

Deletion Mapping of Homoeologous Group 6-Specific Wheat Expressed Sequence Tags

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ABSTRACT

To localize wheat (*Triticum aestivum* L.) ESTs on chromosomes, 882 homoeologous group 6-specific ESTs were identified by physically mapping 7965 singletons from 37 cDNA libraries on 146 chromosome, arm, and sub-arm aneuploid and deletion stocks. The 882 ESTs were physically mapped to 25 regions (bins) flanked by 23 deletion breakpoints. Of the 5154 restriction fragments detected by 882 ESTs, 2043 (loci) were localized to group 6 chromosomes and 806 were mapped on other chromosome groups. The number of loci mapped was greatest on chromosome 6B and least on 6D. The 264 ESTs that detected orthologous loci on all three homoeologs using one restriction enzyme were used to construct a consensus physical map. The physical distribution of ESTs was uneven on chromosomes with a tendency toward higher densities in the distal halves of chromosome arms. About 43% of the wheat group 6 ESTs identified rice homologs upon comparisons of genome sequences. Fifty-eight percent of these ESTs were present on rice chromosome 2 and the remaining were on other rice chromosomes. Even within the group 6 bins, rice chromosomal blocks identified by 1–6 wheat ESTs were homologous to up to 11 rice chromosomes. These rice-block contigs were used to resolve the order of wheat ESTs within each bin.

COMMON wheat (*Triticum aestivum* L.) is an allohexaploid ($2n = 6x = 42$, AABBDD) containing three homoeologous genomes (SEARS 1954). Among

important cereals, the wheat genome with 16,000 Mb is the largest and rice (*Oryza sativa* L.) with 415 Mb is the smallest (ARUMUGANATHAN and EARLE 1991). The wheat genome is ~100 times larger than the model plant Arabidopsis. Even in Arabidopsis only ~45% of the genome represents the gene-containing fraction that is interspersed with noncoding DNA primarily composed of retrotransposon-like repetitive sequences (BARAKAT *et al.* 1997; BENNETZEN *et al.* 1998; SIDHU and GILL 2004).

Estimates for the gene-containing fraction of the wheat genome range from 1 to 5% obtained from the available sequence data comparisons with other plant genomes to 15% by DNA reassociation kinetics experi-

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ments (FLAVELL *et al.* 1974; SANDHU and GILL 2002; SIDHU and GILL 2004). Deletion mapping of ~2000 gene marker loci showed that genes on wheat chromosomes are also unevenly distributed (GILL *et al.* 1996a,b; FARIS *et al.* 2000; SANDHU *et al.* 2001; SANDHU and GILL 2002; AKHUNOV *et al.* 2003a,b). About 30% of the wheat genome appears to contain >85% of the genes (ERAYMAN *et al.* 2004). The remaining 70% of the genome, present as large blocks interspersed by the gene-rich regions, appears to be gene empty. Therefore, targeting the expressed portion of the genome is particularly important for wheat.

Obtaining partial cDNA sequences [expressed sequence tags (ESTs)] from various developmental stages and in response to various biotic and abiotic stresses of the plant is an efficient, economical, and quick approach to target the expressed portion of any genome. About 19 million ESTs representing 600,000 unigenes from >80 organisms are available (<http://www.ncbi.nlm.nih.gov>; ADAMS *et al.* 1991). For wheat, >500,000 ESTs corresponding to ~22,000 unigenes have been isolated. However, the full potential of the utility of ESTs in genomics cannot be realized without revealing their physical location on chromosomes. Physical mapping of ESTs is particularly important in wheat because of the highly uneven distribution of genes on chromosomes.

Physical mapping of DNA markers is relatively easy in wheat because a wealth of aneuploid stocks is available. A complete series of nullisomic-tetrasomic (NT; a line lacking a pair of chromosomes, loss of which is compensated for by an extra pair of one of its homoeologs) and ditelosomic (DT) lines (SEARS 1954) can be used to reveal arm location of markers (ANDERSON *et al.* 1992). In addition, 436 chromosome deletion lines are available for the 21 wheat chromosomes that can be used for intrachromosomal mapping (ENDO and GILL 1996). These stocks have been extensively used to physically map >2000 DNA markers (WERNER *et al.* 1992; GILL *et al.* 1993; KOTA *et al.* 1993; HOHMANN *et al.* 1994; DELANEY *et al.* 1995a,b; MICKELSON-YOUNG *et al.* 1995; GILL *et al.* 1996a,b; FARIS *et al.* 2000; WENG *et al.* 2000; SANDHU *et al.* 2001; DILBIRLIGI *et al.* 2004; ERAYMAN *et al.* 2004).

A National Science Foundation-funded collaborative project was initiated with a goal to physically map 10,000 wheat unigene ESTs using the aneuploid and deletion stocks. Wheat homoeologous group 6 data are reported in this article. Similar data for the other six homoeologous groups are presented in the accompanying articles in this issue. We also report identification of rice chromosomal regions homologous to wheat group 6 chromosomes and the use of rice to reveal EST order within each wheat bin.

