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Flow cytometric chromosome sorting from diploid progenitors of bread wheat, *T. urartu*, *Ae. speltoides* and *Ae. tauschii*

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Abstract

Key message Chromosomes 5A^u, 5S and 5D can be isolated from wild progenitors, providing a chromosomebased approach to develop tools for breeding and to study the genome evolution of wheat.

Abstract The three subgenomes of hexaploid bread wheat originated from *Triticum urartu* (A^uA^u), from a species similar to *Aegilops speltoides* (SS) (progenitor of the B genome), and from *Ae. tauschii* (DD). Earlier studies indicated the potential of chromosome genomics to assist gene transfer from wild relatives of wheat and discover novel genes for wheat improvement. This study evaluates the potential of flow cytometric chromosome sorting in the diploid progenitors of bread wheat. Flow karyotypes obtained by analysing DAPI-stained chromosomes were characterized and the contents of the chromosome peaks were determined. FISH analysis with repetitive DNA probes proved that chromosomes $5A^u$, 5S and 5D could be sorted with purities of 78–90 %, while the remaining chromosomes

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M. Kubaláková · H. Šimková · J. Vrána · J. Doležel Centre of the Region Hana for Biotechnological and Agricultural Research, Institute of Experimental Botany, Šlechtitelů 31, 78371 Olomouc, Czech Republic could be sorted in groups of three. Twenty-five conserved orthologous set (COS) markers covering wheat homoeologous chromosome groups 1–7 were used for PCR with DNA amplified from flow-sorted chromosomes and genomic DNA. These assays validated the cytomolecular results as follows: peak I on flow karyotypes contained chromosome groups 1, 4 and 6, peak II represented homoeologous group 5, while peak III consisted of groups 2, 3 and 7. The isolation of individual chromosomes of wild progenitors provides an attractive opportunity to investigate the structure and evolution of the polyploid genome and to deliver tools for wheat improvement.

Introduction

Bread wheat (Triticum aestivum L.) is the second most widely grown crop and is the primary cereal in the temperate region, serving as a staple food for about 40 % of the world's population (http://faostat.fao.org). Bread wheat has a large genome of about 17,000 Mb with three constituent subgenomes, A, B and D (Devos and Gale 2000). The genome architecture of wheat originated from two successive interspecific hybridizations and polyploidizations. The first hybridization occurred approximately 0.3-0.5 million years ago between the A genome progenitor Triticum *urartu* Thum. ex Gandil. ($A^{u}A^{u}$, 2n = 2x = 14) and a B genome progenitor species which is considered to be similar to Aegilops speltoides Tausch. (SS, 2n = 2x = 14). This resulted in the evolution of wild emmer wheat T. turgidum ssp. dicoccoides (Körn.) Thell. (A^uA^uBB , 2n = 4x = 28) (Dvorak et al. 1993; Maestra and Naranjo 1998). Human selection then created cultivated emmer wheat T. turgidum ssp. dicoccon (Schrank) Thell., which hybridized spontaneously with Ae. tauschii (DD, 2n = 2x = 14) around 9,000 years ago to produce allohexaploid wheat, *T. aestivum* L. $(2n = 6x = 42, A^uA^uBBDD)$ (Dvorak et al. 1998).

Domestication and thousands of years of cultivation narrowed down the genetic variation within bread wheat. The narrow gene pool of wheat relative to that of its wild diploid progenitor species was demonstrated by Lubbers et al. (1991), who reported a polymorphic index of 0.41 after investigating 102 Ae. tauschii accessions at 20 RFLP loci. This was significantly higher than the index determined for hexaploid wheat (0.04) after investigating 21 wheat cultivars originating from different location around the world using 33 genomic probes (Cox 1998). Thus, the genetically diverse diploid progenitors represent a huge reservoir of alleles and genes, which could be used to increase the genetic variation of hexaploid wheat through interspecific hybridization (Cox 1998; Friebe et al. 1996). Several agronomic traits, including resistance to pests and diseases, tolerance to abiotic stresses and traits affecting nutritional and bread making quality or yield components, were identified in accessions of wild diploid progenitors of wheat. Rouse and Jin (2011), for example, identified 86 accessions of T. urartu conferring resistance against the stem rust race TTKSK (Ug99) of Puccinia graminis f. sp. tritici. Diploid T. urartu is also recognized as a source of traits affecting bread-making quality (Guzmán and Alvarez 2012). Ae. speltoides serves as a source of biotic stress tolerance genes, which have already been transferred to wheat (Lr28, Lr35, Lr36, Lr47, Lr51, Sr32, Sr39, Pm12, Pm32, Gb5) (see Kilian et al. 2011). Moreover, this species is a potential source of other important traits such as resistance against pests and diseases (Fusarium head blight, Hessian fly, Septoria tritici), grain hardness protein, heat tolerance and tolerance to manganese toxicity (see Kilian et al. 2011). Ae. tauschii is probably the most frequently studied diploid progenitor. Many resistance genes against pests and diseases have already been transferred from Ae. tauschii to wheat (see Schneider et al. 2008; Kilian et al. 2011) and accessions of Ae. tauschii are also attractive sources of adaptive traits, yield components, bread-making quality and abiotic stress tolerance (Fritz et al. 1995; Cox et al. 1990; Schachtman et al. 1992; Limin and Fowler 1993; Xiu-Jin et al. 1997).

Despite the extensive research efforts, the introgression of favourable agronomic traits from diploid progenitor species to cultivated wheat via interspecific hybridization remains difficult due to the considerable number of undesirable genes located in the targeted genomic regions (Klindworth et al. 2013). A better knowledge of the diploid progenitors of the wheat genomes, including their homology to bread wheat subgenomes, and the application of molecular tools could accelerate the targeted introgression of favourable traits into wheat, as well as providing an invaluable resource for studying the structure and evolution of the wheat genome. Molecular genetic studies on diploid progenitors, including the creation of genetic linkage maps and large-insert bacterial artificial chromosome (BAC) libraries, have already been reported (Gill et al. 1991; Boyko et al. 1999; Akhunov et al. 2005).

The advent of next-generation sequencing (NGS) platforms paved the way for cost-effective large-scale DNA sequencing in plants (Margulies et al. 2005; Schatz et al. 2010; Edwards and Batley 2010). These methods are increasingly being applied in plant genomic research and in breeding major cereal crops and related species, including bread wheat, Ae. tauschii and T. urartu (Brenchley et al. 2012; You et al. 2011; Ling et al. 2013), thus facilitating SNP discovery, physical mapping and whole genome shotgun sequencing (Kumar et al. 2012). However, the successful application of NGS technology for de novo sequencing in species with large and complex genomes may be problematic, as the high proportion of repetitive sequences (57, 61 and 57 % for the A^u, S and D genomes, respectively, in diploid wheats) hampers the assembly of short NGS reads (Kilian et al. 2011; Shangguan et al. 2013).

The analysis of large genomes can be simplified by reducing the sample complexity by isolating single chromosomes and analysing them instead of the whole genome. In recent decades, the development of flow cytometric chromosome sorting in plants has opened the way for the application of genomics tools to chromosomes to obtain shotgun sequences, to develop markers and to construct sequenceready physical maps (Doležel et al. 2012, 2014). The application of the NGS technology (Edwards and Batley 2010) allowed the sequencing of wheat chromosome arm 7DS to $34 \times$ coverage using Illumina and the assembly of low copy and genic regions of this chromosome, representing approximately 40 % of the chromosome arm and all known 7DS genes (Berkman et al. 2011). The Roche 454 technology was applied for the sequencing of barley chromosome 1H and the arms of barley chromosomes 2H-7H to about $2 \times$ coverage and 21,766 barley genes were assigned to individual chromosome arms. Using the conserved synteny with the genomes of rice, sorghum and Brachypodium, the barley genes were arranged in a putative linear order on the individual chromosome arms (Mayer et al. 2009, 2011). In a similar study, Martis et al. (2013) sequenced flow-sorted chromosomes of rye and established linear gene order model (genome zipper) comprising 22,426 or 72 % of the detected set of 31,008 rye genes. Moreover, the study indicated that introgressive hybridizations and/or a series of whole-genome or chromosome duplications played a role in rye speciation and genome evolution.

Chromosome genomics relies on the ability to isolate chromosomes via flow-cytometric sorting. The flowcytometric analysis of mitotic chromosomes has been reported in 24 plant species (Doležel et al. 2014), including hexaploid and tetraploid wheat and their wild relatives in the genus Aegilops and Dasypyrum (Molnár et al. 2011b; Grosso et al. 2012). To date, flow cytometric chromosome analysis and sorting has not been reported for the diploid progenitors of bread wheat. As the technology could greatly aid the transfer of genes from wild relatives to cultivated wheat and the study of the evolution of polyploid wheat, we set out to explore the possibility of isolating individual chromosomes from diploid T. urartu, Ae. speltoides and Ae. tauschii by means of flow sorting. Chromosomes were sorted from the individual peaks of flow karvotypes and were identified by FISH with a set of repetitive DNA probes. DNA amplified from isolated chromosomes was used as a template for PCR using conserved orthologous set (COS) markers with the aim of identifying their genomic location in the diploid progenitors of wheat and of confirming the cytological identification of sorted chromosomes. The results of the present work provide an important step towards analysing the molecular organization of chromosomes in the diploid progenitors of wheat and towards developing tools to support alien gene transfer in wheat improvement programmes.

