GENETIC DIVERSITY OF *GALIUM CRACOVIENSE* EHREND. (Rubiaceae) – THE POLISH ENDEMIC PLANT

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**ABSTRACT**

Genetic diversity of *Galiun cracoviense*, a narrow endemic species, limited to the small area in southern Poland and concentrated on Jurassic limestone outcrops near Częstochowa, was examined using the AFLP marker. Twenty-nine individuals from three spatially isolated populations were used for the study. AFLP analysis yielded 157 bands, of which 110 (70%) were polymorphic. The AMOVA analysis revealed a substantially higher variation within populations (89.35%) than among them (10.65%). Values of parameters describing population genetic diversity, such as Shannon index and gene diversity index estimated for each population, were highly similar. The results indicate a high level of genetic polymorphism as well as a high genetic similarity of the isolated populations of *G. cracoviense* and thus an unconstrained gene flow between them. Based on the results we conclude that additional demographic and genetic studies are necessary to monitor potential decrease of populations size resulting mainly from the mechanical destruction of plants and their habitats caused by intense tourism. Due to the small general range of occurrence, conservation should include the highest possible number of populations of *G. cracoviense*.

**KEY WORDS:** AFLP, *Galiun cracoviense*, gene flow, genetic diversity, isolated populations, narrow endemic species, glacial relicts, Poland.

**INTRODUCTION**

The theory of population genetics predicts that small, isolated populations should experience increased random genetic drift, inbreeding and reduced interpopulation gene flow (Wright 1969). This may in turn result in a depletion of genetic diversity within populations, increased genetic divergence among populations, inbreeding depression (Ellstrand and Elam 1993) and reduced fitness to adapt to changing environmental conditions. Such loss of genetic variability and additionally low levels of genetic diversity have been reported for many rare and/or endemic species (Ellstrand and Elam 1993) and have been considered a consequence of rarity (Gitzendanner and Soltis 2000). In this context endemic taxa, due to their restricted distribution, often small population size and narrow ecological amplitudes, are especially vulnerable to extinction (Piękośl-Mirkowa and Mirek 2005). On the other hand, endemic plants in historical time scale, have always been rare with naturally isolated and small populations because of restricted habitat requirements (Holderegger 1997) (e.g. rocky outcrops). Such species groups which are naturally rare in a specific area, occurring in small, isolated populations are defined as “old rare species” (Huenneke 1991; Oostermeijer et al. 1996). In contrast, “new rare species” are species which were formerly much more common in a particular area, and their populations only became smaller, less abundant, and more isolated because of human influence (Huenneke 1991).

Poland has a small number of endemic plant taxa, due to its geographic location and Quaternary geological history. The majority of endemic species occurs in southern Poland in the Carpathians and the Sudeten, which were not covered by the Scandinavian ice sheet during the Pleistocene glaciations. Although formed in the highest ranges of the Carpathians and Sudeten during successive glaciations, local glaciers did not cover entire ridges, which allowed many plant species to survive in situ. The isolated taxonomic position of some Western Carpathian endemic species indicates their advanced phylogenetic age and suggests they may have survived in the Carpathians not only the latest glaciation.

This situation is different in lowland Poland, repeatedly covered by ice sheets during successive Pleistocene glaciations. Only few endemic taxa, classified as neoendemic species, occur in this area. *Cochlearia polonica* E. Fröhl.
and *Galium cracoviense* Ehrend., occurring in the Kraków-
Częstochowa Upland (the Polish Jura) in southern Poland,
belong to the most widely recognized Polish lowland en
demic species.

The Kraków-Częstochowa Upland is a fairly low range
formed by Jurassic limestone, with maximum altitude 504
m a.s.l., stretching ca. 100 km between Częstochowa and
Kraków. It is one of the areas richest in vascular plant spe
cies in Poland. Particularly many species are associated
with natural saxicolous grasslands Festucetum pallentis
(Baba 2004; Mirek 2004), inhabited also by *Galium craco
viense*, the focus of this study. A number of dealpine spe
cies, such as *Allium montanum* F. W. Schmidt, *Festuca
pallens* Host, *Gymnocarpium robertianum* (Hoffm.) New
man, *Hieracium bifidum* Hornem., *H. caesium* (Fries)
Fries, *Polygala brachyphylla* Chodat, *Saxifraga paniculata*
Mill. and *Viola rupestris* F. W. Schmidt, that reach the lo
cal northern range limit on the limestone rocks in Olszyn,
grow together with *Galium cracoviense* (Szeląg 2000).

