Alloplasmic recombinant lines (H. vulgare)-T. aestivum with 1RS.1BL translocation: initial genotypes for production of common wheat varieties

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Alloplasmic lines are formed when the cytoplasm of one species is replaced by the cytoplasm of another as a result of repeated recurrent crosses of wide hybrids with the paternal genotype. Since the cytoplasm replacement results in new intergenomic interactions between a nucleus and cytoplasm leading to variability of plant characteristics, alloplasmic lines with restored fertility can be an additional source of biodiversity of cultivated plants. Earlier, recombinant alloplasmic lines (H. vulgare)-T. aestivum designated as L-17(1)-L-17(37) were formed from a plant with partially restored fertility of the BC generation of barley-wheat hybrid H. vulgare (cv. Nepolegayushchii) × T. aestivum (cv. Saratovskaya 29). This male-sterile hybrid was consistently backcrossed with wheat varieties Mironovskaya 808 (twice) and Saratovskaya 29, and Mironovskaya 808 had a positive impact on the restoration of fertility. This paper presents the results of investigation into a group of recombinant alloplasmic lines (L-17F1), as well as into doubled haploids (DH) lines – alloplasmic DH-17-lines obtained from another culture of alloplasmic lines (L-17F1). The most productive of these lines were used as initial breeding genotypes. Hybrid form Lutescens 311/00-22 developed from the crossing of the alloplasmic DH-1(1)-17 line (as maternal genotype) with euplasmic line Com37 (CIMMYT), the source of the 1RS.1BL euplasmic line Com37 (CIMMYT), the source of the 1RS.1BL translocation, proved to be successful for breeding. The presence of the 1RS.1BL translocation in the genome of the Lutescens 311/00-22 form and the L-311(1)-L-311(6) alloplasmic lines isolated from it did not lead to a decrease of fertility or sterility in the plants. This indicates that the chromosome of the 1BS wheat does not carry the gene(s) that determine the restoration of fertility in the studied (H. vulgare)-T. aestivum alloplasmic lines. Alloplasmic lines L-311(1)-L-311(6) showed their advantage in comparison with the standard varieties for resistance to leaf and stem rust, yield, and grain quality. The breeding tests performed at Omsk Agricultural Scientific Center, Agrocomplex "Kurgansemena", Federal State Unitary Enterprise "Ishimskoe" (Tyumen Region), using alloplasmic lines L-311(5), L-311(4) and L-311(6) resulted in varieties of spring common wheat Sigma, Uralosibirskaya 2 and Ishimskaya 11, respectively.

Key words: alloplasmic lines (H. vulgare)-T. aestivum; DH-lines; translocation 1RS.1BL; varieties of common wheat.
Alloplasmic lines are formed as a result of repeated recurrent crosses of wide hybrids with the paternal genotype and combine the cytoplasm of the maternal species with the nuclear genome of the pollen parent (Tsunewaki, 1996). The new nuclear-cytoplasmic interactions formed due to cytoplasm replacement can cause epigenetic modifications of nuclear genes (Soltani et al., 2016), leading to changes at the level of transcription and metabolism (Crosatti et al., 2013; Soltani et al., 2016). These processes may lead to plant developmental disorders (Suzuki et al., 1995), changes in resistance to stress factors (Dhitaphichit et al., 1989; Keane, Jones, 1990; Buloychik et al., 2002; Talukder et al., 2015) and changes in the manifestation of agronomically important traits (Ekiz et al., 1998; Liu et al., 2002; Atienza et al., 2013). The most common sign of a nuclear-cytoplasmic conflict included in alloplasmic lines is cytoplasmic male sterility (CMS) (Tsunewaki, 1996; Bentolila et al., 2002).

