Genetic differences among the four *Stipa* species endangered and protected in Poland

Maria Krzakowa*, Marcin Michalak & Maria Judek

Department of Genetics, Adam Mickiewicz University, Umultowska 89, 61-614 Poznań, Poland, *e-mail: krzakowa@amu.edu.pl

Abstract: The distinct character of four species of *Stipa* genus, *S. borysthenica*, *S. capillata*, *S. joannis* and *S. pulcherrima* was manifested in the variability of three enzyme systems, including glutamate-oxalacetate transaminase (GOT), esterase (EST) and peroxidase (PX). The studies were conducted on two-month-old seedlings cultured under the same glasshouse conditions, obtained from seeds from the Botanical Gardens collection. Fifteen loci were described, of which eight proved to be polymorphic. The studies included also the complex species of *S. pennata* and the closely related species of *S. tirsa*.

Key words: Stipa borysthenica, S. capillata, S. joannis, S. pulcherrima, S. pennata, S. tirsa, enzyme, electrophoresis

1. Introduction

In Poland occur four species of the Stipa genus, including S. borysthenica Klokov, S. capillata L., S. joannis Čelak and S. pulcherrima K. Koch. All belong to the group of grasses covered by strict protection and, being endangered, they are included in the Polish Red Book of Plants (Kaźmierczakowa & Zarzycki 2001). Populations of *Stipa* species in Poland have a low frequency due to human activity as they were used them for decoration due to the exceptionally aesthetic habit of their panicles (Kozłowski 2002). For this reason, the species in some regions are endangered by extinction (Żukowski & Jackowiak 1995). Occurrence of the species is linked to xerothermic grasses of a steppe type in the Festuco-Brometea class. These may change their character both in a natural way and due to spontaneous succession of bushes and trees which leads to shadowing and disappearance of heliophilic species (including those of Stipa genus). It may also be due to human activity in the form of excessive pasturage or effects of eutrophic waters flowing down from the neighbouring fields under cultivation (this pertains particularly S. joannis). The increasingly rare biotopes of Stipa genus species are protected in preserves which play the role of living gene banks. The most frequently protected are species of *S*. capillata and S. joannis (Piękoś-Mirkowa & Mirek 2002).

S. borysthenica grows in Poland very infrequently. The species is known to occupy only five xerothermic

stands along the lower Odra River and in Wielkopolska (Ceynowa-Giełdon 2001a). S. capillata is a species typical for the Potentillo-Stipetum capillatae community. It grows in small populations along Lower Odra River, in northwestern Poland, in the Toruń-Eberswalde proglacial streat valley, in Nida Lowland, Małopolska and Lubelska Highlands (Filipek 1974; Górska-Zajączkowska & Weglarski 1993; Piekoś-Mirkowa & Mirek 2002; Towpasz & Mitka 2001). Individual stands of S. joannis can be encountered mainly along the lower Vistula River, in the Notecka proglacial stream valley, in Warta valley, and along the lower Odra River (Górska-Zajączkowska et al. 1989; Michalik 1991; Majtkowska & Majtkowski 2004). In the north of Poland the species forms the Potentillo-Stipetum community. S. pulcherrima is a very rare species in Poland, endangered by extinction. Along the lower Odra River it composes the Linosyridi-Stipetum pulcherrimae complex. It is thought to represent a typical species of xerothermic grasses of the *Festucetalia-valesiaceae* order, together with S. joannis (Matuszkiewicz 2001). It occupies a few stands by the Odra River and on the Sandomierska Highland (Ceynowa-Giełdon 2001b).

The progression of the degradation of biotopes in recent years, constrains botanists to undertake additional measures to preserve the gene pool of endangered species using *ex situ* culture in botanical gardens. Taking into account that inter-specific differences, according to biochemical character, were not investigated earlier

[©] Adam Mickiewicz University in Poznań (Poland), Department of Plant Taxonomy. All rights reserved.

in the genus *Stipa*, we have decided to take advantage of such collections for this preliminary study.

