Cytogeography of *Pilosella officinarum* (Compositae): Altitudinal and Longitudinal Differences in Ploidy Level Distribution in the Czech Republic and Slovakia and the General Pattern in Europe

PATRIK MRÁZ1,2, *, BARBORA ŠINGLIAROVÁ1,2, TOMÁŠ ŠURFUS3,4 and FRANTIŠEK KRAHULEC4

1Institute of Biology and Ecology, P. J. Šafárik University – Faculty of Science, Mnísova 23, SK-041 54 Košice, Slovakia, 2Institute of Botany, Slovak Academy of Sciences, Diárovská cesta 14, SK-845 23 Bratislava, Slovakia, 3Department of Botany, Charles University, Benátská 2, CZ-128 01 Praha, Czech Republic and 4Institute of Botany, Academy of Sciences of the Czech Republic, Práhonice 1, CZ-252 43 Czech Republic

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Background and Aims *Pilosella officinarum* (syn. *Hieracium pilosella*) is a highly structured species with respect to the ploidy level, with obvious cytogeographic trends. Previous non-collated data indicated a possible differentiation in the frequency of particular ploidy levels in the Czech Republic and Slovakia. Therefore, detailed sampling and ploidy level analyses were assessed to reveal a boundary of common occurrence of tetraploids on one hand and higher ploids on the other. For a better understanding of cytogeographic differentiation of *P. officinarum* in central Europe, a search was made for a general cytogeographic pattern in Europe based on published data.

Methods DNA-ploidy level and/or chromosome number were identified for 1059 plants using flow cytometry and/or chromosome counting on root meristem preparations. Samples were collected from 336 localities in the Czech Republic, Slovakia and north-eastern Hungary. In addition, ploidy levels were determined for plants from 18 localities in Bulgaria, Georgia, Ireland, Italy, Romania and Ukraine.

Key Results Four ploidy levels were found in the studied area with a contrasting pattern of distribution. The most widespread cytotype in the western part of the Czech Republic is tetraploid (*4x*) reproducing sexually, while the apomictic pentaploids and mostly apomictic hexaploids (*5x* and *6x*, respectively) clearly prevail in Slovakia and the eastern part of the Czech Republic. The boundary between common occurrence of tetraploids and higher ploids is very obvious and represents the geomorphologic boundary between the Bohemian Massif and the Western Carpathians with the adjacent part of Pannonia. Mixed populations consisting of two different ploidy levels were recorded in nearly 11% of localities. A statistically significant difference in a vertical distribution of penta- and hexaploids was observed in the Western Carpathians and the adjacent Pannonian Plain. Hexaploid populations tend to occur at lower elevations (usually below 500 m), while the pentaploid level is more or less evenly distributed up to 1000 m a.s.l. For the first time the heptaploid level (*7x*) was found on one site in Slovakia. In Europe, the sexual tetraploid level has clearly a sub-Atlantic character of distribution. The plants of higher ploidy level (*penta- and hexa-*) with mostly apomictic reproduction prevail in the northern part of Scandinavia and the British Isles, the Alps and the Western Carpathians with the adjacent part of Pannonia. A detailed overview of published data shows that extremely rare records on existence of diploid populations in the south-west Alps are with high probability erroneous and most probably refer to the closely related diploid species *P. peleteriana*.

Conclusions The recent distribution of *P. officinarum* in Europe is complex and probably reflects the climatic changes during the Pleistocene and consequent postglacial migrations. Probably both penta- and hexaploids arose independently in central Europe (Alps and Carpathian Mountains) and in northern Europe (Scandinavia, Great Britain, Ireland), where the apomictic plants colonized deglaciated areas. We suggest that *P. officinarum* is in fact an amphidiploid species with a basic tetraploid level, which probably originated from hybridizations of diploid taxa from the section *Pilosella*.

Key words: Amphidiploidy, apomixis, Asteraceae, flow cytometry, geographical parthenogenesis, *Hieracium*, postglacial migration, polyploidy.

INTRODUCTION

The genus *Pilosella* Hill., often treated as a subgenus of *Hieracium* L. (syn. *Hieracium* subgen. *Pilosella* (Hill) Gray), is one of the taxonomically most intricate vascular plant groups of the temperate flora. The reticulate pattern of morphological variation reflected in several thousands of taxa described from the species level to the form (Zahn, 1921–1923) complicates taxonomic treatment.

Widespread polyploidy, various modes of reproduction (sexuality, obligate and facultative apomixis of aposporous type, haploid parthenogenesis, vegetative propagation), inter- and intraspecific hybridization within the same and across different ploidy levels are the most important processes involved in microevolution of the genus (Krahulcová et al., 2000). The ploidy level occurring in natural populations varies from diploid (*2n = 2x = 18*) to octoploid (*2n = 8x = 72*). The most frequent cytotypes are tetraploids, pentaploids and hexaploids. Diploids are less frequent, and triploids, heptaploids, octoploids and aneuploids are rather rare (Schuhwerk and Lippert, 1997;...
Krahulcová et al., 2000). Higher ploidy levels up to dodecaploid (2n = 12x = 108) were found in plants obtained by experimental hybridization (Skalińska, 1976). In approximately half of the taxa of the genus Pilosella analysed more than one ploidy level was found, even in plants growing together in one locality (Schuhwerk, 1996 and references therein).