The investigation of population genetic processes or ge
netic structure in glacial relict plant allows to study the con
sequences of long term isolation for the genetic diversity
of plant populations. An investigation of the genetic di
versity within and among glacial relict populations, there
fore, can shed light upon population genetic questions of
general interest. Additionally, estimation of genetic diver
sity for this group of species is useful for optimization of sam
pling strategies and for conservation and research on ge
netic resources (Hamrick et al. 1991; Schaal et al. 1991;

In the present paper, the AFLP fingerprinting was ap
plied to answer the following questions: (1) What is the ef
fect of the long-term isolation (since the end of the last glaci
ation) on partitioning of molecular variance and on level of
genic diversity? (2) Are there any geographic differen
tiation among the populations and different spatial patterns
of genetic variability detectable in isolated patches forming
the range of *Galium cracoviense*?

**MATERIAL AND METHODS**

**Study species**

*Galium cracoviense* belongs to *G.* sect. *Leptogalium*
Lange, which comprises a complex of polymorphic diploid
and polyploid species (Ehrendorfer 1960). The centre of
the section’s range is located in the mountainous areas of
SW Europe. In NE Europe, it is represented only by few
endemic species occurring in small areas. *G. cracoviense* is
a diploid and grows only on a few limestone outcrops near
the village of Olszyn (Fig. 1) in the northern part of the
Kraków-Częstochowa Upland (Piotrowsicz 1958; Ehrendor
fer 1960, 1962). It is morphologically most similar to *G.
obelandicum* Ehrend., a diploid endemic to Œland island
(Ehrendorfer 1960), and the tetraploid *G. sudeticum* Tau
sch, which occurs in the Karkonosze Ms. and in the Sla
vkovsky Les hills in the western Czech Republic (Krahul
cová and Štěpánková 1998).

**Plant material**

After survey of all sites of *Galium cracoviense*, popula
tions from northern (A), central (B) and southern (C) part
of the total geographical range have been selected. This
sampling reflected the distribution pattern of plants within
populations and well represented the distribution of the
species in the whole area. Twenty nine samples were col
lected from the mentioned populations in May 2004 and all
of them were used in molecular analysis (Table 1). Sam
ples were collected only from clearly separated clumps of
*G. cracoviense*. The way of sampling had to be a hard com
promise between optimal statistical approach and conse
vation needs (*G. cracoviense* is placed in Polish Red Data
Book of Vascular Plants, Mirek 2001). Fragments of young
living plants were sampled from 9 to 10 randomly chosen
individuals from each population and kept in plastic tubes
with silica gel. Samples were stored in the laboratory at
-80°C prior to DNA extraction.

**DNA extraction and AFLP fingerprinting**

Total DNA was extracted using the DNeasy Plant Mini
Kit (QiaGen), according to the manufacturer’s protocol,
using ca. 10 mg of dried tissue per sampled plant. DNA
quality was estimated on 1.2% agarose gels. AFLP analysis
followed the procedure described by Vos et al. (1995) with
modification (Cieslak et al. 2007a). DNA was digested
with two restriction enzymes: *Eco* RI and *Mse* I (New En
gland Biolabs, Inc.). Resulting fragments were ligated to
double-strand adapters using T4 DNA Ligase (Roche Diagnos
tics). Restriction success was verified on 1.5% agarose gels.
The samples were then diluted 1:10 with deionized H2O. PCR
amplification was carried out in two steps: preselective and se
lective amplification. Preselective amplification was perfor
med using primers with single selective nucleotides: *Eco* RI + A and *Mse* I + C. PCR products were diluted 1:20.
Selective amplification was performed using primers with
three selective nucleotides (*Eco* RI primers were labelled

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**Fig. 1. General distribution of *Galium cracoviense* Ehrend. (hatched area) and locations of the three studied populations (black points).**
TABLE 1. Origin of plant material of Galium cruciviolense and estimation of genetic diversity in three populations (A, B, C): N – no. of plants in population; P – no. of polymorphic bands; C – no. of private bands; h – Nei’s gene diversity; S – Shannon’s information index.

