Sequence Elimination and Cytosine Methylation Are Rapid and Reproducible Responses of the Genome to Wide Hybridization and Allopolyploidy in Wheat

Hezi Shaked, Khalil Kashkush, Hakan Ozkan, Moshe Feldman, and Avraham A. Levy

Department of Plant Sciences, The Weizmann Institute of Science, Rehovot 76100, Israel

Interspecific or intergeneric hybridization, followed by chromosome doubling, can lead to the formation of new allopolyploid species. Recent studies indicate that allopolyploid formation is associated with genetic and epigenetic changes, although little is known about the type of changes that occur, how rapidly they occur, and the type of sequences involved. To address these matters, we have surveyed F1 hybrids between diploid species from the wheat (Aegilops and Triticum) group and their derived allotetraploids by screening a large number of loci using amplified fragment length polymorphism and DNA gel blot analysis and by assaying the extent of cytosine methylation. We found that sequence elimination is one of the major and immediate responses of the wheat genome to wide hybridization or allopolyploidy, that it affects a large fraction of the genome, and that it is reproducible. In one cross between Ae. sharonensis × Ae. umbellulata, 14% of the loci from Ae. sharonensis were eliminated compared with only 0.5% from Ae. umbellulata, with most changes occurring in the F1 hybrid. In contrast, crosses between Ae. longissima × T. urartu showed that sequence elimination was more frequent after chromosome doubling. Alterations in cytosine methylation occurred in ~13% of the loci, either in the F1 hybrid or in the allotetraploid. For eight of nine bands that were isolated, the sequences that underwent elimination corresponded to low-copy DNA, whereas alterations in methylation patterns affected both repetitive DNA sequences, such as retrotransposons, and low-copy DNA in approximately equal proportions.

INTRODUCTION

The duplication of genomes (polyploidy), either of the same genome (autopolyploidy) or of diverged genomes (allopolyploidy or amphiploidy), is a major force of evolution. It is thought that most living eukaryotes have undergone one or more polyploidization events during their history (Spring, 1997; Wolfe and Shields, 1997). One interpretation of the success of polyploidy is that it releases one set of alleles from selective pressure, thus facilitating the acquisition of new functions and buffering the mutation load. In the case of allopolyploidy, the fixation of permanent heterozygosity (Stebbins, 1971; Allard et al., 1993) has the potential to offer a substantial heterozygote advantage. Despite these potential benefits, polyploidy is an enormous challenge with regard to the orchestration of gene expression, DNA replication, and chromosome pairing. How the new polyploid genome deals with such genomic stress is a significant, albeit poorly understood, question.

In plants, it is estimated that polyploidy occurred in the history of ~70% of flowering species (Masterson, 1994). Genome sequencing data may provide further evidence for polyploidy, as suggested by the recent finding that Arabidopsis, whose genome is among the smallest in angiosperms, is probably an ancient tetraploid (The Arabidopsis Genome Initiative, 2000). Allopolyploidy, brought about by interspecific or intergeneric hybridization followed by chromosome doubling, was found to be associated with genomic alterations, such as sequence elimination, activation of transposons, intergeneric coconversion, methylation, gene silencing, or subtler changes in gene expression (for reviews, see Matzke et al., 1999; Comai, 2000; Wendel, 2000).

Previous works showing allopolyploidy-induced genetic and epigenetic alterations have focused on selected sequences such as rDNA genes (Sasakuma et al., 1995), cDNAs (Song et al., 1995), restriction fragment length polymorphisms (RFLPs), or low-copy sequences from chromosome- or genome-specific DNA (Feldman et al., 1997; Liu et al., 1998a, 1998b) or transgenes (Scheid et al., 1996). In this work, we have analyzed an unbiased set of wheat DNA loci in diploid wheat species, the interspecific F1 offspring, and
their derived synthetic allotetraploids using amplified fragment length polymorphism (AFLP) (Vos et al., 1995) and methylation-sensitive amplification polymorphism (MSAP) (Xiong et al., 1999) to obtain a quantitative estimate of the timing and frequency of allopolyploidy-associated genetic and epigenetic alterations and to determine what type of sequences are affected. The wheat (Aegilops and Triticum) group offers an ideal system in which to study the evolution of polyploids because several species are young, such as the hexaploid bread wheat, which is only ~85,000 years old (Feldman, 2001), most diploid progenitors are known, and allopolyploids can be synthesized easily.

RESULTS

Frequency of Allopolyploidy-Associated Genetic Alterations as Estimated by AFLP

To assess the frequency of allopolyploidy-associated genetic changes in a quantitative and unbiased manner, two different crosses were studied: Ae. sharonensis (SS) × Ae. umbellulata (UU) (cross A) and Ae. longissima (SS) × T. urartu (AA) (cross B). The F1 hybrids and the allopolyploids derived from each cross were analyzed by AFLP. The banding pattern of the hybrids is expected to yield a pattern that is additive of both parents because parents derive from inbred lines (confirmed in this study) and because AFLP markers are dominant. All of the cases of deviation from such additivity scored in this study were reproducible (see Methods). Deviation was observed for bands present in only one of the parents (polymorphic bands) and absent in the F1 hybrid and/or the allotetraploid. Another, albeit more rare, type of deviation from additivity was the appearance of new bands that were not present in the parents, disappeared in F1, and reappeared in the allotetraploid (data not shown). These deviations from additivity were probably caused by changes in methylation of the EcoRI site. Alterations in AFLP patterns caused by methylation changes are described in the MSAP analysis. Here, we have focused on the analysis of bands that disappeared in F1 and/or the allopolyploid.