Sell and West (1976) recognize 63 ‘numbered’ species (including nothospecies) of Pilosella (treated as subgenus of Hieracium) in Europe. Six species having only one capitulum per stem (with the exception of hybrids) are members of the section Pilosellina Zahn. All but one are diploids occurring mainly in western, southern and central parts of Europe: P. argyrocoma (S fores) (southem and central Spain), P. castellana (Boiss. & Reuter) F. W. Schultz & Sch. Bip. (Spain and northern Portugal), P. hoppeana (Schult.) F. W. Schultz & Sch. Bip. (central and eastern Alps), P. pseudopilosella (Ten.) F. W. Schultz & Sch. Bip. (southern Europe, from Portugal and Spain to Bulgaria, Turkey and Romania), P. peleteriana (Mérat) F. W. Schultz & Sch. Bip. (northern and western Europe and the western part of central Europe). In addition, some authors distinguish a lowland form of P. hoppeana as a distinct diploid taxon – P. macrantha (Ten.) F. W. Schultz & Sch. Bip. (central and southern Europe) (e.g. Holub, 1986; Chrttek, 1998, 2002; Gottschlich, 1998; Schuhwerk and Fiscker, 2003). The only almost exclusively polyploid species of section Pilosellina is Pilosella officinarum (F. W. Schultz & Sch. Bip.) (syn. Hieracium pilosella L.). It is distributed much wider than the diploid taxa mentioned above (cf. Hultén and Fries, 1986; Brüttigam, 1992). It extends from the British Isles across the whole of Europe, except the Arctic parts, to western Asia and north-western Siberia. Moreover, it has been introduced into New Zealand, North America and Patagonia, and has become an invasive and troublesome weed (Hultén and Fries, 1986; Chapman et al., 2000; Cárdenas Vergara, 2005; Wilson et al., 2006). Pilosella officinarum usually grows on dry, permeable and nutrient-poor soils from sea level to the sub-Alpine belt. The obligate heliophilous species occurs in tussock grassland communities usually with regular disturbance (graizing, mowing). Due to its low competitive ability, it tends to establish itself on open, sparsely vegetated, sites (roadside dikes, eroded slopes, landslides, etc.). Morphologically, P. officinarum may be distinguished from other representatives of Pilosellina section by long and slender stolons bearing small distant leaves decreasing in size towards the apex, involucral bracts (0.5–1)–2 mm wide, covered by stellate, glandular and eglandular trichomes (Sell and West, 1976). Despite its more or less easy identification in the field, a great phenotypic plasticity has been recorded within the species (Turesson and Turesson, 1960; Gadella, 1987, 1991). A high level of morphological variation is reflected in numerous infraspecific taxa described from the entire distribution range (Nägeli and Peter, 1885; Zahn, 1921–1923). The correlation between some phenotypic characters on one hand (e.g. rosette size, the number and length of stolons) and ploidy level on the other was revealed by Gadella (1991). In total, five cytotypes (2x, 4x, 5x, 6x and 7x) have been recorded in natural populations of P. officinarum (cf. Table 1). The mode of seed reproduction of each particular cytotype depends on the ploidy level. Diploids are sexual. Tetraploid plants reproduce sexually, but several apomictic populations deviate from this general rule (Gadella, 1984, 1987). Pentaploid P. officinarum is almost exclusively

### Table 1. Ploidy levels found in Pilosella officinarum in Europe according to the literature and present records

<table>
<thead>
<tr>
<th>Country</th>
<th>Ploidy level</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Armenia</td>
<td>4x</td>
<td>Nazarova and Ghuksayan, 2004</td>
</tr>
<tr>
<td>Austria</td>
<td>4x, 5x, 6x</td>
<td>Turesson and Turesson, 1960; Gadella, 1972, 1984; Schuhwerk and Lippert, 1997</td>
</tr>
<tr>
<td>Belarus</td>
<td>4x, 5x</td>
<td>Dmitrieva, 1987; Parfenov and Dmitrieva, 1988</td>
</tr>
<tr>
<td>Belgium</td>
<td>4x</td>
<td>Gadella, 1972, 1984</td>
</tr>
<tr>
<td>Bulgaria</td>
<td>4x, 5x, 6x</td>
<td>Mráz et al., this study</td>
</tr>
<tr>
<td>Croatia</td>
<td>6x</td>
<td>Gadella, 1984</td>
</tr>
<tr>
<td>Czech</td>
<td>4x, 5x, 6x</td>
<td>Měšíček and Jarolímová, 1992</td>
</tr>
<tr>
<td>Republic</td>
<td>7x</td>
<td>Krahulcová and Krahulec, 1999</td>
</tr>
<tr>
<td>Denmark</td>
<td>4x</td>
<td>Turesson and Turesson, 1960; Gadella, 1972, 1984</td>
</tr>
<tr>
<td>Finland</td>
<td>4x, 5x</td>
<td>Turesson and Turesson, 1960; Jalas and Pellinen, 1985</td>
</tr>
<tr>
<td>France</td>
<td>2x, 4x, 5x</td>
<td>Delcourt, 1972; Auquier and Renard, 1979; Natarajan, 1981, 1988; Gadella, 1972, 1984</td>
</tr>
<tr>
<td>Germany</td>
<td>4x</td>
<td>Turesson and Turesson, 1960; Gadella, 1972; Bräutigam and Bräutigam, 1996; Schuhwerk and Lippert, 1997, 2002; Albers and Prösting, 1998; Rotreklová et al., 2005</td>
</tr>
<tr>
<td>Hungary</td>
<td>5x, 6x</td>
<td>Mráz et al., this study</td>
</tr>
<tr>
<td>Ireland</td>
<td>4x, 5x, 6x</td>
<td>Gadella, 1972, 1984; Finch, 2005; Watson, 2005; Mráz et al., this study</td>
</tr>
<tr>
<td>Italy</td>
<td>2x, 4x, 5x</td>
<td>Gadella, 1972, 1984</td>
</tr>
<tr>
<td>Luxembourg</td>
<td>4x</td>
<td>Gadella, 1972, 1984</td>
</tr>
<tr>
<td>Macedonia</td>
<td>6x</td>
<td>Gadella, 1972</td>
</tr>
<tr>
<td>Netherlands</td>
<td>4x, 5x, 7x</td>
<td>Gadella and Kliphuis, 1963; Gadella, 1972, 1984</td>
</tr>
<tr>
<td>Norway</td>
<td>4x, 5x</td>
<td>Gadella, 1972</td>
</tr>
<tr>
<td>Poland</td>
<td>4x, 5x, 6x</td>
<td>Skalińska, 1967; Skalińska et al., 1971; Gadella, 1972; Pogan et al., 1987; Pogan and Wcislo, 1989; Rotreklová et al., 2005</td>
</tr>
<tr>
<td>Portugal</td>
<td>4x</td>
<td>Fernades and Queirós, 1971; Gadella, 1972</td>
</tr>
<tr>
<td>Romania</td>
<td>5x, 6x</td>
<td>Gadella, 1972; Mráz et al., this study</td>
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<tr>
<td>Russia</td>
<td>6x</td>
<td>Lavrenko and Sereditov, 1991</td>
</tr>
<tr>
<td>Slovakia</td>
<td>4x, 5x, 6x</td>
<td>Májo vský et al., 1970; Uhříková and Feráková, 1977; Mičieta, 1982; Murin, 1986; Pišt’anský and Mičieta, 2000; Rotreklová et al., 2002, 2005; Mráz et al., this study</td>
</tr>
<tr>
<td>Spain</td>
<td>4x</td>
<td>Gadella, 1984</td>
</tr>
<tr>
<td>Sweden</td>
<td>4x, 5x, 6x</td>
<td>Turesson and Turesson, 1960; Turesson, 1972; Lövkvist and Hultgård, 1999</td>
</tr>
<tr>
<td>Switzerland</td>
<td>4x, 5x, 6x</td>
<td>Gadella, 1972, 1984</td>
</tr>
<tr>
<td>United</td>
<td>4x, 5x, 6x</td>
<td>Turesson and Turesson, 1960; Gadella, 1972, 1984; Morton, 1974; Moore, 1982; Edmonds et al., 2005; Finch, 2005; Grime et al., 2005; Watson, 2005</td>
</tr>
<tr>
<td>Ukraine</td>
<td>4x, 6x</td>
<td>Paschuk, 1987; Mráz et al., this study</td>
</tr>
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apomictic, although a rare sexual seed production was also reported (Turesson and Turesson, 1960; Turesson, 1972; Gadella, 1984). Facultative apomixis in pentaploids was later confirmed embryologically by Pogan and Wcislo (1995). Recently, two accessions of fully sexual pentaploids have been found in the Czech Republic (Krahulcová et al., 2000; Rotreklová et al., 2002). Hexaploids are either sexual or apomictic, while very rare heptaploids are either apomictic or sterile (Gadella, 1984, 1991). Vegetative reproduction by means of over-ground stolons is common for all cytotypes and, together with apomixis, it might contribute to the unicolonial structure of populations.

