Similarities and differences in the nuclear genome organization within Pooideae species revealed by comparative genomic in situ hybridization (GISH)

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Abstract In this paper, we highlight the affinity between the genomes of key representatives of the Pooideae subfamily, revealed at the chromosomal level by genomic in situ hybridization (GISH). The analyses were conducted using labeled probes from each species to hybridize with chromosomes of every species used in this study based on a round robin rule. As a result, the whole chromosomes or chromosome regions were distinguished or variable types of signals were visualized to prove the different levels of the relationships between genomes used in this study. We observed the unexpected lack of signals in secondary constrictions of rye (RR) chromosomes probed by triticale (AABBRR) genomic DNA. We have also identified unlabeled chromosome regions, which point to species-specific sequences connected with disparate pathways of chromosome differentiation. Our results revealed a conservative character of coding sequence of 35S rDNA among selected species of the genera Aegilops, Brachypodium, Festuca, Hordeum, Lolium, Secale, and Triticum. In summary, we showed strong relationships in genomic DNA sequences between species which have been previously reported to be phylogenetically distant.

Keywords Pooideae · Brachypodium · Cereals · Forage grasses · Comparative genomics · In situ hybridization

Introduction

The Poaceae family (Barnh., Gramineae Juss.) is the fourth largest group of plants on Earth and can be found in nearly all regions and climate zones. More than 10,000 species and 600 genera constitute this family, but only three subfamilies, namely Ehrhartioideae, Pooideae, and Panicoideae, play the principal role of providing the human race with nutrition and a source of renewable energy. The Pooideae subfamily includes some economically important cereals, such as bread wheat (Triticum aestivum L.), barley (Hordeum vulgare L.), rye (Secale cereale L.), and triticale (×Triticosecale Wittm.), as well as many lawn and pasture grasses, such as perennial ryegrass (Lolium perenne L.). The genome composition of the species within Pooideae differs in respect to their size, ploidy level, basic chromosome number, and repetitive sequences content. For example, bread wheat (Triticum aestivum L.; 17 Gb; 2n = 6x = 42 chromosomes) is an allopolyploid, composed of six sets of chromosomes belonging to A, B, and D genomes with high content of non-coding, repetitive DNA. Contrasting, Brachypodium distachyon has a small genome (272 Mb; 2n = 2x = 10 chromosomes) and low amount of repetitive sequences (Vogel and Hill 2008; International Brachypodium Initiative (IBI) 2010), making it a good model for structural genomic studies of grasses. It was previously reported that B. distachyon is a comparative and functional genetic model and its potential relevance for research on cereals, on the basis of its physiological and genetic advantages (Draper et al. 2001; Vogel and Bragg 2009). For example, the high degree of genetic synteny between wheat, B. sylvaticum, and B. distachyon enabled the identification of the genes present in the Ph1 (Pairing homeologous 1) locus of wheat (Griffiths et al. 2006; Huo et al. 2009), demonstrating the potential application of the genus Brachypodium in comparative genomics. The phylogenetic relationships between...
B. distachyon and other cereals and grasses have been derived on the basis of the analysis of the internal transcribed spacer (ITS) and rDNA coding sequences (Hsiao et al. 1994), genomic markers, such as random amplified polymorphic DNA (RAPD), restriction fragment length polymorphism (RFLP) (Catalán et al. 1995), chloroplast restriction sites, and ndhF gene sequence (Kellogg 2001; Mochida and Shinozaki 2013). Those approaches showed that B. distachyon can be considered as a kind of missing link between rice (Oryza sativa L.) and temperate grasses.

Exploring the genomes of a wide range of plant species revealed the existence of highly conserved sequences and synteny in gene order. Comparative genomics can distinguish synteny, which means the presence of two or more loci on the same chromosome, and collinearity, which refers to the similarities in the physical order of loci (McCouch 2001). However, recently, the synteny is defined as the conservation of blocks of order within the sets of chromosomes which are being compared to each other. The first study of chromosome evolution and comparative genomics was performed between potato (Solanum tuberosum) and tomato (Lycopersicon esculentum) (Bonierbale et al. 1988). Initially, low copy RFLP markers and low resolution genetic maps were used to determine the phylogenetic relationships between Poaceae species, and established that most of the grass genomes are composed from 30 rice-independent linkage blocks (Gale and Devos 1998; Keller and Feuillet 2000). These methods enabled the macrocollinearity of grass genomes to be determined; however, they also overestimated the level of synteny. Recently, the development of additional genomic resources such as EST marker databases and whole genome sequences, combined with improved software, has enabled comparative genomic studies to reach a higher level of resolution, uncovering the microcollinearity of sequences, which form the basis to establish a model of grass genome evolution (Devos 2005; Wei et al. 2007; Salse et al. 2008; Murat et al. 2014).

One of the main factors leading to chromosome evolution and speciation are changes in the amount and distribution of repetitive DNA sequences (Flavell et al. 1979; Cuadrado and Jouve 2002). It is hypothesized that these sequences play an important role in chromosome organization, stabilization of chromosome structure, recognition and segregation of chromosomes in mitosis and meiosis, and regulation of gene activity (Vershchinin et al. 1995). Repetitive sequences account for up to 90 % of plant genomes (Heslop-Harrison 2000), and show diverse compositions in different genomes. Changes in repetitive sequences could be particularly useful in studies of genomic evolution. Some families of those sequences, e.g., Afa family (Nagaki et al. 1995), are present in more than one genus, which could be an evidence of their genomic relationship. Genomic in situ hybridization (GISH) can distinguish particular genomes at the chromosomal level. GISH can result in chromosome painting and show similarities of repetitive DNA distribution between related species. Moreover, the physical locations of conserved sequences can be visualized on chromosomes as well.