From a practical perspective, alloplasmic lines are obtained from different species of cultivated plants and characterized by CMS that represents a system for obtaining hybrid seeds in heterotic breeding (Cisar, Cooper, 2002). Given that new intergenic interactions arise during cytoplasm replacement, alloplasmic lines with restored fertility can be an additional source of biodiversity in cultivated plants (Liu et al., 2016). For wheat alloplasmic lines, examples of their practical use are not numerous. The VPM1 alloplasmic line carrying the cytoplasm of *Aegilops ventricosa* Tausch was the source of two commercial varieties of wheat Roason and Rendezvous (Jones et al., 1998) resulting from hybridization of the amphiploid (*Ae. ventricosa × T. persicum*) with common wheate variety Marne. This alloplasmic line and the varieties derived from it are resistant to a number of fungal pathogens due to the introgression of *Ae. ventricosa* genes *Pch-1* and *Sr38/Lr37/Yr17* (Delibes et al., 1988; Friebe et al., 1996). Common wheat variety ‘Xiaoshan 2134’ carrying the cytoplasm of *Ae. crusii*, is characterized by high grain quality, resistance to salinity and high yield (Liu et al., 2002). The authors ascribe these traits to the effect of nuclear-cytoplasmic heterosis. The lines of common wheat with cytoplasm *Triticum timopheevii* and *Secale cereale* characterized by tolerance to drought (Semenov et al., 2014) and high-quality gluten characteristics are considered promising for breeding (Klimushina et al., 2013; Semenov et al., 2016). With the use of rye cytoplasm, new forms of triticale have been obtained (Gordey et al., 2011).

Earlier, we reported the production of the recombinant alloplasmic lines of common wheat derived from backcross progenies of barley-wheat hybrids *H. vulgar* × *T. aestivum* (Pershina et al., 1998) and restoration of their fertility (Pershina et al., 1999a). Alloplasmic lines (*H. vulgar* × *T. aestivum* characterized by different fertility levels proved to be valuable models for studying the variability of both nuclear and organellar (mitochondrial and chloroplast) genomes in the process of nuclear-cytoplasmic co-adaptation (Bildanova et al., 2004; Aksyonova et al., 2005; Pershina et al., 2014). Some
of the recombinant alloplasmic lines with full fertility restoration were included in breeding of common wheat (Belan et al., 2017).

The aim of presented research was to generalize and analyse the results of obtaining and studying of recombinant and introgressive alloplasmic lines (H. vulgare)-T. aestivum, which have been successfully used in breeding and have become a source of the new varieties of spring common wheat.

Materials and methods

Production of recombinant and introgressive alloplasmic lines (H. vulgare)-T. aestivum. The Nepolegyayushchii variety of cultivated barley was crossed with spring common wheat Saratovskaya 29, and the pollinated flowers were treated with a solution of gibberellic acid. Using embryo cultivation, hybrid plants H. vulgare × T. aestivum were grown and were characterized by male sterility but female fertility (Pershina et al., 1998). The F1 hybrids were consistently backcrossed with wheat Mironovskaya 808 (twice) and Saratovskaya 29. Among the plants from the BC1 generation, one partially fertile plant was isolated (three seeds were set in one spike, and 34 in another) (Pershina et al., 1999a). Each fertile plants grown from these seeds became the source of recombinant alloplasmic lines designated L-17(1)–L-17(37) (Fig. 1). To form each subsequent generation of alloplasmic lines, the seeds from the main spike of the most productive plant were selected. Alloplasmic DH-lines DH(1)-17F5, DH(1)-17F6, and DH(1)-17F7, were derived by another culture of alloplasmic line L-17F2 (Pershina et al., 1999b).

In this paper, we present the results of a study of seven alloplasmic lines: L-17(3)F4, L-17(4)F5, L-17(9)F6, L-17(12)F7, L-17(18)F8, L-17(21)F9, and L-17(24)F10; three alloplasmic DH lines: DH(1)-17F2, DH(2)-17F3, and DH(3)-17F4; and six introgression alloplasmic lines: L-311(1)F10–L-311(6)F15. The introgression alloplasmic lines were isolated from the hybrid form L-311/00-22F, obtained as a result of the crossing of alloplasmic line DH(1)-17F2 with euplasmic line Com37 (Belan et al., 2010). Using the results of chromosome C-bandng, E.D. Badaeva identified the wheat-rye translocation 1RS.1BL in the alloplasmic lines of group L-311 (Pershina et al., 2013).

