Polyploid evolution: The ultimate way to grasp the nettle

Ludmila Rejlova1,2*, Jindřich Chrtek1,2, Pavel Trávníček1, Magdalena Lučanová1,3, Petr Vít1,4, Tomáš Urfus1,2*

1 Institute of Botany, The Czech Academy of Sciences, Průhonice, Czech Republic, 2 Department of Botany, Faculty of Science, Charles University, Prague, Czech Republic, 3 Department of Botany, Faculty of Science University of South Bohemia, České Budějovice, Czech Republic, 4 Faculty of Environmental Sciences, Czech University of Life Sciences Prague, Prague, Czech Republic

* rejlovalud@gmail.com (LR); tomas.urfus@natur.cuni.cz (TU)

Abstract

Polyploidy is one of the major forces of plant evolution and widespread mixed-ploidy species offer an opportunity to evaluate its significance. We therefore selected the cosmopolitan species Urtica dioica (stinging nettle), examined its cytogeography and pattern of absolute genome size, and assessed correlations with bioclimatic and ecogeographic data (latitude, longitude, elevation). We evaluated variation in ploidy level using an extensive dataset of 7012 samples from 1317 populations covering most of the species’ distribution area. The widespread tetraploid cytotype (87%) was strongly prevalent over diploids (13%). A subsequent analysis of absolute genome size proved a uniform Cx-value of core U. dioica (except U. d. subsp. cypria) whereas other closely related species, namely U. bianorii, U. kiovien-sis and U. simensis, differed significantly. We detected a positive correlation between relative genome size and longitude and latitude in the complete dataset of European populations and a positive correlation between relative genome size and longitude in a reduced dataset of diploid accessions (the complete dataset of diploids excluding U. d. subsp. kurdistanica). In addition, our data indicate an affinity of most diploids to natural and near-natural habitats and that the tetraploid cytotype and a small part of diploids (population from the Po river basin in northern Italy) tend to inhabit synanthropic sites. To sum up, the pattern of ploidy variation revealed by our study is in many aspects unique to the stinging nettle, being most likely first of all driven by the greater ecological plasticity and invasiveness of the tetraploid cytotype.

Introduction

Polyploidy, sometimes referred to as whole-genome multiplication, is generally considered a major force in plant evolution, producing novelties which may eventually lead to single-step speciation, that is, saltation [1–4]. Moreover, the substantial success of angiosperms, the largest clade of land plants, is attributed to polyploidy [5]. Probably 15% (but at least 2–4%) of all speciation events in angiosperms are estimated to have involved polyploidization [6, 7]. Different
ploidy levels can either correspond to already discrete lineages or species [8], or constitute intraspecific variation [9]. Newly established polyploid lineages frequently undergo subsequent diploidization [10–12], which is usually followed by genome downsizing [13–15]. Nevertheless, distinct cytotypes frequently coexist in sympatry and, according to the current state of knowledge, at least 16% of all vascular plant species consist of multiple cytotypes [16].

Polyploidy directly affects a number of key biological features (e.g. cell and plant size and duration of mitosis) ultimately associated with distinct physiology and ecology [17, 18]. Such novelties frequently result in improved adaptation potential, fitness, etc. [19–21], which is further mirrored, for example, by broader ecogeographic and climatic niches of polyploids compared to their diploid or lower-ploidy progenitors [22–24]. Specific features of plants have repeatedly been linked to polyploidy (e.g. phenology, mycorrhizal colonization, pollinator behaviour, herbivore predation, salinity tolerance and migration potential [25–30]). Moreover, polyploid cytotypes tend to inhabit a broad range of synanthropic habitats, in contrast to their diploid congeners (Arabidopsis arenosa (L.) Lawalrée, Cardamine amara L., Centaurea stoebe L., Solidago gigantea Ait., etc. [31–35]), and their greater ecological plasticity and synanthropic affinity can increase their invasive potential [36, 37].

The essential first step when gaining insight into the evolution of polyploid plants is cyto-geography, the study of cytotype diversity and its past and predicted future distribution patterns [13]. Knowledge of the cytotype distribution pattern usually reveals phenomena such as environmental segregation or reproductive isolation of cytotypes [38–40].