Materials and methods

Plant material

Triticum urartu accession MvGB115, *Ae. speltoides* accession MvGB905 and *Ae. tauschii* accession MvGB605, maintained in the Martonvásár Cereal Gene Bank, were used for flow cytometric chromosome analysis and sorting for in situ hybridization experiments and for COS marker analysis.

Preparation of liquid suspensions of chromosomes

The synchronization of the cell cycle of root tip meristem cells using 2 mM hydroxyurea and their accumulation in metaphase using 2.5 µM amiprohos-methyl were carried out as described by Kubaláková et al. (2005). Suspensions of intact chromosomes were prepared from synchronized root tips according to Vrána et al. (2000). Briefly, 50 roots were cut 1 cm from the root tip and fixed in 2 % (v/v) formaldehyde in Tris buffer (10 mM Tris, 10 mM Na₂EDTA, 100 mM NaCl, 0.1 % Triton X-100, pH 7.5) at 5 °C for 20 min. After washing in Tris buffer, the meristem tips were excised and transferred to a tube containing 1 ml of LB01 buffer (15 mM Tris, 2 mM Na2EDTA, 80 mM KCl, 20 mM NaCl, 0.5 mM spermine, 15 mM β-mercaptoethanol, 0.1 % Triton X-100) (Doležel et al. 1989) at pH 9. Metaphase chromosomes were released after homogenization with a Polytron PT1300 homogenizer (Kinematica AG, Littau,

Switzerland) at 20,000 rpm for 13 s. The crude suspension was passed through a 50- μ m pore size nylon mesh to remove large cellular debris and stored on ice until analysis on the same day.

Flow cytometric chromosome analysis and sorting

The samples were analysed using a FACSVantage SE flow cytometer (Becton-Dickinson, San José, USA) equipped with an argon ion laser set to multiline UV and 300 mW output power. Chromosome suspensions were stained with DAPI (4',6-diamidino-2-phenylindole) at a final concentration of 2 μ g/ml and analysed at rates of 200–400 particles per second. DAPI fluorescence was acquired through a 424/44 band-pass filter. Approximately 30, 000 chromosomes were analysed from each sample and the results were displayed as histograms of relative fluorescence intensity (flow karyotypes). In order to verify the chromosome content of individual peaks on the flow karyotypes, 1, 000 chromosomes were sorted from each peak at rates of approximately 5-10 per second into a 15 µl drops of PRINS buffer (100 mM Tris, 50 mM KCl, 2 mM $MgCl_2 \times 6H_2O$) supplemented with 5 % sucrose on a microscope slide (Kubaláková et al. 1997), air-dried and used for FISH with probes for DNA repeats that give chromosome-specific fluorescent labelling patterns.

Fluorescence in situ hybridization

Total genomic DNA was extracted from fresh leaves of Ae. tauschii (D genome), S. cereale and O. sativa using Quick Gene-Mini80 (FujiFilm, Tokyo, Japan) according to the manufacturer's instructions. The repetitive DNA sequences Afa family, pSc119.2 and the 18S unit of the 45S ribosomal RNA gene were amplified using PCR from the genomic DNA of Ae. tauschii, S. cereale and rice, respectively, as described by Nagaki et al. (1995), Contento et al. (2005) and Chang et al. (2010). Afa, pSc119.2 and the 18S rRNA unit were labelled with digoxigenin-11-dUTP (Roche, Manheim, Germany), biotin-16-dUTP (Roche) and a mix of biotin-11-dUTP (50 %) and digoxigenin-11-dUTP (50 %), respectively, by nick-translation using standard kits from Roche following the manufacturer's instructions. Digoxigenin and biotin were detected using anti-digoxigenin-rhodamine Fab fragments (Roche) and streptavidin-FITC (Roche), respectively.

Pretreatments and stringency washes (Schneider et al. 2005) were applied only for slides containing root tip metaphase cells. These steps were omitted in experiments on flow-sorted chromosomes. The hybridization mix (30 μ l per slide), containing 50 % formamide, 2 × SSC, 10 % dextran sulphate, 20 ng of 18S rDNA and 70 ng each of the pSc119.2 and *Afa* family probes in the presence of 6.25 µg Salmon sperm DNA, was denatured at 80 °C for 10 min and stored on ice for 5 min. The chromosome DNA was denatured in the presence of the hybridization mix at 75 °C for 6 min and allowed to hybridize overnight at 37 °C. The detection of the hybridization signals involved the use of 10 μ g ml⁻¹ each of streptavidin-FITC and anti-digoxigenin-Rhodamin. Finally, the slides were counterstained with 2 μ g ml⁻¹ DAPI and examined under a Zeiss Axioskop-2 fluorescence microscope using a Plan Neofluar oil objective 63×, N.A. 1.25 (Zeiss, Oberkochen, Germany) equipped with filter sets appropriate for DAPI (Zeiss filter set 02) and for FITC and Rhodamin (Zeiss filter set 24). Images were acquired with a Spot CCD camera (Diagnostic Instruments, Sterling Heights, USA) and compiled with Image Pro Plus software (Media Cybernetics, Silver Spring, USA). The relative fraction of group 5 chromosomes as a percentage of the whole genome was determined by measuring the lengths of at least 20 chromosomes per homoeologous group on DAPI-stained pictures of T. urartu MvGB115, Ae. speltoides MvGB905 and Ae. tauschii MvGB605.

Amplification of chromosomal DNA

Chromosomes were sorted from each peak on the flow karyotype in batches of 25–50,000 (equivalent to 20–40 ng) into PCR tubes containing 40 μ l of sterile deionized water. The chromosomes were treated with proteinase K and their DNA was amplified by multiple displacement amplification (MDA) using an Illustra GenomiPhi V2 DNA Amplification Kit (GE Healthcare, Chalfont St. Giles, United Kingdom) as described by Šimková et al. (2008). The amplified DNA was used as a template for PCR with microsatellite markers.

COS marker analysis

Genomic DNA was prepared from the accessions of *T. urartu* MvGB115, *Ae. speltoides* MvGB905 and *Ae. tauschii* MvGB605, used for the flow-cytometric analysis and from the wheat (*T. aestivum* L.) genotype Mv9kr1 as described by Cseh et al. (2013). A total of 25 conserved orthologous set (COS) markers (Online Resource 1) specific for wheat homoeologous groups I-VII were chosen from publicly available COS marker collections (the Wheat Genetic Improvement Network: http://www.wgin.org.uk/resources/Markers/TAmarkers.php; Tools and Resources (TR) collections: http://www.modelcrop.org/cgi-bin/gbrowse/brachyv1/; Quraishi et al. 2009).

PCR reactions were performed in a 12- μ l reaction volume in a reaction mix consisting of 1 × PerfectTaq Plus PCR Buffer (5 Prime GmbH, Hamburg, Germany) and 0.4 μ M primers; 50 ng genomic DNA or 1.5 ng of MDA

DNA from flow-sorted chromosomes was used as a template. PCR was carried out in an Eppendorf Mastercycler (Eppendorf, Hamburg, Germany) using the reaction profiles WGIN: 95 °C (15 min), 39 cycles of [95 °C (0.5 min), 58 °C (0.5 min), 72 °C (0.5 min)], hold at 72 °C (5 min) then at 10 °C and TR: 94 °C (10 min), 16 cycles of [95 °C (0.5 min), 58 °C (1 min), decreasing by 0.5 °C per cycle to 50 °C, 72 °C (1 min)], 25 cycles of [94 °C (0.5 min), 50 °C (1 min), 72 °C (1 min)], hold at 15 °C. The annealing temperature and PCR reaction profiles are summarized together with the primer sequences in Online Resource 1. The PCR amplicons were separated with a Fragment AnalyzerTM Automated CE System equipped with a 12-Capillary Array Cartridge (effective length 33 cm) (Advanced Analytical Technologies, Ames, USA). The results were analysed using PROsize v2.0 software.