### Table 1. AFLP Banding Pattern in the Parental Plants of One Interspecific Cross (Ae. sharonensis [SS] × Ae. umbellulata [UU]) and One Intergeneric Cross (Ae. longissima [SS] × T. urartu [AA]) in Their F1 Hybrids and in the First Generation of the Allotetraploids [S'SUU and S'SAA]

<table>
<thead>
<tr>
<th>Crossa</th>
<th>Bands in Parents</th>
<th>Bands That Disappeared</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Polymorphic</td>
<td>Totalb</td>
</tr>
<tr>
<td></td>
<td>♀</td>
<td>♂</td>
</tr>
<tr>
<td>S'S × UU (cross A)</td>
<td>171</td>
<td>202</td>
</tr>
<tr>
<td>S'S × AA (cross B)</td>
<td>80</td>
<td>166</td>
</tr>
</tbody>
</table>

aIn cross A, the maternal genome (SS) was from Ae. sharonensis, whereas in cross B, it was from Ae. longissima (SS).

bThe total number of bands for both parents. Monomorphic bands were scored only once.
DNA Gel Blot Validation of the AFLP-Detected Genetic Alterations

The AFLP-observed deviations from additivity could result from the heterozygosity of the parents, from artifacts of the polymerase chain reaction (PCR)-based AFLP analysis, or from methylation alteration, or they could be the result of random or reproducible genetic alterations induced by allopolyploidy. To distinguish between these various possibilities, we performed DNA gel blot analysis in independently synthesized allotetraploids. This analysis should reveal potential heterozygosity of the parents and indicate the reproducibility of the observed alterations.

Nine bands that disappeared, either in F1 or in the allopolyploid (all from cross B), were isolated from acrylamide AFLP gels (Figure 1) and used as probes for DNA gel blot analysis. The results of this analysis are shown in Table 2 and are described below. Eight of the nine bands originated from low copy DNA, and one was from high-copy DNA. Sequence analysis did not reveal any significant similarity for any of the nine sequences. In two cases, that of AFLP-isolated fragments 4 and 5 (AIF4 and AIF5), validation could not be determined, either because there was no polymorphism between the parents (AIF4) or because the band originated from high-copy DNA (AIF5) and discrete bands could not be analyzed. In all seven cases in which validation could be tested, the band elimination observed by AFLP was confirmed by DNA gel blot analysis (Figure 2). Note that, for all of the probes tested, the pattern of elimination was reproducible in three different S1 plants of each allotetraploid. This finding, as well as the reproducibility of migration patterns in the parents (data not shown), indicates that the deviation from additivity was not caused by the heterozygosity of the parents. Moreover, some of the enzymes used in the DNA gel blot analysis (DrαI and EcoRV) are not particularly sensitive to methylation and band disappearance was not associated with the appearance of new bands, suggesting that for the cases studied here, the band disappearance in AFLP was not caused by cytosine methylation.

In summary, AFLP is a robust and high throughput means to assess the induction of genomic rearrangements. Elimination was the most frequent genomic rearrangement; it was reproducible and could not be attributed to heterozygosity or methylation.

Frequency of Allopolyploidy-Associated Epigenetic Alterations as Estimated by MSAP

The enzyme HpaII is sensitive to methylation of either cytosine residue at the recognition site (5'-CCGG), whereas its isoschizomer MspI is sensitive only to methylation of the external cytosine. Therefore, methylation of the internal cytosine would lead to a differential cleavage by the two isoschizomers and thus to the appearance of different MSAP fragments in the sequencing gel loaded with the amplification products from EcoRI+Mspl versus EcoRI+Hpall digests (Figure 3). Another, albeit less frequent, cause of the differential banding pattern, namely the presence of fragments in the EcoRI+Hpall digests and their absence in the EcoRI+Mspl digests, has been observed (Figure 3) and has been attributed to hemimethylation of
Table 2. DNA Gel Blot Analysis of AIF Probes That Showed AFLP Band Disappearance in the F1 Hybrid and/or in the Allotetraploid Plants of Cross B (S/S' × AA)

<table>
<thead>
<tr>
<th>Probe</th>
<th>Primer Pairs</th>
<th>Fragment Size (bp)</th>
<th>Origin of Fragment</th>
<th>Eliminationa</th>
<th>Copy Number</th>
<th>Sequence Similarityb</th>
</tr>
</thead>
<tbody>
<tr>
<td>AIF1</td>
<td>E+AAC/M+CAG</td>
<td>420</td>
<td>urartu</td>
<td>+</td>
<td>Low</td>
<td>None</td>
</tr>
<tr>
<td>AIF2</td>
<td>E+AAC/M+CAG</td>
<td>202</td>
<td>urartu</td>
<td>+</td>
<td>Low</td>
<td>None</td>
</tr>
<tr>
<td>AIF3</td>
<td>E+AAC/M+CAG</td>
<td>150</td>
<td>urartu</td>
<td>+</td>
<td>Low</td>
<td>None</td>
</tr>
<tr>
<td>AIF4</td>
<td>E+AAC/M+CAG</td>
<td>80</td>
<td>urartu</td>
<td>ND</td>
<td>Low</td>
<td>None</td>
</tr>
<tr>
<td>AIF5</td>
<td>E+AGG/M+CTT</td>
<td>530</td>
<td>longissima</td>
<td>ND</td>
<td>High</td>
<td>None</td>
</tr>
<tr>
<td>AIF6</td>
<td>E+AGG/M+CTT</td>
<td>510</td>
<td>urartu</td>
<td>+</td>
<td>Low</td>
<td>None</td>
</tr>
<tr>
<td>AIF7</td>
<td>E+AGG/M+CTT</td>
<td>450</td>
<td>urartu</td>
<td>+</td>
<td>Low</td>
<td>None</td>
</tr>
<tr>
<td>AIF8</td>
<td>E+AAC/M+CTT</td>
<td>170</td>
<td>longissima</td>
<td>+</td>
<td>Low</td>
<td>None</td>
</tr>
<tr>
<td>AIF9</td>
<td>E+AAC/M+CTT</td>
<td>165</td>
<td>longissima</td>
<td>+</td>
<td>Low</td>
<td>None</td>
</tr>
</tbody>
</table>