Despite the undisputed evolutionary significance of polyploidy, there is a lack of comprehensive cytogeographical studies, with only a few focusing on widespread weedy plants [41] even though they represent highly suitable model taxa for investigating the evolutionary potential of polyploids (e.g. Mercurialis annua L., Tripleurospermum inodorum (L.) Sch. Bip., Senecio inaequidens DC. [42–45]). Surprisingly, the stinging nettle Urtica dioica L., one of the most troublesome polyploid weeds, remains considerably understudied, despite being highly important in agriculture, the textile and cosmetics industries [46–48], and medicine [49, 50]. The species represents a nitrophilous, synanthropic and invasive species with a cosmopolitan distribution [51–53]. Urtica dioica is characterized by huge variation mirrored by a high number of intraspecific taxa distinguished either solely based on morphological characters (e.g. various types of indumentum [54–56]) or with consideration for sexual morphs (predominantly stochastically occurring [54, 55, 57]). Finally, polyploidy is a truly substantial source of variation in U. dioica. Published diploid chromosome counts frequently refer to plants from relict or semi-natural habitats (e.g. alluvial forests [58–60]) whereas tetraploids have been reported to occur in habitats of various types, even highly synanthropic ones [58, 61–63]. However, even though relatively many chromosome counts have been published (e.g. [64]), the distribution pattern has so far only been studied marginally and locally. Moreover, ploidy levels were not directly considered in recent phylogenetic reconstructions [65–68].

We have adopted the only current taxonomic treatment of U. dioica consisting of several subspecies. Apart from the nominate tetraploid and widely distributed subspecies U. d. subsp. dioica, all subspecies are supposed to be diploid and somewhat restricted in their distribution area: U. d. subsp. kurdistanica (found in alpine habitats of Anatolia and Near Eastern mountain ranges [69, 70]), U. d. subsp. pubescens (scattered in lowlands from Italy across the Balkan Peninsula to the delta of the river Volga [71–73]), U. d. subsp. sondenii (tundra marches [59, 73, 74]) and U. d. subsp. subinermis (alluvial forests, floodplain forests [54, 55, 71, 75]). Finally, the unique steno-endemic U. dioica subsp. cypria is treated as a subspecies of U. dioica (a single population in Cyprus, population UP1219; S2 Table) even though its morphology is distinct [72, 76]. However, the infraspecific phylogeny of U. dioica is still largely unresolved [68] and the ranks of its infraspecific taxa also remain a matter of debate [58]. Three recent phylogenies
[65, 67, 68] also place four other taxa within the crown clade of *Urtica* (corresponding to *U. dioica*): *U. atrovirens* Req., *U. bianorii* (Knoche) Paiva, *U. kioviensis* Rogow. and *U. simensis* Hochst. ex A. Rich.

The present study aims to assess the ploidy and genome size variation within *U. dioica* across Europe (with contiguous areas of West Asia). We placed particular emphasis on the following questions: (1) What is the general cytogeographic pattern of *U. dioica* in Europe (with contiguous areas of West Asia)? (2) Is genome size a suitable taxonomic marker for resolving current taxonomic ambiguities? (3) Do certain cytotypes occur in particular habitats?

### Materials and methods

#### Materials

**Plant material.** Plants were collected between 2012 and 2018 at 1317 localities (1305 localities of *U. dioica* and 12 localities of closely related species) across Europe and West Asia (Fig 1, S1 Table, S1, S2 and S6 Figs). Although the sampling was primarily random, we focused partially on relict and semi-natural habitats (e.g. ravine and alluvial forests, alpine vegetation and tundra marches, Mediterranean mountains) because (partly allegedly) diploid taxa (*U. d. subsp. kurdistanica*, subsp. *pubescens*, subsp. *sondenii*, subsp. *subinermis*) were often reported from such habitats [54, 55, 59, 69–75]. In total, 7012 plants (6977 individuals of *U. dioica* and 35 individuals of closely related species) were sampled (5–10 plants per population; the distance between sampled plants was at least 3 m to avoid re-sampling of the same clone). As a rule, fresh leaves were used for flow cytometric analyses, in some cases silica-gel-dried leaves were used (~10% of samples). A subset of plants was transferred to the experimental garden of the Institute of Botany of the Czech Academy of Sciences in Průhonice (N49.99474, E14.56617, 320 m a.s.l.) for further cultivation and chromosome counting. Voucher specimens will be deposited in the Herbarium of the Charles University, Prague (PRC). GPS coordinates, the elevation and type of habitat were recorded for each population (S1 Table). The study did not necessitate any specific permissions and did not involve endangered or protected species.

#### Methods

**Flow cytometry.** Cytotypes were identified by means of flow cytometry, a technique enabling us to analyse large numbers of samples over a short period and to collect appropriate many samples of all taxa and cytotypes [77]. Relative genome size was ascertained for all plants (S1 Table) and absolute genome size was estimated for a subset of samples (Table 1).