In this work, we compare both the model species (Brachypodium distachyon) as well as other important grasses, using genomic DNA probes. Our intention was to verify and classify cytologically visible similarities and differences in repetitive non-coding DNA sequences locations in given Pooidae species. A comparative approach using GISH, instead of comparison of selected sequences or specific chromosome regions, was performed to widen the understanding of the relationships within the Pooidae subfamily at the chromosomal level.

Materials and methods

Plant material

Ten species (Table 1) were selected for comparative GISH analysis: Aegilops speltoides Tausch., Aegilops tauschii Coss., Brachypodium distachyon L., Festuca pratensis L., Hordeum vulgare L., Lolium perenne L., Secale cereale L., Triticosecale Wittm., Triticum aestivum L., and Triticum urartu Tumanian ex Gandilyan. Brachypodium distachyon material was sourced from the University of Silesia in Katowice, Poland. Seeds of Ae. speltoides, S. cereale, T. aestivum, and T. urartu were kindly supplied for the study from the National Small Grains Germplasm Research Facility, National Small Grains Collection (Aberdeen, Idaho, USA). The remaining species were provided by a collection of the Institute of Plant Genetics, Polish Academy of Sciences in Poznan, Poland.

Chromosome preparation

Seeds of each species were germinated on filter paper in Petri dishes for 3–4 days in the dark. The root tips were immersed in ice-cooled water for 26 h, fixed in ethanol and acetic acid (3:1, v/v), and stored at −20 °C until required. Mitotic chromosome preparations were made from root tips digested in a mixture of enzymes, diluted with 0.01 M sodium citric buffer, containing 20 % (v/v) pectinase (Sigma), 1 % (w/v) cellulose (Calbiochem), and 1 % (w/v) cellulase ‘Onozuka R-10’ (Serva). Meristems were dissected from root tips, squashed in drops of 45 % acetic acid, and the good quality preparation was frozen (Hasterok et al. 2006).

DNA isolation and probe labeling

Genomic DNA (gDNA) from young leaves of all selected species were isolated with the C-TAB method (Doyle and
Doyle 1990). DNA from all plants were obtained at the same stage. After the extraction of DNA, samples were labeled by nick translation with tetramethylrhodamine-5-dUTP (Sigma-Aldrich). The ribosomal sequences 35S rDNA and 5S rDNA were labeled with digoxigenin-11-dUTP by nick translation and with tetramethyl-rhodamine-5-dUTP (Sigma-Aldrich) by polymerase chain reaction (PCR), respectively (Kwiatek et al. 2016b).

**In situ hybridization**

The GISH procedures were performed according to the protocol of Kosmala et al. (2006) and Kwiatek et al. (2016a, b), with minor modifications. The GISH protocol was standardized by several repeats to ensure that the obtained results were comparable and reproducible. The hybridization mixture consisted of 50 % deionized formamide, 10 % dextran sulfate, 2 × SSC, 0.5 % SDS, as well as 100–120 ng/slide gDNA probe. The blocking DNA was not used. Several initial GISH experiments were carried out to reveal optimal specifications, such as probe concentrations and wash temperatures. After establishing optimal GISH conditions, we used only those conditions for all experiments described in this paper. Chromosome preparations and hybridization mixture were denatured together at 80 °C for 2 min and then hybridized overnight at 37 °C. The post-hybridization washes were performed, according to Heslop-Harrison (2000), at 42 °C in 2 × SSC buffer. Probes labeled with tetramethylrhodamine-5-dUTP were directly visualized. After the acquisition of images, selected slides were washed off and reprobed with a new set of probes (35S rDNA and 5S rDNA). The reprobing procedures were performed according to the protocol of Heslop-Harrison et al. (1992), with minor modifications. Immunodetection of the digoxigenated probe (35S rDNA) was performed using fluorescein isothiocyanate-conjugated anti-digoxigenin antibody (Sigma-Aldrich). Mitotic cells were examined with an Olympus XM10 CCD camera attached to an Olympus BX61 automatic epifluorescence microscope. Image processing was carried out using Olympus Cell-F imaging software and Micrografx Picture Publisher software.

**Results**

GISH analyses were performed in order to determine the relationships between total genomic DNA of key crops and forage species at the chromosomal level. The experiments were carried out using labeled probes from each species to hybridize with chromosomes of every species used in this study based on a “round robin” rule. Overall, 100 GISH experiments were used (10 probes × 10 species, Table 2). The experiments showed various types of signal locations, including: telomeric and centromeric regions, rDNA loci, subcentromeric and subtelomeric regions, as well as chromosome labeling or signals dispersed along whole chromosomes. Additionally, rDNA-FISH (ribosomal DNA fluorescent in situ hybridization) was carried out when GISH revealed rDNA-like signals.