Growing conditions and study of alloplasmic lines. Alloplasmic lines L-17F5 and DH-17F2 were grown in the field of Institute of Cytology and Genetics near Novosibirsk in 1999 when there was no mass development of leaf rust or stem rust pathogens. Control wheat line Saratovskaya 29 (Sar29) was used to obtain barley-wheat hybrids and alloplasmic lines (H. vulgare)-T. aestivum. The alloplasmic lines of the L-311F10 group were grown in the field of the Institute of Cytology and Genetics in 2017. Euplasmic line Om37 carrying translocation 1RS.1BL and isolated from variety Omskaya 37, and alloplasmic line L-17(3)F12 were used as controls. Plants were grown on plots 50 cm wide in rows of 10, with a distance of 25 cm between rows. The germination rate was determined by the number of seedlings. During harvesting, the height of the plants, the number of productive spikes, the length of the main spike, the number of spikelets per main spike, the number of seeds per main spike, and the frequency of plants with full fertility (more than 35 seeds in the main spike) were estimated.

To confirm the presence of the translocation 1RS.1BL, the SCAR marker iag95 linked to the genes Lr26 and Sr31 localized in the short arm of the rye chromosome 1R (Mago et al., 2002), and genomic in situ hybridization (Mukai, Gill, 1991) were used in the lines included in this study. At least 20 plants were used per treatment. The differences between the average values of the lines were compared using t-tests. Data were analysed using Statistica v.7.0.61.0.

Alloplasmic hybrid form L-311/00-22 and alloplasmic lines L-311(1)–L-311(6) were grown in the field of Laboratory of Spring Common Wheat of Omsk Agrarian Scientific Center, following the methods described in (Belan et al., 2017). Since 2007, the L-311 alloplasmic lines have been tested according the full breeding scheme for BN-1, BN-2, and BN-3 (breeding nurseries of the first, second, and third year of study) and in a competitive variety trial nursery. Assessments of resistance to powdery mildew (Blumeria graminis f. sp. tritici), leaf rust (Puccinia recondita f. sp. tritici), and stem rust (Puccinia graminis f. sp. tritici) were made according to previously described methods (Belan et al., 2017). In addition, yield and grain quality parameters (1000-grain weight, seed protein content) were determined. Using analogous methods, alloplas-
mic line L-311(4) has been studied in the Agrocomplex “Kurgansemena” competitive variety trial nursery since 2013, and alloplasmic line L-311(6) – since 2014 in the Enterprise “Ishinskoe” (Tyumen Region) competitive variety trial nursery.

Results and discussion
In this study recombinant alloplasmic lines (H. vulgare)-T. aestivum were investigated, in which barley variety Nepolegaysuchii (Nep) was a source of the cytoplasm, and the recombinant nuclear genome was formed with the use of common wheat varieties Saratovskaya 29 (Sar29) and Mironovskaya 808 (Mir808) (see Fig. 1) (Pershina et al., 1999a). Use of the Mir808 variety in recurrent crosses of male sterile barley-wheat hybrid H. vulgare (Nep) × T. aestivum (Sar29) had a positive effect on fertility restoration: one of the plants of the BC4 generation set seeds after self-pollination (see Fig. 1).

According to our data, the Mironovskaya yarovaia wheat variety recovered from Mir808 (Dorofeev et al., 1987), along with the Ulyanovka and Pyrotrix 28 varieties (Pershina et al., 1999a, 2014), which also restored full fertility of common wheat in the barley (H. vulgare) cytoplasm. When the backcrossing of barley-wheat hybrids H. vulgare × T. aestivum (Sar29) included only the Sar29 variety in the backcrossed progenies, fixation of complete sterility occurred (see Fig. 1) (Pershina et al., 1999a, 2014). In addition, depending on the genetic diversity of wheat varieties included in the backcrossing of barley-wheat hybrids, numerous alloplasmic lines (H. vulgare)-T. aestivum with different levels of fertility were obtained, some of which, when self-pollinated, segregated into plants with low fertility including those being completely sterile (Pershina et al., 1999a, 2014).