Results

Chromosome analysis using flow cytometry (flow karyotyping)

The analysis of DAPI-stained chromosome suspensions prepared from the diploid progenitors of hexaploid wheat resulted in flow karyotypes with three peaks (Fig. 1). However, there were differences between the species in the degree of resolution of the individual peaks and their position on the flow karyotype. While the second peak on the flow karyotypes of *T. urartu* and *Ae. speltoides* could only partially be resolved from the composite peaks I and III (Fig. 1a, b), in *Ae. tauschii* peak II was clearly discriminated (Fig. 1c). Moreover, the chromosome peaks on the flow karyotype of *T. urartu* were observed at higher fluorescence intensity channels (115–160) than those of *Ae. speltoides* (channels 105–140), while the peak positions in *Ae. tauschii* were located at the lowest fluorescence interval (channels 95–135) (Fig. 1).

FISH on mitotic metaphase spreads

In order to obtain a reference karyotype for the identification of flow-sorted chromosomes, root tip mitotic metaphase spreads from accessions of the diploid progenitor species were investigated using fluorescence in situ hybridization (FISH) with repetitive DNA probes (*Afa* family, pSc119.2, 18S rDNA), which allowed the whole set of chromosomes in *T. urartu* MvGB115, *Ae. speltoides* MvGB905 and *Ae. tauschii* MvGB605 to be identified (Fig. 2). Differences were observed in the FISH hybridization patterns of the genotypes used in this study compared with those used previously or those of durum and hexaploid wheat (Badaeva et al. 1996a, b; Kubaláková et al.

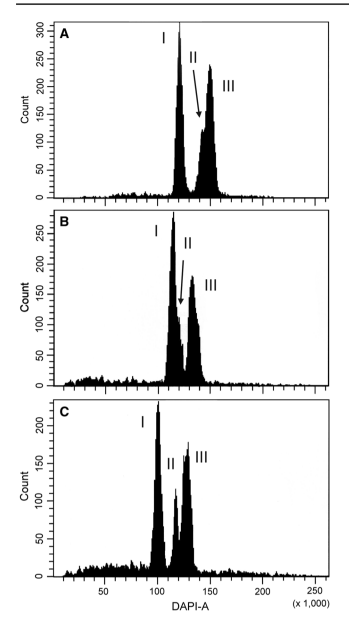


Fig. 1 Flow karyotypes obtained after the analysis of DAPI-stained chromosome suspensions prepared from *T. urartu* MvGB115 $(2n = 2x = 14, A^uA^u)$ (**a**), *Ae. speltoides* MvGB905 (2n = 2x = 14, SS) (**b**) and *Ae. tauschii* MvGB605 (2n = 2x = 14, DD) (**c**). Each flow karyotype consists of two composite peaks I and III, each of them representing a group of three chromosomes. Chromosome 5 is represented by peak II, which is well resolved only in *Ae. tauschii*. In *T. urartu* and *Ae. speltoides* the peak of chromosome 5 is not well resolved, appearing as a shoulder on peak III in *T. urartu* and on peak I in *Ae. speltoides*

2005; Sepsi et al. 2008). Strong 18S rDNA signals were observed on the short arms of $1A^{u}$ and $5A^{u}$, which together with the *Afa* hybridization patterns (diagnostic bands on the chromosome arms $1A^{u}S$ and $5A^{u}L$) allowed the discrimination of the two chromosomes (Fig. 2a). Distinctive *Afa* labelling patterns were also observed on chromosome $2A^{u}$

and chromosome arm $4A^{u}S$. Chromosomes $3A^{u}$, $6A^{u}$ and $7A^{u}$ could be identified from their hybridization patterns, which were similar to those of the wheat chromosomes. The chromosomes of *Ae. speltoides* MvGB905 showed hybridization patterns similar to those of hexaploid wheat, which allowed the identification of the whole set of chromosomes (Fig. 2b).

The idiogram representing the genomic distribution of the *Afa* family, pSc119.2 and 18S rDNA sequences in *T. urartu* MvGB115, *Ae. speltoides* MvGB905 and *Ae. tauschii* MvGB605 is shown in Fig. 3. Differences in the fluorescence hybridization patterns facilitated the identification of each of the seven chromosomes of the diploid progenitors in the wheat background.

The chromosomes that showed different hybridization patterns relative to wheat were 2S, where a weak subtelomeric signal was detected relative to wheat 2B, 3S, where strong pSc119.2 subtelomeric signals were detected on the long arm, 4S, where the subtelomeric region of the long arm had only one weak pSc119.2 signal, and 7S, which showed a weak telomeric pSc119.2 band on the long arm relative to the corresponding wheat chromosomes. Only minor differences were found on chromosomes 3D and 4D (weak telomeric 18S rDNA and pSc119.2 signals, respectively, on the short arms) in Ae. tauschii relative to the corresponding chromosomes of wheat (Fig. 2c). However, the identification of the chromosomes was not hindered by these differences and all chromosomes of the diploid progenitor species could be distinguished on the basis of their fluorescence labelling patterns.

Description of flow karyotypes

The chromosome content of the individual peaks was determined after FISH on flow-sorted chromosomes with probes for Afa family, pSc119.2 and 18S rDNA repeats (Fig. 4; Table 1). The strong fluorescent signals given by the FISH probes were similar to those observed on mitotic metaphase spreads and allowed the unambiguous identification of the sorted chromosomes. In T. urartu, peaks I and III contained chromosomes 1A^u, 4A^u and 6A^u, and 2A^u, 3A^u and 7A^u, respectively. Peak II corresponded to chromosome 5A^u, representing 15.23 % of the *T. urartu* genome, which could be sorted at a purity of >78.53 % (Fig. 4; Table 1). Peaks I and III were also composite in Ae. speltoides, containing chromosomes 1S, 4S and 6S, and 2S, 3S and 7S, respectively (Fig 4; Table 1). As expected, the partially resolved peak II corresponded to chromosome 5S (~13.88 % of the S genome), which was sorted at a purity of 89.8 %. The distribution of the D genome chromosomes in Ae. tauschii between the peaks of the flow karyotypes was shown by FISH analysis to be similar to that observed in T. urartu and Ae. speltoides.

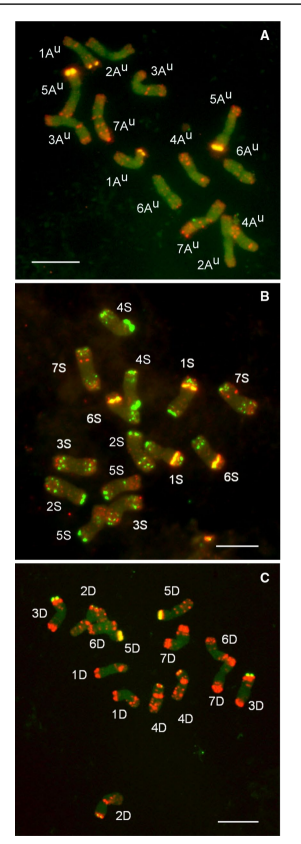


Fig. 2 Fluorescence in situ hybridization (FISH) with probes for repetitive DNA sequences pSc119.2 (green), Afa family (red) and 18S rDNA (yellow) on mitotic metaphase spreads of T. urartu MvGB115 (a), Ae. speltoides MvGB905 (b) and Ae. tauschii MvGB605 (c). Scale bars 10 μm

The composite peaks I and III contained the chromosome groups 1D, 4D and 6D, and 2D, 3D and 7D, respectively. The well-resolved peak II corresponded to chromosome 5D (~15.56 % of the D genome), which represented 87.3 % of the sorted particles (Fig. 4; Table 1).

Assignment of COS markers to peaks on flow karyotypes

A set of conserved orthologous set (COS) markers specific for wheat homoeologous groups 1-7 were mapped to subgenomic DNA samples from individual peaks on the flow karyotypes to confirm the cytological assignment of chromosomes to the flow karyotype peaks. Of the 25 COS markers investigated, all 25 gave PCR products in wheat (genotype Mv9kr1), 24 in T. urartu MvGB115 and Ae. tauschii MvGB605 and 23 in Ae. speltoides MvGB905 (Online Resource 2). The 25 markers resulted in a total of 68 PCR products (range 1-7 PCR products/marker/ genotype, mean 2.72 PCR products) in wheat (genotype Mv9kr1), while 137 products were amplified in the diploid progenitors (49, 42 and 46 products in the T. urartu, Ae. speltoides and Ae. tauschii genotypes, respectively). Of the 137 PCR products detected in the diploid species, 49 (35.8 %) were non-polymorphic, while 88 (64.2 %) were polymorphic relative to hexaploid wheat (the proportion of non-polymorphic amplicons was 38.8, 33.3 and 34.8 % in T. urartu, Ae. speltoides and Ae. tauschii, respectively).

Because each chromosome of the wild progenitors has a major location in one of the peaks on the flow karyotype (Table 1), the yield of PCR products differed between the peaks and the highest amount of PCR products was observed in the peak where the locus-carrying chromosome had its major location (Fig. 5). For example, the marker $X_{GPL:C:731424}$, specific for group 4 chromosomes of wheat, produced a 260-bp PCR amplicon with continuously decreasing yield in the T. urartu flow karyotype peaks I, II and III (no amplicon in peak III) (Fig. 5, Online Resource 2), where the 4A chromosome contents were 38.61, 0.56 and 0 %, respectively (Table 1). With this marker a similar relationship could be observed in Ae. speltoides and Ae. tauschii between the yield of group 4 chromosome-specific PCR amplicons and the relative content of group 4 chromosomes determined by cytomolecular methods for the flow karyotype peaks.