<table>
<thead>
<tr>
<th>Population</th>
<th>Collection site</th>
<th>Coordinates</th>
<th>N</th>
<th>P (%)</th>
<th>C</th>
<th>h</th>
<th>S</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Olsztyn, Towarne hill, Poland</td>
<td>50°46'N 19°16'E</td>
<td>10</td>
<td>59 (54%)</td>
<td>1</td>
<td>0.1479</td>
<td>0.2175</td>
</tr>
<tr>
<td>B</td>
<td>Olsztyn, Zamkowa hill, Poland</td>
<td>50°45'N 19°16'E</td>
<td>10</td>
<td>56 (51%)</td>
<td>1</td>
<td>0.1549</td>
<td>0.2317</td>
</tr>
<tr>
<td>C</td>
<td>Olsztyn, Biaklo hill, Poland</td>
<td>50°44'N 19°16'E</td>
<td>9</td>
<td>50 (45%)</td>
<td>4</td>
<td>0.1529</td>
<td>0.2242</td>
</tr>
</tbody>
</table>

with fluorescent marker FAM 6): Eco RI AAT/Mse I CTA; Eco RI AGT/Mse I CAC; Eco RI ACG/Mse I CAG.

Amplification products were separated in POP 4 polymer with an internal size standard (GeneScan Rox 500) on the ABI Prism 3100-Avant automated sequencer (Applied Biosystems). Three samples were used in duplicates for each analysis as controls, to assess possible genotyping errors (Bonin et al. 2004). Primer screening was conducted on three individuals. Twenty-two selective primer combinations were tested, three of which were chosen for the final analysis (see above). The selection of primer combinations was based on the number of polymorphic fragments and high repeatability. Data were analysed using the GeneScan 1.3 Analysis Software (Applied Biosystems). Good quality fragments were scored in the range of 50-500 bp using Genographer, version 1.6 (Montana State University; http://hordeum.oscs.montana.edu). Finally, data were assembled in a binary 0/1 matrix.

**Data analysis**

Within-population genetic diversity for each population was quantified as (1) percentage of polymorphic loci, (2) Shannon’s information index (Lawton 1972) and (3) gene diversity index (Nei 1978). Additionally, private and discriminating markers were assessed in populations following Cieslak et al. (2007a). Private markers were those specific only to one population, while discriminating markers were private markers present in all samples from the population.

The degree of genetic isolation among populations was estimated by $N_{st}$, the number of migrations per generation. $N_{st}$, reflecting the effective migration rate among populations, was calculated as follows: $N_{st} = (1 - F_{st})/4 F_{st}$ (Wright 1969; Slatkin and Barton 1989) where $F_{st}$ is the proportion of the total genetic diversity among populations (calculated using POPGENE version 1.32; Yeh et al. 1997). The UPGMA dendrogram representing genetic relationships among individuals of G. cruciviolense was constructed on the basis of Jaccard’s similarity coefficient (Jaccard 1908). Group support was assessed by bootstrap analysis with 1000 replications (Treecon program, ver. 1.3; van de Peer and de Wachter 1994). Spatial representation of relative similarities between individuals was provided by Principal Coordinates Analysis (PCO). It was performed using MVSP 3.10b software (Kovach 1999).

Data were also analysed with the Bayesian model in Baps 4.14 (Corander et al. 2006). Both a nonmixture model with uncorrelated allele frequencies and an admixture one with correlation of allele frequencies were used. This Bayesian method detects population structure by clustering individuals into panmictic groups assuming Hardy–Weinberg equilibrium and linkage equilibrium within clusters. Both the number of populations in the sample and their allele frequencies are treated as unknown parameters and are jointly estimated by the program. The method does not make use of geographical information for detecting clusters. The procedure was run 70 times each for K=1 to 10 as the assumed maximum number of populations present in the sample. Note that the choice of K is not equal to the number of clusters that is sought in the sample. Instead, the program considers all values equal to or smaller than K to be a plausible number of clusters.

Genetic structure of populations and variation levels were assessed by the analysis of molecular variance (AMOVA; Excoffier et al. 1992). This analysis was based on the pairwise square Euclidean distance among molecular phenotypes. Significance levels were determined by 1023 permutations. The analysis was conducted at two levels: among groups representing populations and within them. AMOVAAs and values of $F_{st}$ were calculated using ARLEQUIN version 2.0 (Schneider et al. 2000; http://arlequin.ch/arlequin/).

**RESULTS**

Three AFLP primer combinations yielded a total of 157 bands. Of these bands, 110 (70%) were polymorphic. The number of polymorphic bands in particular populations ranged between 50 (population C) and 59 (population A). None of the populations had discriminating markers, but private markers were present: one per population in A and B and four in population C. The values of parameters describing genetic diversity estimated for each population were highly similar (Table 1). Calculated values of the gene flow ($N_{e}$) between populations were relatively high and similar to one another indicating a substantial interchange of genes among populations. The values were as follows: $N_{eAB} = 3.78; N_{eAC} = 3.35; N_{eBC} = 3.73$. Moreover, AMOVA indicated a substantially higher genetic variation within populations (89.35%) than among populations (10.65%); Table 2) and a low value of fixation index $F_{st} = 0.10646$ (Table 3).