a Elimination in F1 and/or the allotetraploid was validated (+) or could not be determined (ND). AIF1 to AIF7 were eliminated in the allotetraploid only, and AIF8 and AIF9 were eliminated in both F1 and the allotetraploid.
b Similarity was considered significant for e values ≤ 0.005.

The two parental lines, the F1 hybrid, and the allotetraploid were compared for their methylation patterns (Table 3). Seven pairs of selective primers produced 501 clear bands in the two parental lines Ae. sharonensis (genome S/S') and T. monococcum ssp aegilopoioi (genome A') (Table 3). Each of the bands represented a recognition site cleaved by one or both of the isoschizomers. Of the 501 resolved bands, 159 were methylated in one or both parental lines (Table 3). The two parents showed essentially the same degree of methylation, namely 30.2% for Ae. sharonensis and 33.2% for T. monococcum ssp aegilopoioi (data not shown). Therefore, on average, 31.7% (159 of 501) of the 5'-CGG sites in the parental line genome were cytosine methylated in the leaves (Table 3).

The two parental lines, the F1 hybrid, and the allotetraploid were compared for their methylation patterns (Table 4, Figure 3). As mentioned for the AFLP analysis of genetic alterations, we expected that F1 and the allotetraploid would have the combined methylation pattern of both parents. Any deviation from additivity was considered an alteration in methylation pattern related to the F1 and/or the allotetraploid situation. Five major classes were identified among the methylated fragments (Table 4): (1) 21 bands (13.2% of all 159 methylated sites) were monomorphic, that is, they had the same pattern of methylation among parental lines, the F1 hybrid, and the allotetraploid; (2) 117 bands (73.5% of all methylated sites) showed differential cytosine methylation between the parents but no deviation from additivity in F1 or the allotetraploid; (3) three bands (1.8% of all methylated sites) showed differential cytosine methylation patterns between parents and F1 hybrids, but methylation was restored to the parental type in the allotetraploid; (4) 11 bands (6.9% of all methylated sites) showed alteration in cytosine methylation starting in the F1 hybrid that was maintained in the allotetraploid (interestingly, of those, 10 were from aegilopoioi and one was from sharonensis); and (5) seven bands (4.4% of all loci) showed methylation alteration in the allotetraploid but not in the F1 hybrid.

DNA Gel Blot Validation of the MSAP-Detected Epigenetic Alterations

DNA gel blot analysis was performed to confirm the methylation patterns detected by MSAP using 12 MSAP-isolated fragments (MIFs) as probes. These included five fragments (of 11 total) that showed methylation alteration in both the F1 hybrid and the allotetraploid and all seven fragments that showed methylation alteration in the allotetraploid only (Table 4). Total DNA from leaves was digested with either HpaII or MspI. All 12 fragments were sequenced and analyzed for similarity to known sequences in the database. Three of the five fragments that showed methylation alteration in the F1 hybrid (MIFs 1 to 5) showed high similarity to repetitive DNA sequences and highly repeated retrotransposons, and two fragments showed no significant similarity to known sequences (Table 4).

Sequence analysis of the seven fragments that showed methylation alteration in the allotetraploid (MIFs 6 to 12) showed no significant similarity to known sequences (Table 5). DNA gel blot analysis of these fragments showed that three of these seven fragments were high-copy number and the rest were low-copy number (Table 5). Methylation alteration in the allotetraploid was validated for all four low-copy fragments using DNA gel blot analysis. An example is shown in Figure 4A for MIF8 as a probe (Table 5). Furthermore, the four low-copy fragments were hybridized to membranes containing the same DNA samples digested with methylation-insensitive restriction enzymes. In these cases, there was no alteration in the DNA gel blot hybridization pattern. An example is shown in Figure 4B using MIF8 as a probe. Similarly, altered methylation in F1 and/or the allo-
polyploid was validated for all five low-copy probes tested. These results show perfect agreement between DNA gel blot analysis and methylation alteration in the allopolyploid detected by MSAP. The high-copy probes led to a smear on the autoradiogram, leaving us unable to assess methylation alteration (data not shown).

Note that using MSAP, we also recovered fragments that were not differentially methylated but that showed band disappearance in F1 and/or the allopolyploid, as was found using AFLP (Figures 1 and 2). DNA gel blot analysis of two such bands showed elimination in the allopolyploid using both membranes that included DNA samples digested by isoschizomers (Figure 5A) and membranes that included the same DNA samples digested with nonsensitive restriction enzymes (Figure 5B). These two bands were both low copy and showed no significant similarity to known sequences.