2. Material and methods

The species were represented by the following samples of seeds:

S. borysthenica – 2 samples: Botanical Garden, Adam Mickiewicz University; the seeds originated from exchange with Botanical Garden in Budapest;

S. capillata – 10 samples: Botanical Garden, Adam Mickiewicz University – the seeds originated from populations in the vicinity of Węgrzyce near Gorzów, from the Ojcowski National Park and St. Laurent's Mountain, and from the Botanical Garden, Frankfurt am Main; Botanical Garden, University of Łódź; Botanical Garden, University of Marie Skłodowska-Curie in Lublin – the seeds originated from natural populations in Wiślica and from an exchange program; Botanical Garden, University of Warsaw; Botanical Garden, Jagiellonian University in Cracow;

S. joannis – 4 samples: Botanical Garden, Adam Mickiewicz University – the seeds originated from collection of our own, from the Barbarka population near Toruń and Skorocice population near Busko; Botanical Garden, Institute of Plant Culture and Acclimation in Bydgoszcz: the seeds originated from Choragiewki population;

S. pennata – 2 samples: Botanical Garden, University of Warsaw; Botanical Garden, Adam Mickiewicz University: the seeds originated from Frankfurt am Main;

S. pulcherrima – 2 samples: Botanical Garden, Institute of Plant Culture in Bydgoszcz – the plants were developed from seeds obtained by exchange with the Botanical Garden in Berlin-Dahlem; Botanical Garden, Adam Mickiewicz University: seeds of plants originating from Ukraine;

S. tirsa - 2 samples: Botanical Garden, Adam Mickiewicz University: the seeds originated from exchange with the Botanical Garden in Berlin and our own collection.

Seedlings originating from Botanical Gardens seed collections were cultivated in identical glasshouse conditions. A crude extract of individual plants was subjected to electrophoresis in 11% starch gel in

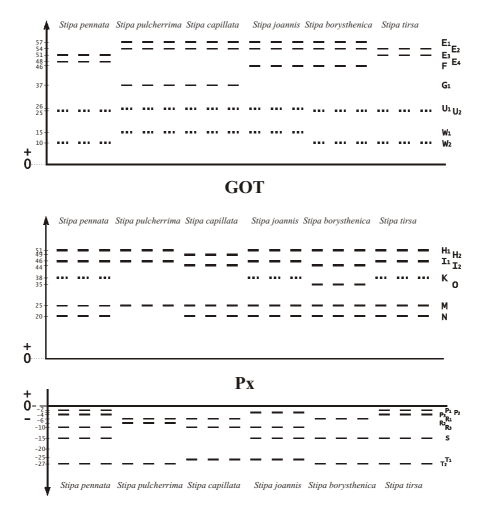


Fig. 1. Schematic diagrams of electrophoretically detected allozymes for the examined 6 species



lithiumborate buffer system, pH 8.3. The gels were stained specifically (Show-Prasad 1970) for three enzymes: glutamate-oxalacetic transaminase (GOT, EC.2.6.1.1), esterase (EST, EC 3.1.1.2), peroxidase (PX, EC 1.11.1.7). Interspecific differences were illustrated by Sphi coefficient (Leuschner 1974), principal component analysis and aglomerative clustering using the unweighted-pair-group method (UPMGA). In this way, six species were compared, using the nomenclature of the Botanical Gardens from which the seeds came.

3. Results and discussion

For each enzyme system, band patterns on the gels permitted the establishment of separate loci and their allozymes (Fig. 1).

GOT. Assuming that, similarly to other plants, the glutamate-oxalacetic transaminase is a dimer (Hillis & Moritz 1990), the two-banded phenotypes reflect expression in two separate loci. Each of bands containing two allozymes (in cases of H and I loci) are

documented by separate, single-banded loci as well as for M and N (showing no hybrid bands).