**Brachypodium distachyon chromosomes**

The GISH experiments on *B. distachyon* (2n = 2x = 10) chromosomes resulted in the 35S rDNA loci being identified using all probes of gDNA (genomic DNA) probes tested. The 5S
Table 2 Specification of comparative mapping of the Pooideae subfamily according to a “round robin” rule

| Species          | B. distachyon                  | H. vulgare                      | S. cereale                    | T. aestivum                   | Ae. tauschii                  |
|------------------|--------------------------------|--------------------------------|-------------------------------|-------------------------------|-------------------------------|
| No. of rDNA loci | 35S 2 loci; 5S 2 loci           | 35S 4 loci; 5S 8 loci           | 35S 2 loci; 5S 4 loci          | 35S 4 loci; 5S 12 loci        | 35S 2 loci; 5S 4 loci          |
| **B. distachyon**| Centromeric regions + 2 signals of 35S rDNA | 4 signals of 35S rDNA (NOR chromosomes 6 and 7) | Telomeric regions + 2 signals of 35S + 4 signals of 5S rDNA | Telomeric regions + 4 signals of 35S rDNA | Telomeric regions + 2 signals of 35S rDNA |
| **H. vulgare**   | 35S rDNA                       | Chromosome labeling + centromeric regions | Chromosome labeling + 2 pericentromeric signals in chromosomes without rDNA loci | Chromosome labeling, in some chromosomes lack of signals in centromeric and terminal regions | Labeling of chromosome segments |
| **S. cereale**   | Strong dispersed signals + 2 signals of 35S + 2 signals of 5S rDNA | Centromeric regions | Chromosome labeling + telomeric regions | Chromosome labeling and signals in chromosomes without rDNA loci | Chromosome labeling |
| **T. aestivum**  | 35S rDNA + 5S rDNA + telomeric regions + centromeric regions | Pericentromeric regions | Very weak chromosome labeling without NORs and telomeric regions | Chromosome labeling | Chromosome labeling |
| **Ae. tauschii** | 35S rDNA + telomeric regions + centromeric regions | Chromosome labeling | Chromosome labeling without telomeric regions | Chromosome labeling | Chromosome labeling + telomeric regions + signals in chromosomes without rDNA loci |
| **Ae. speltoides** | 2 signals of 35S + 2 signals of 5S rDNA + centromeric regions | Pericentromeric regions + centromeric regions | Chromosome labeling without terminal regions | 14 signals of 5S rDNA, centromeric regions in 14 chromosomes of B genome | Chromosome labeling + 2 terminal signals in 1 pair without rDNA loci |
| **T. urartu**    | 2 signals of 35S rDNA           | Chromosome labeling without terminal regions | Chromosome labeling without terminal regions | Chromosome labeling, 14 chromosomes with stronger intensity (genome A) | Chromosome labeling without terminal regions |
| **×Triticosecale** | 35S rDNA + 5S rDNA + strong dispersed signals | Pericentromeric regions | Chromosome labeling without 2 NORs + telomeric regions | Chromosome labeling without terminal regions, centromeric regions and 2 NORs | Chromosome labeling without terminal regions and NORs |
| **F. pratensis** | 35S rDNA + centromeric regions | Chromosome labeling + centromeric regions | Chromosome labeling without terminal regions | Weak dispersed signals | Dispersed signals + telomeric regions (not in all chromosomes) |
| **L. perenne**   | 2 signals of 35S + 2 signals of 5S rDNA + telomeric regions | Chromosome labeling + 4 signals of 35S rDNA | Chromosome labeling without terminal regions + 2 signals of 35S rDNA | 4 signals of 35S rDNA + telomeric regions | 2 signals of 35S + 2 signals of 5S rDNA + telomeric regions + weak dispersed signals |

| Species          | Ae. speltoides                  | T. urartu                      | ×Triticosecale                | F. pratensis                   | L. perenne                    |
|------------------|--------------------------------|--------------------------------|-------------------------------|-------------------------------|-------------------------------|
| No. of rDNA loci | 35S 4 loci; 5S 2 loci           | 35S 4 loci; 5S 4 loci          | 35S 6 loci; 5S 10 loci        | 35S 2 loci; 5S 2 loci          | 35S 7 loci; 5S 2 loci          |
| **B. distachyon**| 4 signals of 35S + 2 signals of 5S rDNA | Telomeric regions + 4 signals of 35S rDNA | Telomeric regions + 6 signals of 35S + 10 signals of 5S rDNA | Telomeric regions + 2 signals of 35S rDNA | Telomeric regions + 7 signals of 35S rDNA |
| **H. vulgare**   | Chromosome labeling + pericentromeric signals | Chromosome labeling + 2 pericentromeric signals | Pericentromeric signals in 14 chromosomes of B genome | Chromosome labeling + 2 signals of 35S rDNA | Chromosome labeling + 7 signals of 35S rDNA |
| **S. cereale**   | Chromosome labeling without terminal regions, 4-5 signals in chromosomes without rDNA loci | Chromosome labeling | Labeling of 14 chromosomes (genome R) | Weak chromosome labeling + 2 signals of 35S rDNA | Strong, dispersed signals |
| Species                  | Ae. speltoides          | T. urartu               | ×Triticosecale         | F. pratensis            | L. perenne               |
|-------------------------|-------------------------|-------------------------|------------------------|-------------------------|--------------------------|
| No. of rDNA loci        | 35S 4 loci; 5S 2 loci   | 35S 4 loci; 5S 4 loci   | 35S 6 loci; 5S 10 loci | 35S 2 loci; 5S 2 loci   | 35S 7 loci; 5S 2 loci    |
| T. aestivum             | Chromosome labeling     | Chromosome labeling     | Labeling of 28...     | Chromosome labeling     | Chromosome labeling      |
|                         | without terminal regions + pericentromeric signals | without terminal regions (1 arm in 2 chromosomes) | (genomes A and B, very weak labeling of 14 R-genome chromosomes) | + 2 signals of 35S rDNA | + 7 signals of 35S rDNA |
| Ae. tauschii            | Chromosome labeling     | Chromosome labeling     | Chromosome labeling, 14 chromosomes with stronger intensity (genome B) | Weak chromosome labeling + 2 signals of 35S rDNA | Weak chromosome labeling + 7 signals of 35S rDNA |
|                         | without terminal regions | (8 chromosomes with stronger intensity) | | | |
| Ae. speltoides          | Chromosome labeling     | Chromosome labeling     | Labeling of 14 chromosomes (genome B) | Chromosome labeling     | Weak chromosome labeling + 7 signals of 35S rDNA |
|                         | + pericentromeric regions | without terminal regions in 1 pair | | | |
| T. urartu               | Chromosome labeling     | Chromosome labeling     | Chromosome labeling (A genome - strong signals, B - weaker intensity, R - very weak labeling) | Weak chromosome labeling + 2 signals of 35S rDNA | Weak chromosome labeling + 7 signals of 35S rDNA |
| ×Triticosecale          | Chromosome labeling     | Chromosome labeling     | Chromosome labeling | 2 signals of 35S rDNA | Chromosome labeling      |
|                         | of selected regions + pericentromeric signals | (6 chromosomes with stronger intensity; 2 chromosomes without labeling of terminal regions) | | | |
| F. pratensis            | Chromosome labeling     | Chromosome labeling     | Chromosome labeling | Chromosome labeling | Chromosome labeling      |
|                         | without terminal regions and interstitial block in chromosome 5 | | | | |
| L. perenne              | 4 signals of 35S + 2 signals of 5S rDNA | 4 signals of 35S rDNA + terminal signals in 2 chromosomes + dispersed signals | Chromosome labeling + 6 signals of 35S rDNA | Strong, distinct signals | Chromosome labeling without terminal regions in 1 pair |