Diploid plants of *P. officinarum* are rare and their distribution is considered to be of a relict character (e.g. Gadella, 1984). They were reported from the Valley of Aosta (Italy) (Gadella, 1972) and south-eastern France (Delcourt, 1972), respectively (but see the Discussion below). In most of Europe, the tetraploid and pentaploid populations of *P. officinarum* are by far the most common cytotypes (Gadella, 1984). Tetraploids are widespread in the lowlands of west and central Europe (e.g. Turesson and Turesson, 1960; Gadella, 1972, 1984; Pogan and Wcislo, 1989; Schuhwerk and Lippert, 1997, 2002; Krahulcová and Krahulec, 1999), while the pentaploids occur chiefly in regions that were covered by the Pleistocene glaciation – Scandinavia, the British Isles (Turesson and Turesson, 1960; Gadella, 1972, 1984, 1987; Finch, 2005; Watson, 2005). Several hexaploid populations of *P. officinarum* were found mainly in the Alps, Scandinavia, Balkan Peninsula (e.g. Turesson and Turesson, 1960; Gadella, 1972, 1984, 1991; Lavrenko and Seredtov, 1991; Schuhwerk and Lippert, 1997) and the Western Carpathians (see below). The rare occurrence of heptaploids was reported from only three localities in Sweden (Turesson and Turesson, 1960), one site in the Netherlands (Gadella, 1984) and one population in the Czech Republic (Košt’álová, 2004).

Four ploidy levels (tetra-, penta-, hexa- and heptaploid) have been recorded in the Czech Republic and Slovakia (Májovsky et al., 1970; Uhríková and Feráková, 1977; Mičieta, 1982; Murin, 1986; Mešiček and Jarolímova, 1992; Krahulcová and Krahulec, 1999; Pšt’anský and Mičieta, 2000; Krahulcová et al., 2001; Rotreklová et al., 2002, 2005; Košt’álová, 2004). Recently, Pšt’anský and Mičieta (2000) recorded tetraploids in approx. 30 localities mainly in southern and western Slovakia, while other authors reported pentaploid and hexaploid plants mostly from eastern, northern and central Slovakia. Most of the chromosome counts coming from the Czech Republic that had been published indicated that the plants analysed were tetraploids.

Almost all published data on ploidy level of *P. officinarum* are based on classical chromosome counting. This precise method is, however, considerably time-consuming. Since routine introduction of the flow cytometry in plant science in the nineties of the last century (Doležal, 1991), this approach has rapidly become popular for estimating DNA-ploidy level (Doležel, 1991). This is mainly due to the very easy sample preparation and the possibility of screening large numbers of individuals in a very short time. Here, the search which was carried out for a boundary between the area of distribution of the tetraploid cytotype and the range of pentaploid and hexaploids of *P. officinarum* in the territory of Slovakia and the Czech Republic, using mostly a flow cytometric approach, is reported. Moreover, an attempt was made to find out if there was a correlation between the distribution of particular ploidy levels on one hand and the altitude on the other. To understand better the cytogeographic differentiation of *P. officinarum* in central Europe, a search, based on published data, was made for a general cytogeographic pattern in Europe.