The UPGMA cluster analysis demonstrated lack of genetic discontinuities among the populations. None of the spatially isolated groups formed a separate, homogeneous cluster (Fig. 2). PCO scatter diagram (with axes 1 and 2 explaining 12.67% and 8.8% of variability, respectively; Fig. 3) showed also that populations formed one complex, although there was a trend for plants growing in population
TABLE 2. AMOVA analysis results for three populations (groups) and 29 individuals of *Galium cracoviense*.

<table>
<thead>
<tr>
<th>Source of variation</th>
<th>d.f.</th>
<th>Sum of squares</th>
<th>Variance components</th>
<th>Variation percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Among populations within groups</td>
<td>2</td>
<td>66,610</td>
<td>1.84530</td>
<td>10.65</td>
</tr>
<tr>
<td>Within populations</td>
<td>26</td>
<td>402,700</td>
<td>15.48846</td>
<td>89.35</td>
</tr>
<tr>
<td>Total</td>
<td>28</td>
<td>469,310</td>
<td>17.33376</td>
<td>100.00</td>
</tr>
</tbody>
</table>

TABLE 3. Pairwise *F<sub>ST</sub>* values among study populations. All values are significant at the p<0.001 level.

<table>
<thead>
<tr>
<th>Populations</th>
<th>A</th>
<th>B</th>
<th>C</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>0.00000</td>
<td></td>
<td></td>
</tr>
<tr>
<td>B</td>
<td>0.10691</td>
<td>0.00000</td>
<td></td>
</tr>
<tr>
<td>C</td>
<td>0.13361</td>
<td>0.07686</td>
<td>0.00000</td>
</tr>
</tbody>
</table>

Fig. 2. UPGMA dendrogram based on Jaccard’s Coefficient for 29 individuals and three populations (A, B, C) of *Galium cracoviense*.

A to be located on the right side of the axis 2 (this population is situated in the northernmost part of the geographical range of *Galium cracoviense* (Fig. 1)), and for plants growing in populations B and C to be located at the other extreme in the plot. The weak differentiation was confirmed by the Bayesian analysis. Especially in admixture analysis the highest average likelihood was obtained for K=2 i.e. identifying two groups. However, one of these groups was represented by a single individual (population B, no. 18). Because of its strong genetic divergence from the rest of individuals, it was not included in subsequent analyses. All remaining individuals from A, B and C populations were located in one genetic group.

DISCUSSION

Our results clearly demonstrate that populations of *Galium cracoviense* are characterised by a high level of genetic variability. In spite of the fact that investigated populations occur as isolated groups growing on clearly separated rocky outcrops (Fig. 1), they are comparably polymorphic. They are also genetically similar and did not form divergent clusters, showing the distinct spatial isolation of populations. The values of pairwise genetic distances (*F<sub>ST</sub>*) between populations were very low and highly significant (p<0.001, Table 3) indicating high genetic similarity of all populations. Additionally, the value of gene flow between the populations, calculated on the base of the genetic distances (*F<sub>ST</sub>*) within the same localities expressed as the number of individuals migrating between populations per generation (Table 3), showed a high level of migration.

Results of the AMOVA analysis showed that substantially more genetic variation of *G. cracoviense* is distributed within populations than between them (Table 2), indicating a relatively restricted population differentiation, as was expected from outcrossing species. Also the evaluation of *F<sub>ST</sub>* is close to the values of this parameter reported by authors of the papers analyzing the population structure (though those were often based on data derived both from isozymes and DNA analysis) in mixed and outcrossing species (Loveless and Hamrick 1984; Smith and Pham 1996; Gaudeul et al. 2000; Wróblewska et al. 2003). High genetic homogeneity of the studied populations was also confirmed by the Bayesian analysis.

Such a pattern of genetic variability in small populations and for species with a narrow range has been noted before, for instance in *Cochlearia macrorhiza* (Koch et al. 2003), *Digitalis minor* (Sales et al. 2001), *Gentianella austriaca* (Greimler and Dobeš 2000), *Iris aphylla* (Wróblewska and Brzosko 2006) or *Seseli jarenyni* (López-Pujol et al. 2002). High levels of diversity were demonstrated in the narrow endemic *Allium aaseae* (Smith and Pham 1996), in the rare and endangered south African shrub *Leucadenron elimense* (Tansley and Brown 2000), also rare and endangered *Leucopogon obtectus* (Zawko et al. 2001) and in small populations of endemic *Erodium paularense* (Martin et al. 1997).