Columns = chromosome preparations; rows = gDNA probes
rDNA loci were labeled by *S. cereale*, *T. aestivum*, *Ae. speltoides* (Fig. 1a), ×*Triticosecale*, and *L. perenne* gDNA probes only. Pericentromeric signals were observed when *B. distachyon*, *T. aestivum*, *Ae. tauschii*, *Ae. speltoides* (Fig. 1a), and *F. pratensis* gDNA probes were used. Rye and triticale probes also gave strong and clear signals dispersed in all *B. distachyon* chromosomes. Furthermore, *B. distachyon* telomeric regions were labeled when *T. aestivum*, *Ae. tauschii*, and *L. perenne* DNA were used as genomic probes.

**Hordeum vulgare chromosomes**

All barley (*H. vulgare*; 2n = 2x = 14) chromosomes were labeled by *H. vulgare*, *Ae. tauschii*, *F. pratensis*, and *L. perenne* gDNA probes. By contrast, *T. urartu* probe marked every barley chromosome but left the telomeric regions unlabeled. Clear signals in the centromeric regions were observed when *H. vulgare*, *S. cereale*, *Ae. speltoides* (Fig. 1b), and *F. pratensis* gDNA probes were hybridized. In addition, pericentromeric regions were identified when *T. aestivum*, *Ae. speltoides*, and ×*Triticosecale* (Fig. 2b) gDNA probes were used. Hybridization using gDNA probes generated from *B. distachyon* and *L. perenne* resulted in four strong signals, which corresponded to the 35S rDNA loci.

**Secale cereale chromosomes**

All rye (*S. cereale*; 2n = 2x = 14) chromosomes were labeled by *H. vulgare*, *S. cereale*, and ×*Triticosecale* probes. The triticale probe did not hybridize with two 35S rDNA loci (Fig. 3c). The *H. vulgare* probe also identified an additional two signals located on the pericentromeric regions of the chromosomes lacking rDNA loci. Hybridizations with the rest of probes, except for *B. distachyon* probe, resulted in labeling of all the rye chromosome except for the telomeric regions. Similarly to triticale, the wheat probe also did not label two
35S rDNA loci on rye chromosomes. In contrast, *B. distachyon* gDNA probe gave signals in two 35S and four 5S rDNA loci, as well as in terminal regions. Finally, the *L. perenne* probe highlighted an additional two regions which corresponded to the 35S rDNA loci.

**Triticum aestivum** chromosomes

All wheat chromosomes (*T. aestivum; 2n = 6x = 42*) were labeled by *T. aestivum*, *T. urartu*, *Ae. tauschii* (Fig. 1c), and *S. cereale* gDNA probes, although the *T. urartu* probe resulted in 14 chromosomes of A genome being marked stronger. The rye gDNA probe showed strong and dispersed signals in most of the wheat chromosomes. Triticale and barley gDNA probes also labeled all chromosomes; however, the centromeric and telomeric regions as well as two 35S rDNA loci (only in the case of triticale probe) remained unlabeled. Contrastingly, the *Ae. speltoides* gDNA probe gave only 14 signals in 5S rDNA loci and 14 centromeric regions in B genome chromosomes. The *B. distachyon* probe resulted in labeled telomeric sites and four signals in 35S rDNA loci, while *L. perenne* yielded four 35S rDNA loci and telomeric regions. Finally, weak and dispersed signals were observed using *F. pratensis* gDNA probe.