Sample preparation followed a simplified two-step protocol [78]. A part of a petiole was chopped together with the internal reference standard *Bellis perennis* L. (2C = 3.38 pg; [79]) using a sharp razor blade in a plastic Petri dish containing 500 μl of the ice-cold buffer Otto I (0.1-M monohydrate citric acid and 0.5% Tween 20). The suspension was filtered through a 42-μm nylon mesh and the isolated nuclei were stained for 5 minutes with 1 ml of the buffer Otto II (0.4-M Na₂HPO₄ · 12H₂O) supplemented with the fluorochrome 4’,6-diamidino-2-phenylindole (DAPI; final concentration 4 μg · ml⁻¹) and β-mercaptoethanol (final concentration 2 μl · ml⁻¹).

Absolute genome size was estimated using the intercalating fluorochrome propidium iodide (PI) supplemented with RNase IIA (both at final concentrations of 50 μg · ml⁻¹). Each sample were analysed three times on three consecutive days to rule out diurnal fluctuation. If the deviation among all particular measurements of the same individual exceeded the threshold of 3%, additional analyses were conducted [78].

To assess heteroploid hybridization, seeds from the mixed population (i.e. population UP0466) were also analysed. Achenes were removed from the pericarp and chopped in the
Fig 1. Distribution of two dominant cytotypes of *Urtica dioica* in Europe and West Asia. Map of all samples based on flow cytometric analyses of 1305 populations. The size of the circles reflects the number of populations.

https://doi.org/10.1371/journal.pone.0218389.g001

Table 1. Summary of absolute genome size of *Urtica dioica* and closely related species (2C-values in pg) and detected numbers of (somatic) chromosomes.

| Taxon                                | No. of individuals analysed/No. of populations | Mean Cx-value (pg) ± SD | 2C-value range (pg) | Chrom. number (2n) | Difference compared to 2x (%) | Difference compared to 4x (%) |
|--------------------------------------|-----------------------------------------------|------------------------|---------------------|---------------------|------------------------------|------------------------------|
| *U. d. subsp. dioica*                | 32/27                                         | 0.55 ± 0.04²           | 2.08-2.20           | 52                  | –                            | –                            |
| *U. d. subsp. dioica* – 3x           | 1/1                                           | 0.54                   | –                   | 39                  | –                            | –                            |
| *U. d. subsp. kurdistanica*          | 6/3                                           | 0.59 ± 0.01²⁴          | 1.15-1.20           | –                   | –                            | –                            |
| *U. d. subsp. pubescens*             | 16/14                                         | 0.58 ± 0.03²⁴          | 1.10-1.21           | 26                  | –                            | –                            |
| *U. d. subsp. sondenii*              | 4/2                                           | 0.57 ± 0.01²⁴          | 1.12-1.15           | 26                  | –                            | –                            |
| *U. d. subsp. subinermis*            | 19/13                                         | 0.58 ± 0.03²⁴          | 1.10-1.25           | 26                  | –                            | –                            |

**Closely related species:**

| Taxon                                | No. of individuals analysed/No. of populations | Mean Cx-value (pg) ± SD | 2C-value range (pg) | Chrom. number (2n) | Difference compared to 2x (%) | Difference compared to 4x (%) |
|--------------------------------------|-----------------------------------------------|------------------------|---------------------|---------------------|------------------------------|------------------------------|
| *U. atrovirens*                      | 4/3                                           | 0.60 ± 0.004²³         | 1.18-1.19           | –                   | 3.5                          | 45.4                         |
| *U. bianorii*                        | 1/1                                           | 0.83²⁴                 | –                   | –                   | 43.5                         | 24.3                         |
| *U. d. subsp. cypria*                | 6/1                                           | 0.55 ± 0.005²⁴         | 1.65-1.67           | –                   | 44.4                         | 23.9                         |
| *U. kiovienis*                       | 5/4                                           | 0.71 ± 0.025²³         | 1.36-1.43           | –                   | 22.6                         | 35.3                         |
| *U. simensis*                        | 1/1                                           | 0.74²³                 | –                   | –                   | 28.7                         | 32.1                         |

* Different letters indicate groups of taxa that are significantly different in Tukey HSD test.

https://doi.org/10.1371/journal.pone.0218389.t001
single-phase seeds buffer LBO1 (15M Tris, 2M Na$_2$EDTA, 0.5M spermine tetrahydrochloride, 80M KCl, 20M NaCl, 0.1% Triton X-100, stored at −20˚C [80]) together with the fluorochrome 4',6-diamidino-2-phenylindole (DAPI) and β-mercaptoethanol.

