Genome constitution and differentiation of subgenomes in Siberian and Far Eastern endemic species of the genus *Elymus* (Poaceae) according to the sequencing of the nuclear gene *waxy*  

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Fifty-three species of perennial grasses in the genus *Elymus* (Poaceae), which are widespread in Russia, are generally assumed to have three haplome combinations: StH, StY and STY. The StH-genome species, endemic to Russia, remain the least studied. R. Mason-Gamer and co-authors have previously shown in a series of studies that a molecular phylogenetic analysis of the low-copy gene *waxy* (GBSS1) sequences significantly complements cytogenetic data on the genomic constitution and evolutionary relationships among both North American and Asian species of the genus *Elymus*. To determine the species’ genomic constitution and to evaluate the level of phylogenetic differentiation, we examined the GBSS1 gene in 18 species of *Elymus* from Siberia and the Russian Far East, including the following 14 endemics: *E. charkeviczii*, *E. jacutensis*, *E. kamczadalorum*, *E. komarovii*, *E. kronokensis*, *E. lenensis*, *E. macrourus*, *E. margaritae*, *E. subfibrosus*, *E. sajanensis*, *E. transbaicalensis*, *E. peschkovae*, *E. uralensis*, and *E. viridiglumis*. PCR amplification products of GBSS1 gene fragments (including exons 9–14) were cloned and 6–8 clones per accession were sequenced. It appears that all the species studied have St and H subgenomic gene variations. The most significant differences between the subgeneric variants St and H were found in intron 13. The H subgenome contains a 21-bp-long deletion in intron 13 in all *Elymus* genotypes, probably derived from a common ancestor of the H and P genomes. Instead of this deletion, all St subgenomes have a relatively conservative sequence similar to that of the genus *Pseudoroegneria*, whose ancestor is considered to be the donor of the modern St subgenome for all *Elymus* species. Cluster phylogenetic analysis revealed differentiation in St and H subgenome sequences into two evolutionarily variant species: St1 vs. St2 and H1 vs. H2, respectively. Variants of the St and H subgenomes were found homologous to various modern species of the ancestral genera *Pseudoroegneria* and *Hordeum*: St1 to *P. strigosa*, St2 to *P. spicata*, H1 to *H. jubatum*, and H2 to *H. californicum*. The details of the relationships between Russian and North American species of the genus, as well as a number of microevolutionary interconnections in the group of boreal endemic species of Siberia and the Russian Far East were revealed. The new results obtained here are essential for the development of a phylogenetically oriented taxonomic system for the genus *Elymus*.  

Key words: *Elymus*; phylogeny; allopolyploids; genome constitution; GBSS1.

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Геномная конституция и дифференциация субгеномов эндемичных сибирских и дальневосточных видов рода *Elymus* (Poaceae) по данным секвенирования ядерного гена *waxy*  

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В России распространены 53 вида многолетних трав рода *Elymus L.* (Poaceae), предположительно трех гаплоидных комбинаций: StH, StY и STY. Наименее изученными остаются бореальные StH-геномные виды — эндемики РФ. Ранее R. Mason-Gamer с соавторами в серии исследований было показано, что молекулярно-филогенетический анализ последовательностей низкокопийного гена *waxy* (GBSS1) существенно дополняет цитогенетические данные по геномной конституции и эволюционным взаимоотношениям как среди североамериканских, так и среди азиатских видов рода *Elymus*. Мы исследовали ген GBSS1 у 18 видов *Elymus*.
Introduction