Alloplasmic lines L-17(1)-L-17(37) formed from a partially fertile BC3 generation plant with a 2n = 42 did not contain barley chromosomes and were fertile in F4 generations (Bildanova et al., 2004). The germination of the recombinant L-17F4 alloplasmic lines was similar to the control level (not less than 98 %), and all the plants were fertile (Table 1).

In the majority of alloplasmic lines, the frequency of plants with full fertility (seed set higher than 35 seeds in the main spike) was at the level of the Sar29 parent line, varying from 65 % in L-17(18)F4 to 95 % in L-17(21)F4 and L-17(12)F4 (see Table 1). The full fertility was significantly lower than that of the control (60 %) in alloplasmic line L-17(9)F4. The analysis of the productivity characteristics (the number of productive spikes, the length of the main spike, the number of spikelets and seeds per main spike, and the number of seeds per plant) differed between different alloplasmic lines. So, in L-17(9)F4 and L-17(18)F4, the values of some of these parameters were significantly lower than in the Sar29 control line, while in L-17(4)F4 they reached the control level. Alloplasmic lines L-17(21)F4, L-17(12)F4, L-17(3)F4 and L-17(24)F4 exceeded line Sar29 in the number of seeds per plant. The increase in L-17(21)F4 was due to an increase in the number of seeds per main spike; an increase of a number of productive spikes in alloplasmic lines L-17(3)F4 and L-17(12)F4; and an increase in the number of productive spikelets per plant and the number of spikelets per main spike in alloplasmic line L-17(24)F4. For 1000-grain weight, the values of this parameter were the same at the
control level for the majority of the lines studied and was lower for lines F-17(9)F₄, L-17(18)F₄, and L-17(21)F₄ (see Table 1).

These data show that the new nuclear genome that formed as a result of recombination between two genomes of wheat varieties Saratovskaya 29 and Mironovskaya 808 when interacting with the cytoplasm of barley had a positive effect not only on the restoration and maintenance of fertility of alloplasmic lines (*H. vulgare*-*T. aestivum* but also on the traits that determine the productivity of plants. This is mostly due to the high ecological plasticity and high productivity of Saratovskaya 29 and Mironovskaya 808, which have been grown for a long time in many regions with different soil and climatic conditions and hybridized with each other and with other genotypes of wheat to produce many varieties (Dorofeev et al., 1987).

Another important result of our study is that fertility and the signs of high productivity were also evident in the DH lines derived from alloplasmic line L-17F₂. Earlier we had been able to obtain 20 green plants grown from anther culture of alloplasmic line L-17F₂ on a modified medium of P-II (Pershina et al., 1999b). Eleven of these plants with 2n = 42 were fertile. These plants were used for development of a separate alloplasmic DH line. In the current paper we present the results of a study of the self-pollinated progeny of three alloplasmic DH lines: DH(1)-17F₂, DH(2)-17F₂, and DH(3)-17F₂ (see Table 1). In these lines, in comparison with line Sar29, the number of productive spikes, the number of spikelets and seeds per main spike, and the number of seeds per plant were significantly higher. Due to their high level of fertility and productivity, these alloplasmic DH lines, along with the original alloplasmic lines L-17(3)F₄ and L-17(24)F₄, were included in the breeding. In addition, among these alloplasmic lines, we found three more undersized lines such as L-17(3)F₄, DH(2)-17F₂, and DH(3)-17F₂.