DISCUSSION

In previous works, we have shown that a limited set of loci could undergo rapid elimination in newly synthesized allopolyploids from the wheat group (Feldman et al., 1997; Liu et al., 1998b; Ozkan et al., 2001). These findings were based on the analysis of fragments that had the unexpected property of not being present in all genomes of polyploid wheat (chromosome-specific or genome-specific sequences). In this study, we have addressed the frequency of genetic and epigenetic changes in interspecific and intergeneric F1 hybrids and in newly synthesized allopolyploids and have sequenced some of the loci involved. This was done by analyzing a large and unbiased sample of wheat loci using the AFLP method (Vos et al., 1995) or the derived MSAP method (Xiong et al., 1999). The banding patterns obtained by both methods were found to be reproducible; moreover, AFLP or MSAP bands showing unexpected inheritance in the F1 hybrid or the allopolyploid were isolated, sequenced, and used as probes for DNA gel blot analysis. In all cases, the DNA gel blot data corroborated the AFLP or MSAP data, indicating that AFLP and MSAP provide reliable and high throughput methods to test for genetic and epigenetic changes, respectively.

The Observed Genetic and Epigenetic Alterations Are Not the Result of Heterozygosity

The unexpected inheritance of AFLP and MSAP bands in the F1 hybrids and the allopolyploids was not attributable to the heterozygosity of the parents or to PCR artifacts, as supported by the following evidence. The diploid parents of the allopolyploid were collected as single plants and selfed, and a single plant was used for each cross. The accessions used to produce the hybrids were checked by DNA gel blot for heterozygosity and were found to be homozygous at 50 loci (H. Ozkan, unpublished data). The independently made F1 and allopolyploids had similar AFLP and MSAP banding patterns. Bands that were present in F1 and disappeared in the allopolyploid could not disappear as a result of genetic segregation because chromosome doubling from dihaploid parents should produce homozygous lines. Moreover, band disappearance was not caused by chromosome loss because plants were checked cytologically and found to be euploid, as expected. Finally, the DNA gel blot validation of

Figure 2. DNA Gel Blot Hybridization with Three Different AIF Probes That Showed Band Disappearance with the AFLP Method.

Genomic DNA of Ae. longissima (TL; genome S/S), T. urartu (TMU; genome AA), and three independently made newly synthesized allopolyploids (S1 generation; genome S'S/AA) was digested and blotted on a nylon membrane. (A) AIF1 probe (Table 2) was isolated from T. urartu and hybridized to EcoRV-digested genomic DNA. Sequence elimination was found in both Ae. longissima (bottom arrow) and T. urartu (top arrow). (B) AIF2 probe (Table 2) was isolated from T. urartu and hybridized to BamHI-digested genomic DNA. This was the only probe that showed a single-copy fragment in both parents. The T. urartu band, shown by the arrow, disappeared in the allotetraploid. (C) AIF6 (Table 2) was isolated from T. urartu and hybridized to HindIII-digested genomic DNA. Two bands from T. urartu, shown by arrows, disappeared in the allotetraploid. This pattern of elimination also was shown with enzymes that are not sensitive to methylation, such as DraI, with all of the probes (data not shown).
band elimination (Figures 2 and 5) did not provide evidence for heterozygosity. Therefore, the observed reproducibility in band elimination or methylation suggests that some sequences in the genome have unknown features that make them the target for sequence elimination or for methylation or demethylation in interspecific F1 hybrids and/or allopolyploids. These data are consistent with the findings of Ozkan et al. (2001), who analyzed a limited number of loci in 35 different interspecific and intergeneric hybrids and their derived amphiploids.

**Frequency and Timing of Genetic and Epigenetic Alterations**

The results shown in Table 1 indicate that a high fraction of the genome can be eliminated in newly synthesized allopolyploids and that the parental genomes are not affected equally. Note that DNA loss frequencies can be determined only for polymorphic bands, whereas the appearance of new bands, which was very rare, could be determined irrespective of the polymorphism of the parents. In cross A, 14% of the polymorphic bands (24 of 171) from *Ae. sharonensis* disappeared compared with only 0.5% (one of 202) for *Ae. umbellulata*. In cross B, there was less difference between the two parents, but overall the elimination rates were similar. The finding that one genome can eliminate up to 14% of its loci in a single generation (assuming that loss frequency is similar for monomorphic and polymorphic bands) shows that allopolyploidy leads not only to the establishment of new species in one step but also to the rapid evolution of the individual genomes in the allopolyploid background. Such rapid evolution could explain the elusive nature of the B genome of wheat (Talbert et al., 1995). These findings also suggest that genomes do not have a one-way ticket to obesity (Bennetzen and Kellogg, 1997) but that mechanisms exist for DNA elimination as discussed below. The timing of elimination was dependent on the genomic combination: most of the elimination occurred in the F1 hybrid of cross A, whereas in cross B, most of the elimination occurred after chromosome doubling. These data show that large-scale genomic rearrangements can occur extremely rapidly, perhaps as early as in the zygote of interspecific and intergeneric F1 plants. Indeed, when deletion occurred in F1 (Ozkan et al., 2001; this work), it was usually an all-or-nothing event, and there was no weak hybridization signal that could suggest the chimeric nature of the F1 tissues from which the genomic DNA was extracted. This suggests that elimination was a very early event. Note that there may be a difference between sequences that disappear in the F1 hybrid and those that disappear after chromosome doubling, as found by Ozkan et al. (2001) for chromosome-specific versus genome-specific sequences. Our work extends previous works on wide hybrids in cereals and other species showing that genomic rearrangements could occur, usually in repetitive DNA, and that new RFLP fragments could be observed (Lin et al., 1985; Lapitan et al., 1988; Svitashev et al., 1995).