The area studied

Research has been carried out in the area of the Czech and Slovak Republics with an adjacent part of north-eastern Hungary. The area studied belongs to two different biogeographic regions, the mountain range of the Western Carpathians and the Bohemian Massif. The border between both regions is situated in the eastern part of the Czech Republic, lying north-north-east to south-south west. These two regions differ in a variety of environmental and historical parameters. In this respect, differences in the cytotype distribution cannot be explained in any easy way. On the other hand, this area covering their border can show that the pattern in cytotype distribution can be very contrasting even across a very narrow zone.

The Bohemian Massif has an old Paleogenic relief, younger areas being only canyons, those areas with Tertiary volcanism in the northern part of Bohemia, and glacial cirques in the Sudetes and the Šumava Mountains. The highest point is Mík Sněžka (1602 m), the lowest is the valley of the River Elbe on the German border at 115 m. Mostly acid Varisian parts were later covered with Permnian-Carboniferous or Mesozoic sediments. Base-rich bedrocks are concentrated at lower altitudes. Vegetation cover has a coarser grain (homogeneous on a larger scale) in comparison with the Carpathian Mountains.

The Western Carpathians, including the Intra-Carpathian (Pannonian) Basin, represent the north-west part of the Carpathian are extending from north-east Austria and south-east Czech Republic to north-east Slovakia and south-east Poland. The relief is young, of Tertiary age, similar to the Alps. The highest point is Gerlach Peak (2655 m). The bedrock is more complicated, mostly of Mesozoic and Tertiary ages. Calcium-rich substrates occur from lowland to the high mountains; e.g. in central Slovakia almost consistent limestone substrates can be found from the xerothermic Slovak Karst to the highest altitudes of the Belaer Tatra with altitudes above 2000 m. Some areas are very continental, with climatic conditions which do not allow the growth of Fagus sylvatica as in the area between the High and Low Tatra Mountains. On the other hand, some not distant areas are more oceanic, as in north-west Slovakia. For all these reasons, the vegetation cover is fine-grained (homogeneous in small areas but, on a larger scale, heterogeneous). Large regions with homogeneous vegetation are rare.

The area of the Czech Republic has a rather uniform climate; the warmest month is July and it is also the
month of highest rainfall. This contrasts with Slovakia, 
where the same condition applies only at higher altitudes. At 
alower altitudes, the warmest month is also July, but the 
highest rainfall is distributed from May to September, 
depending on the exact geographic position (Vesecký, 
1961). In this way, the same area is rather oceanic in 
May–June and more continental in September and vice versa. Slovakia (the Carpathian Mountains with the 
Pannonian Plain) is therefore fine-grained and more diverse 
with respect to relief, bedrock and climate.

MATERIALS AND METHODS

Material collection

Plants of Pilosella officinarum F. W. Schultz & Sch. Bip. 
(syn. Hieracium pilosella L.) for the present study were 
collected in 2003–2006 in their natural habitats throughout 
Slovakia and the Czech Republic, to a lesser extent also in 
the north-eastern part of Hungary to cover all geographic 
regions. They were cultivated in pots in the Botanical 
Garden of P. J. Šafárik University, Košice and in the exper-
imental field of the Institute of Botany, Academy of 
Sciences of the Czech Republic, Průhonice. For the com-
plete list of localities see Supplementary Information 1 
available online). Besides the plants from the region 
above mentioned, some plants from a further 18 localities 
different parts of Europe have also been analysed 
(Supplementary Information 4, available online).

As a rule, three or five plants from each population were 
sampled, three from the pure populations, five in the case of 
co-occurrence of other potentially hybridizing species of 
the genus Pilosella. Efforts were made to avoid collecting 
samples originating from one clone. If it was apparent 
that plants at the collecting site did originate from one 
plant, they belonged to one clone (usually plants growing 
very close together in a very small area of several square 
cm), only one individual plant per locality (population) 
dug up. In some cases however, several cultivated 
plants died before analysis. For both these reasons, some 
populations are represented by only one plant. To determine 
the proportion of mixed cytotypes in populations, only 
those populations with two and more plants analysed 
were involved. Despite the fact that the ploidy level of 
only one plant had been estimated by us, some localities 
marked in Supplementary Information 1, available online) 
can be considered as collecting sites with two or 
more analysed plants because the chromosome number of 
other plant/plants from the same locality was published 
elsewhere (see Supplementary Information 2, available 
online). Therefore, in addition to the data collected for 
this research, a few previous accounts from the literature 
(Roterklova et al., 2005) were used to search for some 
localities consisting of two different ploidy levels. These 
plants were not included in the total number of plants ana-
lysed in this present study. The voucher specimens have 
been deposited recently in the herbarium of Patrik Mráz, 
at the Institute of Biology and Ecology, P. J. Šafárik 
University, Košice and in the herbarium of the Institute of 
Botany, Průhonice (PRA).

Chromosome counts

The chromosome counts are based on the somatic mitosis 
in the root-tip cuttings of pot-cultivated plants. The material 
was pre-treated at room temperature with a 0.5% solution of 
colchicine for 1.5–3 h and then fixed in a cold mixture of 
etanol and acetic acid (3 : 1) for at least 1 h. The fixed 
material was stored in 70% ethanol at 4 °C until processed. 
The root tips were macerated in 1 n HCl at 60 °C for 7– 
10 min. The squash and smear method with cellophane 
replacing the glass covers (Murín, 1960) and with Giemsa 
solution in a phosphate buffer was used. Selected perma-