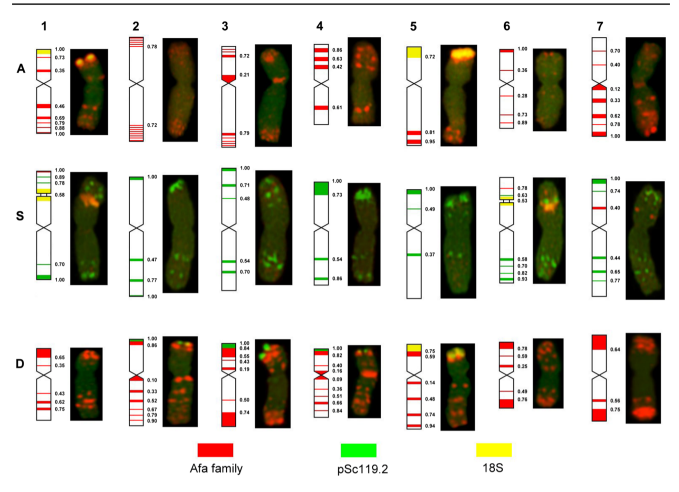


Fig. 3 Idiogram and karyotype of the chromosomes of *T. urartu* MvGB115 (a), *Ae. speltoides* MvGB905 (s) and *Ae. tauschii* MvGB605 (d) showing genomic distribution of repetitive DNA

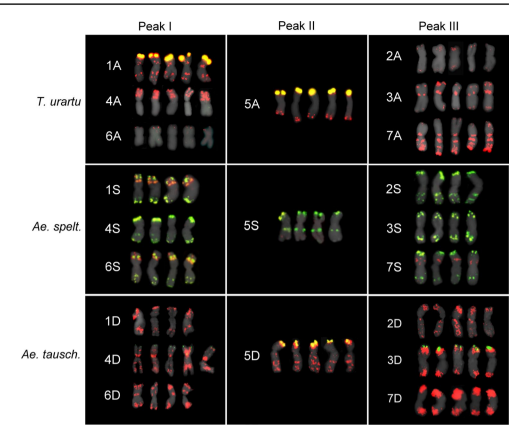
sequences pSc119.2 (green), Afa family (red) and 18S rDNA (yellow). Fraction lengths were determined as means of 15–20 measurements for each band

Based on the differences in the yield of PCR amplicons between the flow karyotype peaks, all the PCR amplicons could be assigned to peaks on flow karyotypes of the diploid species (Table 2, Online Resource 2). The COS markers specific for different ESTs and representing the group 1–7 chromosomes of hexaploid wheat showed similar results to the cytomolecular determination of the chromosome content of the flow karyotype peaks. In the first flow karyotype peak of T. urartu, Ae. speltoides and Ae. tauschii, the highest yield was detected for the PCR amplicons of markers specific for group 1, 4 and 6 chromosomes, while the markers specific for group 2, 3 and 7 chromosomes were assigned to peak III in the diploid progenitors. The markers representing the group 5 chromosomes were assigned unambiguously to peak II of T. urartu, Ae. speltoides and Ae. tauschii and with four exceptions (group 3-specific X_{BM134465} in Ae. tauschii, group 4-specific X_{4S} and group 6-specific X_{6A} in Ae. speltoides and group 7-specific X_{GPI:C:767323} in T. urartu) no markers specific for other homoeologous groups were assigned to peak II in any of the three species. These results indicate that flow karyotype peak II represents only the group 5 chromosomes of the diploid progenitors, consistently with the cytological results.

Discussion

Flow cytometric chromosome analysis and sorting has been developed in species with high socio-economic importance, such as bread wheat, durum wheat, barley and rye (Vrána et al. 2000; Kubaláková et al. 2003, 2005; Lysák et al. 1999). Chromosome genomics, which has been described as the application of genomic tools to flow-sorted chromosomes (Doležel et al. 2007), facilitates the molecular analysis of chromosome structure, the high-throughput development of markers, the construction of ready-to-sequence physical maps and positional gene cloning. The present work extends the potential of chromosome genomics to the diploid progenitors of bread wheat: *T. urartu, Ae. speltoides* and *Ae. tauschii*.

Fig. 4 Mitotic metaphase chromosomes sorted from individual peaks of flow karyotypes of T. urartu MvGB115, Ae. speltoides MvGB905 and Ae. tauschii MvGB605. Sorted chromosomes were identified after FISH with probes for repetitive DNA sequences pSc119.2 (green), Afa family (red) and 18S rDNA (yellow). Chromosomes were assigned to the peaks where their frequencies were the highest. Four or five representative examples are given for each chromosome



The fact that the chromosome peaks on the flow karyotype of T. urartu were shifted towards higher fluorescence channels as compared to Ae. speltoides and Ae. tauschii suggests that the average size of the A^u genome chromosomes, and hence the size of the A^u genome is larger than that of the S and D genomes. These results agree with the 1C values published for T. urartu (5.784 and 5.88 pg DNA), Ae. speltoides (5.15 pg DNA) and Ae. tauschii (5.08 and 5.1 pg DNA) (Özkan et al. 2003, 2010; Furuta et al. 1986; Rees and Walters 1965) and indicate that their genomes underwent different evolutionary changes. As the number of genes per monoploid genome is similar in plant species, differences in genome size are mainly due to repetitive DNA sequences, of which retroelements are considered to play a dominant role in genome expansion (Bennetzen 2007; Feuillet and Keller 2002; Lisch 2009). Charles et al. (2008) showed that specific types of transposable elements underwent differential proliferation in various wheat genomes during their evolution. The activation of Copia and CACTA transposable elements has occurred in the wheat A genome, while the proliferation of Gypsy elements was observed in the B genome. The differential proliferation of these retroelements in the A and B genomes of wheat occurred prior to allopolyploidization events (Charles et al. 2008) and hence in the diploid wheat progenitors.

Differences in the molecular organization of the A, S and D genomes of *T. urartu*, *Ae. speltoides* and *Ae. tauschii*

relative to tetraploid and hexaploid wheat have been demonstrated at the chromosome level using FISH. The FISH analysis performed in the present work involved probes for pSc119.2, Afa family and 18S rDNA repeats, whose genomic distribution was previously described in tetraploid and hexaploid wheat as well as in their diploid progenitors (Badaeva et al. 1996a, b; Kubaláková et al. 2005; Molnár et al. 2009). The most pronounced differences in the chromosomal structure concern the A genome of T. urartu relative to bread wheat, including the intensively labelled NOR regions of chromosomes 1 and 5, which were eliminated in durum and bread wheat after polyploidization (Jiang and Gill 1994). The inactivation of major NORs was also observed after the formation of the allopolyploid Aegilops species (Badaeva et al. 2004; Molnár et al. 2011a; Feldman et al. 2012). The difference in the hybridization pattern of chromosome 4A between T. urartu and bread wheat is due to the pericentric inversion of 4A and translocations involving chromosomes 5A and 7B, which occurred at the polyploid level (Hernandez et al. 2012). Smaller differences in the FISH karyotype were detected between Ae. speltoides and the B genome of wheat, while almost the same karyotype was observed in Ae. tauschii relative to the D genome chromosomes of wheat.

