Metabolomic approach to search for fungal resistant forms of Aegilops tauschii Coss. from the VIR collection

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Abstract. Broadening of the genetic diversity of donors of resistance to biotic environmental factors is a challenging problem concerning Triticum L., which can be solved by using wild relatives of wheat, in particular, Aegilops tauschii Coss., in breeding programs. This species, believed to be the donor of D genome of common wheat (T. aestivum L.), is a source of some traits important for breeding. This greatly facilitates the possibility of crossing Ae. tauschii with common wheat. Aegilops L. species are donors of effective genes for resistance to fungal diseases in wheat. For instance, genes that determine resistance to rust agents in common wheat were successfully introgressed from Ae. tauschii into the genome of T. aestivum L. The aim of our study was to identify differences in metabolomic profiles of Ae. tauschii forms (genotypes), resistant or susceptible to such fungal pathogens as Puccinia triticina f. sp. tritici and Erysiphe graminis f. sp. tritici. These indicators may be used as biochemical markers of resistance. A comparative analysis of groups of Ae. tauschii accessions showed that metabolomic profiles of the forms with or without resistance to fungal pathogens differed significantly in the contents of nonproteinogenic amino acids, polyols, phyto sterols, acylglycerols, mono- and oligosaccharides, glycosides, phenolic compounds (hydroquinone, kempferol), etc. This fact was consistent with the previously obtained data on the relationship between Fusarium resistance in oats (Avena sativa L.) and certain components of the metabolomic profile, such as acylglycerols, nonproteinogenic amino acids, galactinol, etc. Thus, our studies once again confirmed the possibility and effectiveness of the use of metabolomic analysis for screening the genetic diversity of accessions in the VIR collection, of Ae. tauschii in particular, in order to identify forms with a set of compounds in their metabolomic profile, which characterize them as resistant. Ae. tauschii accessions with a high content of pipecolic acids, acylglycerols, galactinol, stigma sterol, glycerol, azelaic and pyrogallic acids, campesterol, hydroquinone, etc., can be used for creating wheat and triticale cultivars with high resistance to fungal pathogens causing powdery mildew, brown rust, and yellow rust.

Key words: Aegilops tauschii Coss.; metabolomic approach; disease resistance; fungal pathogens.

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Использование метаболомного подхода для поиска форм Aegilops tauschii Coss. из коллекции ВИР им. Н.И. Вавилова, устойчивых к грибным патогенам

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Аннотация. Расширение генетического разнообразия доноров устойчивости Triticum L. к биотическим факторам среды – актуальная задача, которую возможно решить благодаря использованию в селекционных программах дикорастущих родичей пшеницы, в частности Aegilops tauschii Coss. По существующим представлениям, последний – донор генома D мягкой пшеницы T. aestivum L. и носитель ряда ценных селекционных признаков. Это значительно облегчает скрещивание Ae. tauschii с мягкой пшеницей. Виды рода Aegilops L. являются донорами эффективных генов устойчивости пшеницы к грибным болезням. Так, от видов Ae. tauschii в геном T. aestivum L. успешно интродрессированы гены, детерминирующие устойчивость мягкой пшеницы к возбудителям ржавчины. Целью нашего исследования было выявление различий по метаболомным профилям форм (генотипов) Ae. tauschii, устойчивых и неустойчивых к грибным патогенам.
Introduction

Wheat (Triticum L.) is one of the most significant crops in the world, including the Russian Federation. For the majority of the world’s population, it is one of the staple foods. The yield and quality of wheat largely depend on the resistance of cultivars to environmental stress factors, including fungal diseases. Most of the cultivated foreign and domestic cultivars are susceptible to diseases caused by agents of stem rust (Puccinia graminis) and mildew (Blumeria graminis (DC.) Speer f. sp. tritici Marchal.) and septoria leaf blotch (Mycosphaerella graminicola (Fuckel) J. Schroet. (=Septoria tritici; Phaeosphaeria nodorum (E. Muell.) Hedjar. (=Leptosphaeria nodorum (E. Muell., =Septoria nodorum (Berk.)). Crop losses can reach up to 40 % (Afanaseko, 2010; Kolomiets et al., 2017). The creation of wheat cultivars resistant to the most harmful fungal pathogens is one of the effective ways to combat them.