**Aegilops tauschii** chromosomes

*Aegilops tauschii* (*2n = 2x = 14*) chromosomes were marked in GISH experiments using *T. aestivum*, *Ae. tauschii*, *S. cereale*, and *Ae. speltoides* gDNA probes. Additionally, *Ae. tauschii* probe hybridization resulted in strong telomeric signals and clear dispersed signals along all chromosomes. Six telomeric sites were detected using rye probe. The *Ae. speltoides* gDNA probe gave also two terminal signals in *Ae. tauschii* chromosomes. The ×*Triticosecale* and *T. urartu* gDNA probes also marked all chromosomes; however, the telomeric regions and two 35S rDNA loci (only in the case of triticale probe) remained unlabeled. Labeled segments of *Ae. tauschii* were observed when *H. vulgare* gDNA probe was used in the GISH experiments. In contrast, *L. perenne* (Fig. 1d) and *F. pratensis* gDNA probes resulted in weak dispersed sites observed in all

![Figure 2](image_url) Hybridization patterns for comparative mapping of gDNA (red): a *B. distachyon* gDNA probe to *Ae. speltoides* chromosomes showing centromeric regions, 5S and 35S rDNA; b ×*Triticosecale* gDNA probe to *H. vulgare* chromosomes: pericentromeric and peritelomeric signals; c *L. perenne* gDNA probe to *F. pratensis* chromosomes: other distinct, strong signals (white arrows); d *S. cereale* gDNA probe to *T. urartu* chromosomes: chromosome labeling and dispersed signals. Scale bar = 5 μm

![Figure 3](image_url) Labeling of all chromosomes using gDNA (red) of selected species beyond: a subtelomeric regions: *T. urartu* gDNA probe to *Ae. speltoides* chromosomes; b centromeric regions: *H. vulgare* gDNA probe to *T. urartu* chromosomes (the arrows indicate distinct pericentromeric signals); c NORs: triticale gDNA probe to *S. cereale* chromosomes (the arrows indicate NORs)
chromosomes. Additionally, the L. perenne probe gave signals in two 35S and two 5S rDNA loci, and also labeled all telomeric regions (Fig. 1d). Two 35S rDNA loci and telomeric sites were also labeled when GISH was carried out using B. distachyon gDNA probe.

_Aegilops speltoides_ chromosomes

The experiments with labeled DNA of _Ae. tauschii, T. urartu_ (Fig. 3a), _S. cereale, T. aestivum_, and _F. pratensis_ (Fig. 1e) revealed labeling of all _Ae. speltoides_ (2n = 2x = 14) chromosomes with the exception of the telomeric regions, whereas gDNA probe of _S. cereale_ revealed strong, distinct signals on several chromosomes. GISH with _Ae. speltoides_ gDNA probe resulted in chromosome labeling and brighter signals in pericentromeric regions. Similar results for all chromosomes of _Ae. speltoides_ were obtained when gDNA of × _Triticosecale, T. aestivum_, and _H. vulgare_ were used for GISH. What is more, the subtelomeric region of a pair of chromosomes possessing the 5S loci remained unlabeled when hybridization with _F. pratensis_ gDNA (Fig. 1e) was performed. Four 35S and two 5S rDNA loci were obtained for the DNA of _L. perenne_ and _B. distachyon_ (Fig. 2a) and centromeric regions were highlighted using _B. distachyon_ only.

_Triticum urartu_ chromosomes

Labeled gDNA of five species, _S. cereale_ (Fig. 2d), _Ae. tauschii, T. urartu, F. pratensis_, and _H. vulgare_ lead to chromosome labeling on their entire length. Additionally, distinct signals in the pericentromeric regions in two chromosomes and unlabeled centromeric regions in all chromosomes were observed in experiments using _H. vulgare_ DNA (Fig. 3b). Chromosome painting with lack of signals in the terminal part of one pair of chromosomes was obtained for the DNA of _Ae. speltoides_ (Fig. 1f), _T. aestivum_, and × _Triticosecale_. Additionally, _Ae. speltoides_ probe resulted in some strong, clear signals in pericentromeric regions (Fig. 1f). _Brachypodium distachyon_ probe resulted in strong subtelomeric signals and gave four signals of 35S rDNA loci. The genomic DNA of _L. perenne_ revealed only the 35S rDNA sequences.

× _Triticosecale_ chromosomes

Hybridization with gDNA of _B. distachyon_ and _H. vulgare_ to chromosomes of triticale (× _Triticosecale_, 2n = 6x = 42) provided contrasting results to the remaining experiments, where chromosome labeling was observed. The gDNA of _B. distachyon_ was mapped in telomeric sites and gave six signals of 35S rDNA loci and ten signals of 5S rDNA loci. The dispersed, strong signals located near the centromeric and telomeric regions in genome B were visible for the experiment with gDNA of _H. vulgare_. GISH with _S. cereale_ probe enabled to distinguish the R genome chromosomes, while the DNA probe of _Ae. speltoides_ visualized the B genome, and DNA of _T. urartu_ the A genome chromosomes. What is more, the hybridization with DNA of _T. aestivum_ enabled to identify A and B genome as well as R genome chromosomes in terms of negative discrimination (very weak labeling). Probing of homologous gDNA to triticale chromosomes resulted in entire chromosomes being labeled. Finally, probing with _Ae. tauschii_ gDNA resulted in chromosome labeling, but with more intensity of the B genome of × _Triticosecale_.

_Festuca pratensis_ chromosomes

GISH with gDNA probes of selected species, with the exception of _F. pratensis_ and _L. perenne_ (Fig. 2c), revealed two signals of 35S rDNA in the karyotype of _F. pratensis_ (2n = 2x = 14). Furthermore, in nearly all cases where 35S rDNA signals were detected, complete chromosome labeling was also observed; the exceptions were _B. distachyon_ and × _Triticosecale_. However, _B. distachyon_ probe produced weak signals in telomeric regions. Chromosome labeling was clearly visible when _F. pratensis_ gDNA was used to probe chromosomes of _F. pratensis_. Hybridization of _L. perenne_ DNA to _F. pratensis_ revealed strong, distinct signals in some chromosomes (Fig. 2c).