Alterations in cytosine methylation also were associated with the hybrid and allopolyploid condition (Table 4, Figure 3). Few events (1.8%) were found whereby the methylation pattern was altered in F1 but was restored to the parental
type in the allopolyploid. Most events of methylation pattern alteration had occurred in the F1 hybrid (6.9%) and were maintained in the allopolyploid (4.4%). These results are higher than reported for methylation alterations in F1 of intraspecific crosses; in rice, Xiong et al. (1999) reported, using MSAP, that methylation patterns were altered for 4.1% of the sites analyzed in the F1 hybrids of crosses between cultivars. An interesting observation, which needs to be substantiated by a larger sample, is that 10 of 11 bands that showed methylation alteration in the F1 hybrid and the allopolyploid were from the genome of one parent (*aegilopoides*; Table 4), suggesting

Table 3. Number of Bands Amplified Using Seven MSAP Selective Primer Combinations in the Parental Lines *Ae. sharonensis* (SlSl) and *T. monococcum* ssp *aegilopoides* (AmAm), the F1 Hybrid (SlAm), and the Allotetraploid (SlSlAmAm)

<table>
<thead>
<tr>
<th>Primer (HM +4/E+3)</th>
<th>Total Number of Bands in the Two Diploid Parents&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Polymorphic Sites&lt;sup&gt;b&lt;/sup&gt;</th>
<th>Total Number of Methylated Sites&lt;sup&gt;c&lt;/sup&gt;</th>
<th>Methylation Alteration in the F1 Hybrid</th>
<th>Methylation Alteration in the Amphiploid</th>
</tr>
</thead>
<tbody>
<tr>
<td>E + AAG</td>
<td>76</td>
<td>2</td>
<td>31</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>E + AGC</td>
<td>65</td>
<td>4</td>
<td>28</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>E + AGG</td>
<td>64</td>
<td>3</td>
<td>18</td>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>E + ACG</td>
<td>72</td>
<td>7</td>
<td>20</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>E + ACT</td>
<td>78</td>
<td>4</td>
<td>22</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>E + ACA</td>
<td>70</td>
<td>9</td>
<td>15</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>E + ACC</td>
<td>76</td>
<td>8</td>
<td>25</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td>Total</td>
<td>501</td>
<td>37</td>
<td>159</td>
<td>14</td>
<td>18</td>
</tr>
</tbody>
</table>

<sup>a</sup> The selective primer HM +4 (5'-CATGAGTCTGCTGGTCAA [HM + TCCA]) was used in combination with each of the seven E+3 primers listed. The core sequence of EcoRI is identical to that used in the AFLP protocol by Vos et al. (1995).

<sup>b</sup> Bands from four lanes were analyzed in each selective primer combination, two lanes for each parent (digested with either HpaII or MspI).

<sup>c</sup> Bands that were monomorphic (identical in both parents) were counted only once.

<sup>d</sup> Sites showing conventional AFLP polymorphism, not related to methylation.

<sup>e</sup> Bands were considered methylated if they showed polymorphism between the two isoschizomers. Bands with the same methylation pattern in both parents were scored only once.

Table 4. Patterns of Cytosine Methylation in the Parental Lines, the F1 Hybrid, and the Allotetraploid<sup>a</sup>

<table>
<thead>
<tr>
<th>Methylation Patterns</th>
<th><em>sharonensis</em> (SlSl)</th>
<th><em>aegilopoides</em> (AmAm)</th>
<th>F1 Hybrid (SlAm)</th>
<th>Allotetraploid (SlSlAmAm)</th>
<th>Number of Sites</th>
<th>Total (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Monomorphic</td>
<td>–</td>
<td>+</td>
<td>–</td>
<td>–</td>
<td>21</td>
<td>21 (13.2)</td>
</tr>
<tr>
<td>Polymorphic</td>
<td>+</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>47</td>
<td></td>
</tr>
<tr>
<td>between parents, no change in</td>
<td>–</td>
<td>+</td>
<td>–</td>
<td>–</td>
<td>35</td>
<td></td>
</tr>
<tr>
<td>F1 and amphiploid</td>
<td>–</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>24</td>
<td>117 (73.5)</td>
</tr>
<tr>
<td>Methylation alteration in F1 only</td>
<td>+</td>
<td>+</td>
<td>–</td>
<td>+</td>
<td>1</td>
<td>3 (1.8)</td>
</tr>
<tr>
<td>Methylation alteration in F1 and amphiploid</td>
<td>–</td>
<td>–</td>
<td>+</td>
<td>–</td>
<td>1</td>
<td>11 (6.9)</td>
</tr>
<tr>
<td>Methylation alteration in amphiploid</td>
<td>–</td>
<td>+</td>
<td>–</td>
<td>+</td>
<td>6</td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup> +, band present; –, band absent. Each row represents a different MSAP banding pattern that was observed in the parents-F1-allotetraploid combination.
that there may be a difference between genomes in their ability to be modified (demethylated or hypermethylated) in the polyploid background. Methylation alterations in F1 involved mostly repetitive DNA (Table 5), with some examples from retrotransposons. Although we did not find examples of known genes, the methylation alterations found here might be related to the allopolyploidy-associated changes in methylation of coding sequences (Liu et al., 1998a) and to the gene silencing recently described for Arabidopsis using cDNA-AFLP (Comai, 2000). The demethylation of retrotransposons shown in this work (Table 5) might be consistent with transposon activation observed as the spread of dispersed repetitive DNA, including retroelements, to new genomes since polyploid formation in cotton (Zhao et al., 1998). Similarly, demethylation may affect DNA transposon activity, as was shown for several maize elements, such as Mutator, Activator, and Suppressor-mutator (Chandler and Walbot, 1986; Dennis et al., 1986; Banks et al., 1988). On