Estimation of ploidy level

Flow cytometry was used to detect the DNA-ploidy level 
(Suda et al., 2006) for most of the plants. An analysis of 
relative DNA content was performed with a PA II ploidy 
analyser (Partec GmbH, Münster, Germany) equipped 
with an HBO-100 mercury arc lamp in the Flow 
Cytometry Laboratory, Institute of Botany, Academy of 
Sciences, Průhonice, Czech Republic and FACScalibur 
instrument (Becton Dickinson, USA) equipped with an 
argon-ion laser excitation at 488 nm in the Flow 
Cytometry Laboratory, Institute of Biology and Ecology, 
P. J. Šafárik University, Košice. Sample preparations were 
carried out in a two-step procedure (Otto, 1990; Doležel 
and Gohde, 1995). Approximately 1 cm² of leaf tissues 
from both the sample and the reference internal standard 
were ground together for about 30 s in a Petri dish contain-
ing 1 ml of ice-cold Otto I buffer (4·2 g citric acid 
monohydrate + 1 mL 0·5 % Tween 20 adjusted to 200 mL 
and filtered through a 42·µm filter). Filtration through a 
42-µm nylon mesh was followed by centrifugation at 
150 g for 5 min. The supernatant was removed 
and 100 µL of fresh Otto I buffer was added. The nuclei in 
the pellet were resuspended and stored for 30 min at room 
temperature for incubation. For DNA staining 1 mL of 
Otto II buffer (0·4 n sodium hydrogencarbonate dodeka-
hydrate) including 50 µL of propidium iodide, 50 µL ribo-
nuclease, 2 µL mercaptoethanol (FACScalibur, Becton 
Dickinson) or DAPI (4',6-diamidino-2-phenylindole) at a 
concentration of 4 µg mL⁻¹ (PA II flow cytometer, Partec 
GmbH) was used. The clones of previously cytologically 
studied diploid (2n = 2x = 18) plants of Pilosella lactu-
cella (Wallr.) P. D. Sell & C. West (Rotreklová et al., 
2002, 2005) were used as an internal reference standard 
for the relative DNA content measurements. Moreover, 
one tetraploid and several pentaploid and hexaploid 
plants of P. officinarum with known chromosome 
numbers were used in separate and mixed flow cytometry 
analysis to determine the exact position of peaks of 
known polyploids in relation to the diploid standard peak 
(Fig. 1). Histograms were accumulated at a flow rate of 
about 20–50 particles per second for a total count of 
3000–5000 nuclei. The resulting values were expressed 
as a peak ratio, which is a ratio of the mean position of 
the G1 peak in the DNA histogram of the tested plant
to the mean position of the G0/G1 peak in the histogram of the reference plant.

Maps

The distribution maps of cytotypes/ploidy levels in the Czech Republic and Slovakia are based on the co-ordinates determined by a GPS receiver, or found ex post facto from the tourist maps at a scale of 1 : 50 000 (usually old literature data). For most references from Europe for which the appropriate geographical co-ordinates were not given in original sources, the geographical position of collecting sites was estimated using Microsoft Encarta World Atlas (1998 Edition) and GeoNet Name Server (http://gns.ww.nga.mil/geonames/GNS/index.jsp; accessed in December 2005). However, for approx. 10 % of references, estimation of co-ordinates failed (marked by an asterisk in Supplementary Information 3, available online) usually due to the absence of the name of the nearest village/town, or the existence of two or more villages/towns with the same name. Most of the chromosome numbers of the plants from the British Isles were obtained from the online version of Cytological database of the Botanical Society of the British Isles (accessed in February 2005). Distributional maps were prepared using distribution mapping software DMAP (Morton, 2004).

Statistical analysis

One-way analysis of variance (ANOVA) and Tukey’s pairwise comparison (using Minitab for Windows Release 11) were applied to determine the significance (P < 0.05) of the difference in the altitudinal distribution between pentaploid, hexaploid and mixed populations in the Western Carpathians and adjacent Pannonia (Slovakia and the eastern part of the Czech Republic).

RESULTS

Ploidy level distribution in the Czech Republic and Slovakia

The DNA-ploidy levels and/or chromosome numbers were detected for 1059 plants sampled at 336 localities throughout the Czech Republic and Slovakia. Some plants were sampled also in north-eastern Hungary, along the Slovak–Hungarian state border. In total, 1055 plants were analysed by flow cytometric analysis. For eight plants the ploidy level was found using two approaches – by classical counting and by flow cytometry – while another four plants were counted only (cf. Supplementary Information 1, available online).

Altogether, four ploidy levels, tetra-, penta-, hexa- and heptaploid, were revealed in the area on which the study focused. The tetraploid level (4x; altogether 426 plants which represent 40.2 % of all plants analysed) was found to be the most common, followed by pentaploid (5x; 389 plants, 36.7 %) and hexaploid (6x; 241 plants, 22.8 %). Three heptaploid plants (7x) were discovered in a mixed population with one pentaploid plant at only one site in western Slovakia (Fig. 2). The record of heptaploid ploidy level is the first for P. officinarum in the territory of the Western Carpathians. The effort made to determine the chromosome number of heptaploid plants was not successful (the plants died), thus the new ploidy level should be considered merely as a DNA-ploidy level, i.e. not based on an exact chromosome count. Estimations of ploidy levels given for the plants from the Hungarian part of the Western Carpathians are the first records of ploidy level for P. officinarum for this area. In 32 localities out of 302 (10.6 %), from which at least two plants were analysed, mixed populations consisting of two different ploidy levels were found.

The distribution of ploidy levels in the Czech Republic is not proportional to that in Slovakia. While tetraploids are the most widespread in the Czech Republic, specifically in its western part, penta- and hexaploids predominated in Slovakia and in the eastern part of the Czech Republic (Fig. 2). The boundary between a common occurrence of the tetraploid cytotype and higher ploids is very conspicuous and corresponds well with the natural geological and geomorphological boundary between the Bohemian Massif and the Western Carpathians with the adjacent Pannonian Plain (Král, 1999). If the proportion of the particular ploidy level for each geographic region is taken into account separately, i.e. the Bohemian Massif on one hand and the Western Carpathians with Pannonia on the other, then the differences are very striking (Fig. 3).

Apart from latitudinal differentiation in ploidy level distribution in the Czech Republic and Slovakia, a statistically significant difference was also found between the proportion of pentaploids and hexaploids across the altitudes in the territory of the Western Carpathians and adjacent Pannonia (Slovakia, north-east Hungary and the eastern part of the Czech Republic) (Table 2). Generally, pure hexaploid populations tend to occur at lower elevations (usually below 500 m), while the pentaploids are very common above 500 m a.s.l. Mixed populations consisting...
Fig. 2. Distribution of ploidy levels of *Pilosella officinarum* in the Czech Republic, Slovakia and north-east part of Hungary based on present data.
of two different ploidy levels were found relatively evenly along the altitudinal gradient up to 1000 m (Fig. 4).