_Lolium perenne_ chromosomes

The hybridization of genomic DNA of _B. distachyon, H. vulgare, T. aestivum, Ae. tauschii, Ae. speltoides_, and _T. urartu_ to chromosomes of _L. perenne_ (2n = 2x = 14) identified the 35S rDNA loci. What is more, the _B. distachyon_ gDNA probe labeled the telomeric sites of _L. perenne_. Strong, dense signals were observed for _S. cereale_ probe. _Lolium perenne_ chromosomes probed with gDNA from _H. vulgare, Ae. tauschii, T. urartu_, and _Ae. speltoides_ resulted in both 35S rDNA signals and chromosome labeling. Probing gDNA of × _Triticosecale_ (Fig. 1g), _F. pratensis_, and _L. perenne_ resulted in the labeling of all chromosomes. GISH with gDNA of _L. perenne_ to _L. perenne_ chromosomes showed an unlabeled terminal part of a pair of chromosomes that lacked rDNA loci.

Discussion

In this study, we detected and described the hybridization patterns of various genomic DNA probes generated from key representatives of cereals, forage grasses, and _B. distachyon_ on mitotic chromosomes of those species. The aim of this approach was to score the affinity of the given
species by using gDNA as probes, without blocking DNA from the species chosen for chromosome spreads preparations. As a result, the whole chromosomes or chromosome regions were distinguished or variable types of signals were visualized to prove the different levels of the relationships between genomes used in this study.

A part of the presented results are either as they were expected or confirm the results reported in other papers. However, our study also reveals some novel insights, which require more specific analysis in the future. The most characteristic DNA motifs in the Pooidae subfamily are telomeric and rDNA sequences. Our results confirmed the previous study of Hasterok et al. (2004), who reported labeled pericentromeric regions and 35S rDNA loci on B. distachyon chromosomes using B. distachyon genomic probe. This phenomenon could be explained by the small content of repetitive sequences in this genome. Besides, Mayer et al. (2011) compared the gene order of B. distachyon with H. vulgare and reported that the number of conserved syntenic loci was high (14,422), reflecting a closer phylogenetic affinity. What is more, Ma et al. (2010) carried out GISH reciprocal experiments between B. distachyon and H. vulgare and observed mainly 35S ribosomal DNA loci. In the present study, the same type of affinity was observed in GISH runs with all of the gDNA probes, indicating that this high copy DNA sequence is conserved between analyzed grasses. Ribosomal DNA loci, which are essential for all organisms, can be detected at the cytogenetic level; however, homogenization operates very strongly in these regions in all Poaceae genomes (Sallares and Brown 1999). Signals observed in the centromeric regions of B. distachyon led to the assumption that this species is more closely related to Ae. speltoides and Ae. tauschii than with T. urartu. Moreover, we showed a certain degree of genomic relationship between B. distachyon and F. pratensis. This point of view could also be confirmed by the phylogenetic analysis of grass species based on the ndfH chloroplast gene sequence made by Mochida and Shinozaki (2013).

Considering their importance in agriculture, the most analyzed temperate cereal species in comparative genomics are barley and wheat, along with their relatives. Based on comparison to the previously constructed high-density physical marker map of wheat (Qi et al. 2004), Mayer et al. (2011) reported that barley contains an archetypal Triticeae genome. Most of the chromosome arms showed well-conserved synteny with previously reported chromosomal translocations (Qi et al. 2006). In the present work, we observed labeled pericentromeric regions on H. vulgare chromosomes using Ae. speltoides, T. aestivum, and ×Triticosecale gDNA probes. These observations are correlated with the results of Icsó et al. (2015), where gDNA of H. vulgare was used for the chromosome probing of T. aestivum. They reported that gDNA of barley produced hybridization bands mainly in the pericentromeric and intercalary chromosome regions of B genome chromosomes and it corresponded to the GAA FISH pattern. Surprisingly, we observed differences in the labeling of wheat chromosomes using barley genomic probe in comparison to chromosomes of wheat labeled with gDNA probes of its ancestral species. The three genomes of wheat share similar repetitive sequence types, with the D genome specific repetitive sequences being the most frequent (Nagaki et al. 1995; Jia et al. 2013). In addition, the existence of tandemly repeated sequence dpTa1 presented in 58 species of the Triticeae tribe have been reported (Vershinin et al. 1994). The occurrence of signals detected in phylogenetically distant species may indicate strong relationships between repetitive sequences in selected grasses. For example, the pSc119.2 sequence from rye is widely used in the identification of wheat chromosomes and its relatives (Cuadrado and Jouve 1994; Schneider et al. 2003, 2005; Wiśniewska et al. 2013).

On the other hand, we have observed an unexpected lack of signals in secondary constrictions of rye chromosomes probed by triticale genomic DNA. This may be connected with the differences in rDNA sequences between 1R chromosomes of rye and 1R chromosomes of triticale. In general, unlabeled chromosome regions can be related with species-specific sequences or disparate pathways of chromosome differentiation, which was exposed by the lack of labeled segments of Ae. speltoides chromosomes probed with F. pratensis gDNA. The most likely explanation for the unlabeled telomeric regions could be the fact that the Ae. speltoides genome carries subtelomeric sequences Spelt1 and Spelt52 (Salina et al. 2006, 2009). On the other hand, the lack of signals from the 5S chromosome segment of Ae. speltoides remains unknown but indicates that this region could have gone under structural changes during S genome evolution. What is more, we observed single signals in pericentromeric locations in some chromosomes of T. urartu probed with Ae. speltoides gDNA. Similarly, Belyayev et al. (2000) reported pericentromeric signals in the chromosomes of T. urartu and suggested that those regions are conserved in A and B genomes. Moreover, comparative GISH resulted in a lack of signals in the subtelomeric regions in the chromosomes of T. aestivum, Ae. speltoides, Ae. tauschii, and H. vulgare. This could be determined by the different constitution of subtelomeric regions. For example, the composition of the termini of S. cereale chromosomes is complex and, besides the basic telomeric sequence (T/A)1-4 G1-8, it also contains tandemly organized DNA families such as pSc119.2, pSc200, and pSc250 (Salina et al. 2009).