MATERIALS AND METHODS

Genetic stocks: Chromosome arm locations of the selected ESTs were revealed using 21 NT and 24 DT lines (SEARS 1954,

1966; SEARS and SEARS 1978). For sub-arm localization of ESTs, 101 deletion lines carrying 120 breakpoints were selected (ENDO and GILL 1996). Of these, 17 were for homoeologous group 6 that, along with 6 DT breakpoints, divided the group 6 chromosomes into 26 bins. All the aneuploid and deletion stocks used for the study were in cultivar Chinese Spring (CS) background. The deletion breakpoints were expressed as a fraction length (FL) value of the arm retained in the deletion chromosome. The stocks were provided by the Wheat Genetics Resource Center (WGRC), Kansas State University, Manhattan, KS.

EST selection: As of February 2, 2004, 8318 singletons from ~117,000 ESTs derived from ~37 cDNA libraries were mapped by the whole project. Details concerning the cDNA libraries, ESTs, and singletons are given elsewhere (<http://wheat.pw.usda.gov/NSF>; LAZO *et al.* 2004; ZHANG *et al.* 2004). For the analysis presented here and in the accompanying articles in this issue, the March 17, 2003, data set of 4485 mapped and verified ESTs was used. From this data set, 882 ESTs mapped to homoeologous group 6.

Deletion mapping: Genomic DNA isolation, restriction enzyme digestion, and gel-blot analysis were performed as described by SANDHU *et al.* (2001). Gel-blot analysis was performed using 15 µg of genomic DNA digested with *EcoRI* enzyme. The NT, DT, and the deletion lines were used in a single hybridization reaction on a set of five filters. The lane order for the filters is provided at <http://wheat.pw.usda.gov/NSF>. Each fragment band (locus) was mapped to a chromosome region (bin) flanked by breakpoints of the largest deletion possessing the fragment and the smallest deletion lacking it. The chromosome size data of CS were taken from B. S. GILL *et al.* (1991). The number of expected loci per arm was calculated on the basis of its physical length. The mapping data along with the gel-blot analysis images are available at http://wheat.pw.usda.gov/cgi-bin/westsq1/map_locus.cgi.

Consensus physical map: A consensus physical map of homoeologous group 6 chromosomes was constructed using the criteria described in GILL *et al.* (1996a,b) except that only the ESTs that detected orthologous loci on all three chromosomes were used. The breakpoints of all group 6 deletions were placed on a hypothetical chromosome drawn to scale on the basis of the mean length of group 6 chromosomes. The deletion mapping data from the three chromosomes were then combined to position each EST to the shortest possible chromosome interval. In case of a discrepancy, a location consistent with two homoeologs was used.

Wheat-rice comparisons: The 882 group 6 ESTs were compared with the rice genomic sequence using "blast" (<http://www.ncbi.nlm.nih.gov/>; ALTSCHUL *et al.* 1997). A cutoff *E*-value of $E < 10^{-20}$ and sequence length >100 bases (for *E*-values $< E < 10^{-20}$) were used to identify rice homologs that were equivalent to >65% nucleotide sequence homology. Rice bacterial artificial chromosome (BAC) and P1-derived artificial chromosome (PAC) contigs (<http://rgp.dna.affrc.go.jp>) corresponding to each group 6 bin were identified and used to order 385 group 6 wheat ESTs.

RESULTS

Distribution of ESTs: Of the March 7, 2003 data set of 4485 mapped and verified project ESTs, 882 ESTs mapped to homoeologous group 6 chromosomes, using only the *EcoRI* enzyme. These group 6 ESTs detected 5154 restriction fragment bands, and each fragment band was considered as a locus. The NT and DT analyses mapped 2849 (55%) of these loci to specific chromosome arms. The remaining 2305 (45%) fragment bands