The genus *Elymus* L. (wildrye) is the largest genus of the tribe *Triticeae* Dumort in the family Poaceae Barn. It contains exclusively amphiploid self-pollinating perennial grasses (Dewey, 1984; Löve, 1984). Species of the genus are widespread on all continents from the Holarctic to the tropics, with more than half populations growing in Central Asia (Lu, 1994). The genomic constitution of all species formed by haplotomes from the ancestors of several modern genera: *Pseudoroegneria* (Nevski) A. Löve (St haplome), *Hordeum* L. (H haplome), *Agropyron* Gaertn. (P haplome) and *Australopyrum* (Tzvelev) A. Löve (W haplome), as well as the Y haplome from an unknown ancestor. The St haplome is common for all species of the genus. After institution and recognition of the genomic classification system for the tribe *Triticeae* (Dewey, 1984), a taxonomic system began to spread, in the broadest sense is divided into different species. This information is essential for the construction of a phylogenetically oriented taxonomic system for the species grown in Russia. The availability of numerous *GBSS1* gene sequences in the NCBI Nucleotide database (http://www.ncbi.nlm.nih.gov/nuccore) enables a comparative analysis of selected locus sequences in combination with other molecular markers allow phylogenetic relationships between taxa to be reconstructed. Moreover, our studies proved that data on the genomic constitution of species and their microevolutionary relationships are to be taken as a fundamental basis to construct phylogenetically-oriented genus systematics for the species grown in Russia. The availability of numerous *GBSS1* gene sequences in the NCBI Nucleotide database (http://www.ncbi.nlm.nih.gov/nuccore) enables a more detailed assessment of the relationships between a large number of genotypes of each species in comparative studies. Here we present a comparative analysis of nucleotides sequences of an ~1300-bp-long fragment of the *GBSS1* gene including exons 9 to 14 in 18 *Elymus* species (including 14 endemics) growing in Siberia and in the Russian Far East in order to establish or confirm their genomic constitution and to assess the evolutionary differentiation levels of subgenomes in different species. This information is essential for the construction of a phylogenetically oriented taxonomic system of the genus species growing in Russia.

Materials and methods

The analyzed accessions included genus *Elymus* species widespread in the Asian part of Russia, mainly with unknown or unconfirmed genomic constitutions (Supplementary markers were obtained by R. Mason-Gamer with collaborators (Helfgott, Mason-Gamer, 2004; Mason-Gamer, 2001, 2004, 2008, 2013; Mason-Gamer et al., 1998; 2010a, b). In particular, their studies have shown that comparative data on nucleotide sequences of the low-copy gene *waxy* (granule-bound starch synthase 1, *GBSS1*) are consistent with cytogenetic data in regard to the genomic constitution and evolutionary origin of North American (Mason-Gamer, 2001) and Asian (Mason-Gamer et al., 2010a) species of the genus *Elymus*.

We have analyzed the applicability of the nuclear low-copy genes *bmy2* and *waxy*, as well as ITS rRNA clusters as genetic markers for the assessment of phylogenetic relationships between species of the genus growing in Siberia and in the Russian Far East (Shmakov et al., 2015). We confirmed that comparative analysis of selected locus sequences in combination with other molecular markers allow phylogenetic relationships between taxa to be reconstructed. Moreover, our studies proved that data on the genomic constitution of species and their microevolutionary relationships are to be taken as a fundamental basis to construct phylogenetically-oriented genus systematics for the species grown in Russia. The availability of numerous *GBSS1* gene sequences in the NCBI Nucleotide database (http://www.ncbi.nlm.nih.gov/nuccore) enables a more detailed assessment of the relationships between a large number of genotypes of each species in comparative studies. Here we present a comparative analysis of nucleotides sequences of an ~1300-bp-long fragment of the *GBSS1* gene including exons 9 to 14 in 18 *Elymus* species (including 14 endemics) growing in Siberia and in the Russian Far East in order to establish or confirm their genomic constitution and to assess the evolutionary differentiation levels of subgenomes in different species. This information is essential for the construction of a phylogenetically oriented taxonomic system of the genus species growing in Russia.
Material 1). Known GBSSI gene sequences of the St and H genomes deposited in the NCBI database were used as references for comparative analysis (Supplementary Material 2). Genomic DNA was extracted from fresh or dried leaves as previously described by Khanuja et al. (1999) with modifications, or by using Nucleospin Plant II kits (Macherey-Nagel, Germany) according to the manufacturer’s recommendations.

The previously described (Mason-Gamer et al., 1998) primers F-for (TGCGAGCTCGACAACATCATGC) and M-bac (GGCGAGCGCGCGCGACCGATCCCCTCGC) were used for PCR-amplification of an ~1300-bp-long GBSSI fragment overlapping gene exons from 9 to 14. The PCR reaction mixture of a total volume of 15 µl contained Taq buffer, 0.2 mM of each dNTP, 1.5 mM MgCl2, 1 µM of each primer, 20 ng genomic DNA, and 1.0 unit of HS Taq DNA polymerase (Eurogen, RF). The following temperature profile was used (a C-1000 thermal cycler, Bio-Rad, USA): primary denaturation for 4 min at 94 °C, then 38 three-step cycles with denaturation for 25 sec at 94 °C, annealing for 30 sec at 65 °C and elongation for 1 min at 72 °C, followed by final elongation for 20 min at 72 °C to terminate the terminal non-matrix addition of deoxyadenosine at the 3'-end of the PCR product (Mason-Gamer et al., 1998). Amplification products were analyzed by 1.7 % agarose gel electrophoresis in TAE buffer at an electric field strength of 4 V/cm.