Our comparative analysis of genomic relationships within crucial Pooidae species revealed some intriguing results, such as unexpected chromosome labeling between distant species (e.g., ×Triticosecale vs. L. perenne), dispersed signals (e.g., L. perenne vs. F. pratensis), or labeling of selective chromosome regions (e.g., S. cereale vs. Ae. speltoides). The
strong, distinct signals observed for *L. perenne* and *F. pratensis* can be useful for the karyotyping of *F. pratensis* chromosomes. Till now, only two pairs of chromosomes, 2F with 355 rDNA and 3F with 5S rDNA loci, can be easily identifiable (Jones et al. 2002; Thomas 1981).

In conclusion, our results are in parallel with the comparative studies made on model species and key representatives of the Pooidae subfamily. However, some results indicated unknown intergeneric dependencies that need further analysis. Moreover, reciprocal GISH analysis on *Aegilops*, *Hordeum*, and *Triticum* species could shed light on the chromosome structure and evolution within the Triticeae tribe.

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**Compliance with ethical standards**

**Ethical approval** This article does not contain any studies with human participants or animals performed by any of the authors.

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**References**

Belyayev A, Raskina O, Korol A, Nevo E (2000) Coevolution of A and B genomes in allotetraploid *Triticum dicoccoides*. Genome 43:1021–1026

Bennett MD, Leitch IJ (2012) Plant DNA C-values database (release 6.0, December 2012). Available online at: http://data.kew.org/cvalues/

Bonierbale MW, Plaisted RL, Tanksley SD (1988) RFLP maps based on a common set of clones reveal modes of chromosomal evolution in potato and tomato. Genetics 120:1095–1103

Catalán P, Shi Y, Armstrong L, Draper J, Stace CA (1995) Molecular phylogeny of the grass genus *Brachypodium* P. Beauv. based on RFLP and RAPD analysis. Bot J Linn Soc 117:263–280

Cuadrado A, Jouve N (1994) Highly repetitive sequences in B chromosomes of *Secale cereale* revealed by fluorosence in situ hybridization. Genome 37:709–712

Cuadrado A, Jouve N (2002) Evolutionary trends of different repetitive DNA sequences during speciation in the genus *Secale*. J Hered 93:339–345

Devos KM (2005) Updating the ‘crop circle’. Curr Opin Plant Biol 8:155–162

Doyle JJ, Doyle JL (1990) A rapid total DNA preparation procedure for fresh plant tissue. Focus 12:13–15

Draper J, Mur LA, Jenkins G, Ghosh-Biswas GC, Bablak P, Hasterok R et al (2001) *Brachypodium distachyon*. A new model system for functional genomics in grasses. Plant Physiol 127:1539–1555

Flavell RB, O’Dell M, Smith DB (1979) Repeated sequence DNA comparisons between *Triticum* and *Aegilops* species. Heredity 42:309–322

Gale MD, Devos KM (1998) Comparative genetics in the grasses. Proc Natl Acad Sci U S A 95:1971–1974

Griffiths S, Sharp R, Foote TN, Bertiin I, Wanous M, Reader S et al (2006) Molecular characterization of *Ph1* as a major chromosome pairing locus in polyploid wheat. Nature 439:749–752

Hasterok R, Draper J, Jenkins G (2004) Laying the cytotoxiconomic foundations of a new model grass, *Brachypodium distachyon* (L.) Beauv. Chromosome Res 12:397–403

Hasterok R, Dulaiva J, Jenkins G, Leggett M, Langdon T (2006) Multisubstrate chromosome preparations for high throughput comparative FISH. BMC Biotechnol 6:20

Heslop-Harrison JS (2000) Comparative genome organization in plants: from sequences and markers to chromatin and chromosomes. Plant Cell 12:617–636

Heslop-Harrison JS, Harrison GE, Leitch IJ (1992) Reprobing of DNA: DNA in situ hybridization preparations. Trends Genet 8:372–373

Hsiao C, Chatterton NJ, Assay KH, Jensen KB (1994) Phylogenetic relationships of 10 grass species: an assessment of phylogenetic utility of the internal transcribed spacer region in nuclear ribosomal DNA in monocytes. Genome 37:112–120

Huo NY, Vogel JP, Lazo GR, You FM, Ma Y, McMahon S et al (2009) Structural characterization of *Brachypodium* genome and its syntetic relationship with rice and wheat. Plant Mol Biol 70:47–61

Icsó D, Molnár-Láng M, Linc G (2015) Constructing an alternative wheat karyotype using barley genomic DNA. J Appl Genet 56:45–48

International Brachypodium Initiative (IBI) (2010) Genome sequencing and analysis of the model grass *Brachypodium distachyon*. Nature 463:763–768

Jia J, Zhao S, Kong X, Li Y, Zhao G, He W et al (2013) *Aegilops tauschii* draft genome sequence reveals a gene repertoire for wheat adaptation. Nature 496:91–95