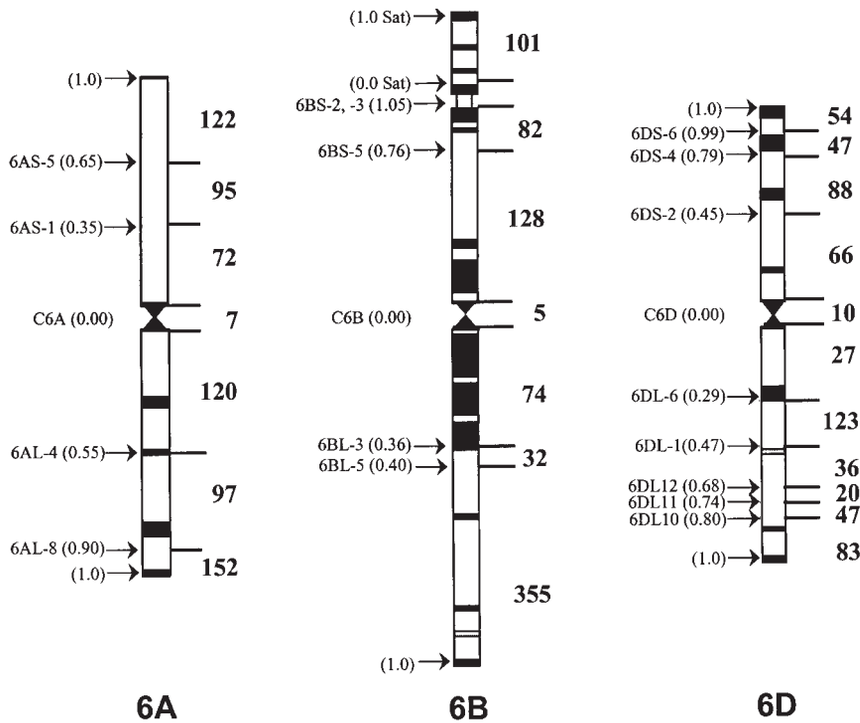


FIGURE 1.—Deletion maps of wheat chromosomes 6A, 6B, and 6D. The chromosomes are drawn to scale. Each centromere is marked by a constriction and the C-bands are shown as solid boxes. The deletion breakpoints along with fraction length (FL) are marked by arrows on the left of the chromosomes and the number of EST loci in each bin is on the right.

were not mapped, as these were monomorphic among the NT lines. The 882 ESTs, mapped on homoeologous group 6, detected 2043 loci on group 6 and 806 loci on other homoeologous groups. Among the homoeologs, chromosome 6B had the highest number of loci and 6D had the lowest. Of the 882 ESTs, 518 detected 665 loci on chromosome 6A, 601 detected 777 loci on 6B, and 488 ESTs detected 601 loci on chromosome 6D (Figure 1).

Distribution of the group 6-specific EST loci on the three homoeologous chromosomes showed that 264 EST probes detected 873 loci on all three homoeologs, 192 detected 472 on two, and the remaining 426 detected 698 loci on only one of the chromosomes (Figure 2). The number of loci mapping on the long arms was significantly higher than that on the short arms. The numbers of short-arm loci were 289, 311, and 255 for

6A, 6B, and 6D, respectively, compared to 369, 461, and 336 for the long arms (Figure 3). Twenty-two loci mapped in the centromeric region, of which 7 were on 6A, 5 on 6B, and 10 on 6D (Figure 1). A comparison of these numbers with the expected number based on the physical length of the arms showed significant differences (χ^2 , $P = 0.012$) between the observed and the

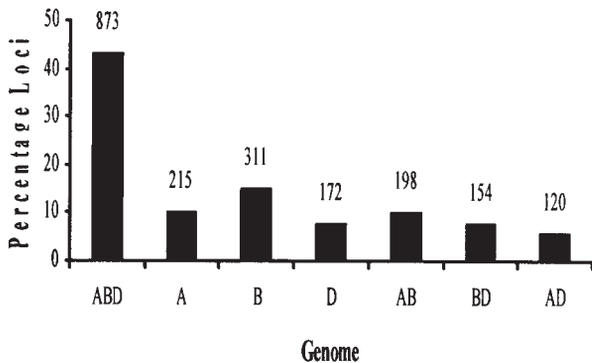


FIGURE 2.—Distribution of 2043 homoeologous group 6-specific EST loci by genome. The actual number of loci is shown at the top of each bar.

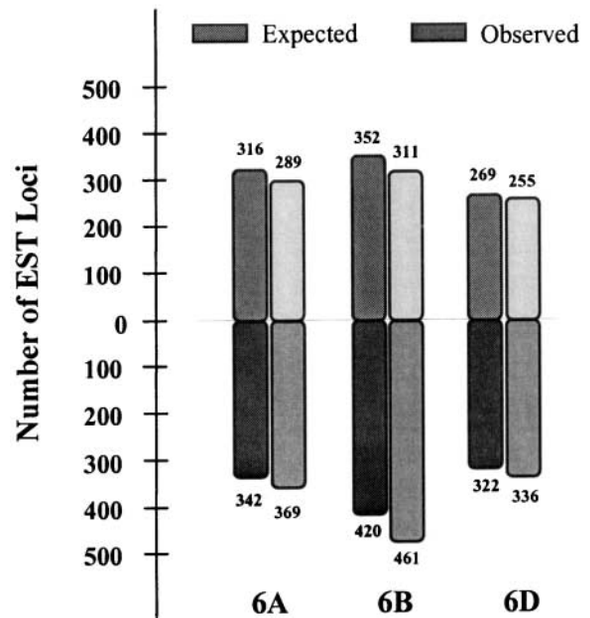


FIGURE 3.—Comparison of observed and expected numbers of homoeologous group 6-specific EST loci. The shaded and solid bars represent the short and long arms, respectively. The actual numbers are given at the top of the bars; not counted are the 22 loci that mapped to centromere bins.