EST. The band pattern pointed to the monomeric character of the enzyme. The five distinguished loci well characterized the species. The most polymorphic proved to be the E locus with four alleles. At this locus the species demonstrated fixation of the heterozygous condition, frequently termed ,,the fixed heterozygosity" which, nevertheless, was typical for the species. Thus, four species (S. pulcherrima, S. capillata, S. joannis and S. borysthenica) demonstrated E1E2 phenotype, S. tirsa showed E2E3 phenotype while S. pennata demonstrated E3E4 phenotype or differed from each other even if, as closely related, they shared the common E3 allele. It should be recalled that S. pennata as a morphologically variable species served as a common species for a few other species the taxonomic identity of which underwent periodic modifications. Material for our studies originated from botanical garden collections, included small numbers of seeds and, therefore, monomorphism of the samples. Even if they

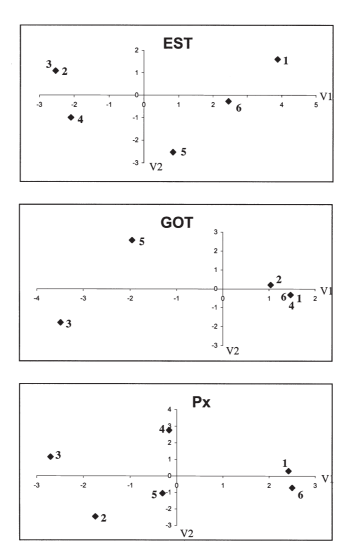


Fig. 2. Scatter diagram for the species investigated on the plane of the first two axes of the Principal Components. 1 - S. *pennata*, 2 - S. *pulcherima*, 3 - S. *capillata*, 4 - S. *joannis*, 5 - S. *borysthenica*, 6 - S. *tirsa*

are heterozygotic, they might occasionally result from their original form, a single plant. Ambiguities of the type may be resolved only by appropriate population studies.

Persistence of "fixed heterozygosity" might reflect also the biology of the studied species since xerophytic groups of species: the first including *S. pennata* and *S. tirsa* with *S. borysthenica* attached to them and the other formed by *S. pulcherrima* and *S. joannis*, while *S. capillata* proved its separate character.

In all the schemes a noticeable tendency appears for links between *S. pennata* (1) and *S. tirsa* (6) samples.

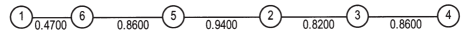


Fig. 3. Dendrite of the shortest connections between *Stipa* species constructed on the basis of the summarised data obtained from all enzyme systems Explanations: 1 - S. *pennata*, 2 - S. *pulcherima*, 3 - S. *capillata*, 4 - S. *joannis*, 5 - S. *borysthenica*, 6 - S. *tirsa*

tuft grasses of genera such as *Stipa* and of other stable species of grasses with the patchy growth, e.g., those of *Bromus* genus, are mostly autogamic due to the presence of cleistogamic flowers (Stebbins 1958).

PX. Band patterns indicated that, similarly to other grasses (Krzakowa & Kraupe 1981; Krzakowa 1996; Krzakowa & Mikulski 1997; Krzakowa *et. al.* 2003, 2005) including the studied species of *Stipa* genus, the enzyme system shows its monomeric behaviour. Two loci could be distinguished, each with three alleles, including loci P and R as well as locus T with two allozymes. Locus S, not detected in studied samples of *S. capillata* and *S. pulcherrima*, in the remaining species was present in the form of a single band.

In order to illustrate reciprocal relationships between the species, the technique of principal components was applied for individual enzymes (Fig. 2). In respect to esterases (EST), *S. pennata*, *S. borysthenica* and *S. tirsa* formed a single species group. The most pronounced similarity was manifested by two species: *S. pulcherrima* and *S. capillata*, which occupied the same space, while *S. joannis* was quite distinct.

In respect to glutamate-oxalacetate-transaminase (GOT), here species including *S. pennata*, *S. joannis* and *S. tirsa*, proved similar. Worth noting, *S. capillata* and *S. borysthenica* manifested a distinct character. The species variability pattern in the space formed for peroxidases (PX) by the two principal components confirmed the distinct character of *S. capillata*, linked *S. pennata* and *S. tirsa* and pointed to a similar character of *S. pulcherrima*, *S. joannis* and *S. borysthenica*.

The total dendrite constructed for all the populations (Fig. 3) on the basis of the lowest values of the Sphi coefficient demonstrated close taxonomic similarity between *S. pennata* and *S. tirsa* while the other species are connected by similar taxonomic distances.

