<td>28.8***</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>F₂ (7A ++ 7C) × self</td>
<td></td>
<td>1</td>
<td>19</td>
<td>16</td>
<td>10</td>
<td>6</td>
<td>44</td>
<td>118</td>
<td>1:2:1</td>
<td>12.6**</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>BC ++ × (7A ++ 7B)</td>
<td></td>
<td>--</td>
<td>15</td>
<td>--</td>
<td>30</td>
<td>14</td>
<td>82</td>
<td>141</td>
<td>1:1</td>
<td>46.8***</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>BC ++ × (7A ++ 7C)</td>
<td></td>
<td>--</td>
<td>9</td>
<td>--</td>
<td>29</td>
<td>9</td>
<td>40</td>
<td>87</td>
<td>1:1</td>
<td>18.4***</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*The introgressed segments in each line are shown on the maps in Figure 4, where ‘A’ denotes the segment on the short arm, and ‘B’ the segment on the long arm of the same chromosome. The 7C segment on chromosome 7 is considered the ‘B’ segment. The wild type or *L. esculentum* alleles at the corresponding marker loci are indicated by ‘+’. Note: the control, single segment populations were grown separately.

*bSegregation data are the number of plants in each genotypic class; underlined values indicate genotypes with crossovers between the A and B segments.

*c=n=sample size, excluding individuals with recombination within the A or B segments

*dChi-square values test for goodness-of-fit to expected Mendelian ratios, and are based on data in the parental (nonrecombinant) classes only. Significance levels are: ns, not significant; *, P < 0.05; **, P < 0.01; ***, P < 0.001.
Figure Legends

Figure 1. Genetic map of *S. lycopersicoides* introgression lines. The introgressed segments in each line are indicated by solid bars to the right of the chromosome maps. Thicker lines indicate regions homozygous for *S. lycopersicoides* markers. Numbers above the line are the accession identifiers (all ‘LA’ numbers unless otherwise indicated). Dashed lines connecting marker loci to the introgressed segments indicate which markers were evaluated in each line. The location of recombination events detected in progeny are indicated by ‘×’. (An absence of ×’s in a given interval indicates no cross-overs were detected). Recombination data are summarized in Table 1. The distance between markers are from TANKSLEY *et al.* (1992), the position of centromeres (0) from PILLEN *et al.* (1996). The location of a paracentric inversion on chromosome 10 that distinguishes cultivated tomato from *S. lycopersicoides* is shown by a dashed line with double arrow heads (from PERTUZE *et al.* 2002).

Figure 2. Correlation between observed frequencies of homeologous recombination in *S. lycopersicoides* introgression lines (expressed as a percentage of the expected value) and the genetic length of introgressed segments on the tomato map. Observed recombination rates represent the combined map units of all marker intervals in each segment. The expected genetic lengths are the distances between corresponding markers on the RFLP map of tomato (TANKSLEY *et al.* 1992). Included in the correlation are four data points (open circles) representing recombination frequencies in substitution lines containing *S. lycopersicoides* chromosome 7 or 8 (SL-7, SL-8). Note: data on LA4276 are not included because this line carries an inverted segment that does not recombine.
Figure 3. Diagram illustrating the construction of double *S. lycopersicoides* introgression lines, and their possible use for increasing recombination frequency in the progeny. The ‘target’ introgression line contains a homeologous segment with a gene of interest, such as a hypothetical disease resistance factor (R). The ‘driver’ line contains a homeologous segment on the opposite arm of the same chromosome. The ‘bridging’ introgression is a line containing a donor segment from *L. pennellii*. Each double introgression line is heterozygous for two alien segments, initially in repulsion linkage phase, on the same chromosome. The locations of crossover events, predicted to occur preferentially in homologous stretches, are indicated with an ‘×’. Representative recombinant chromosomes that are obtainable in the progeny are shown.

Figure 4. Recombination in ‘target / driver’ double introgression lines for chromosomes 1, 2 and 7. For each chromosome, the location of the two introgressed segments is shown to the right of the reference map, based on recombination in F2 *L. esculentum × L. pennellii* (*F2 esc × pen*). The total map units in each interval is shown to the left of the reference map. Recombination in single introgressed segments served as controls for the corresponding double introgression lines. Linkage estimates were obtained from both F2 and backcross (BC) progeny. The rate of recombination was measured both within the introgressed segments (solid regions, indicating homeology), and in the interval between them (open segments, indicating homology). The total number of individuals genotyped in each progeny array is indicated by ‘n’. The asterisk on the IL-7C map is to indicate that the control data for this marker interval was from a line with a slightly longer segment, extending beyond TG199 to marker TG216 (not shown on the map).
Figure 5. Genetic maps of recombination frequencies within *S. lycopersicoides* introgression lines for chromosome 2 in different genetic backgrounds. (A) Recombination in F2 progeny of introgression line LS38-11 crossed to *L. pennellii*. (B) Genetic distances between the same markers taken from the RFLP map of tomato (TANKSLEY et al. 1992). (C) Recombination in F2 progeny of LA4239 in the background of *L. esculentum* (data from Table 1). Chromosomes are shaded to indicate the genetic distances are in cM, based on the Kosambi mapping function. Map (A) consisted of two linkage groups at the threshold of LOD $\geq 3.0$.

Figure 6. Recombination in ‘bridging’ introgression lines containing introgressed segments from *S. lycopersicoides* and *L. pennellii* on chromosome 7. Dotted lines indicate the marker loci used to measure recombination within each segment and their positions on the RFLP map (TANKSLEY et al. 1992). Recombination frequencies are expressed as the percentage of the expected values for the same marker intervals. An ‘×’ indicates the region to which each recombination value applies. The positions of the *L. pennellii* segments are from LIU and ZAMIR (1999). The controls are the lines containing a single introgressed segment, either from *S. lycopersicoides* (LA4259) or *L. pennellii*. The single segment *L. pennellii* controls were genotyped with the same markers as the corresponding segments in the double introgression stocks. Recombination estimates are based on F2 populations (of size n) corresponding to the crosses indicated above the chromosomes. Single introgression lines were crossed to *L. esculentum* cv.s M82 or VF36 so that all recombination tests were carried out in a constant genetic background, equivalent to F1 VF36 × M82.
Chromosome 2

IL-2A × IL-2B

BC ♂ 2.8 1.7 n=179

BC ♀ 1.0 0 0 n=97

F2 43.0 0 0.6 n=84

unlinked

F2 IL-2A 0 16.9 n=94

F2 IL-2B 2.2 43.1 n=117

F2 esc × pen

38.0

IL-2A

LA4237

TG554

TG308

TG48

TG507

IL-2B

LA4239

n=94

n=179

n=84

n=97

n=117
(A) $F_2$ $IL \times pen.$

(B) $F_2$ $esc. \times pen.$

(C) $F_2$ $IL \times esc.$

$L. esculentum / L. pennellii$

$L. esculentum / S. lycopersicoides$

$L. pennellii / S. lycopersicoides$