To date, from a practical point of view, the hybrid combination obtained in Laboratory of Spring Common Wheat Breeding (Omsk Agrarian Scientific Center) by crossing alloplasmic line DH(1)-17F₂ (maternal genotype) with line Com37 (pollinator, provided for the study by A.I. Morgunov from the CIMMYT collection) has been the most effective. In the early breeding nurseries, hybrid alloplasmic form Lutescens 311/00-22 was reproduced as a population, from which, after selection based on agronomical valuable traits, six promising alloplasmic lines were identified: L-311(1), L-311(2), L-311(3), L-311(4), L-311(5), and L-311(6).

Using C-banding, E.D. Badaeva revealed in these alloplasmic lines a wheat-rye translocation identified as 1RS.1BL (Pershina et al., 2013), which was inherited in a number of self-pollinated generations. Figure 2 shows the spikes of alloplasmic lines L-311(4)F₁₀, L-311(5)F₁₀, L-311(6)F₁₀, and the presence of translocation of 1RS.1BL in these lines proved by the results of GISH and PCR analysis. Since translocation 1RS.1BL was not detected the alloplasmic line of group L-17, it can be assumed that Com37 used as a pollinator in production of hybrid Lutescens 311/00-22, had this translocation. (Currently,
the Com37 line and alloplasmic lines DH(1)-17F$_2$ have been lost.)

The fact that the substitution of chromosome 1BS for 1RS did not result in decrease of fertility or sterility in alloplasmic lines (H. vulgare)-T. aestivum indicates that the restoration and maintenance of fertility of these lines does not depend on the effect of $Rf$-genes controlling the restoration of wheat fertility in the alien cytoplasm and localized in chromosome 1BS (Tsunewaki, 2015).

In our previous work, other genotypes of alloplasmic lines (H. vulgare)-T. aestivum were obtained in which the introduction of wheat-rye translocation 1RS.1BL into the genome, including a combination with wheat-Agropyrum translocation 7DL-7Ai, did not have a negative effect on the fertility (Pershina et al., 2014).

The results of a study into the morphobiological traits of alloplasmic lines L-311(1)F$_{10}$-L-311(6)F$_{10}$ in comparison with alloplasmic line L-17(3)F$_{12}$ (has no the1RS.1BL translocation), and euplasmic line Om37 carrying this translocation, are presented in Table 2.

In all the six alloplasmic lines of group L-311F$_{10}$ carrying the 1RS.1BL translocation the lengths of the main spike, the number of spikelets and the number of seeds per main spike were significantly higher than in control line L-17(3)F$_{12}$ and line Om37. In addition, alloplasmic lines L-311(3)F$_{10}$, L-311(4)F$_{10}$, L-311(5)F$_{10}$, and L-311(6)F$_{10}$ exceeded the control line in the number of seeds per plant, and the lines L-311(4)F$_{10}$, L-311(5)F$_{10}$, and L-311(6)F$_{10}$, in the number of productive spikes.

Since rye chromosome arm 1RS carries a complex of genes ($Lr26/Sr31/Yr9/Pm8$) controlling resistance to fungal pathogens (Singh et al., 1990), the varieties with 1RS.1BL have become widespread throughout the world (Rabinovich, 1998; Schlegel, 2018). Mass cultivation of homogeneous varieties of wheat with this translocation has reduced the effectiveness of the gene resistance to fungal pathogens, including appearance of the highly aggressive $Ug99$ race virulent to the Sr31 gene (Pretorius et al., 2000).

At the same time, the effect of 1RS.1BL translocation on the manifestation of agronomically important and adaptive

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### Table 2. Characteristics of alloplasmic lines L-311(1)F$_{10}$-L-311(6)F$_{10}$ in comparison with euplasmic line Om37-1RS.1BL and alloplasmic line L-17(3)F$_{12}$