Wild relatives of cultivated forms of wheat, rye, barley, oats, etc. serve as an inexhaustible source of resistance genes for creating cultivars combining high yields and resistance to environmental factors. Evolutionary, Aegilops L. species are close to those of the genus Triticum L. (Dorofeev, 1971; Migushova, 1975, Konarev, 1980; Liu et al., 2015; Arora et al., 2012). However, the donors used in breeding programs, in most cases are characterized by the same resistance genes. With time, it leads to the appearance of “adapted” forms of pathogens that infect cultivars previously considered to be resistant. Many known resistance genes from Ae. tauschii Coss. are not used in practice to improve wheat cultivars, as their protective effect is considered low (Pretorius, 1997; Kolmer, Anderson, 2011). Expansion of the genetic diversity of donors of resistance to wheat fungal diseases will help breeders solve this problem, and the global collection of wheat wild relatives at the N.I. Vavilov All-Russian Research Institute of Plant Industry (VIR) plays a crucial role in this task (Vavilov, 1919).

The VIR collection of the genus Aegilops L. contains over 5,000 accessions of various ecological and geographical origins, and it includes thirteen diploid, ten tetraploid, and five hexaploid species. Since 1956, species possessing complex immunity to fungal diseases have been identified in the collection: diploid Ae. mutica Boiss., Ae. speltoides Tausch., Ae. aecheri Boiss., Ae. bicornis (Forsk.) Jaub. et Spach., Ae. comosa Sibth. & Sm., Ae. uniaristata Vis., Ae. heldreichii Hozm.; tetraploid Ae. ovata L., Ae. triaristata Wild., Ae. ventricosa Tausch., Ae. variabilis Eig. Such a diversity of genetic material makes it possible to select appropriate genotypes for the subsequent production of wheat cultivars with improved biological characteristics. Ae. tauschii is a carrier of D genome, which is close to the polyploid wheat genome, and this fact greatly facilitates the crossing of Ae. tauschii with common wheat when transmitting effective disease resistance genes (Dobrotvorskaia et al., 2017). Besides, flour of such cultivars has high baking quality (Semenova et al., 1973). By now, most of the effective genes that determine resistance to rust and wheat spot blotch agents have been successfully introgressed into the genome of T. aestivum L. (McIntosh et al., 1995; Mujeeb-Kazi et al., 2001; Yang et al., 2003; Adonina et al., 2012).

Recently, nonspecific metabolomic analysis has found wide application for plant species phenotyping and resistance studies, which provides a unique opportunity to scan a wide range of compounds that make up the metabolomic profile in the source material and give an objective assessment (using metabolic markers) of the plant’s response to environmental factors (Konarev et al., 2015). This approach is increasingly used to identify individual metabolites or their groups that can characterize the protective status of the studied object, which makes it possible to identify accessions resistant to environmental stressors (Taji et al., 2006; Chakraborty, Newton, 2011; Valitova et al., 2016; Loskutov et al., 2017). Currently, metabolomic profiles of various crops from the VIR collection are investigated. Wild and cultivated forms of oats resistant and susceptible to Fusarium were studied, and significant differences among them were shown for a number of compounds (acylglycerols, nonproteinogenic amino acids, galactinol, etc.) (Loskutov et al., 2017, 2019). Total screening of wild forms of various crops will make it possible to shape a model “metabolomic profile of a resistant cultivar”.

The aim of this study was the identification of metabolic markers to be used in screening the genetic diversity of wild relatives for the forms with effective resistance genes, which can be efficiently introduced into the common wheat genome.
for the use in breeding programs. The research objectives included the study of metabolomic profiles of *Ae. tauschii* accessions with and without resistance to leaf rust and powdery mildew pathogens, in order to detect metabolites marking the susceptibility of *Ae. tauschii* to fungal pathogens. The results of the study can shorten and optimize the selection of source material when creating wheat cultivars highly resistant to fungal pathogens.

**Materials and methods**

Fifty *Ae. tauschii* accessions from the VIR collection (see the Table), grown at the Dagestan Experimental Station of VIR (DES VIR) in 2017 and harvested at full ripeness, were used as the material for the study. The sample was composed from the main of *Ae. tauschii* botanical varieties in the VIR collection, taking into account the most complete ecogeographical representation.