Grouping by the closest neighbourhood technique (UPGMA) pointed to phylogenetic relationships between the species (Fig. 4). It resulted in the formation of two Inclusion in our studies of samples representing *S*. *pennata* complex, which still persists in the world

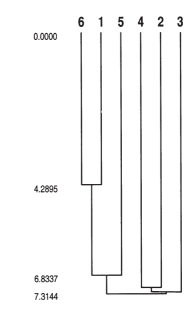


Fig. 4. Dendrogram depicting hierarchial structure of genetic relatedness among *Stipa* species

Explanations: 1 - S. pennata, 2 - S. pulcherima, 3 - S. capillata, 4 - S. joannis, 5 - S. borysthenica, 6 - S. tirsa

nomenclature (Strid & Tan 1991), allows the recollection that races of *S. joannis*, *S. tirsa* and *S. pulcherrima* have been isolated from the *S. pennata* species (Hegi 1906), and later reached the status of species (Hegi 1992). Also *S. borysthenica* has originated from *S. pennata* subsp. *sabulosa* (Rybcov 1972; Hegi 1992). The inter-specific differences demonstrated in the present study are sufficiently interesting to prompt further studies on natural populations as long as they are still available.

Acknowledgements. The authors are thankful to Mrs. Barbara Malchrowicz for her technical assistance. Scientific work financed from the resources earmarked for science in years 2005-2007 as Research Project no. 2 P04C 095 28.

References

- CEYNOWA-GIEŁDON M. 2001a. *Stipa borysthenica* Klokov -Ostnica piaskowa. In: R. KaźMIERCZAKOWA & K. ZARZYCKI (eds.). Polska czerwona księga roślin. Paprotniki i rośliny kwiatowe, wyd. 2, pp. 463-465. PAN, Instytut Botaniki im. W. Szafera, Instytut Ochrony Przyrody, Kraków.
- CEYNOWA-GIEŁDON M. 2001b. Stipa pulcherrima Koch Ostnica powabna. In: R. KaźMIERCZAKOWA & K. ZARZYCKI (eds.). Polska czerwona księga roślin. Paprotniki i rośliny kwiatowe, wyd. 2, pp. 459-460. PAN, Instytut Botaniki im. W. Szafera, Instytut Ochrony Przyrody, Kraków.
- FILIPEK M. 1974. Kserotermiczne zespoły murawowe nad dolną Odrą i Wisłą na tle zbiorowisk pokrewnych. Bad. Fizjogr. Pol. Zach. seria B-Botanika 27: 45-82.
- Górska-ZAJĄCZKOWSKA M. & WĘGLARSKI K. 1993. Ostnica włosowata *Stipa capillata* L. – rzadki i zagrożony gatunek flory północno-zachodniej Polski. Biuletyn Ogrodów Botanicznych, Muzeów i Zbiorów 2: 5-14.
- Górska-Zajączkowska M., Jańczyk J. & Węglarski K. 1989. Ostnica Jana (*Stipa joannis* Cel.). Rzadki i zagrożony gatunek flory Polski. Wiad. Bot. 33(3): 95-110.
- HEGI G. 1906. Illustrierte Flora von Mittel-Europa. Band I, pp. 203-206. Verlag von J. F. Lehmann, München.
- HEGI G. 1992. Illustrierte Flora von Mittel-Europa. Band I, Teil 3, pp. 397-426. Verlag Paul Parey, Berlin-Hamburg.
- HILLIS D. M. & MORITZ C. 1990. Molecular systematics. 588 pp. Sinauer Associates Inc. Publishers, Sunderland, Massachusetts, USA.
- KAŹMIERCZAKOWA R. & ZARZYCKI K. (eds.). 2001. Polska czerwona księga roślin. Paprotniki i rośliny kwiatowe, wyd. 2, 664 pp. PAN, Instytut Botaniki im. W. Szafera, Instytut Ochrony Przyrody, Kraków.
- KozŁowski S. 2002. Trawy w polskim krajobrazie. In: L. FREY (ed.). Polska Księga Traw, pp. 301-322. Instytut Botaniki im. W Szafera, PAN, Kraków.
- KRZAKOWA M. 1996. Genetic diversity of *Phragmites* australis (Cav)Trin.ex Stued. revealed by electrophoretically detected differences in peroxidases. In: C. OBINGER, U. BURNER, R. EBERMANN, C. PENEL & H. GREPPIN (eds.). Plant Peroxidases: Biochemistry and Physiology, pp. 184-189. University of Geneva.
- KRZAKOWA M., CELKA Z. & DRAPIKOWSKA M. 2005. Genetic variability of *Calamagrostis arundinacea* populations growing in *Calamagrostio arundinaceae-Quercetum petraeae* community. In: L. FREY (ed.) Biology of Grasses, pp. 23-30. W. Szafer Institute of Botany, Polish Academy of Sciences, Krakow.
- KRZAKOWA M., JAŃCZYK-WĘGLARSKA J. & ŚLIWIŃSKA E. 2003. Peroxidase polymorphism in *Calamagrostis epigejos* (Poaceae) indicating interspecific hybridization. In: Z. ZWIERZYKOWSKI, M. SURMA & P. KACHLICKI (eds.).