General pattern of ploidy level distribution in Europe

*Pilosella officinarum* is the European taxon most examined by karyology. Chromosome numbers were counted in plants originating from 655 localities, excluding present data, across the whole continent (refer to Supplementary Information 3, available online, and Table 1). It was possible to localize geographically nearly 600 sites (Fig. 5). The most common cytotype, tetraploid (without present data), was reported in 284 localities (43 %), followed by pentaploid found at 257 collecting sites (40 %). The hexaploid ploidy level is obviously rarer, i.e. detected in 74 cases (11 %). Mixed populations consisting of two or more different cytotypes were found on 40 sites (6 %). However, in most publications the number of plants analysed per population was not given and therefore the proportion of mixed populations would be probably higher if only populations with at least two or more analysed plants per locality were taken in consideration. Tetraploids are distributed mostly in western Europe and the western part of central Europe, being the only cytotype detected in Denmark and Germany. It prevails considerably in the Netherlands (82 %), Poland (71 %) and France (65 %). The pentaploids have two main centres of distribution: at high latitudes in northern Europe (Sweden, 70 %; the British Isles, 64 %) and in major orophytic systems in Europe – the Alps (30 %) and the Carpathian Mountains with the adjacent part of Pannonia (present data for the Western Carpathians indicate 66 % of pentaploids). The predominant ploidy level in the Alps is hexaploid (59 %, in Switzerland even 84 %). The records on diploids and heptaploids are extremely scarce. The former

<table>
<thead>
<tr>
<th>Ploidy level</th>
<th>N</th>
<th>$X \pm $ s.d.</th>
</tr>
</thead>
<tbody>
<tr>
<td>5x</td>
<td>86</td>
<td>546 ± 292$^a$</td>
</tr>
<tr>
<td>6x</td>
<td>40</td>
<td>370 ± 189$^b$</td>
</tr>
<tr>
<td>5x + 6x</td>
<td>21</td>
<td>513 ± 253$^{ab}$</td>
</tr>
</tbody>
</table>

Only populations with two or more analysed plants were included. The tetraploids and heptaploids were, according to their comparatively low abundance, excluded from this analysis. Altitudinal ranges and means are given in metres above sea level. N, Total number of populations; X, mean; s.d., standard deviation. Means in columns sharing the same superscript letters are not significantly different (Tukey’s pairwise comparisons, $P = 0.003$, $F = 6.19$).

Fig. 3. Proportions of the ploidy level of *Pilosella officinarum* in the Czech Republic, north-east Hungary and Slovakia based on present data, expressed as a portion of plants of a particular ploidy level compared with the total plants analysed. The proportions over the whole area studied are given (total), together with the proportions found in two different geomorphological regions – the Bohemian Massif and the Western Carpathians with the adjacent part of Pannonia.

Fig. 4. Proportions of pure pentaploid (5x), pure hexaploid (6x) and mixed populations (5x + 6x) of *Pilosella officinarum* found at different altitudinal ranges in the Western Carpathians and the adjacent part of Pannonia, expressed as a number of populations of the particular ploidy level compared with the total number of populations analysed in the particular altitudinal range.
were found only in the south-west Alps (but see Discussion), the latter mostly in northern and central Europe (Sweden, Netherlands and the Czech Republic; heptaploids are not included in Fig. 5). The new data from 18 European localities confirm this general pattern: Bulgaria 4x, 5x and 6x; Georgia 4x and 5x; Ireland 5x and 6x; Italy 6x; Romania 5x, 6x and mixed 5x + 6x; Ukraine 6x (cf. Table 1 and Supplementary Information 4, available online).

**DISCUSSION**

**Ploidy level distribution in the Czech Republic and Slovakia**

According to the results of the present research, the tetraploids strongly prevail in the western part of the Czech Republic, while the pentaploids and hexaploids represent two main cytotypes in Slovakia and the eastern part of the Czech Republic. This corresponds well with scattered data published in previous studies, with the exception of the tetraploids. Our results contradict the data published by Přístanský and Míčeta (2000), who reported a significant predominance of tetraploid populations in the Western Carpathians, but only a rare occurrence of pentaploid and hexaploid plants (cf. Supplementary Information 2, available online). As preliminary results did not confirm any common occurrence of tetraploids in Slovakia (Šingliaráová and Mráz, 2004), in 2004 five localities were visited from which the tetraploids were published by Přístanský and Míčeta (2000). However, no tetraploid was detected in any of them. Based on these observations, all data of Přístanský and Míčeta (2000) were considered as dubious.

The present study revealed a new heptaploid ploidy level in *P. officinarum* in the territory of the Western Carpathians. So far, the heptaploids had been detected only in three localities in Sweden (Turesson and Turesson, 1960), in one site in the Netherlands (Gadella, 1984) and one plant in a population of hexaploid plants near Prague in the Czech Republic (Košt’álova, 2004). In the Western Carpathians locality, from four analysed plants three were heptaploid and only one pentaploid. Although pentaploids reproduce the most often via aposporic apomixis, there are some data on facultative apomixis (Turesson and Turesson, 1960; Turesson, 1972) or even full sexuality (Krahulcová et al., 2000; Rotreklová et al., 2002). Moreover, apomictic pentaploids usually produce 2x to 3x pollen grains (Gadella, 1987; Krahulcová and Krahulec, 2000). A possible explanation of an increased ploidy level may be the fusion of reduced and unreduced gametes, as was suggested in the case of a large Dutch heptaploid population situated between two localities – the first occupied by tetra- and the second by pentaploid plants (Gadella, 1988).
A sympatry of two ploidy levels within one population was confirmed in nearly 11% of populations. The presence of cytotype mixtures is pronounced especially in the Western Carpathians (16-4%), whereas in the Bohemian Massif a co-occurrence of different ploidy levels is rarer (5-9%) and confined only to the warmest regions in relict river valleys or in the zone adjacent to the Western Carpathians and the Pannonian Plain. Higher numbers of mixed populations in the Western Carpathians might be explained by a high presence of two different ploidy levels (5x and 6x) in this territory. However, this is not the case for the Bohemian Massif which has only one completely dominating tetraploid cytotype. Whether the presence of mixed cytotypes is mainly due to the more or less stochastic co-existence of different clones with different ploidy levels or to the local formation from one dominant ploidy level is yet unknown. Undoubtedly, the production of fully or partially reduced or unreduced gametes and gene flow between plants in the locality may contribute to the presence of a cytotype mixture in populations (cf. Krahulcová et al., 2000). Mixed cytotype populations were found previously also in other parts of Europe in 37 localities (cf. Supplementary Information 3, available online), as well as in the Western Carpathians (Skalniška, 1967).