Field studies of *Ae. tauschii* accessions were conducted at DES VIR on irrigated 1-m² plots according to the method accepted in VIR (Merezhko et al., 1999). During the growing season, the air temperature averaged +20.4 °C, the amount of precipitation was 15.4–16.3 mm, and the total of active temperatures amounted to 3400–4500 °С. Field assessment of the infection of *Ae. tauschii* accessions by fungal pathogens was carried out in fields of DES VIR, and the laboratory evaluation of the degree of infection was carried out at the VIR Department of Genetics in compliance with methods employed at VIR (Tyryshkin et al., 2004). Resistance was determined on a 9-point scale, where 9 means the absence of disease symptoms or presence of small necrotic spots; 7, microscopic pustules surrounded by necrotic zone; 5, small pustules surrounded by a wide necrotic zone; 3, medium-size pustules surrounded by chlorotic tissue; and 1, large pustules forming continuous lesion zones. Plants scoring 9–7 were classified as resistant, and those with 5–1 points as susceptible. Plants were examined at 5-day intervals throughout the growing season. For each accession, an integral score was derived from the highest infection points (see the Table).

Metabolomic profiles of grain from *Ae. tauschii* accessions were studied at the Department of Biochemistry and Molecular Biology of VIR (in 5 biological and 3 analytical replications) (Loskutov et al., 2017). The grains were cleaned of glumes and ground; 50 mg of the flour of an accession were homogenized (Loskutov et al., 2017). The grains were cleaned of glumes and ground; 50 mg of the flour of an accession were homogenized by mortar and pestle, and the sample was kept at −20 °C for 3 days. A 100-µL portion of the extract was evaporated to dryness using a Labconco CentriVap Concentrator (USA). The dry residue was silylated using bis(trimethylsilyl)trifluoroacetamide at 100 °C for 40 minutes. The separation of trimethylsilyl ethers of metabolites was carried out using an HP-5MS 5 % phenyl–95 % methylpolysiloxane capillary column (30.0 m, 250.00 µm, 0.25 µm) on an Agilent 6850 gas chromatograph with an Agilent 5975B VL quadrupole mass selective detector MSD (Agilent Technologies, USA). The analysis was performed at 1.5 mL/min inert gas flow rate through a column. The column was heated from +70 up to +320 °C at 4 °C/min heating rate. The temperature of the mass spectrometer’s detector was +250 °C, and that of the injector was +300 °C. The injected sample volume was

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**Leaf disease resistance in *Ae. tauschii* Coss. accessions**

| VIR Catalog No. | Subspecies (ssp.), variety (var.) | Mildew | Rust | brown | yellow |
|----------------|-----------------------------------|--------|------|-------|--------|
| k-101          | var. typica                       | 9      | 3    | 3     |
| k-108          | ssp. strangulata                  | 9      | 7    | 7     |
| k-112          |                                   | 9      | 9    | 9     |
| k-113          |                                   | 9      | 9    | 9     |
| k-291          |                                   | 9      | 3    | 7     |
| k-315          |                                   | 9      | 9    | 3     |
| k-336          |                                   | 9      | 9    | 3     |
| k-338          |                                   | 9      | 3    | 3     |
| k-340          |                                   | 9      | 3    | 3     |
| k-520          |                                   | 9      | 9    | 9     |
| k-617          |                                   | 9      | 5    | 3     |
| k-1098         | var. typica                       | 9      | 5    | 9     |
| k-1102         | var. meyeri                       | 9      | 9    | 9     |
| k-1111         | ssp. strangulata                  | 9      | 9    | 9     |
| k-1155         |                                   | 9      | 9    | 1     |
| k-1723         |                                   | 9      | 9    | 9     |
| k-1783         | var. meyeri                       | 9      | 3    | 3     |
| k-527          | ssp. strangulata                  | 9      | 9    | 5     |
| k-1520         | var. typica                       | 9      | 9    | 7     |
| k-2271         |                                   | 9      | 9    | 7     |
| k-3187         |                                   | 9      | 9    | 9     |
| k-994          | var. typica                       | 9      | 3    | 3     |
| k-1619         |                                   | 9      | 3    | 5     |
| k-1659         |                                   | 9      | 9    | 9     |
| k-1740         |                                   | 9      | 9    | 9     |
| k-608          | var. meyeri                       | 9      | 3    | 3     |
| k-1216         |                                   | 9      | 9    | 9     |
| k-1022         | var. typica                       | 9      | 5    | 3     |
| k-1770         | ssp. strangulata                  | 9      | 9    | 9     |
| k-1657         | ssp. strangulata                  | 9      | 3    | 3     |
| k-1662         |                                   | 9      | 9    | 9     |
| k-1958         | var. typica                       | 9      | 9    | 9     |
| k-1966         | ssp. strangulata                  | 9      | 9    | 7     |
| k-4043         |                                   | 9      | 9    | 7     |
| k-4049         |                                   | 9      | 9    | 7     |
| k-4056         |                                   | 9      | 9    | 7     |
| k-967          | var. meyeri                       | 9      | 5    | 1     |
| k-677          |                                   | 9      | 5    | 1     |
| k-553          | var. meyeri                       | 9      | 3    | 1     |
| k-560          |                                   | 9      | 3    | 1     |
| k-1172         |                                   | 9      | 9    | 1     |
| k-1498         |                                   | 9      | 1    | 3     |
| k-494          | ssp. strangulata                  | 9      | 9    | 7     |
| k-497          |                                   | 9      | 9    | 9     |
| k-4564         | ssp. strangulata                  | 9      | 3    | 3     |
| k-423          | ssp. strangulata                  | 9      | 7    | 3     |
| k-394          | var. meyeri                       | 9      | 5    | 3     |
| k-396          |                                   | 9      | 3    | 3     |
| k-663          |                                   | 9      | 1    | 3     |
1.2 μL. Pyridine solution of tricosan (1 μg/μL) served as the internal standard.