Genomic asymmetry, i.e. the alteration of wheat subgenomes relative to the diploid progenitors, may be related to the time that passed between the two polyploidization

 Table 1
 Assignment of chromosomes to peaks of flow karyotypes for *T. urartu* MvGB115, *Ae. speltoides* MvGB905 and *Ae. tauschii.*

 MvGB605. The numbers represent the percentage of the chromosome type in the whole peak content

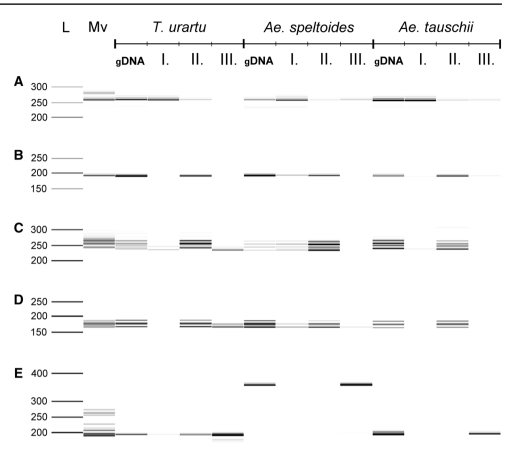
Species Genome		Chromosome	Peak I	Peak II	Peak III	
T. urartu spel- toides	А	1	25.64	1.41	_	
		2	_	0.28	32.69	
		3	_	2.25	34.39	
		4	38.61	0.56	-	
		5	0.28	78.53	1.69	
		6	35.44	0.84	_	
		7	_	16.1	31.21	
No. of chromosomes analysed			347	354	471	
Ae. speltoides	S	1	29.69	5.19	0.19	
		2	_	-	36.59	
		3	_	-	36.39	
		4	31.51	2.28	-	
		5	11.31	89.8	0.19	
		6	27.47	2.7	0.38	
		7	_	-	26.24	
No. of chromosomes analysed			495	481	522	
Ae. tauschii	D	1	34.17	0.82	_	
		2	_	0.13	31.02	
		3	_	11.0	29.67	
		4	36.46	0.68	_	
		5	_	87.3	0.84	
		6	29.35	_	_	
		7	-	_	38.44	
No. of chromos analysed	436	727	593			

events of hexaploid wheat (the formation of T. dicoccoides occurred approximately 0.3-0.5 million years ago, while hexaploid wheat evolved around 9,000 years ago) and to the process of genetic diploidization (Dvorak et al. 1998). Genetic diploidization causes non-random elimination of coding and non-coding sequences in parental genomes (Özkan et al. 2001; Feldman and Levy 2005) and could explain the observation of strong Afa FISH signals in the A genome of T. urartu and the lower number and size of Afa loci in hexaploid wheat. Diploidization affects both the expression of rRNA encoding genes, storage protein genes and other genes related to plant morphology, and the adaptation to environmental and biological factors, leading to the unequivocal partition of tasks between the constituent genomes of durum and bread wheat (Feldman et al. 2012). While the A genome of hexaploid wheat preferentially controls morphological traits, the B and D genomes control the reactions to biotic and abiotic factors (Feldman et al. 2012).

The finding that the 5A^u, 5S and 5D chromosomes can be isolated from the diploid progenitors of bread wheat may facilitate the dissection of many important traits. The group 5 chromosomes harbour loci ensuring genetically stable inheritance and determining the morphological traits responsible for widespread cultivation and adaptation to different agroecological conditions and abiotic stresses (Cattivelli et al. 2002). Among them, the *Ph1* locus, the dominant factor in the genetic system ensuring diploid-like meiotic chromosome pairing in tetra- and hexaploid wheat, is located on the long arm of chromosome 5B (Riley and Chapman 1958). The O gene, a predominant domestication gene in bread and durum wheat, conferring the free-threshing (naked grain) phenotype and pleiotropically affecting spike morphology, rachis fragility, glume tenacity, plant height and heading characters (Leighty and Boshnakian 1921; Mackey 1954; Muramatsu 1986; Kato et al. 1999; Faris et al. 2003; Simons et al. 2006) has been located on chromosome 5A. The VRN1 gene, which is critical for the adaptation of polyploid wheats to autumn sowing and divides wheat varieties into the winter and spring market classes, has been mapped to the colinear regions of the long arm of chromosomes 5A, 5B and 5D (Galiba et al. 1995; Snape et al. 1997). The group 5 chromosomes of hexaploid wheat also carry major loci affecting winter hardiness and freezing tolerance, designated as FROST RESISTANCE-1 (FR-1; Sutka and Snape 1989) and FR-2, which is approximately 30 centimorgans proximal to VRN-1 and includes a cluster of 11 (or more) C-REPEAT BINDING FACTOR (CBF) genes (Vágújfalvi et al. 2003; Dhillon et al. 2010). Other loci affecting drought and salt tolerance and the crossability of wheat have also been mapped to group 5 chromosomes (Quarrie et al. 1994; Koebner et al. 1996; Riley and Chapman 1967; Sitch et al. 1985; Krolow 1970).

The identification of novel alleles of important genes in the wild diploid wheat species could facilitate their use for coping with the pleiotropic effects of climate change (Feuillet et al. 2008). The production of synthetic wheat by crossing tetraploid durum wheat and Ae. tauschii is an important pathway for the utilization of wild genetic diversity in breeding programmes. As reported by Dreisigacker et al. (2008), 1,100 synthetic wheat lines were produced by CIMMYT from a collection of ~900 Ae. tauschii accessions and showed significant variation for morphological and agronomic traits and for tolerance to biotic and abiotic stresses. As the draft genome sequences of T. urartu and Ae. tauschii became available (Ling et al. 2013; Jia et al. 2013), the isolation of group 5 chromosomes from various diploid wheat species will provide a cost-effective means for the targeted re-sequencing of the chromosomes using next generation technology (Vitulo et al. 2011). This will support comprehensive gene content analysis and allele discovery, and facilitate the development of gene-based markers for

Fig. 5 PCR products of conserved orthologous set (COS) markers amplified from the wheat genotype Mv9kr1 (Mv), from total genomic DNA (gDNA) and from subgenomic DNA samples derived from chromosomes of particular peaks (I, II, III) on flow karyotypes of T. urartu MvGB115, Ae. speltoides MvGB905 and Ae. tauschii. MvGB605. The pictures represent markers amplified from flow karyotype peak I $(X_{GPL:C:731424})$ (a), peak II (X_{GPI:C:726959}, X_{5A}, X_{GPI:C:729592}) (**b**–**d**) and peak III (X_{tr150}) and those specific for homoeologous group 4 ($X_{GPI:C:731424}$), group 5 (X_{GPI:C:726959}, X_{5A} and X_{GPI:C:729592}) and group 2 (X_{tr150})



specific genomic regions (Wenzl et al. 2010; Berkman et al. 2011; Mayer et al. 2009, 2011; Wicker et al. 2011). Moreover, the availability of chromosome sequences will permit the validation of genome assemblies at the chromosome level.

Our findings confirm and expand the results of previous studies indicating the high transferability of COS markers between species (Parida et al. 2006; Burt and Nicholson 2011; Howard et al. 2011; Molnár et al. 2013). The fact that 64.2 % of the products obtained with chromosomes isolated from wild relatives were polymorphic relative to those obtained in hexaploid wheat suggested that the substantial genetic diversity of wild progenitors is due to the variability of intron regions (Yu et al. 2005). Thus, COS markers can be used to identify the chromatin of wild progenitors in the wheat background and for marker-assisted selection to facilitate the transfer of useful agronomic traits (Quraishi et al. 2009). The 24 COS markers assigned to chromosomes of the diploid progenitors of wheat can be used in pre-breeding programmes to facilitate gene introgression. In a previous study, 100 COS markers were assigned to the U and M genome chromosomes of diploid and tetraploid Aegilops species (Molnár et al. 2013) and used for the analysis of macrosynteny between Aegilops, wheat and the model species Brachypodium and rice. These results indicate that the markers used in the present study and the COS markers that are assigned to the chromosomes of the diploid progenitors of wheat in the future will be suitable tools to study the genome rearrangements and synteny perturbations that occurred during the polyploidisation and domestication of bread wheat.

The FISH analysis of flow-sorted chromosome fractions showed that chromosomes 5A^u, 5S and 5D could be isolated at high purity (78.5, 89.8 and 89.3 %, respectively) from T. urartu, Ae. speltoides and Ae. tauschii. Further improvements in the protocol might lead to the increased purity of these chromosome fractions. However, as demonstrated by Mayer et al. (2011), the level of chromosome contamination observed in the present study does not compromise the bioinformatic analysis of the sequence data obtained by sequencing DNA amplified from single chromosome fractions. Other chromosomes from the diploid wheat species formed composite peaks on the flow karyotypes and could be sorted into groups of three. This limitation could be overcome in the future using the multiparametric flow cytometric analysis of chromosomes labelled by FISH with fluorescent probes for microsatellites, as shown by Giorgi et al. (2013). The FISH karyotypes of diploid wheat species developed in this study and by Megyeri et al. (2012) might serve as a guide when choosing probes for labelling chromosomes prior to flow-sorting.