Surprisingly, the boundary between a common occurrence of tetraploids and higher ploids of P. officinaria is rather sharp and corresponds well to the natural boundary of two geomorphological units: the Western Carpathians with the adjacent Pannonian Plain and the Bohemian Massif. From the cytological point of view, a similar boundary between two cytotypes of P. bauhini was recorded by Rotreklová (2004), albeit, with the reverse pattern in comparison with P. officinarum. The tetraploids of P. bauhini are more frequent in Slovakia and Hungary and rare in the Czech Republic, Poland or Germany. On the other hand, pentaploid populations prevail in the Czech Republic and Germany. The border between Hercynian (including the Bohemian Massif) and Carpathian regions seems to be an important biogeographical barrier might have contributed to the different floristic and cytological patterns of these regions.

A significant difference in the proportion of penta- and hexaploid populations across altitude was found within the Western Carpathians and adjacent Pannonia. While pentaploids are more or less evenly present up to 1000 m, the hexaploids usually grow in regions with a warmer climate, usually below 500 m a.s.l. However, there are several regional deviations. Prevailing populations of the pentaploid level are present in the Zemplínske vrchy Mountains (south-east Slovakia) belonging to the warmest region of the Western Carpathians situated in the neighbouring zone with the Pannonian Plain. On the other hand, several hexaploid populations were recorded at a high elevation with a cold and humid climate in the Oravška Magura Mountains. In the Bohemian Massif, rare penta- and hexaploids or mixed populations are confined mostly to the warm, low-elevated, regions, such as river valleys, or to the adjacent zone with the Pannonian Plain (Fig. 2). One hexaploid population was found on the top of the Hrubý Jeseník Mountains (eastern part of the Sudetes range, Czech Republic). It seems that there are at least two hexaploid types in the area studied, differing in distribution and breeding systems (T. Uršu, unpubl. res.). The first one is confined mostly to thermophilous vegetation in the Carpathian Mountains and has an apomictic breeding system; it is probably related to apomictic hexaploids occurring throughout the Carpathian Mountains to the Balkan. The second hexaploid type is confined to relict river valleys in the Bohemian Massif and is sexual; this type is probably related to sexual hexaploids of the Alps (Gadella, 1984).

Amphidiploid origin of Pilosella officinarum?

Diploids of P. officinarum that are considered to have a relict distribution were found by Delcourt (1972) and by Gadella (1972) only in the south-western Alps. However, 16 records of data from the French Alps published by the former author were doubted later due to mis-identification as a closely related but different diploid species from the section Pilosellina – P. peleteriana (Gadella, 1984). Nevertheless, two diploid plants of P. officinarum counted by Gadella might belong to this species, as it is obvious from the photograph of these plants (Gadella, 1972: 362). These diploids originating from a very widely defined locality ‘the valley of Aosta’ (north-west Italy) have long stolons with decreasing leaf size towards the stolon apex. On the other hand, these plants could also represent the hybrids between true P. officinarum and some diploid taxon from P. section Pilosellina. These questionable data on existence of diploids of P. officinarum may suggest that a well-established diploid cytotype within P. officinarum does not really exist in nature. Another fact supports this hypothesis: most of the diploid species of Pilosella that had been counted up to the present, including the closely related taxa from the section Pilosellina, have been found usually in several if not many localities and occupied much wider ranges (e.g. Zahn, 1921–1923; Bräutigam, 1992; Schuhwerk, 1996). Moreover, polyploidy
in diploid taxa of section *Pilosellina* is either unknown or very scattered records of polyploids might be regarded as mis-identifications with *P. officinarum* or interspecific hybrids. The almost exclusive presence of polyploid populations with the tetraploid ones being the commonest leads to the hypothesis that *P. officinarum* is likely to be an amphidiploid species originated from one or more crosses between diploid members of section *Pilosellina*. Both place and time of this hybridization are difficult to estimate. Analysis of ITS sequences showed low differentiation between diploid taxa (Fehrer et al., 2007a), which suggests a relatively low age of the particular members of this group. With respect to chloroplast haplotypes, diploid members of the section *Pilosellina* share both main types. *Pilosella hoppeana*, *P. macrantha* and *P. peleteriana* share the haplotype typical of steppe and mountain species such as *P. onigenesis*, *P. alpícola*, *P. glacialis*, *P. echioides*, etc., i.e. those species, which occurred together during the Glacial Period in steppic and tundra-like habitats in the area of central Europe. The other diploids confined to southern Europe (*P. castellana*, *P. argyrocoma* and *P. pseudopilosella*) share the haplotype with *P. lactucella*, *P. vahlii* and *P. breviscapa* (Fehrer et al., 2007a). Central European populations of *P. officinarum* exhibit the same haplotype as *P. hoppeana* and *P. macrantha* (Krahulec et al., 2004, Fehrer et al., 2007a, b). The close relationship of diploid *P. peleteriana* and *P. officinarum* has been proven by an allozyme pattern observed in the plants originating from Scandinavia (Tyler, 2005). The possible polyphyletic and polytopic origin of polyploid populations of *P. officinarum* is supported by its enormous morphological variation. Zahn, a monographer of the genus, distinguished about 600 subspecies (Zahn, 1921–1923, 1922–1930). Morphologically, the tetraploid plants of *P. officinarum* found recently in Bulgaria resemble hybrids between hexaploid *P. officinarum* and diploid *P. macrantha*. To understand the origin of *P. officinarum* it is necessary to know more about the detailed distribution of haplotypes in the whole distribution area of *P. officinarum* and its diploid relatives. Recently Trewick et al. (2004) included several plants from their natural European range of distribution in their study on the origin of the introduction of *P. officinarum* into New Zealand. They found a mixed distribution of two common chloroplast haplotypes with no clear geographic pattern. However, three rare haplotypes were distributed mainly in the Alps, Sudeten Mountains, the Carpathian Mountains and Finland.