<table>
<thead>
<tr>
<th>Genotypes</th>
<th>Number of productive spikes</th>
<th>Length of main spike, cm</th>
<th>Number of spikelets per main spike</th>
<th>Number of seeds per main spike</th>
<th>Number of seeds per plant</th>
</tr>
</thead>
<tbody>
<tr>
<td>Om37</td>
<td>5.0 ± 0.5</td>
<td>10.5 ± 0.2</td>
<td>18.6 ± 0.3</td>
<td>41.5 ± 1.2</td>
<td>183.2 ± 18.1</td>
</tr>
<tr>
<td>L-311(1)F$_{10}$</td>
<td>5.2 ± 0.3</td>
<td>11.5 ± 0.2**</td>
<td>19.6 ± 0.3*</td>
<td>47.9 ± 1.2**</td>
<td>194.3 ± 18.5</td>
</tr>
<tr>
<td>L-311(2)F$_{10}$</td>
<td>6.0 ± 0.4</td>
<td>11.8 ± 0.1***</td>
<td>19.9 ± 0.2***</td>
<td>49.5 ± 0.8***</td>
<td>231.8 ± 18.3</td>
</tr>
<tr>
<td>L-311(3)F$_{10}$</td>
<td>6.1 ± 0.5</td>
<td>12.4 ± 0.1***</td>
<td>20.3 ± 0.3***</td>
<td>49.5 ± 1.2</td>
<td>257.4 ± 16.2**</td>
</tr>
<tr>
<td>L-311(4)F$_{10}$</td>
<td>6.8 ± 0.4*</td>
<td>12.4 ± 0.1***</td>
<td>20.9 ± 0.3***</td>
<td>56.8 ± 1.0***</td>
<td>263.5 ± 19.3**</td>
</tr>
<tr>
<td>L-311(5)F$_{10}$</td>
<td>6.5 ± 0.5*</td>
<td>12.7 ± 0.2***</td>
<td>20.3 ± 0.4***</td>
<td>50.9 ± 2.1***</td>
<td>252.1 ± 22.2*</td>
</tr>
<tr>
<td>L-311(6)F$_{10}$</td>
<td>6.3 ± 0.4*</td>
<td>12.8 ± 0.2***</td>
<td>21.2 ± 0.4***</td>
<td>53.8 ± 1.2***</td>
<td>260.3 ± 22.6*</td>
</tr>
<tr>
<td>L-17(3)F$_{12}$</td>
<td>4.7 ± 0.4</td>
<td>8.5 ± 0.2(***</td>
<td>14.6 ± 0.2(***</td>
<td>32.6 ± 1.2(***</td>
<td>153.7 ± 13.4</td>
</tr>
</tbody>
</table>

Note: The difference in comparison with euplasmic line Om37 is significantly greater at $^* p < 0.05$; $^{**} p < 0.01$; $^{***} p < 0.001$; and significantly less at $^{***} p < 0.001$.

### Table 3. Results of studying promising alloplasmic lines of group L-311 in a competitive variety trial nursery, Omsk, 2010

<table>
<thead>
<tr>
<th>Lines</th>
<th>Growing season, days</th>
<th>Productivity, t/ha</th>
<th>1000-grain weight</th>
<th>Protein content in the grain (N×5.7), %</th>
<th>Lesion of rust disease</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>lines ± of Om33</td>
<td>lines ± of Om33</td>
<td>lines ± of Om33</td>
<td>lines ± of Om33</td>
<td>lines ± of Om33</td>
</tr>
<tr>
<td>L-311(1)</td>
<td>98</td>
<td>4.75 ± 0.16</td>
<td>47.7 ± 4.6</td>
<td>16.56 ± 1.48</td>
<td>0 ± 0.00</td>
</tr>
<tr>
<td>L-311(2)</td>
<td>97</td>
<td>5.00 ± 0.41</td>
<td>44.9 ± 1.8</td>
<td>18.24 ± 3.16</td>
<td>1 ± 0.02</td>
</tr>
<tr>
<td>L-311(3)</td>
<td>98</td>
<td>5.29 ± 0.7</td>
<td>43.0 ± 0.1</td>
<td>18.78 ± 3.7</td>
<td>1 ± 0.02</td>
</tr>
<tr>
<td>L-311(4)</td>
<td>98</td>
<td>5.05 ± 0.46</td>
<td>44.4 ± 1.3</td>
<td>17.78 ± 2.7</td>
<td>5 ± 0.05</td>
</tr>
<tr>
<td>L-311(5)</td>
<td>94</td>
<td>5.25 ± 0.66</td>
<td>45.8 ± 2.7</td>
<td>17.64 ± 5.6</td>
<td>5 ± 0.05</td>
</tr>
<tr>
<td>L-311(6)</td>
<td>93</td>
<td>5.00 ± 0.4</td>
<td>45.5 ± 2.4</td>
<td>17.53 ± 2.4</td>
<td>10</td>
</tr>
<tr>
<td>Om33-St</td>
<td>93</td>
<td>4.59 ± 0.0</td>
<td>43.1 ± 15.08</td>
<td>80.0 ± 5.0</td>
<td>50</td>
</tr>
</tbody>
</table>