The results were processed using UniChrom and AMDIS software. The peaks were identified using the NIST 2010 mass spectra library and as well as libraries of the St. Petersburg University Research Park and of the V.L. Komarov Botanical Institute of the Russian Academy of Sciences (Puzanskiy et al., 2015). The biochemical parameter values are given in ppm (μg/g).

Statistical data were evaluated with the Statistica 7.0 software package. The initial set of characters was screened by the method of one-way analysis of variance to identify the characters (metabolites) whose contents reliably discriminated Ae. tauschii accessions resistant and susceptible to the studied pathogens. The discriminant analysis was used to assess informativeness of resistance characters (metabolites) of Ae. tauschii accessions.

Results

Metabolomic profiles of caryopses of Ae. tauschii accessions with and without resistance to fungal pathogens differed in several indicators. Higher contents of organic acids were observed in the resistant forms of Ae. tauschii (1190 ppm) compared with the susceptible ones (1090 ppm). The dominating organic acids in the metabolomic profile of Ae. tauschii caryopses were malic and methylmalonic acids (203 vs. 164 and 168 vs. 153 ppm, respectively). The caryopses of resistant and susceptible forms were found to contain, respectively, the following contents of organic acids (ppm): 102 and 79 galacturonic acid, 78 and 76 lactic, 43 and 48 gulonic, 25 and 44 glucolic, 30 and 47 azelaic, 28 and 27 oxalic, 42.8 and 41.0 fumaric, 19.9 and 13.0 ribonic, and 11.5 and 10.0 glyceric. The sum of minor acids with concentrations no higher than 10 ppm each was 24.0 ppm in both resistant and susceptible forms of Ae. tauschii. All metabolic reactions in plants occur with the participation of phosphoric acid and its derivatives, which explains its rather high content in the studied accessions (320.2 in resistant and 277.0 ppm in susceptible forms), the amounts of methylphosphate being 64.0 and 56.0 ppm, respectively.

Three nonproteinogenic amino acids constituted 60 % of free amino acids in resistant and susceptible Ae. tauschii accessions: 3-hydroxyipipolic (137.2 and 116.6), pipolic (0.5 and 0.3), and 5-hydroxyipipolic (0.7 and 0.7 ppm, respectively). The remaining amino acids were represented by essential (valine, isoleucine, threonine, phenylalanine, and tryptophan) and non-essential ones (α-alanine, glycine, serine, proline, hydroxyproline, asparagine, glutamine, tyrosine, aspartic acid, and glutamic acid). The dominating ones were asparagine (18.7 and 13.4 ppm), valine (15.6 and 16.0), α-alanine (10.6 and 13.1), glutamine (12.3 and 9.0) and glutamic acid (5.7 and 5.5 ppm), respectively. The amounts of the remaining amino acids did not exceed 4.0 ppm. The sums of free amino acids (except nonproteinogenic) did not differ significantly between the studied forms of Aegilops resistant and susceptible to fungal pathogens: 88.4 and 85.3 ppm, respectively.