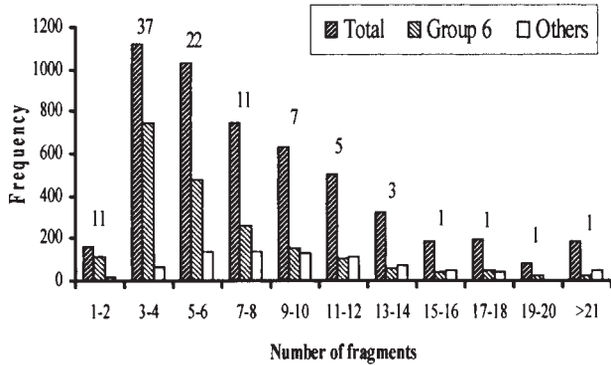


FIGURE 4.—Comparison of the frequency of the bands (fragments) detected by ESTs mapping on group 6 with that of those mapping on other wheat homoeologous groups. The percentage of ESTs detecting the number of fragments is given at the top of the bars.

expected numbers (Figure 3). In general, the observed number of loci for the short arms was lower than expected. For example, 46% of the loci were expected to be present on 6BS whereas only 40% were observed. Consequently, the long arm had ~11% more loci than expected. Similar observations were made for the other two chromosomes.

The distribution of ESTs was uneven along the group 6 chromosomes. Distal bins had more EST loci per unit size compared to the proximal bins (Figure 1). The bin proximal to deletion C-6DL-6 had the lowest EST density. This bin is ~1.6 μ m in length and that translates to ~106 Mb DNA (B. S. GILL *et al.* 1991). The calculated EST density based on the 27 loci mapped to the bin was ~0.25 loci/Mb. The bin distal to deletion 6DS-6 had the highest density, 16 loci/Mb, with 47 loci and a size of ~3 Mb. Similarly, EST density in the 6A bin distal to deletion 6AL-8 was 4.38 loci/Mb compared to 0.63 loci/Mb for the bin proximal to deletion 6AL-4. Detailed mapping information for all the ESTs in each group 6 bin can be accessed from the GrainGenes database website (<http://wheat.pw.usda.gov/wEST>).

Copy number of expressed sequences: On average, each EST detected 5.8 fragment bands with a range from 1 to 39. Frequency of loci detected by group 6 ESTs in comparison with other homoeologous groups showed that only 37% of the ESTs detected the expected 3 or 4 fragment bands (Figure 4). About 11% of the ESTs detected only 1–2 loci, suggesting deletion of homoeologous sequences. The remaining 52% of the ESTs detected 5 or more fragment bands. From the total 2849 loci, 2043 mapped to group 6 chromosomes and the remaining 806 mapped on the other chromosomes. Approximately 30% of loci were duplicated. Of these, 6% were intrachromosomal duplications and the remaining were on other chromosomes. Among the 30 intrachromosomal duplications, 15 were on opposite arms. No difference was observed among homoeologs for the rate of intrachromosomal duplications. However, >75% of

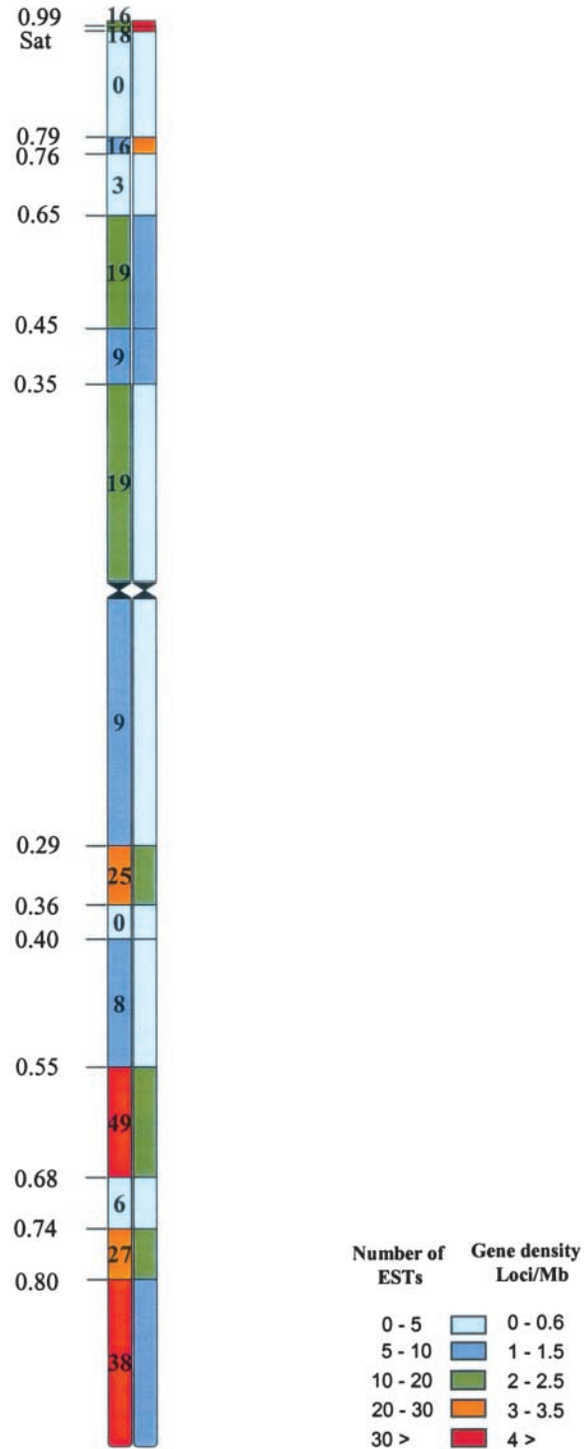


FIGURE 5.—A consensus physical map of homoeologous group 6 chromosomes. The physical map on the left was generated on the basis of the actual number of ESTs per bin, whereas the map on the right was based on EST density. Deletion breakpoints are indicated by lines and FL values on the left. The numbers in the boxes and color scheme for the left side of the consensus physical chromosome represent the number of ESTs present in that region and the right side represents the gene density based on the percentage of the chromosome arm.

these duplications were observed in the proximal 50% of the chromosomes. The proximal bins C-6AL4, C-6DS2, and C-6AS1 showed the greatest number of duplications.