Since allopolyploid Elymus genomes contain at least two subgenomic variations of the GBSSI gene, amplification was performed in three replicates to minimize the “PCR drift” effect due to stochastic fluctuations at the initial stages of PCR (Wagner et al., 1994). The combined PCR product was ligated into vector pAL2-T (Eurogen, RF) and then cloned in chemically competent XL1-Blue E. coli cells. Positive colonies containing recombinant plasmids with a GBSSI insert were selected by blue/white coloring of E. coli grown on Amp(+) LB-Agar containing X-gal/IPTG. Twenty white colonies for each accession were tested for an insert of the expected length by PCR-amplification with universal M13 primers (Eurogen, RF) followed by electrophoresis analysis. Since allotetraploid Elymus species are believed to be the donor of the modern St genome, and its ancestor is Agropyron-like, our results on the presence of only the St and H genomes support the allopolyploidy hypothesis. Elymus species from the genera Agropyron and Hordeum species) and H (genus Hordeum species) are clearly indicated the presence of only the St and H genomes in all the studied species from Siberia and the Russian Far East, thus confirming that these species belong to the tetraploid StH genome group. Interestingly, the allotetraploid group of Elymus species from North America is also represented mainly by StH genome species (Mason-Gamer, 2001). Only rare individuals of several alien Asian StHY and StY genome species were reported there (Barkworth et al., 2007).

The most notable differences between the St and H subgenome fragments are located in intron 13 (Fig. 1). The H subgenome sequences of this intron in all Elymus genotypes analyzed contained a 21-bp-long deletion, most likely coming from a common ancestor of the H and P subgenomes, since it is also present in modern representatives of related monogenic species from the genera Hordeum and Agropyron. However, all St and Y subgenomes had at the very site of this deletion a relatively conservative sequence, which largely matches a sequence in the genus Pseudoroegneria, whose ancestor is believed to be the donor of the modern St genome. Small deletions are also common for other regions of this intron, but are less frequent in the other GBSSI fragment regions analyzed.

In addition, our analysis of the accessions did not confirm the previously published data on the existence of conservative sites that are absolutely specific to the H and St haplomes (Smakov et al., 2015). This was true only partially of some sequences belonging to different haplomes.

Cluster analysis of the whole GBSSI region from 9 to 14 exon sequences, as well as separate sequences of introns or

\[ \text{Supplementary Materials 1 and 2 are available in the online version of the paper: https://vavilov.elpub.ru/jour/manager/files/SupplAgafonov_engl.pdf} \]
exons, showed common patterns with certain nuances of phyletic connections both within and between related groups of the *Elymus* taxa analyzed. The analysis of the most conservative sites (exons 9–14) showed uniformity within the same subgenomes and at the same time distinction among different subgenomes (Fig. 2).

In the species studied the two subgenomes were found clearly differentiated. For instance, the sequences of the St subgenome were divided into two groups: St 1 and St 2. The sequences of the St1 subgenome for Siberian species are probably older since they were found not only in the northern biotypes *E. macrourus*, *E. jacutensis*, *E. kamczadalorum* and more southern StY genome species *E. gmelini* and *E. pen-
Sequences of a greater number of the *Elymus* species from North American natural accessions initially were subdivided according to the same principle (Mason-Gamer, 2001), therefore we have built a dendrogram that included the endemic species of Asian Russia in comparison with some sequences of North American species. The sequences of the St subgenome including exons 9 to 14 with introns were used. The results are shown in Fig. 4.

In this version of the dendrogram, Asian species were also distributed among two clades with the same composition as in...
the dendrogram constructed using exons alone. Some of the North American species (marked by dots on the dendrogram) together with the Asian species P. strigosa formed a joint clade with the group of the St1 subgenome, while the others met in the St2 group along with all accessions of the North American species P. spicata. GBSS1 sequences of the Y subgenome in E. gmelinii and E. pendulinus showed a closer relationship with the St2 group, which does not contradict the data on the evolutionary origin of this subgenome (Mason-Gamer et al., 2010a).