**General pattern of ploidy level distribution in Europe and its relationship with polyploidy and apomixis**

Gadella (1984, 1987, 1991) studied the distribution of particular cytotypes of *P. officinarum* in Europe and tried to explain its pattern. The revision presented in this paper (Fig. 5) showed that this pattern is more complex and fine grained (as was shown in the area of the Slovak and Czech Republics) than that suggested by Gadella. Despite the fact that *P. officinarum* is the most karyologically studied vascular plant species, it is realized that the published data cover only some parts of its natural distribution range sufficiently (western, central and northern Europe). Large areas in southern, south-eastern and eastern Europe have scarce or almost no data.

In total, four different cytotypes of *P. officinarum* were found in Europe (see Table 1 and Fig. 5). The records on diploids are highly questionable (see above). The most common ploidy levels are 4x, 5x and 6x. The range of sexual tetraploid cytotypes clearly separates the higher ploidy levels (5x and 6x) into two groups occurring in geographically different regions – into northern Europe and the mountains of central and south-east Europe (the Alps and the Carpathian Mountains, mountains in Bulgaria). Such a pattern of ploidy level distribution suggests an independent origin of penta- and hexaploids. Concerning the results from the Czech Republic and Slovakia, it seems that they match the general pattern in central Europe. While the prevailing tetraploid cytotype in the Bohemian Massif shows linkage to the tetraploid populations in the western part of central Europe, penta- and hexaploid populations in the Western Carpathians and the Pannonian Plain are likely to be related to the high ploids found in the Alps and in the Balkans (cf. Fig. 5).

It was hypothesized that the prevailing occurrence of high ploids (5x and 6x) correlates with either high latitudes or high altitudes and that their common distribution in northern Europe and in the Alps may be the result of the last Pleistocene glaciation (Gadella, 1984, 1987, 1991). The detailed map given in Fig. 5 shows that the tetraploid level is confined to western Europe and the western part of central Europe and that it has a sub-Atlantic distribution character. Northwards, sexual tetraploids are rare or completely missing and they are replaced by apomictically reproducing penta- and hexaploids (cf. Turesson and Turesson, 1960). The boundary of tetraploids and high ploids matches well with the border of the ice-sheet during the Last Glacial Maximum (cf. Adams, 1997) in the British Isles and it is very close to this geographic position in Scandinavia. Interestingly, the same pattern of cytotype distribution was found in *Parnassia palustris* L. (Parnassiaceae), where the boundary between diploids and tetraploids more or less correlates with the limit of Last Glacial Maximum (Gornall and Wentworth, 1993; Borgen and Hultgård, 2003). It therefore seems that the relationship between the presence of high ploid apomictic plants (5x and 6x) in northern Europe and glaciations might have a real basis and suggests evolutionary advantages of polyploidy associated with apomixis in the colonization of deglaciated areas in Scandinavia (cf. Asker and Jerling 1992). Merxmüller (1975) pointed out that diploid, sexually reproducing taxa of the closely related genera *Hieracium* and *Pilosella* are mostly confined to the southern latitudes, while there was a tendency for polyploids, mostly apomorphic species, to prevail in northerly situated regions. Such geographically limited parthenogenesis is known also in other sexual–apomictic genera and was summarized by Bierzychudek (1985) (for thorough recent revision on complex causality of geographical parthenogenesis, see Hörandl, 2006). The diploid members of section *Pilosellina* have a more restrained range of distribution in...
comparison with polyploid *P. officinarum* and are confined mostly to southern and central Europe (see Introduction). This recalls the situation of several other groups of polyploid vascular plants associated with apomixis, e.g. *Antennaria* L. (Bayer and Stebbins 1987), *Ranunculus auricomus* group (Hörandl, 2006) and *Taraxacum* (den Nijs et al. 1990), where polyploid apomicts tend to have larger ranges than sexuals. In the present case, the colonizing success of *P. officinarum* might be attributed to the combinations of different factors, such as its probable allopolyploid origin (see above), increased heterozygosity and the existence of a high number of genetically different clones, the presence of an apomorphic mode of reproduction in high ploids (5x and 6x) with the occurrence of residual sexuality, vegetative reproduction via above-ground stolons, the possibility of long-distance dispersal via achenes with a pappus, and the opportunity of recurrent formation of novel genotypes via hybridization. It is possible that all these factors have played an important role in shaping the present cytogeographic patterns of *P. officinarum*.

**SUPPLEMENTARY INFORMATION**

Supplementary information is available online at www.aob.oxfordjournals.org and contains the lists of Pilosella officinarum localities accompanied by geographical co-ordinates, ploidy levels and/or chromosome numbers (and references) for data (1) presented in this study for the areas of the Czech Republic, Slovakia and Hungary; (2) for previously reported data from the Czech Republic and Slovakia; (3) for previously published data from the rest of European area; and (4) for new data from Europe outside of Slovakia, Czech Republic and Hungary.

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**LITERATURE CITED**