Consensus physical map: The consensus physical map containing 262 ESTs is presented in Figure 5. Location of 223 ESTs was consistent among the three homoeologs (Table 1). For the remaining 39 that showed discrepancies (marked by * in Table 1), the physical location consistent with two homoeologs was used. Of the 39 ESTs with discrepant map locations among homoeologs, 21 mapped to the adjacent bins, suggesting that the discrepancy was due to FL value variation among the homoeologs. The remaining 18 ESTs mapped to nonadjacent bins, of which 13 mapped on the opposite arm.

Uneven distribution of ESTs was even more distinct on the consensus physical map. The distal 60% of the consensus map contained ~80% of the ESTs (Figure 5). The number of ESTs per bin ranged from zero (in the long-arm bin 0.36-0.40 and the short-arm bin 0.79-Sat) to 49 (in the long-arm bin 0.55-0.68). Because of significant size differences among bins, EST density was calculated per unit size. The EST density per unit chromosome length ranged from 0% in consensus regions 0.36-0.40 of the long arm and 0.79-Satellite (Sat) of the short arm to 23% in the short-arm region Sat-0.99 (Figure 5). In general, the smaller-sized bins had a higher EST density. For example, the short-arm regions Sat-0.99 and 0.99-1.00 (~3 Mb each) accounted for 20-23% of the ESTs.

Comparative mapping: To find rice regions corresponding to each of the group 6 bins, 882 group 6 ESTs were compared against the available rice sequences. At the level of stringency used for comparison, only 385 (43%) of the 882 wheat ESTs identified rice homologs; of these, 225 (58%) showed homology to rice chromosome 2, whereas the remaining 160 (42%) corresponded to regions on the other 11 rice chromosomes. The percentage of the wheat ESTs mapping on the other rice chromosomes ranged from 1.3% on chromosome 11 to 8.3% on chromosome 3. With a mean of 6.5, the number of rice chromosomes represented in each wheat bin ranged from 1 (in the long-arm bin 0.36-0.40) to 11 (in the long-arm bin 0.80-1.00) (Figure 6). Within each wheat bin, rice chromosomes other than chromosome 2 were identified by ESTs ranging from 1 to 6 with an average of 2.7. However, chromosome-specific rice contigs corresponding to each wheat bin were discontinuous as homologs were not present for all BACs/PACs present in rice contigs.

To examine differences among wheat bins for rice homology, the number of ESTs per consensus region (Figure 6, blue bar chart) was compared with that of ESTs showing homology to rice sequence (Figure 6, red bar chart). The width of the blue bar chart was drawn to scale on the basis of the location of 225 ESTs that were present on the consensus physical map. The width

of the red bar chart was drawn to scale using 385 ESTs that identified rice homologs. Significant differences were observed among the wheat bins for their homology to rice. For example, wheat regions 0.29-0.36 and 0.55-0.68 on the long arm and 0.76-0.79 on the short arm showed the highest levels of homology with the rice chromosomes (Figure 6). On the other hand, the long-arm regions 0.36-0.40 and 0.68-0.74 and short-arm region 0.79-Sat possessed the least homology. Wheat ESTs mapping on all homoeologs identified rice homologs more frequently than others did. Of the 262 ESTs present on the consensus map, 143 (54%) detected rice homologs compared to 39% for the remaining ESTs.

Rice BAC/PAC contigs corresponding to each of the group 6 bins were used for intrabin ordering of ESTs (Table 1). The order of the 385 wheat ESTs present in the 16 bins of the consensus map was revealed using rice sequences. Of these, 219 were homologous to rice chromosome 2, 32 to chromosome 3, 17 to chromosome 1, and the remaining were homologous to other rice chromosomes.

DISCUSSION

Major cereal crops including wheat (*T. aestivum* L.), maize (*Zea mays* L.), barley (*Hordeum vulgare* L.), and rice (*O. sativa* L.) belong to the grass family Poaceae. Comparisons of genetic maps and DNA sequences have suggested that these grass genomes originated from a common ancestor 50-60 million years ago (BENNETZEN and FREELING 1993; KELLOGG 1998) and have similar gene composition and colinearity (AHN and TANKSLEY 1993; AHN *et al.* 1993). The number of functional genes in these crop plants is not known. The number of genes in rice estimated from genome sequence analysis ranges from 32,000 to 50,000 (GOFF *et al.* 2002). In hexaploid wheat, the gene number estimates range from 75,000 to 150,000, or ~10,000-20,000 gene loci per homoeologous group (SIDHU and GILL 2004). Here we report physical mapping of >2000 loci (10-20% of the total) for wheat homoeologous group 6. We also show the general distribution of genes on the chromosomes.