Higher concentrations of polyols and phytosterols were recorded in the susceptible forms of Ae. tauschii (341.8 and 336.5 ppm, respectively). Glycerol, xylitol, dulcitol, myo-inositol and inositol derivatives (59.5, 84.4, 70.4, 23.4, 9.6 ppm, respectively) prevailed in caryopses of the susceptible forms, while galactol (92.5 ppm) dominated in the resistant ones. Phytosterols were mainly represented by sitosterol, stigmasterol, and campesterol; their contents were 219.8, 85.8, and 30.9 ppm in the susceptible forms, and 160.3, 44.5, and 22.7 ppm in the resistant ones, respectively.

The dominating fatty acids in caryopses of both resistant and susceptible forms of Ae. tauschii were found to have higher contents of acylglycerols (954.0 vs. 745.6 ppm), mainly due to diacylglycerol (DAG) amounting to 631.0 and 464.0 ppm, respectively. The amounts of monoacylglycerols, i.e., MAG-2 C18:3 and MAG-1 C18:1, were also higher in resistant forms: 188.0 vs. 152.7, 54.0 vs. 34.2 ppm. On the contrary, the content of MAG-1 C16:0 was higher in the susceptible forms: 94.7 vs. 81.0 ppm in resistant.

Monosugars in Ae. tauschii caryopses were represented mainly by hexoses (over 80 %). These values were higher (1164.8 ppm) for resistant forms of Aegilops than for susceptible ones (1026.7 ppm). Of the hexoses, glucose (819.8) was predominant in resistant forms, while in the susceptible ones these were glucose (455.4) and fructose (318.1 ppm). The total pentose contents (ribose and xylose) did not exceed 40 ppm. The contents of glycerol-3 phosphate did not differ significantly between resistant and susceptible forms of Aegilops. They amounted to 42.8 and 37.4 ppm, respectively.

Oligosaccharides in caryopses of resistant and susceptible forms of Ae. tauschii amounted to 13480.6 and 14920.0 ppm, respectively. They were mainly represented by sucrose and raffinose. The content of sucrose was higher in susceptible forms (11604.9), while that of raffinose was higher (4882.0 ppm) in resistant ones.

Derivative sugars, identified as methyl-D-galactopyranoside, were found in both resistant and susceptible forms of Ae. tauschii (169.2 and 84.4 ppm, respectively).

In the group of phenolic compounds, the contents of hydroquinone (93.6 in resistant and 74.0 ppm in susceptible accessions) were found to be the highest. The values for kempferol; pyrogallic, 2,3-dihydroxybenzoic, salicylic, and caffeic acids; and α-tocopherol were 29.1, 1.5, 0.2, 0.3, 0.4, 0.4 ppm in resistant and 14.0, 1.3, 0.2, 0.2, 0.4 and 0.3 ppm in susceptible forms.

The above data reflect the activity of metabolic processes in Ae. tauschii caryopses. This activity characterizes primary and secondary metabolism, i.e., exchange of nitrogen-containing compounds, including amino acids; the Krebs cycle; carbohydrate metabolism; glycolysis; pentose phosphate cycle; exchange of signal (inositol) compounds; shikimate and glyoxylate pathways; etc.

Statistical processing of the data showed that the metabolome of susceptible forms of Ae. tauschii differs with varying degrees of significance from that of resistant ones in a number of indicators. Resistant forms of Ae. tauschii were divided into
The results of our work show that the metabolomic profiles of resistant and susceptible forms of *Ae. tauschii* differ significantly. When using the metabolomic analysis data, there is a high probability (up to 98 %) of identifying forms resistant to fungal pathogens among the accessions taken into the study without additional tests. We recommend that this approach be used to optimize the breeding process.