Application of Novel Cytological and Molecular Techniques in Genetics and Breeding of the Grass, pp. 109-114. Institute of Plant Genetics PAS, Poznań.

- KRZAKOWA M. & KRAUPE A. 1981. Isozyme investigations of natural populations of the cheatgrass (*Bromus tectorum* L.). Bot. Jahrb. Syst. 103(1-4): 393-399.
- KRZAKOWA M. & MIKULSKI W. 1997. Peroxidase as marker in pure lines of perennial ryegrass (*Lolium perenne* L.). In: Z. STASZEWSKI, W. MŁYNIEC & R. OSINSKI (eds.). Ecological aspects of breeding fodder crops and amenity grasses, pp. 322-326. Proceedings of 20th Meeting of EUCARPIA, Fodder and Ammenity Grass Section, Radzików, Poland, October 1996.
- LEUSCHNER D. 1974. Einführung in die numerische Taxonomie. 139 pp. Gustav Fischer Verlag, Jena.
- MAJTKOWSKA G. & MAJTKOWSKI W. 2004. Stanowisko ostnicy Jana (*Stipa joannis*) w Wielkich Lniskach k. Grudziądza. In: VI Ogólnopolskie Spotkanie Naukowe, Biologia Traw, p. 58. Kraków.
- MATUSZKIEWICZ W. 2001. Przewodnik do oznaczania zbiorowisk roślinnych Polski. In: J. B. FALIŃSKI (ed.). Vademecum Geobotanicum 3, 537 pp. Wyd. Nauk. PWN, Warszawa.
- MICHALIK S. 1991. Ochrona czynna stanowisk ostnicy Jana Stipa joannis Cel. na skale Jonaszówka w Ojcowskim Parku Narodowym. Prądnik. Prace Muz. Szafera 3: 175-181.
- PIĘKOŚ-MIRKOWA H. & MIREK Z. 2002. Zagrożenie i ochrona gatunkowa traw. In: L. FREY (ed.). Polska Księga Traw, pp. 209-234. Instytut Botaniki W. Szafera, Polska Akademia Nauk, Kraków.
- Ryвcov H. I. 1972. Opriedielitiel vysshih rastenij Krima. 44-45 pp. Nauka, Leningrad.
- SHOW C. R. & PRASAD R. 1970. Starch gel electrophoresis of enzymes, a compilation of recipes. Bioch Gen. 4: 297-320.
- STEBBINS G. L. 1958. Zmienność i ewolucja roślin. 252 pp. Wyd. Nauk. PWN, Warszawa.
- STRID A. & TAN K. 1991. Mountain flora of Greece. Vol. 2, pp. 825-830. Edinburgh University Press.
- TOWPASZ K. & MITKA J. 2001. Grasses in xerothermic grassland on the Proszowice Plateau, southern Poland. In: L. FREY (ed.). Studies on Grasses in Poland, pp. 303-311. W. Szafer Institute of Botany, Polish Academy of Sciences, Kraków.
- ŻUKOWSKI W. & JACKOWIAK B. 1995. List of endangered and threatened vascular plants in Western Pomerania and Wielkopolska (Great Poland). In: W. ŻUKOWSKI & B. JACKOWIAK (eds.). Endangered and threatened vascular plants of Western Pomerania and Wielkopolska. Publications of the Department of Plant Taxonomy of the Adam Mickiewicz University of Poznań 3: 9-96. Bogucki Wyd. Nauk., Poznań.