Deletion mapping revealed significant differences among group 6 homoeologs for the number of loci. The comparison between the expected and observed numbers of loci indicated that the number of loci is not always proportional of the size of the chromosome arm. Maximum number was observed for the 6B and minimum for 6D. This difference may partly be due to the variable sizes of the homoeologs, which are predicted to be 863, 673, and 667 Mb for 6B, 6D, and 6A, respectively (B. S. GILL *et al.* 1991). Another factor explaining this difference may be the number of duplicated loci that may differ among homoeologs.

Dramatic differences were observed for the number of loci per bin. These differences were more pronounced on the consensus physical map mainly because there

TABLE 1
Order of wheat ESTs mapped on group 6 physical map determined from rice BAC/PAC comparison

										Long-arm bins									
										Short-arm bins									
0.99-1.00	Sat-0.99	0.79-Sat	0.76-0.79	0.65-0.76	0.45-0.65	0.35-0.45	0.0-0.35	C	0.0-0.29	0.29-0.36	0.36-0.40	0.40-0.55	0.55-0.68	0.68-0.74	0.74-0.80	0.80-1.00			
BE403550	BE591939	BF291990	BE500816	BF429432	BE446568	BE426401	BG606725	BE443164	BE424119	BF429374	BE490777	BF484678	BF483643	BF202810	BG313802				
BE498351	BE499467	BE498152	BE446453	BE426819	BE499423	BE482663	BE518418	BE637333	BF202312		BE591279	BE405016	BF482687	BE490152*	BF485033				
BE496826	BE490604	BE443439	BF474168	BF484831	BF474648	BE443016	BF202721	BE518064	BE442341		BE493784	BE442611	BE490200	BG263703*	BE403221				
BF201083	BF292168		BF483884	BE497808	BF483648	BF291554	BG312637	BE489789	BE406523		BF472980	BE590521	BM138382	BF483564	BF293225				
BG313246*	BE517858		BE443401	BE403562	BG274266	BG607450	BE445667	BM134356	BE638038		BF483993	BE591696	BE490226	BE442605	BE494057				
BE404947	BE517715		BE498242	BE490111	BE445603	BE443113	BF293628	BE490805	BE500104		BF201435	BF291790	BE637570	BF293956	BM138088				
BE495217	BE592000		BG262500	BF483580	BE500818	BE498099	BF478625	BE496986	BF473299		BE4046439	BE403397	BE405224	BG604547	BF145273				
BE606222	BE495913		BF482700	BE403679	BF482824	BE426591	BE422743	BE444313	BF201426		BE636841	BG274398	BF484968	BG604865	BE442865				
BG263579*	BF200529		BE422443	BE489882	BF428701	BE473139	BE498294	BE498480	BE404912		BG313281	BF485173	BE498182	BE405516	BE591777				
BE500191	BE424523*		BE494189	BF291693	BF200644	BE488526	BE426814	BE404530*	BE517763		BE426413	BG274508	BE498303	BE499203	BF200773				
BE490130	BM134392		BF474461	BE405089	BE445201	BE497343*	BE442709	BE404553*	BE517763		BF202329	BF202738	BE498303	BE495143	BF483267				
BE591957	BE445952		BE488206*	BE425932*	BE446153	BF293341	BE403510	BF428553*	BE494381		BE604721	BM138686	BE498092	BE489894					
BE499652	BF429160		BG313503	BE442576	BF428905	BF428905	BG263085	BG312780*	BE443951		BE443744	BE500611	BE498419	BE637891					
BG262275	BE490715		BE424039	BF291478	BM138480	BM138480	BE425967	BE425967	BE425967		BF482926	BE490512	BG606839*	BE518379					
BE498456	BE498110		BG263145	BE498988	BE442933	BE442933	BE4426594	BE426594	BE426594		BM135343	BE591931	BE6637911	BF294007					
BE605139	BM140482		BE591244	BF429355	BE493888	BE493888	BE493888	BF484691	BF484691		BF482473	BE426214	BE426310	BF475120					
BF473001	BE499685		BF291471	BE606541	BE500541	BE500541	BE500541	BE490786	BE490786		BG274229	BE500686	BE637963	BF474812					
BG262851	BF201597		BE591588	BG275060	BG312802	BG312802	BE494056	BF482566	BF482566		BE607043	BE403938	BE407066	BG274882					
BG274742*	BE398695*		BF473535	BF200758	BF200758	BF200758	BE406474	BE495607	BE495607		BG313765	BM140440	BE425207	BE406840					
BE497450	BF145263		BG262421*	BG275014	BG275014	BG275014	BE498678	BE495949	BE495949		BE425067	BE443342	BE489612	BE403564					
BF145253	BM138339		BE604816	BF201409	BF201409	BF201409	BE498678	BE518352	BF474134		BE518352	BF474134	BE403154	BE423933					
BF293402*	BF428770		BF145263	BM136709	BM136709	BM136709	BE445239	BE403421	BE403421		BE606591	BM138139	BE426017	BE500840					
BF428770	BM138681		BE445324	BE499711	BE499711	BE499711	BF483150	BE591155	BE591155		BE444509	BM136727	BE445641*	BE489573					
			BE424920	BE404539	BE404539	BE404539	BF201649	BE404354	BE404354		BF428572	BE403577	BE494036*	BF483091					
			BF293613	BF478393	BF478393	BF478393	BE405195*	BE442905	BE442905		BE499625	BE497701	BE590638*	BM1137547					
			BE490365	BF483695	BF483695	BF483695	BE406602*	BE637763	BE637763		BE490286	BE490147	BE591788	BF474277					
			BE490565	BE444631*	BE444631*	BE444631*	BE497874	BF293311	BF293311		BE443191	BE403818	BF484672	BE500543					
			BE404723	BE496059	BE496059	BE496059	BE606884	BF478945	BF478945		BE604759	BE442546	BF485398	BF485407					
			BG607042	BE592027	BE592027	BE592027	BF483246	BF482982	BF482982		BF482668	BF201071	BG263812	BE490082					
			BE406943	BE473852*	BE473852*	BE473852*	BF485238	BF473461	BF473461		BF484181	BE443929	BG604504	BE443156					
			BE426362	BF473852*	BF473852*	BF473852*	BG263023	BE444256	BE444256		BF203145	BF293263	BE443711	BE443711					
			BE424657	BG263237	BG263237	BG263237	BE605218	BF428665	BF428665		BG607393*	BE489623	BE443643	BE443643					
							BE443589	BE443589	BE443589		BE424097	BE498001	BE498785	BE498785					
											BE442732	BE498001	BE591228	BE591228					