Our results reveal high genetic variability that protects *Galium cracoviense* against a genetic erosion and potential decrease in genetic diversity. On the other hand, spatially isolated populations (local gene pools) are highly homogeneous (not differentiated) which reduces the species’ overall genetic richness and may cause that changes of environmental conditions may have grave consequences both in evolutionary and ecological contexts. This low differen-
terization within the whole species range reduces potential ability of the plant to adapt to environmental changes compared to species characterized by having diversified gene pool across their geographical ranges (Ellstrand and Elam 1993). Probably, biogeographical history of the species indicates its stability.

Ehrendorfer (1962) treats Galium cracoviense as a glacial relict that survived at its present locality at least the last glaciation. We share this opinion, however, we believe that the term ‘pre-glacial’ would be more appropriate. It is highly probable that some of saxicolous calciphilous species, regarded so far as glacial relics, such as Saxifraga paniculata, Allium montanum, Festuca pallens, Gymnocarpium robertianum, Hieracium bifidum, H. caesium and Polygala brachyptera, which grow together with Galium cracoviense, may have also survived in situ at least the last Pleistocene glaciation. The phenomenon of co-occurrence of saxicolous calciphilous glacial and pre-glacial relics in the same localities in the Apuseni Mountains in Romania was analyzed by Csergö (2002).

The authors base their hypothesis of the ancient, pre-glacial age of Galium cracoviense populations on following premises: 1) the relatively high genetic polymorphism despite very narrow distribution range and small population size could suggest a long-term persistence of such populations in the area and a naturally limited distribution to ecological “islands” of rocky outcrops (Holderegger 1997); 2) the present distribution range of G. cracoviense was not covered by the continental ice sheet during the last two glaciations: Wartanian (Riss) and Vistulian (Würm) (Lang 1994; Ber 2005); 3) habitats preferred by G. cracoviense are widely distributed throughout the whole Kraków–Częstochowa Upland, however the species occurs exclusively on the northern edge of the Upland, which even during the older and most southward Krznanian (Mindel) glaciation presumably remained free from ice cover and established the local, southern border of the continental ice sheet.

This would be a situation analogous to that observed in Alps, where the stations of many saxicolous alpine endemics are found along the line delimiting maximal range of the lost glaciation’s ice sheet (Pitschmann and Reisigl 1959; Prosser and Scortegagna 1998) i.e. places regarded as hypothetical glacial refugia (Schönswetter et al. 2005).

Little is known about the genetic diversity within and among populations of glacial relics in central Europe (Dannemann 2000; Lutz et al. 2000; Reisch 2002). Numerous studies from North America provide substantial evidence that putative relict plant populations harbour high levels of genetic diversity (Lewis and Crawford 1995; Soltis et al. 1997; Allphin et al. 1998) which suggests, that “old rare species” are better adapted to processes connected with small population size and isolation (Schmid and Jensen 2000). It seems to be a typical phenomenon, that plants from glacial refugia show high levels of diversity. This allows the species to respond to the selection pressure imposed by pests and disease and to facilitate adaption to future environmental changes (Barrett and Kohn 1991; Holsinger and Gottlieb 1991). The high levels of diversity of glacial relics in central Europe are present e.g. in Saxifraga paniculata, which also belongs to “old rare species” (Reisch et al. 2003; Reisch and Poschlod 2004; Reisch 2008).

Data on genetic diversity within and among populations of rare and endangered species play a significant role in the formulation of appropriate management strategies directed
towards their conservation and development, besides being advantageous in the understanding of their structure, evolutionary relationships, taxonomy and demography (Milligan et al. 1994). Consequently, the study of population genetics has been identified as one of the priorities for conservation (Holsinger and Gottlieb 1991).

The present survey was the first attempt to understand the genetic structure of populations of Galium cracovicum. Such data are particularly important for species with narrow ranges and low number of populations, additionally endangered by various external threats, as it is in the case of another Polish endemic plant, Cochlearia polonica where molecular data allowed to confirm the narrow endemic status and estimate the genetic resources of the only existing stable population (Cieslak et al. 2007a, b). In the case of Galium cracovicum additional demographic and molecular studies are necessary to further monitoring in detail of this species’ genetic condition (Schwartz et al. 2006). The repeated monitoring of population size indicate a threat mainly by the mechanical destruction of plants and their habitats caused by intense tourism. The decreasing number of individuals in population or number of populations may initiate genetic processes influencing the variability and differentiation of the whole population.

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