Table 5. DNA Gel Blot Analysis of the Fragments That Showed Methylation Alteration in the F1 Hybrid of the Cross Ae. sharonensis (S/S) and T. monococcum ssp aegilopoides (A^nA^?) and in the Derived Allotetraploid

<table>
<thead>
<tr>
<th>Probe</th>
<th>Primer (HM+4/E+3)^a</th>
<th>Fragment Size (bp)</th>
<th>Origin of the Fragment</th>
<th>Methylation Alterationb</th>
<th>Copy Number</th>
<th>Sequence Similarityc</th>
</tr>
</thead>
<tbody>
<tr>
<td>MIF1</td>
<td>E+ AAG</td>
<td>210</td>
<td>sharonensis</td>
<td>+</td>
<td>High</td>
<td>Barley retrotransposon</td>
</tr>
<tr>
<td>MIF2</td>
<td>E+ AAG</td>
<td>260</td>
<td>aegilopoides</td>
<td>+</td>
<td>Low</td>
<td>None</td>
</tr>
<tr>
<td>MIF3</td>
<td>E+ AGG</td>
<td>100</td>
<td>sharonensis</td>
<td>+</td>
<td>High</td>
<td>Barley retrotransposon</td>
</tr>
<tr>
<td>MIF4</td>
<td>E+ AGG</td>
<td>150</td>
<td>aegilopoides</td>
<td>+</td>
<td>High</td>
<td>Ceratitis capitata unknown repetitive DNA fragment</td>
</tr>
<tr>
<td>MIF5</td>
<td>E+ ACA</td>
<td>134</td>
<td>aegilopoides</td>
<td>+</td>
<td>High</td>
<td>None</td>
</tr>
<tr>
<td>MIF6</td>
<td>E+ AAG</td>
<td>350</td>
<td>aegilopoides</td>
<td>–</td>
<td>Low</td>
<td>None</td>
</tr>
<tr>
<td>MIF7</td>
<td>E+ ACT</td>
<td>110</td>
<td>aegilopoides</td>
<td>–</td>
<td>High</td>
<td>None</td>
</tr>
<tr>
<td>MIF8</td>
<td>E+ ACT</td>
<td>240</td>
<td>sharonensis</td>
<td>–</td>
<td>Low</td>
<td>None</td>
</tr>
<tr>
<td>MIF9</td>
<td>E+ ACA</td>
<td>270</td>
<td>aegilopoides</td>
<td>–</td>
<td>High</td>
<td>None</td>
</tr>
<tr>
<td>MIF10</td>
<td>E+ ACC</td>
<td>120</td>
<td>aegilopoides</td>
<td>–</td>
<td>Low</td>
<td>None</td>
</tr>
<tr>
<td>MIF11</td>
<td>E+ ACC</td>
<td>405</td>
<td>aegilopoides</td>
<td>–</td>
<td>Low</td>
<td>None</td>
</tr>
<tr>
<td>MIF12</td>
<td>E+ ACC</td>
<td>275</td>
<td>aegilopoides</td>
<td>–</td>
<td>High</td>
<td>None</td>
</tr>
</tbody>
</table>

^a The selective primer HM+4 (5’-CATGAGTCTGCTCGGTCAA [HM+TCAA]) was used in combination with each of the four E+3 primers listed. The core sequence of EcoRI is identical to that used in the AFLP protocol by Vos et al. (1995).
b +, methylation alteration; –, no alteration.
c Similarity was considered significant for e values ≤ 0.005.

Figure 4. DNA Gel Blot Analysis Using MIF8 Probe That Showed an Alteration in MSAP Pattern in the Allotetraploid (S1) of the Cross between Ae. sharonensis (TH) and T. monococcum ssp aegilopoides (TMB).

(A) Each DNA sample was digested with EcoRI-HpaII (H) and EcoRI-MspI (M), and the two digests were loaded on the gel side by side (HpaII digest on the left and MspI digest on the right). The arrow points to the fragment whose methylation is altered in the allotetraploid.
(B) DNA samples were digested with DraI (not sensitive to cytosine methylation) using the same hybridization probe. MIF8 was obtained using primer combination HM+TCAA/E+ACT (Table 5).
the other hand, we did not find new AFLP bands corresponding to retroelements or DNA transposons (Table 2), suggesting that there was no large-scale occurrence of new transposon insertion in the F1 hybrid or the allopolyploid.

Underlying Mechanism of Rapid and Reproducible Allopolyploidy-Associated Sequence Elimination

Our finding that a significant fraction of the genome can undergo elimination in F1 and in newly synthesized allotetraploids suggests that both interspecific and intergeneric hybridity and allopolyploidy are shocks that trigger a rapid and massive genomic response in wheat. Surprisingly, this response was reproducible; that is, the same loci were always involved in the same patterns of sequence elimination (Table 2; Figure 5; see also Ozkan et al., 2001). We did not find evidence for a random mutator effect, for example, through point mutations, recombination, and transposable element activities. Such rearrangements would have caused new RFLPs or AFLP fragments (compared with the parents) to appear in the F1 hybrid and the allopolyploids. Elimination occurred rapidly and in a reproducible manner. A number of mechanisms could explain the type of elimination observed here: (1) gene conversion between the homoeoalleles (with rather long conversion tracts); (2) excision of the eliminated sequence and subsequent loss, via mechanisms such as transposon excision, V(D)J recombination, or site-specific recombination, as was shown in *Tetrahymena* (Godiska and Yao, 1990); and (3) crossing over between direct repeats that flank the eliminated sequence and loss of the excised circle. Gene conversion is the least likely mechanism because it would require very long conversion tracts and recombination between homologous chromosomes. Moreover, mapping of some eliminated sequences using nullisomic-tetrasomic lines (e.g., WPG90 used by Feldman et al., 1997) showed that these sequences were present in only one set of chromosomes and were absent in the homoeologs.