Table 2Assignment of COSmarkers specific for wheathomoeologous groups 1–7 topeaks I–III on flow karyotypesof T. urartu MvGB115, Ae.speltoides MvGB905 and Ae.tauschii MvGB605

Marker	Wheat homoeologous group	T. urartu		Ae. speltoides			Ae. tauschii			
		I	II	III	I	II	III	I	II	III
X _{BE443103}	1	X			Х			X		
X _{GPI:C:732519}	1	Х			Х			Х		
X _{GPI:C:746781}	1	Х			Х			Х		
X _{BE445693}	2			Х			Х			Х
X _{GPI:C:719382}	2			Х			Х			
<i>X</i> _{tr150}	2			Х			Х			Х
X _{2N}	2			Х			Х			Х
X _{3B}	3			Х			Х			Х
X _{BM134465}	3			Х			Х		Х	
X _{BE404709}	3			Х						Х
X _{GPI:C:748004}	4	Х			Х			Х		
X _{GPI:C:731424}	4	Х			Х			Х		
X _{GPI:C:725135}	4	Х			Х			Х		
X _{4S}	4	Х				Х		Х		
X _{GPI:C:748166}	5								Х	
X _{GPI:C:729592}	5		Х			Х			Х	
X _{5A}	5		Х			Х			Х	
X _{GPI:C:726959}	5		Х			Х			Х	
X _{GPI:C:740549}	6	Х			Х			Х		
X _{6A}	6	Х				Х		Х		
X _{6N}	6	Х			Х			Х		
X _{BE352570}	7			Х			Х			Х
X _{GPI:C:771171}	7			Х			Х			Х
X _{GPI:C:770073}	7			Х			Х			Х
X _{GPI:C:767323}	7		Х				X			Х

To conclude, this study represents an important step forward in developing chromosome genomics for the wild genetic resources of wheat. The flow karyotypes of T. urartu, Ae. speltoides and Ae. tauschii were characterized and the chromosome content of all the peaks on the karyotypes was determined for the first time. The ability to purify group 5 chromosomes in the wild diploid progenitors of wheat paves the way for the rapid re-sequencing of single chromosomes isolated from diverse populations, the physical mapping of DNA sequences to particular chromosomes using PCR, the construction of chromosome-specific BAC libraries and next generation sequencing to identify low-copy and genic sequences and to develop new markers. The COS markers assigned to the chromosomes of diploid wheats could be used in pre-breeding programmes to select chromosome segments carrying agronomically useful genes in T. aestivum—wild wheat recombinant lines.

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Conflict of interest The authors declare that they have no conflict of interest.

Ethical standards The authors declare that the experiments comply with the current laws of the countries (Czech Republic and Hungary) in which they were performed.

References

- Akhunov ED, Akhunova AR, Dvorak J (2005) BAC libraries of *Triticum urartu*, Aegilops speltoides and Ae. tauschii, the diploid ancestors of polyploidy wheat. Theor Appl Genet 111:1617–1622. doi:10.1007/s00122-005-0093-1
- Badaeva ED, Friebe B, Gill BS (1996a) Genome differentiation in *Aegilops*. 1. Distribution of highly repetitive DNA sequences on chromosomes of diploid species. Genome 39:293–306. doi:10.1139/g96-040

- Badaeva ED, Friebe B, Gill BS (1996b) Genome differentiation in *Aegilops*. 2. Physical mapping of 5S and 18S-26S ribosomal RNA gene families in diploid species. Genome 39:1150–1158. doi:10.1139/g96-145
- Badaeva ED, Amosova AV, Samatadze TE, Zoshchuk SA, Shostak NG et al (2004) Genome differentiation in *Aegilops*. 4. Evolution of the U-genome cluster. Plant Syst Evol 246:45–76. doi:10.1007/s00606-003-0072-4
- Bennetzen JL (2007) Patterns in grass genome evolution. Curr Opin Plant Biol 10:176–181. doi:10.1016/j.pbi.2007.01.010
- Berkman PJ, Skarshewski A, Lorenc MT, Lai K, Duran C et al (2011) Sequencing and assembly of low copy and genic regions of isolated *Triticum aestivum* chromosome arm 7DS. Plant Biotechnol J 9:768–775. doi:10.1111/j.1467-7652.2010.00587.x
- Boyko EV, Gill KS, Mickelson-Young L, Nasuda S, Raupp WJ et al (1999) A high-density genetic linkage map of *Aegilops tauschii*, the D genome progenitor of bread wheat. Theor Appl Genet 99:16–26. doi:10.1007/s001220051204
- Brenchley R, Spannag M, Pfeifer M, Barker GLA, D'Amore R et al (2012) Analysis of the bread wheat genome using wholegenome shotgun sequencing. Nature 491:705–710. doi:10.1038/ nature11650
- Burt C, Nicholson P (2011) Exploiting co-linearity among grass species to map the *Aegilops ventricosa*-derived *Pch1* eyespot resistance in wheat and establish its relationship to *Pch2*. Theor Appl Genet 123:1387–1400. doi:10.1007/s00122-011-1674-9
- Cattivelli L, Baldi P, Crosatti C, Di Fonzo N, Faccioli P et al (2002) Chromosome regions and stress-related sequences involved in resistance to abiotic stress in *Triticeae*. Plant Mol Biol 48:649– 665. doi:10.1023/A:1014824404623
- Chang KD, Fang SA, Chang FC, Chung MC (2010) Chromosomal conservation and sequence diversity of ribosomal RNA genes of two distant *Oryza* species. Genomics 96:181–190. doi:10.1016/j.ygeno.2010.05.005
- Charles M, Belcram H, Just J, Huneau C, Viollet A et al (2008) Dynamics and differential proliferation of transposable elements during the evolution of the B and A genomes of wheat. Genetics 180:1071–1086. doi:10.1534/genetics.108.092304
- Contento A, Heslop-Harrison JS, Schwarzacher T (2005) Diversity of a major repetitive DNA sequence in diploid and polyploid *Triticeae*. Cytogenet Genome Res 109:34–42. doi:10.1159/000082379
- Cox TS (1998) Deepening the wheat gene pool. J Crop Prod 1:1–25. doi:10.1300/J144v01n01_01
- Cox TS, Hatchett JH, Gill BS, Raupp WJ, Sears RG (1990) Agronomic performance of hexaploid wheat lines derived from direct crosses between wheat and *Aegilops squarrosa*. Plant Breed 105:271–277. doi:10.1111/j.1439-0523.1990.tb01285.x
- Cseh A, Soós V, Rakszegi M, Türkösi E, Balázs E, Molnár-Láng M (2013) Expression of *HvCslF9* and *HvCslF6* barley genes in the genetic background of wheat and their influence on the wheat β -glucan content. Ann Appl Biol 163:142–150. doi:10.1111/aab.12043
- Devos KM, Gale MD (2000) Genome relationships: the grass model in current research. Plant Cell 12:637–646. doi:10.1105/ tpc.12.5.637
- Dhillon T, Pearce SP, Stockinger EJ, Distelfeld A et al (2010) Regulation of freezing tolerance and flowering in temperate cereals: the VRN-1 connection. Plant Physiol 153:1846–1858. doi:10.1104/pp.110.159079
- Doležel J, Binarová P, Lucretti S (1989) Analysis of nuclear DNA content in plant cells by flow cytometry. Biol Plant 31:113–120. doi:10.1007/BF02907241
- Doležel J, Kubaláková M, Paux E, Bartoš J, Feuillet C (2007) Chromosome based genomics in the cereals. Chromosome Res 15:51–66. doi:10.1007/s10577-006-1106-x

- Doležel J, Vrána J, Safář J, Bartoš J, Kubaláková M et al (2012) Chromosomes in the flow to simplify genome analysis. Funct Integr Genomics 12:397–416. doi:10.1007/s10142-012-0293-0
- Doležel J, Vrána J, Cápal P, Kubaláková M, Burešová V, Šimková H (2014) Advances in plant chromosome genomics. Biotechnol Adv 32:122–136. doi:10.1016/j.biotechadv.2013.12.011
- Dreisigacker S, Kishii M, Lage J, Warburton M (2008) Use of synthetic hexaploid wheat to increase diversity for CIMMYT bread wheat improvement. Aust J Agric Res 59:413–420. doi:10.1071/AR07225
- Dvorak J, Di Terlizzi P, Zhang H-B, Resta P (1993) The evolution of polyploid wheat: identification of the A genome donor species. Genome 36:21–31. doi:10.1139/g93-004
- Dvorak J, Luo MC, Yang ZL (1998) Restriction fragment length polymorphism and divergence in the genomic regions of high and low recombination in self-fertilizing and cross-fertilizing *Aegilops* species. Genetics 148:423–434
- Edwards D, Batley J (2010) Plant genome sequencing: applications for crop improvement. Plant Biotechnol J 8:2–9. doi:10.1111/j.1467-7652.2009.00459.x
- Faris JD, Fellers JP, Brooks SA, Gill BS (2003) A bacterial artificial chromosome contig spanning the major domestication locus Q in wheat and identification of a candidate gene. Genetics 164:311–321
- Feldman M, Levy AA (2005) Allopolyploidy: a shaping force in the evolution of wheat genomes. Cytogenet Genome Res 109:250– 258. doi:10.1159/000082407
- Feldman M, Levy AA, Fahima T, Korol A (2012) Genomic asymmetry in allopolyploid plants: wheat as a model. J Exp Bot 63:5045–5059. doi:10.1093/jxb/ers192
- Feuillet C, Keller B (2002) Comparative genomics in the grass family: molecular characterization of grass genome structure and evolution. Ann Bot 89:3–10. doi:10.1093/aob/mcf008
- Feuillet C, Langridge P, Waugh R (2008) Cereal breeding takes a walk on the wild side. Trends Genet 24:24–32. doi:10.1016/j.tig.2007.11.001
- Friebe B, Jiang J, Raupp WJ, McIntosh RA, Gill BS (1996) Characterization of wheat-alien translocations conferring resistance to diseases and pests: current status. Euphytica 91:59–87. doi:10.1 007/BF00035277
- Fritz AK, Cox TS, Gill BS, Sears RG (1995) Marker-based analysis of quantitative traits in winter wheat by *Triticum tauschii* populations. Crop Sci 35:1695–1699. doi:10.2135/cropsci1995.0011 183X003500060031x
- Furuta Y, Nishikawa K, Yamaguchi S (1986) Nuclear DNA content in diploid wheat and its relatives in relation to the phylogeny of tetraploid wheat. Jpn J Genet 61:97–105. doi:10.1266/jjg.61.97
- Galiba G, Quarrie SA, Sutka J, Morgounov A, Snape JW (1995) RFLP mapping of the vernalization (Vrn1) and frost resistance (Fr1) genes on chromosome 5A of wheat. Theor Appl Genet 90:1174–1179. doi:10.1007/BF00222940
- Gill KS, Lubbers EL, Gill BS, Raupp WJ, Cox TS (1991) A genetic linkage map of *Triticum tauschii* (DD) and its relationship to the D genome of bread wheat (AABBDD). Genome 34:362– 374. doi:10.1139/g91-058
- Giorgi D, Farina A, Grosso V, Gennaro A, Ceoloni C et al (2013) FISHIS: fluorescence in situ hybridization in suspension and chromosome flow sorting made easy. PLoS One 8:e57994. doi:10.1371/journal.pone.0057994
- Grosso V, Farina A, Gennaro A, Giorgi D, Lucretti S (2012) Flow sorting and molecular cytogenetic identification of individual chromosomes of *Dasypyrum villosum* L. (*H. villosa*) by a single DNA probe. PLoS One 7:e50151. doi:10.1371/ journal.pone.0050151
- Guzmán C, Alvarez JB (2012) Molecular characterization of a novel waxy allele (Wx-A^ula) from Triticum urartu Thum. ex