Analysis of metabolomic data for resistant and susceptible forms of *Ae. tauschii* allows for a substantiated conclusion about the degree and nature of fungal pathogen influence on the main stages of primary and secondary metabolism in caryopses of accessions differing in the degree of resistance. To a greater or lesser degree, almost all pathogens affected metabolic processes, i.e., the Krebs cycle; glycolysis; and metabolism of fatty acids, acylglycerols, polyols, phytosterols, mono- and oligosaccharides. With all this, fungal pathogens had practically no effect on the contents of free amino acids (except for threonine and tyrosine). Since tyrosine is a precursor in the synthesis of many bioactive compounds that perform various functions – structural (lignin), protective (phenolic compounds, alkaloids, etc.), transport (electron transfer), and others – the effect of pathogens on the content of this amino acid in a caryopsis may be due to activation of defense mechanisms in response to the penetration of the pathogen into plant tissues, as confirmed by studies conducted outside Russia (Schenck, Maeda, 2018). Changes in the content of another amino acid, threonine, are associated by a number of authors with the influence of adverse biotic environmental factors, for example, insect pests. A plant reduces the concentration of substances necessary for the nutrition of the parasite, thereby affecting its population. The same mechanism may also act in cases of leaf rust and powdery mildew pathogens (Gonzales-Vigil et al., 2011). Azelaic acid, a product of oleic acid oxidation, and pipopecolic (lysine catabolites) are intensely produced in response to the invasion of pathogens, in particular, fungi, into plant tissues; therefore, changes in the concentration of these compounds are well justified and
confirmed by other researchers (Navarova et al., 2012; Zoeller et al., 2012).

Among free phenolic compounds, a significant effect on the resistance to fungal pathogens was detected for hydroquinone (the dominant compound of this group) and pyrogallic acid. Phenolic compounds are known to be actively involved in the formation of plant immunity. The presence of free forms of phenolic compounds most often indicates intensity of glycoside synthesis, where they function as aglycones. According to our data, mainly hydroquinone and pyrogallic acid accumulate in the caryopsis of resistant forms of Ae. tauschii. Their presence in the free state is most likely associated with the destruction of active forms, glycosides, exemplified by abutin. As for pyrogallic acid, its accumulation may also be due to the plant/fungal pathogen interaction (the role of signaling substances) (Seigler, 1998; cit.: Gilbert, 2001).

The data from the present study confirm those we obtained earlier investigating the relationship between the resistance of oat (Avena sativa L.) forms to the Fusarium and such components of the metabolomic profile as acylglycerols, nonproteinogenic amino acids, and galactinol (Loskutov et al., 2019).

**Conclusion**

The results of our study confirm the pertinence and effectiveness of the use of nonspecific metabolomic analysis for the search for and identification of plant forms with a set of compounds proposed as markers of resistance to certain pathogens. Ae. tauschii accessions with high contents of pipecolic acids, acylglycerols, galactinol, stigmasterol, maltose, tyrosine, sorbose, glycerol, azelaic and pyrogallic acids, methyl-D-galactopyranoside, etc. can be included in breeding programs for cultivars of main cereal crops with high resistance to mildew, brown rust, and yellow rust.

**References**

Adonina I.G., Petrah N.V., Timonova E.M., Christov Yu.A., Salina E.A. Construction and study of leaf rust resistant common wheat lines with translocations of Aegilops speltoides Tausch. genetic material. Russ. J. Genet. 2012;48:404-409. DOI 10.1134/S1027295412020020.

Afanasenko O.S. Problems of creating cultivars with long-term disease resistance. Zaschita i Karantin Rastenii = Plant Protection and Quarantine. 2010;3:4-10. (in Russian)

Arora S., Singh N., Kaur S., Bains N.S., Uaey C., Poland J., Chupea A-L., Blinova E.V.-B., Konarev A.V., Khoreva V.I., Shavarda A.L., Blinova E.V., Gnutikov A.A. Biochemical aspects of interactions between fungi and plants: a case study of Fusarium in oats. Selskokhozyaystvennaya Biologiya = Agricultural Biology. 2019;54(3):575-588. DOI 10.10538/agrobiology.2019.3.575eng.

Loskutov I.G., Shelenga T.V., Konarev A.V., Shavarda A.L., Blinova E.V., Dzubenko N.I. The metabolomic approach to the comparative analysis of wild and cultivated species of oats (Avena L.). Russ. J. Genet.: Appl. Res. 2017;7(5):501-508. DOI 10.1134/S2079059717050136.