At this time, we have no support for or against the other mechanisms, and experiments are under way in our laboratory to determine the deletion or recombination breakpoints and thus gain more insight into the underlying elimination mechanism. In any case, this work shows that sequence elimination is the major and immediate response of the wheat genome to wide hybridization and allopolyploidy, that it can affect a large fraction of the genome (up to 14% of the polymorphic fragments), and that it is reproducible. In this respect, allopolyploidy somehow fits McClintock’s definition of a genomic shock that “initiates a highly programmed sequence of events within the cell that serves to cushion the effects of the shock” (McClintock, 1984). Future studies may reveal whether the well-orchestrated and reproducible sequence elimination reported here is an example of preprogrammed adaptation of the genome to amphiploidy or a fortuitous response with no adaptive significance.

Figure 5. DNA Gel Blot Analysis of a Band Isolated from MSAP That Did Not Show Alteration in Methylation but Rather Showed Band Disappearance.

The bands obtained by both isoschizomers in parent *T. monococcum* ssp *aegilopoides* (TMB) were eliminated in the first generation of the allotetraploid (S1). TH, *Ae. sharonensis*.

(A) Each DNA sample was digested with EcoRI+HpaII (H) and EcoRI+MspI (M), and the two digests were loaded on the gel side by side (HpaII digest on the left and MspI digest on the right).

(B) DNA samples were digested with EcoRV using the same hybridization probe. The probe was obtained using primer combination HM+TCAA/E+AAG. Arrows indicate the band of TMB that disappeared in the allotetraploid.

METHODS

Plant Material

The plant material for this study consists of three different combinations of newly synthesized allotetraploids, F1 hybrids, and their homozygous diploid parents. The first cross involved *Aegilops sharonensis* (genome SSS) × *Ae. umbellulata* (genome UU), the second cross involved *Ae. longissima* (genome SSS) × *Triticum urartu* (genome AA), and the third cross involved *Ae. sharonensis* (genome SSS) × *T. monococcum* ssp *aegilopoides* (genome A′A′). The synthetic allotetraploids were obtained after colchicine treatment of the F1 plants. Chromosome number was determined in root tip mitosis of the F1 hybrids and the allotetraploids. The parental lines and the synthetic allopolyploids are maintained in the plant collection of our laboratory, and their production is described by Ozkan et al. (2001).

Amplified Fragment Length Polymorphism (AFLP) Analysis

The AFLP procedure was performed according to the protocol of Gibco BRL according to Vos et al. (1995), with minor modifications. Genomic DNA was extracted from young fresh leaves by the cetyltrimethyl-ammonium bromide procedure (Kidwell and Osborn, 1992). One-half microgram of genomic DNA was digested at 37°C for 2 hr. The restriction-ligation reaction contained 12 units of Msel, 10 units of EcoRI, 1 unit of T4-DNA ligase, 5 pmol of EcoRI adaptor, 50 pmol of Msel adaptor, 0.5 M NaCl, 5 µg of BSA, and 10× DNA ligase buffer (1× DNA ligase buffer is 50 mM Tris-HCl, pH 7.5, 10 mM MgCl2, 10 mM dithiothreitol, 1 mM ATP, 25 mg/ml bovine serum albumin) in a final volume of 10 µL. The digested-ligated DNA was diluted 1:10 with 90 µL of double-distilled water.
The adaptor sequences were as follows: MseI adapters, 5’-TAC-TCAAGGACTCAT-3’ and 5’-GACGATGAGTCCTGAG-3’; EcoRI adapters, 5’-CTCGTAGTACGCGTACC-3’ and 5’-AATTGGTAGACGATC-TAC-3’. Preselective amplification was performed with primers complementary to the core of the adaptor sequences; the EcoRI preselective primer was 5’-GACTGCGTACCAATCTA-3’, and the MseI preselective primer was 5’-GATGAGTCCTGAGTAC-3’. The polymerase chain reaction (PCR) contained 50 ng of restricted-ligated DNA, 50 ng of EcoRI preselective primer, 50 ng of MseI preselective primer, 1 unit of Taq DNA polymerase, 2 μL of 10 × Taq DNA polymerase buffer (1 × Taq DNA polymerase buffer is 50 mM Tris HCl, pH 8.0, 100 mM NaCl, 0.1 mM EDTA, 1 mM DDT, 50% glycerol, 2 μL of 25 mM MgCl₂, and 2.5 mM deoxynucleotide triphosphates in a final volume of 20 μL). The PCR conditions were 20 cycles of 30 sec at 94°C, 60 sec at 56°C, and 60 sec at 72°C. After preamplification, the PCR products were diluted 1:20 with 190 μL of double-distilled water.

Selective amplification was performed with the 32P-end-labeled EcoRI selective primers and the unlabeled MseI selective primers. The labeling reaction contained 50 ng of EcoRI selective primer, 100 μCi of γ-32P-ATP, 10 units of T4 polynucleotide kinase, and 10 μL of 5 × polynucleotide kinase buffer (1 × polynucleotide kinase buffer is 50 mM Tris HCl, 1 mM DDT, 0.1 mM EDTA, 50% glycerol [v/v], pH 7.5) in a final volume of 50 μL. Samples were incubated at 37°C for 1 hr and then heated to 70°C for 10 min. The sequence of the selective primers is similar to that of the preselective primers, with the addition of two variable nucleotides at the 3’ end. The AFLP primers are identified by the last three specific nucleotides of the selective primers, namely EcoRI + ANN primers and MseI + CNN primers.