- Hernandez P, Martis M, Dorado G, Pfeifer M, Gálvez S et al (2012) Next-generation sequencing and syntenic integration of flowsorted arms of wheat chromosome 4A exposes the chromosome structure and gene content. Plant J 69:377–386. doi:10.1111/j.1365-313X.2011.04808.x
- Howard T, Rejab NA, Griffiths S, Leigh F, Leverington-Waite M et al (2011) Identification of a major QTL controlling the content of B-type starch granules in *Aegilops*. J Exp Bot 62:2217–2228. doi:10.1093/jxb/erq423
- Jia J, Zhao S, Kong X, Li Y, Zhao G et al (2013) *Aegilops tauschii* draft genome sequence reveals a gene repertoire for wheat adaptation. Nature 496:91–95. doi:10.1038/nature12028
- Jiang J, Gill BS (1994) New 18S-26S ribosomal RNA gene loci: chromosomal landmarks for the evolution of polyploid wheats. Chromosoma 103:179–185. doi:10.1007/BF00368010
- Kato K, Miura H, Sawada S (1999) QTL mapping of genes controlling ear emergence time and plant height on chromosome 5A of wheat. Theor Appl Genet 98:472–477. doi:10.1007/ s001220051094
- Kilian B, Mammen K, Millet E, Sharma R, Graner A et al (2011) Aegilops. In: Kole C (ed) Wild crop relatives: genomic and breeding resources, cereals. Springer, Berlin, pp 1–76. doi:10.1007/978-3-642-14228-4_1
- Klindworth DL, Hareland GA, Elias EM, Xu SS (2013) Attempted compensation for linkage drag affecting agronomic characteristics of durum wheat 1AS/1DL translocation lines. Crop Sci 53:422–429. doi:10.2135/cropsci2012.05.0310
- Koebner RMD, Martin PK, Orford SM, Miller TE (1996) Responses to salt stress controlled by the homeologous group 5 chromosomes of hexaploid wheat. Plant Breed 115:81–84. doi:10.1111/j.1439-0523.1996.tb00878.x
- Kofler R, Bartoš J, Gong L, Stift G, Suchánková P et al (2008) Development of microsatellite markers specific for the short arm of rye (*Secale cereale* L.) chromosome 1. Theor Appl Genet 117:915–926. doi:10.1007/s00122-008-0831-2
- Krolow KD (1970) Untersuchungen über die Kreuzbarkeit zwischen Weizen und Roggen. Z Pflanzenzücht 64:44–72
- Kubaláková M, Macas J, Doležel J (1997) Mapping of repeated DNA sequences in plant chromosomes by PRINS and C-PRINS. Theor Appl Genet 94:758–763. doi:10.1007/s001220050475
- Kubaláková M, Valárik M, Bartoš J, Vrána J, Číhalíková J et al (2003) Analysis and sorting of rye (*Secale cereale* L.) chromosomes using flow cytometry. Genome 46:893–905. doi:10.1139/ g03-054
- Kubaláková M, Kovářová P, Suchánková P, Číhalíková J, Bartoš J et al (2005) Chromosome sorting in tetraploid wheat and its potential for genome analysis. Genetics 170:823–829. doi:10.1534/genetics.104.039180
- Kumar A, Simons K, Iqbal MJ, de Jiménez M, Bassi FM et al (2012) Physical mapping resources for large plant genomes: radiation hybrids for wheat D-genome progenitor *Aegilops tauschii*. BMC Genomics 13:597. doi:10.1186/1471-2164-13-597
- Leighty CE, Boshnakian S (1921) Genetic behaviour of the spelt form in crosses between *Triticum spelta* and *Triticum aestivum*. J Agric Res 7:335–364
- Limin AE, Fowler DB (1993) Inheritance of cold hardiness in *Triticum aestivum* x synthetic hexaploid wheat crosses. Plant Breed 110:103–108. doi:10.1111/j.1439-0523.1993.tb01220.x
- Ling H-Q, Zhao S, Liu D, Wang J, Sun H et al (2013) Draft genome of the wheat A-genome progenitor *Triticum urartu*. Nature 496:87–90. doi:10.1038/nature11997
- Lisch D (2009) Epigenetic regulation of transposable elements in plants. Ann Rev Plant Biol 60:43–66. doi:10.1146/annurev.arp lant.59.032607.092744

- Lubbers EL, Gill KS, Cox TS, Gill BS (1991) Variation of molecular markers among geographically diverse accessions of *Triticum tauschii*. Genome 34:354–361. doi:10.1139/g91-057
- Lysák MA, Číhalíková J, Kubaláková M, Šimková H, Künzel G et al (1999) Flow karyotyping and sorting of mitotic chromosomes of barley (*Hordeum vulgare* L.). Chromosome Res 7:431–444. doi:10.1023/A:1009293628638
- MacKey J (1954) Neutron and X-ray experiments in wheat and a revision of the speltoid problem. Hereditas 40:65–180
- Maestra B, Naranjo T (1998) Homoeologous relationships of Aegilops speltoides chromosomes to bread wheat. Theor Appl Genet 97:181–186. doi:10.1007/s001220050883
- Margulies M, Egholm M, Altman WE, Attiya S, Bader JS et al (2005) Genome sequencing in microfabricated high-density picolitre reactors. Nature 437:376–380. doi:10.1038/nature03959
- Martis MM, Zhou R, Haseneyer G, Schmutzer T, Vrána J et al (2013) Reticulate evolution of the rye genome. Plant Cell 25:3685–3698
- Mayer KFX, Taudien S, Martis M, Šimková H, Suchánková P et al (2009) Gene content and virtual gene order of barley chromosome 1H. Plant Physiol 151:496–505. doi:10.1104/ pp.109.142612
- Mayer KFX, Martis M, Hedley PE, Simková H, Liu H et al (2011) Unlocking the barley genome by chromosomal and comparative genomics. Plant Cell 23:1249–1263. doi:10.1105/ tpc.110.082537
- Megyeri M, Farkas A, Varga M, Kovács G, Molnár-Láng M, Molnár I (2012) Karyotypic analysis of *Triticum monococcum* using standard repetitive DNA probes and simple sequence repeats. Acta Agron Hung 60:87–95. doi:10.1556/AAgr.60.2012.2.1
- Molnár I, Benavente E, Molnár-Láng M (2009) Detection of intergenomic chromosome rearrangements in irradiated *Triti*cum aestivum-Aegilops biuncialis amphiploids by multicolour genomic in situ hybridization. Genome 52:156–165. doi:10.1139/G08-114
- Molnár I, Cifuentes M, Schneider A, Benavente E, Molnár-Láng M (2011a) Association between simple sequence repeat-rich chromosome regions and intergenomic translocation breakpoints in natural populations of allopolyploid wild wheats. Ann Bot 107:65–76. doi:10.1093/aob/mcq215
- Molnár I, Kubaláková M, Šimková H, Cseh A, Molnár-Láng M, Doležel J (2011b) Chromosome isolation by flow sorting in Aegilops umbellulata and Ae. comosa and their allotetraploid hybrids Ae. biuncialis and Ae. geniculata. PLoS One 6:e27708. doi:10.1371/journal.pone.0027708
- Molnár I, Šimková H, Leverington-Waite M, Goram R, Cseh A et al (2013) Syntenic relationships between the U and M genomes of *Aegilops*, wheat and the model species *Brachypodium* and rice as revealed by COS markers. PLoS One 8:e70844. doi:10.1371/ journal.pone.0070844
- Muramatsu M (1986) The vulgare super gene, Q: its universality in durum wheat and its phenotypic effects in tetraploid and hexaploid wheats. Can J Genet Cytol 28:30–41. doi:10.1139/g86-006
- Nagaki K, Tsujimoto H, Isono K, Sasakuma T (1995) Molecular characterization of a tandem repeat, Afa family, and its distribution among *Triticeae*. Genome 38:479–486. doi:10.1139/g95-063
- Özkan H, Levy AA, Feldman M (2001) Allopolyploidy-induced rapid genome evolution in the wheat (*Aegilops-Triticum*) group. Plant Cell 13:1735–1747. doi:10.1105/TPC.010082
- Özkan H, Tuna M, Arumuganathan K (2003) Nonadditive changes in genome size during allopolyploidization in the wheat (*Aegilops-Triticum*) group. J Hered 94:260–264. doi:10.1093/jhered/ esg053
- Özkan H, Tuna M, Kilian B, Mori N, Ohta S (2010) Genome size variation in diploid and tetraploid wild wheats. AoB Plants 2010:plq015. doi:10.1093/aobpla/plq015