Mcintosh R.A., Wellings C.R., Park R.F. Wheat Rust: An Atlas of Resistance Genes. Australia: CSIRO Publ., 1995.

Merezhko A.F., Udachin R.A., Zuev E.V., Filatenko A.A., Serbin A.A., Liapunova O.A., Kosov V.I., Kurkiev U.K., Okhotnikova T.V., Navruzbekov N.A., Boguslavskii R.L., Abdulaev A.K., Chikha N.N., Mitrofanova O.P., Potokina S.A. Enriching, Con­educting, and Con­erving, and Studying the Global Collections of Wheat, Aegilops, and Triticale. St. Petersburg, 1999. (in Russian)

Migushova E.F. On the origin of wheat genomes. Trudy po Priklad­noy Botanike, Genetike i Selektssi = Proceedings on Applied Botany, Genetics, and Breeding. 1975:55(3):3-26. (in Russian)

Mujeeb-Kazi A., Cano S., Rosas V., Cortes A., Delgado R. Regis­tration of five synthetic hexaploid wheat and seven bread wheat lines resistant to wheat spot blotch. Crop Sci. 2001;41(5):1653-1654. DOI 10.2135/cropsci2001.4151653x.

Navarova H., Bernsdorff F., Doring A.C., Zeier J. Pipecolic acid, an endogenous mediator of defense amplification and priming, is a critical regulator of inducible plant immunity. Plant Cell. 2012;24(12):5123-5141. DOI 10.1105/tpc.112.103564.

Pretorius Z.A. Detection of virulence to Lr41 in a South African pathotype of Puccinia recondita f. sp. tritici. Plant Dis. 1997;81(4):423. DOI 10.1094/PDIS.1997.81.4.423A.

Puzanskiy R.K., Shavarda A.L., Tarakhovskaya E.R., Shishova M.F. Analysis of metabolic profile of Chlamydomonas reinhardii cultured under autotrophic conditions. Appl. Biochem. Microbiol. 2015;51(1):83-94. DOI 10.1134/S0003683815010135.

Schenck C.A., Maeda H.A. Tyrosine biosynthesis, metabolism, and catabolism in plants. Phytochemistry. 2018;149:82-102. DOI 10.1016/j.phytochem.2018.02.003.
Metabolomic approach to search for fungal resistant forms of \textit{Aegilops tauschii} Coss. from the VIR collection

Semenova L.V., Migushova E.F., Devyatkin E.P. Grain quality of \textit{Aegilops} grain, ancestor of wheat. Trudy po Prikladnoy Botanike, Genetike i Selektii = Proceedings on Applied Botany, Genetics, and Breeding. 1973;50(1):216-226. (in Russian)

Taji T., Takahashi S., Shinozaki K. Inositol and their metabolism in abiotic and biotic stress responses. Subcell. Biochem. 2006;39:239-264. DOI 10.1007/0-387-27600-9_10.

Tyrshkin L.G., Kolesova M.A., Chikida N.N. \textit{Aegilops tauschii} Coss. Characteristics of accessions for juvenile leaf disease resistance. In: Catalogue of the VIR Global Collection. St. Petersburg, 2004;763:3-15. (in Russian)

Valitova J.N., Sulkarmayeva A.G., Minibayeva F.V. Plant sterols: diversity, biosynthesis, and physiological functions. Biochemistry (Moscow). 2016;81(8):819-834. DOI 10.1134/S0006297916080046.

Vavilov N.I. Plant Immunity to Infectious Diseases. Moscow, 1919. (in Russian)

Yang W.-Y., Yu Y., Zhang Y., Hu X.-R., Wang Y., Zhou Y.-C., Lu B.-R. Inheritance and expression of stripe rust resistance in common wheat (\textit{Triticum aestivum}) transferred from \textit{Aegilops tauschii} and its utilization. Hereditas. 2003;139:49-55. DOI 10.1111/j.1601-5223.2003.01671.x.

Zoeller M., Stingl N., Krischke M., Fekete A., Wallter F., Berger S. Lipid profiling of the \textit{Arabidopsis} hypersensitive response reveals specific lipid peroxidation and fragmentation processes: biogenesis of pimelic and azelaic acid. Plant Physiol. 2012;160(1):365-378. DOI 10.1104/pp.112.202846.

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