Jones ES, Mahoney NL, Hayward MD, Armstead IP, Jones JG, Humphreys MO et al (2002) An enhanced molecular marker based genetic map of perennial ryegrass (*Lolium perenne*) reveals comparative relationships with other Poaceae genomes. Genome 45:282–295

Keller B, Feuillet C (2000) Colinearity and gene density in grass genomes. Trends Plant Sci 5:246–251

Kellogg EA (2001) Evolutionary history of the grasses. Plant Physiol 125:1198–1205

Komarala A, Zwireckykowski Z, Gasiör D, Rapacz M, Zwireckywska E, Humphreys MW (2006) GISH/FISH mapping of genes for freezing tolerance transferred from *Festuca pratensis* to *Lolium multiflorum*. Heredity (Edinb) 96:243–251

Kwiatek M, Belter J, Majka M, Wiśniewska H (2016a) Allocation of the S-genome chromosomes of *Aegilops variabilis* Eig. carrying powdery mildew resistance in triticale (*×Triticosecale* Wittm.). Protoplasma 253:329–343. doi:10.1007/s00709-015-0813-6

Kwiatek M, Majka M, Szlaskiewicz-Jarzina A, Ponińska A, Pudelska H, Belter J, Wiśniewska H (2016b) Transmission of the *Ph1* locus in *Aegilops ovata* chromosomes carrying gametocidal factors in hexaploid triticale (*×Triticosecale* Wittm.) hybrids. J Appl Genet 57:305–315

Ma L, Vu GTH, Schubert V, Watanabe K, Stein N, Houwen A et al (2010) Synteny between *Brachypodium distachyon* and *Hordeum vulgare* as revealed by FISH. Chromosome Res 18:295–299

Mayer KFX, Martis M, Hedley PE, Šimková H, Liu H, Morris JA et al (2011) Unlocking the barley genome by chromosomal and comparative genomics. Plant Cell 23:1249–1263
McCouch SR (2001) Genomics and synteny. Plant Physiol 125:152–155
Mochida K, Shinozaki K (2013) Unlocking Triticeae genomics to sustainably feed the future. Plant Cell Physiol 54:1931–1950
Murat F, Zhang R, Guizard S, Flores R, Armero A, Pont C et al (2014) Shared subgenome dominance following polyploidization explains grass genome evolutionary plasticity from a seven protochromosome ancestor with 16K protogenes. Genome Biol Evol 6:12–33
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
Qi LL, Echalier B, Chao S, Lazo GR, Butler GE, Anderson OD et al (2004) A chromosome bin map of 16,000 expressed sequence tag loci and distribution of genes among the three genomes of polyploid wheat. Genetics 168:701–712
Qi L, Fribe B, Gill BS (2006) Complex genome rearrangements reveal evolutionary dynamics of pericentromeric regions in the Triticeae. Genome 49:1628–1639
Salina EA, Lim KY, Badaeva ED, Shcherban AB, Adonina IG, Amosova AV et al (2006) Phylogenetic reconstruction of Aegilops section Sitopsis and the evolution of tandem repeats in the diploids and derived wheat polyploids. Genome 49:1023–1035
Salina EA, Sergeeva EM, Adonina IG, Shcherban AB, Afonnikov DA, Belcram H et al (2009) Isolation and sequence analysis of the wheat B genome subtelomeric DNA. BMC Genomics 10:414
Sallares R, Brown TA (1999) PCR-based analysis of the intergenic spacers of the Nor loci on the A genomes of Triticum diploids and polyploids. Genome 42:116–128
Salse J, Bolot S, Throude M, Jouffe V, Piegu B, Quraishi UM et al (2008) Identification and characterization of shared duplications between rice and wheat provide new insight into grass genome evolution. Plant Cell 20:11–24
Schneider A, Linc G, Molnár-Láng M, Graner A (2003) Fluorescence in situ hybridization polymorphism using two repetitive DNA clones in different cultivars of wheat. Plant Breed 122:396–400
Schneider A, Linc G, Molnár I, Molnár-Láng M (2005) Molecular cytogenetic characterization of Aegilops biuncialis and its use for the identification of 5 derived wheat–Aegilops biuncialis disomic addition lines. Genome 48:1070–1082
Thomas HM (1981) The Giemsa C-banding karyotypes of six Lolium species. Heredity 46:263–267
Vershinin AV, Svitashev S, Gummesson PO, Salomon B, von Bothmer R, Bryngelson T (1994) Characterization of a family of tandemly repeated DNA sequences in Triticeae. Theor Appl Genet 89:217–225
Vershinin AV, Schwarzacher T, Heslop-Harrison JS (1995) The large-scale genomic organization of repetitive DNA families at the telomeres of rye chromosomes. Plant Cell 7:1823–1833
Vogel JP, Bragg JN (2009) Brachypodium distachyon, a new model for the Triticeae. In: Feuillet C, Muehlbauer GJ (eds) Genetics and genomics of the Triticeae, vol 7. Springer, New York, pp 427–449
Vogel J, Hill T (2008) High-efficiency Agrobacterium-mediated transformation of Brachypodium distachyon inbred line Bd21-3. Plant Cell Rep 27:471–478
Wei F, Coe E, Nelson W, Bharti AK, Engler F, Butler E et al (2007) Physical and genetic structure of the maize genome reflects its complex evolutionary history. PLoS Genet 3:e123
Wiśniewska H, Kwiatek M, Kulak-Książczyk S, Apolinski B (2013) Introgression of A- and B-genome of tetraploid triticale chromatin into tetraploid rye. J Appl Genet 54:435–440