The H subgenome showed a similar pattern of differentiation. Figure 5 shows a dendrogram constructed from complete sequences of the H genome introns and exons from Russian and North American species (the latter are marked with dots in the figure). Two perennial Hordeum species (marked with asterisks) were taken as references. Gene copies from the H genome appeared to be divided into two main clades (designated as H1 and H2). Clade H1 included exclusively Russian species, while clade H2 was formed by Russian northeastern and all North American species. Each of these clades has its own ancestral taxon from the contemporary genus Hordeum: widespread in Eurasia H. jubatum for the Russian H1 group and North American H. californicum Covas & Stebbins for the H2 group.

Russian species from clade H1 showed significantly greater differentiation than the species from heterogeneous clade H2. Clade H1 appeared to be divided into 3 subclades. Primarily three clones of the northeastern species E. kronokensis and E. peschkovae went to a separate group. This fact indirectly confirms the significant isolation of the latter from Siberian E. confusum, although they are similar in spike morphology. E. confusum, in turn, clustered most closely with the reference H. jubatum. The most distant cluster was formed by all accessions of E. caninus with an addition of the clone of South Ural endemic E. uralensis. The largest cluster was formed by the Siberian mountain species E. komarovii, E. transbaicalensis and E. margaritae, which an addition of a pair of reference clones of E. sibiricus and, as the most unexpected fact, a clone of E. subfibrosus accession from Chukotka.Remarkably, the different reference accessions of E. mutabilis fell into different H subgenome clades.

Mixed clade H2 included not only all North American clones of Elymus species together with H. californicum, but also clones from different regions of Russia: E. kamezadalorum and E. charvevicsii (species from the Kamchatka Peninsula), E. macrourus, E. jacutensis, E. lenensis (northern accessions from the Taymyr Peninsula), E. sajanensis (a Siberian mountain species) and two of three E. mutabilis (reference Chinese

Fig. 3. Differentiation of St subgenomes on the basis of differences in nucleotide sequences in different parts of the GBSS1 gene in Elymus species from the Asian part of Russia in comparison with sequences in reference accessions of the Eurasian species.

| St1 | St2 |
|-----|-----|
| $\text{P.str}_{323}$ | $\text{P.str}_{323}$ |
| $\text{gme}_{7726}_{5}$ | $\text{gme}_{7726}_{5}$ |
| $\text{pen}_{7731}_{5}$ | $\text{pen}_{7731}_{5}$ |
| $\text{kam}_{7726}_{3}$ | $\text{kam}_{7726}_{3}$ |
| $\text{mac}_{0135}_{7}$ | $\text{mac}_{0135}_{7}$ |
| $\text{jac}_{0443}_{1}$ | $\text{jac}_{0443}_{1}$ |
| $\text{tmg}_{1530}_{5}$ | $\text{tmg}_{1530}_{5}$ |
| $\text{tmg}_{0030}_{4}$ | $\text{tmg}_{0030}_{4}$ |
| $\text{mac}_{1520}_{1}$ | $\text{mac}_{1520}_{1}$ |
| $\text{mac}_{1520}_{2}$ | $\text{mac}_{1520}_{2}$ |
| $\text{can}_{5270}_{5}$ | $\text{can}_{5270}_{5}$ |
| $\text{can}_{5272}_{5}$ | $\text{can}_{5272}_{5}$ |
| $\text{can}_{5274}_{5}$ | $\text{can}_{5274}_{5}$ |
| $\text{mut}_{9311}_{5}$ | $\text{mut}_{9311}_{5}$ |
| $\text{mut}_{9328}_{5}$ | $\text{mut}_{9328}_{5}$ |
| $\text{vir}_{1508}_{3}$ | $\text{vir}_{1508}_{3}$ |
| $\text{una}_{0228}_{3}$ | $\text{una}_{0228}_{3}$ |
| $\text{can}_{1502}_{2}$ | $\text{can}_{1502}_{2}$ |
| $\text{cont}_{1502}_{1}$ | $\text{cont}_{1502}_{1}$ |
| $\text{pes}_{1504}_{2}$ | $\text{pes}_{1504}_{2}$ |
| $\text{sib}_{1502}_{6}$ | $\text{sib}_{1502}_{6}$ |