The selective amplification reaction contained 50 ng of template DNA from the preselective amplification, 1 ng of labeled EcoRI selective primer, 5 ng of MseI selective primer, 1 unit of Taq DNA polymerase, 2 μL of Taq DNA polymerase buffer, 2 μL of 25 mM MgCl₂, and 2.5 mM deoxynucleotide triphosphates in a final volume of 20 μL. The PCR cycles were one cycle of 2 min at 94°C, 30 sec at 65°C, and 2 min at 72°C, followed by 10 cycles each of annealing temperature of 1°C less than the former one, and 25 cycles of 1 sec at 94°C, 30 sec at 56°C, and 2 min at 72°C. The PCR products were mixed with 20 μL of dye (98% formamide, 10 mM EDTA, 0.1% bromphenol blue, and 0.1% xylene cyanol). The denatured PCR products were separated on 6% denaturing polyacrylamide (20:1 acrylamide:bisacrylamide, 7.5 M urea, and 1 × Tris-borate-EDTA buffer, pH 7.8) standard sequencing gel (43 cm in length) at 55 W for 1.5 hr. Gels were dried and exposed to x-ray film (Fuji Photo Film, Tokyo, Japan) for 6 to 12 hr at room temperature.

Methylation-Sensitive Amplification Polymorphism (MSAP) Analysis

The original protocol for detecting AFLP provided by Vos et al. (1995) was modified to incorporate the use of methylation-sensitive restriction enzymes. The modified protocol involved the use of the isoschizomers HpaII and MspI in combination with EcoRI. The adapter and the basic primer sequences for the EcoRI end were the same as those provided in the original protocol (Vos et al., 1995). A new double-strand fragment, referred to as the HpaII-MspI adaptor, was designed for each of the isoschizomer digestions using oligonucleotides 5’-GATGAGTCCTGAGTAC-3’ and the complementary strand. The basic primer sequence for the HpaII-MspI digestions (HM-0) was designed accordingly as 5’-ATCATGAGTCTCGGTCCG-3’.

Primer sequences for the EcoRI ends (E+3) and four selective nucleotides for the HpaII-MspI ends (HM+4) were designed as shown in Table 3. All of the adaptor and primer sequences were synthesized by the nucleic acid synthesis unit at the Weizmann Institute of Science.

Scoring of AFLP and MSAP bands

Each AFLP and MSAP gel was run twice. For the AFLP analysis, the replicate gel was run from the same DNA sample but from a different restriction-ligation-amplification reaction. For the MSAP analysis, the replicate gel was derived from an independent DNA extraction. Moreover, with both AFLP and MSAP gels, the upper part and the lower part of the gel, where resolution is not satisfactory, was not used for band scoring. The summary of the various types of bands shown in Tables 1, 3, and 4 is based only on the high-resolution, middle part of the gel, which was reproducible in all cases.

Cloning of Fragments Subjected to Genetic or Epigenetic Alterations and Validation by DNA Gel Blot Analysis

Fragments that showed evidence of polyploidy-associated alterations were excised from AFLP gels and reamplified using the following PCR conditions: 3 min at 94°C, 30 sec at 94°C, 1 min at 56°C, 1 min at 72°C, followed by 34 cycles. The AFLP-isolated fragments (AIFs) and the MSAP-isolated fragments (MIFs) were ligated into pGEM-T easy vector (Promega), transformed to Escherichia coli strain XL-1 Blue, and sequenced using universal T7 and/or T3 primers. The AIFs and the MIFs were labeled with phosphorus-32 and used as hybridization probes for DNA gel blot analysis. The procedure of DNA gel blot analysis followed essentially the method described by Liu et al. (1997), except that methylation analyses used either HpaII or MspI.

Sequence Analysis

Sequences obtained in this study were analyzed for similarity to known sequences in public databases using the BLAST package 2.0 on the National Center for Biotechnology Information server (http://www.ncbi.nlm.nih.gov/BLAST/).

ACKNOWLEDGMENTS

We thank an anonymous referee for constructive comments. This work was supported by a United States–Israel Binational Science Foundation fund grant. H.O. was supported by the Turkish Council of Higher Education and by a short-term European Molecular Biology Organization fellowship. K.K. and H.S. were supported by a fellowship from the Feinberg Graduate School for Ph.D. and M.Sc., respectively.

Received March 1, 2001; accepted June 8, 2001.
REFERENCES


Sequence Elimination and Cytosine Methylation Are Rapid and Reproducible Responses of the Genome to Wide Hybridization and Allopolyplody in Wheat
Hezi Shaked, Khalil Kashkush, Hakan Ozkan, Moshe Feldman and Avraham A. Levy
Plant Cell 2001;13;1749-1759
DOI 10.1105/TPC.010083

This information is current as of October 20, 2017

References
This article cites 27 articles, 11 of which can be accessed free at:
/content/13/8/1749.full.html#ref-list-1

Permissions

eTOCs
Sign up for eTOCs at:
http://www.plantcell.org/cgi/alerts/ctmain

CiteTrack Alerts
Sign up for CiteTrack Alerts at:
http://www.plantcell.org/cgi/alerts/ctmain

Subscription Information
Subscription Information for The Plant Cell and Plant Physiology is available at:
http://www.aspb.org/publications/subscriptions.cfm

© American Society of Plant Biologists
ADVANCING THE SCIENCE OF PLANT BIOLOGY