(continued)

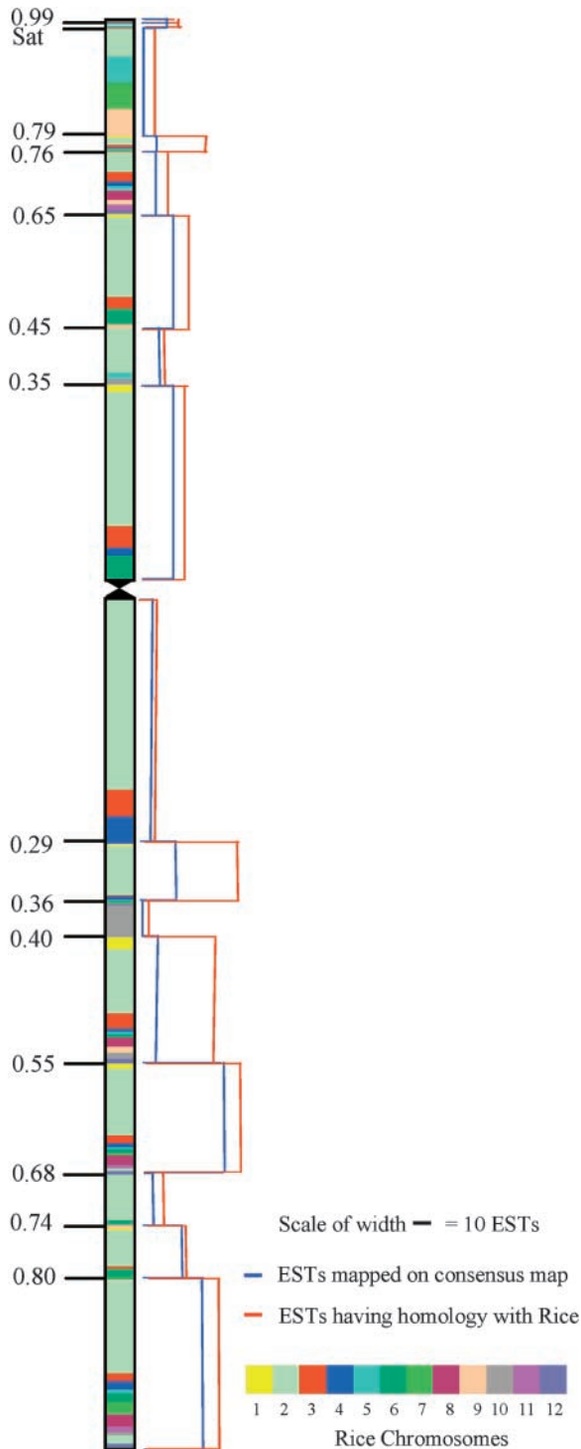


FIGURE 6.—Comparative analysis of wheat group 6 ESTs with rice BAC/PAC sequences. The wheat consensus chromosome 6 is shown with deletion breakpoints and FL values on the left. Each deletion bin is color coded according to the rice chromosome with matching ESTs mapped to that bin. Each bin is spanned by red and blue bar charts on the right of the chromosome. The bar charts are drawn to scale using actual number of ESTs. The width of the red bar chart shows the number of wheat ESTs with homology to the rice chromosomes and the blue bar chart shows the number of ESTs on the consensus wheat group 6 map.