- 1104
- Parida SK, Kumar ARK, Dalal V, Singh NK, Mohapatra T (2006) Unigene derived microsatellite markers for the cereal genomes. Theor Appl Genet 112:808–817. doi:10.1007/ s00122-005-0182-1
- Požárková D, Koblížková A, Román B, Torres AM, Lucretti S et al (2002) Development and characterization of microsatellite markers from chromosome 1-specific DNA libraries of *Vicia faba*. Biol Plant 45:337–345. doi:10.1023/A:1016253214182
- Quarrie SA, Gulli M, Calestani C, Steed A, Marmiroli N (1994) Location of a gene regulation drought-induced abscisic acid production on the long arm of chromosome 5A of wheat. Theor Appl Genet 89:794–800. doi:10.1007/BF00223721
- Quraishi UM, Abrouk M, Bolot S, Pont C, Throude M et al (2009) Genomics in cereals: from genome-wide conserved orthologous set (COS) sequences to candidate genes for trait dissection. Funct Integr Genomics 9:473–484. doi:10.1007/ s10142-009-0129-8
- Rees H, Walters MR (1965) Nuclear DNA and the evolution of wheat. Heredity 20:73–82
- Riley R, Chapman V (1958) Genetic control of the cytologically diploid behaviour of hexaploid wheat. Nature 182:713–715. doi:10.1038/182713a0
- Riley R, Chapman V (1967) The inheritance in wheat of crossability with rye. Genet Res Camb 9:259–267. doi:10.1017/ S0016672300010569
- Román B, Satovic Z, Požárková D, Macas J, Doležel J et al (2004) Development of a composite map in *Vicia faba*, breeding applications and future prospects. Theor Appl Genet 108:1079–1088. doi:10.1007/s00122-003-1515-6
- Rouse MN, Jin Y (2011) Stem rust resistance in A-genome diploid relatives of wheat. Plant Dis 95:941–944. doi:10.1094/P DIS-04-10-0260
- Šafář J, Šimková H, Kubaláková M, Číhalíková J, Suchánková P et al (2010) Development of chromosome-specific BAC resources for genomics of bread wheat. Cytogenet Genome Res 129:211– 223. doi:10.1159/000313072
- Schachtman DP, Lagudah ES, Munns R (1992) The expression of salt tolerance from *Triticum tauschii* in hexaploid wheat. Theor Appl Genet 84:714–719. doi:10.1007/BF00224174
- Schatz M, Langmead B, Salzberg S (2010) Cloud computing and the DNA data race. Nat Biotechnol 28:691–693. doi:10.1038/ nbt0710-691
- Schneider A, Linc G, Molnár I, Molnár-Láng M (2005) Molecular cytogenetic characterization of *Aegilops biuncialis* and its use for the identification of five derived wheat-*Aegilops biuncialis* disomic addition lines. Genome 48:1070–1082. doi:10.1139/ g05-062
- Schneider A, Molnár I, Molnár-Láng M (2008) Utilisation of Aegilops (goatgrass) species to widen the genetic diversity of cultivated wheat. Euphytica 163:1–19. doi:10.1007/s10681-007-9624-y
- Sepsi A, Molnár I, Szalay D, Molnár-Láng M (2008) Characterization of a leaf rust-resistant wheat–*Thinopyrum ponticum* partial amphiploid BE-1, using sequential multicolor GISH and FISH. Theor Appl Genet 116:825–834. doi:10.1007/ s00122-008-0716-4
- Shangguan L, Han J, Kayesh E, Sun X, Zhang C et al (2013) Evaluation of genome sequencing quality in selected plant

species using expressed sequence tags. PLoS One 8:e69890. doi:10.1371/journal.pone.0069890

- Šimková H, Svensson JT, Condamine P, Hřibová E, Suchánková P et al (2008) Coupling amplified DNA from flow-sorted chromosomes to high-density SNP mapping in barley. BMC Genom 9:294. doi:10.1186/1471-2164-9-294
- Simons KJ, Fellers JP, Trick HN, Zhang Z, Tai Y-S et al (2006) Molecular characterization of the major wheat domestication gene Q. Genetics 172:547–555. doi:10.1534/genetics.105.044727
- Sitch LA, Snape JW, Firman SJ (1985) Intrachromosomal mapping of crossability genes in wheat (*Triticum aestivum*). Theor Appl Genet 70:309–314. doi:10.1007/BF00304917
- Snape JW, Semikhodskii A, Fish L, Sarma RN, Quarrie SA et al (1997) Mapping frost resistance loci in wheat and comparative mapping with other cereals. Acta Agron Hung 45:265–270
- Sutka J, Snape JW (1989) Location of a gene for frost resistance on chromosome 5A of wheat. Euphytica 42:41–44. doi:10.1007 /BF00042613
- Sutka J, Galiba G, Vágújfalvi A, Gill BS, Snape JW (1999) Physical mapping of the Vrn-A1 and Fr1 genes on chromosome 5A of wheat using deletion lines. Theor Appl Genet 99:199–202. doi:10.1007/s001220051225
- Vágújfalvi A, Galiba G, Cattivelli L, Dubcovsky J (2003) The cold regulated transcriptional activator *Cbf3* is linked to the frosttolerance gene *Fr-A2* on wheat chromosome 5A. Mol Genet Genomics 269:60–67. doi:10.1007/s00438-003-0806-6
- Vitulo N, Albiero A, Forcato C, Campagna D, Dal Pero F et al (2011) First survey of the wheat chromosome 5A composition through a next generation sequencing approach. PLoS One 6:e26421. doi:10.1371/journal.pone.0026421
- Vrána J, Kubaláková M, Šimková H, Číhalíková J, Lysák MA et al (2000) Flow-sorting of mitotic chromosomes in common wheat (*Triticum aestivum* L.). Genetics 156:2033–2041
- Wenzl P, Suchánková P, Carling J, Šimková H, Huttner E et al (2010) Isolated chromosomes as a new and efficient source of DArT markers for the saturation of genetic maps. Theor Appl Genet 121:465–474. doi:10.1007/s00122-010-1323-8
- Wicker T, Mayer KFX, Gundlach H, Martis M, Steuernagel B et al (2011) Frequent gene movement and pseudogene evolution is common to the large and complex genomes of wheat, barley, and their relatives. Plant Cell 23:1706–1718. doi:10.1105/ tpc.111.086629
- Xiu-Jin L, Deng-Cai L, Zhi-Rong W (1997) Inheritance in synthetic hexaploid wheat 'RSP' of sprouting tolerance derived from *Aegilops tauschii* Coss. Euphytica 95:321–323. doi:10.102 3/A:1003078801358
- You FM, Huo N, Deal KR, Gu YQ, Luo M-C et al (2011) Annotationbased genome-wide SNP discovery in the large and complex *Aegilops tauschii* genome using next-generation sequencing without a reference genome sequence. BMC Genomics 12:59. doi:10.1186/1471-2164-12-59
- Yu J, Wang J, Lin W, Li S, Li H et al (2005) The genomes of Oryza sativa: a history of duplications. PLoS Biol 3:e38. doi:10.1371/ journal.pbio.0030038