Note: Om33-St is a variety of spring common wheat Omskaya 33 used as a reference.
traits depends on the genetic background and plant growing regions (Hoffmann, 2008; Tahmasebi et al., 2015). The gene Sr31 remains effective for protection against stem rust in different regions of Russia (Gultyaeva, 2017).

From the data obtained a conclusion can be made that the genetic background of the alloplasmic lines of group L-311 was favourable for realization of the positive effect of the 1RS.1BL translocation on both agronomically important and adaptive traits in the breeding trials regions including Omsk, Kurgan and Tyumen regions.

The study of alloplasmic lines L-311(1)–L-311(6) for consecutive self-pollinated generations in breeding nurseries and competitive variety trial nurseries was initially carried out in the fields of the Omsk Agrarian Scientific Center. It should be emphasized that leaf rust expansion has been noted there every year since 2001, and in 2007 the epiphytoty of this pathogen was recorded. Since 2009, the threat of stem rust has increased. In addition, powdery mildew is observed in this region every year.

Studying alloplasmic lines L-311(1)–L-311(6) demonstrated their moderate resistance to powdery mildew. In comparison with variety-standard Omskaya 33 susceptible to leaf and stem rust pathogens, the alloplasmic lines showed complex resistance to local populations of leaf and stem rust (Table 3).

Moreover, these alloplasmic lines exceeded the variety-standard for yield, 1000-grain weight, and seed protein content. Table 3 presents the results of studying alloplasmic lines L-311(1)–L-311(6) in 2010 in a competitive variety trial nursery. According to the tests results obtained from 2010 to 2012, middle-ripened alloplasmic line L-311(5) was identified as the most promising and was transferred in 2013 to the State Committee for Testing of New Varieties as variety Sigma. This variety was included in State Register of Selection Achievements of the Russian Federation in West Siberia in 2016.

The middle-ripened alloplasmic line L-311(4) was studied in the competitive variety trial nursery of Agro-complex “Kurgansemena”. Based on the three year results (2012–2015), variety Uralosibirskaya 2 was transferred to the State Committee for Testing of New Varieties in 2016. This variety was tested in the Urals and West Siberian region. In 2014, middle-early alloplasmic line L-311(6) was studied in the competitive variety trial nursery of Enterprise “Ishimskoe”, Tyumen Region. Based on the four-year data this line was transferred to the State Committee for Testing of New Varieties as variety Ishimskaya 11 in West Siberia in 2017.

In addition, the efficiency of using alloplasmic lines from group L-311 for obtaining hybrids and the alloplasmic DH lines combining resistance genes to fungal pathogens localized in wheat-rye translocation 1RS.1BL with complexes of effective resistance genes from other sources has been shown (Pershina et al., 2013).

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Conflict of interest

The authors declare they have no conflict of interest.

References


Talukder S.K., Vara Prasad P.V., Todd T., Babar M.A., Poland J., Bowden R., Fritz A. Effect of cytoplasmic diver-
Alloplasmic lines (*H. vulgare*)-*T. aestivum* with 1RS.1BL for production of wheat varieties

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