Sequencing, annotation and characterization of 17 Mb of chromosome 3B contigs provide novel insights into the wheat genome organization and evolution
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Sequencing, annotation and characterization of 17 Mb of chromosome 3B contigs provide novel insights into the wheat genome organization and evolution

Our current knowledge of the wheat genome structure is limited to the analysis of individual BAC sequences and of random BAC end and plasmid sequences. To improve our understanding of the composition and evolution of this large and complex genome, we have sequenced and completely annotated 136 BAC clones assembled into 12 contigs of 0.5 to 3.1 Mb in size originating from different regions of chromosome 3B. This first set of large wheat contiguous sequences account for 17 Mb and contains 128 genes, 21 pseudogenes and 14 gene relics as well as 2963 transposable elements, representing 83.5% of the sequences. The average gene density is of 1 gene/114 kb with important local variations ranging from 0 to 10 genes per BAC. Gene distribution is not random with 77% of the genes being clustered into several small islands containing 3 genes in average and a slight gradient of gene density towards the telomeric regions. Interestingly, all Mb-sized contigs carry genes irrespective of their chromosomal location. Only 32% of the BACs are devoid of genes and the largest gene-free region is 709 kb indicating that gene islands are small and scattered all along the chromosome, thus supporting the idea that the whole genome needs to be sequenced to ensure access to the entire gene space. Only half of the predicted wheat genes are orthologous and syntenic with rice chromosome 1. The other half consists of genes present in rice but on different chromosomes. Non syntenic genes are found in almost all (9/10) sequenced gene-containing regions with large variations depending on their composition and chromosomal location. The highest level of rearrangement was observed for a disease resistance region with 2/3 of the genes at non orthologous positions whereas other regions exhibit >90% of syntenic genes. We also precisely annotated the 2963 transposable elements and identified 271 different families, including 155 new ones (at least 2 members), and 356 unknown single copy elements. TE composition shows large variations along the chromosomes with some families strictly found in centromeric or telomeric regions. Compared to isolate BAC sequencing, analyses of Mb-sized contigs give new insights into the genome structure by revealing, for instance, that most of TEs are complete and fragmented by nested insertions that can reach more than 200 kb.

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