Sample accessions from the Siberian and Far Eastern Elymus species.
Fig. 4. The maximum likelihood tree constructed from multiple alignment of St subgenome GBSS1 intron and exon (9–14) sequences in Elymus species from the Asian part of Russia in comparison with sequences in Eurasian and North American (marked by dots) reference species (St and Y subgenomes). Asterisks indicate Pseudoroegneria species carriers of the St genome. SH-aLRT (%)/UFboot (%) support values are shown.

mut_5279_H and South Ural ABZ06_2). The third reference, mut_9330_H E. mutabilis, from Altai fell into clade H1. Thereby, only some tendency toward relations between the North American accessions and northern or eastern accessions of Russian species can be derived from the H subgenome sequence analysis. The close relationship between American and Kamchatka species is easier to understand, taken into account the historical connections of these flora with each other, as well as with the species from the wide northern distribution areas of E. macrauris and E. jacutensis. It is more difficult to explain the close proximity of Chinese and South Ural accessions of E. mutabilis to this group. Nevertheless, GBSS1 gene variability provides a tool to trace evolutionary relations of species and local geographical
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races from Siberia and the Russian Far East. If we consider the relative position of the accessions inside the clades of the subgenomes, we will see that the clusters combined the species accessions according to their perceived relationship. E. jucutensis and E. macrourus species, for instance, united into the common clusters in both H and St clades (see Fig. 2), as well as on separate dendrograms of these subgenomes (see Fig. 4, 5), thereby confirming the earlier assumptions about E. jucutensis being an aristate subspecies of E. macrourus (Tsvelyov, 1964). This fact is consistent with data on comparative morphological and peptide electrophoretic analyses and hybridization of these species’ particular biotypes (Agafonov, 2008).

A comparative sequence analysis confirmed the isolation of E. kamczadalorum from the Kamchatka species E. charkeviczii, which was previously established using data on comparative morphology, electrophoresis of seed endosperm storage proteins, sexual hybridization (Agafonov, Gerus, 2008) and molecular ISSR analysis (Kobozeva et al., 2017). The species E. komarovii and E. transbaicalensis formed indistinguishable branches inside clade H, together with the Altai species E. margaritae, while E. transbaicalensis and E. margaritae clones were grouped inside clade St1. The phylogenetic proximity of the first two species has been repeatedly experimentally confirmed previously (Agafonov et al., 2019), while the degree of E. margaritae isolation is currently being studied using biosystematic methods. The most unexpected data were obtained regarding the relationships in the group of South Ural biotypes of E. uralensis, E. viridiglumis, E. caninus, and E. mutabilis. These data are currently being verified in the field and laboratory experiments.

Conclusion

Therefore, despite a complicated reticulate evolution in parallel with various related allopolyploid genera and constantly ongoing active microevolutionary transformations, basic genomes seem to have retained unique ancestral features. This makes it relatively easy to identify the genomic composition and to classify modern species within the framework of a phylogenetically oriented taxonomic model of the genus. In our opinion, the integrity of the genus ought to be preserved, because some species in the independent genus Roegneria with the genomic formula StY (Baum et al., 1991) are similar
in morphology to species in the newly proposed genus *Campeostachys* with the genomic formula StHY (Baum et al., 2011), the species of which are significantly different from each other in morphology. The St genome originating from the ancestors of the genus *Pseudoroegneria* seems to be an anchoring constant for a genetic unification of all members of the genus.

We suppose that differentiation of the genus should be based on a model of microevolutionary complexes representing an aggregate of taxa evolving through hybridization and introgression. The degree of taxa relationship within the complex should be confirmed using biosystematic methods with the obligatory determination of the ability to cross, i.e. taking into account the position in the system of recombination (RGP) and introgressive (IHP) gene pools (Agafonov, Salomon, 2002). In fact, the microevolutionary complex is a projection of the RGP collection onto the taxonomic model of the genus, considering the genomic constitution of the species. Each microevolutionary complex should be thought of as a branched system of different ranks of taxa (species and subspecies), remaining therefore a phylogenetically confirmed structure.

In the future, it is necessary to determine the taxonomic rank of microevolutionary complexes, which can be sections or aggregates of the same species in a broad sense, as shown by the example of a revision of *Pendulini* (Nevski) Tzvelev sub-section of the *Gouldaria* (Huss.) Tzvelev section (Kobozeva, Agafonov, 2015).

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