were three times more breakpoints resulting in a finer resolution. The estimated bin size on the individual deletion maps ranged from ~ 3 (the bin distal to deletion 6DS-6) to 299 Mb (C-6BS-5) with a mean of 88 Mb (Figure 1). On the other hand, consensus-map bin size ranged from 3 (for short-arm bin 0.99–1.00) to 119 Mb (for short-arm bin 0.0–0.35) with a mean of 43 Mb (Figure 5). More than a 30-fold difference was observed for gene density among bins on the consensus map compared to a 14-fold difference in size among individual bins. This comparison suggests that the difference in gene density may be even greater if additional breakpoints were available (GILL *et al.* 1996b; ERAYMAN *et al.* 2004). The limitations of the consensus map construction are pointed out by the fact that $\sim 15\%$ of the ESTs had discrepant locations among the homoeologs. Although order and colinearity are conserved among the three genomes for most of the genes, significant differences may be present due to differential amplification of the three genomes, chromosomal rearrangements, and gene copy number.

Using only one restriction enzyme, $\sim 30\%$ of the ESTs detected loci on all three chromosomes, 22% detected loci on two, and the remaining ESTs detected loci only on one of the three homoeologs. This large number of ESTs mapping to only one of the chromosomes can be attributed to the use of only one restriction enzyme. By using two restriction enzymes for a similar physical mapping experiment, $\sim 81\%$ (61/75) wheat group 1 gene markers detected loci on all three homoeologs, 12% (9/75) detected loci on two, and only 7% (5/75) detected loci on one of the three genomes (SANDHU *et al.* 2001). In the present study, $\sim 45\%$ (2305/5154) of the fragments detected by group 6-specific ESTs were monomorphic and that may be resolved with the use of additional restriction enzymes.

The extent and distribution of EST duplication on group 6 chromosomes was similar to that reported for the wheat genome as a whole (QI *et al.* 2004). About 21% of the wheat sequences have paralogous loci (AKHUNOV *et al.* 2003a). In this study, 32% (287) of the ESTs detected paralogous loci on other chromosomes ranging from 13% for group 4 to 20% for groups 2 and 7. About 4% of the ESTs detected paralogous loci on group 6, of which one-half were on the same chromosome. In barley, 20–30% of probes detected duplicated loci (GRANER *et al.* 1991; KLEINHOFES *et al.* 1993). Some of the other similar estimates obtained from the genetic linkage analysis were 28% of the cDNA clones and 34% of the *Pst*I genomic clones in *T. monococcum* L. (DUBCOVSKY *et al.* 1996), and 31% in *Aegilops tauschii* Coss. (K. S. GILL *et al.* 1991). These duplicated loci could have resulted from interchromosomal exchanges, intergenomic invasions, and dispersion of specific DNA elements during genome evolution through polyploidization (WENDEL 2000; AKHUNOV *et al.* 2003a).

Several wheat-rice comparisons have been made and

slightly different results were observed, depending on methods and criteria. Genetic linkage-map comparisons between wheat and rice identified syntenic chromosomes between the two genomes (AHN *et al.* 1993; KURATA *et al.* 1994; DEVOS *et al.* 1995; SHERMAN *et al.* 1995; VAN DEYNZE *et al.* 1995; SAGHAI-MAROOF *et al.* 1996; DEVOS and GALE 1997). Depending upon stringency, sequence comparisons between wheat and rice showed that 50–98% of the rice genes are similar to those of wheat. For example, GOFF *et al.* (2002) reported ~98% protein sequence homology among rice, maize, wheat, and barley. On the other hand, 65% of wheat ESTs identified rice homologs at $E < 10^{-15}$ (SORRELLS *et al.* 2003). Similar comparisons in the present study at a slightly higher stringency showed that only ~43% of group 6 ESTs have rice homologs. Even at a liberal cut-off value ($E < 10^{-1}$), only 67% (593/882) of the ESTs detected rice homologs. Therefore, we conclude that at least 33% of the wheat ESTs do not have rice homologs.

Previous studies using RFLP markers (GALE and DEVOS 1998) and wheat ESTs (SORRELLS *et al.* 2003) have reported that wheat homoeologous group 6 chromosomes illustrate substantial homology to rice with the best conservation of gene order and content with rice chromosome 2 (SORRELLS *et al.* 2003). In the present study, however, of the 43% group 6 ESTs that identified rice homologs, only 58% were on rice chromosome 2. The remaining were present as small blocks on the other rice chromosomes.

Individual bin comparisons with rice sequences showed that, in addition to chromosome 2, other rice chromosome segments were present in all the bins as 1–10 blocks of varying sizes. Each rice chromosomal block was identified by 1–6 wheat ESTs, suggesting that these blocks are not paralogous loci but are true homologs of wheat group 6 scattered on other rice chromosomes (Figure 6). Since most of the rice chromosomal blocks homologous to wheat group 6 bins have been identified, it should be possible to use the rice sequence information efficiently and accurately for wheat genomics. Furthermore, rice BAC/PAC contigs can be used to order wheat ESTs within bins. In this study, we resolved the order of 385 wheat ESTs within 16 bins (Table 1). However, the accuracy of this order needs to be determined.

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