All samples were incubated for 5–10 minutes at room temperature before being run through each of two flow cytometers (relative genome size: CyFlow ML equipped with a 365-nm UV LED as the light source; absolute genome size: CyFlow SL with a diode-pumped 532-nm solid-state green laser; both Partec GmbH, Münster, Germany). The resulting histograms were evaluated in Partec FloMax 2.3 software (Partec GmbH, Münster, Germany). Only analyses providing peaks with a coefficient of variation of less than 3% for fresh and 5% for silica-dried material were processed further.

One-way analysis of variance (ANOVA), followed by Tukey’s honest significant difference (HDS) test, was used to test the significance of genome size differences between the taxa analysed. Values of genome size were log-transformed before the analysis. All statistical analyses were performed and all plots were produced in the R statistical environment [81].

Chromosome counts. Chromosome counts were determined from root tips of germinating seeds and cultivated individuals. Selected samples were processed according to the modified protocol of Mandáková & Lysak [82].

Fresh roots (~1 cm long) were put into 1.5-ml Eppendorf tubes with distilled water and placed into a container with ice-flakes for 24 hours. Afterwards they were put into a freshly prepared fixative (ethanol: acetic acid, 3: 1, v: v) and stored overnight in a refrigerator (~4˚C). The material was stored at −20˚C in the fixative until further use.

The root tips were washed twice in distilled water (each time for 5 min), then a citrate buffer was applied and roots were washed in an orbital shaker (twice for 5 min). Subsequently, the buffer was sucked out of the sample and a 0.3% mixture of pectolytic enzymes (pectolyase, cellulase, cytohelicase) was added, followed by incubation in an incubator (37˚C, 120 min). Then the enzyme mixture was replaced with the same citrate buffer.

The white tip of the root meristem was cut under a stereomicroscope, excess buffer was removed, and the sample was sprinkled with 60% acetic acid with an incubation time of 1–2 min. The root meristem was disintegrated using dissecting needles and the obtained meristematic suspension was covered by a cover-slip. The slide was moved 2–3 times above a flame and then the material was carefully squashed.

The slides were placed into a freezer (~−80˚C) and after 10 minutes in the freezer the cover-slips were separated from the slides by razor. The samples were subsequently dyed with 15 μl of Vectashield with 4’,6-diamidino-2-phenylindole (DAPI). The preparations were covered with new cover-slips and fixed with nail polish.

Chromosomes were observed under a Nikon Eclipse E600 microscope equipped with a Nikon DS-Qi1Mc camera, and images were acquired using NIS-Elements AR software.

Ecological relations. We used exactly recorded locations of all populations to get a basic grasp of the ecological preferences of major cytotypes (the diploid and tetraploid cytotypes of U. dioica). To evaluate the ecological relations of major cytotypes, we applied simple modelling using the Bioclim algorithm according to Chumová et al. [83]. In the first step, georeferenced data were spatially stratified to avoid discrepancies caused by unequal sampling (R package ‘spThin’, Aiello-Lemmens et al. [84]; a 20-km and 5-km threshold distance for tetraploid and diploid population, respectively, was used). The resulting 576 localities were used for the extraction of bioclimatic data. Data from raster layers for all 19 bioclimatic variables were extracted using the ‘extract’ function in the ‘raster’ R package [85]. Principal trends in the variation of bioclimatic variables were detected by PCA. Mutually uncorrelated variables were identified by stepwise forward selection and subjected to linear discriminant analysis. All analyses were conducted using the ‘MorphoTools’ R package for multivariate data handling [86].
In addition, correlations of relative fluorescence intensity value with elevation (dataset divided into two elevation ranges: 0–500 and >500 m above sea level), latitude and longitude were quantified by fitting a linear or quadratic function. To assess the affinity of both cytotypes to human-affected habitats, we adopted a four-level scale of synanthropy (sensu Tunxen 1956 [87]), which was arbitrarily assigned to each sampling locality. Subsequently, Pearson’s chi-square test [88] was used to determine the dependence between the degree of synanthropy and ploidy level.

Results
Flow cytometry
We determined the relative genome size of 6836 plants from 1295 populations (176 individuals were excluded due to poor-quality flow-cytometric histograms). Our results confirm the occurrence of two dominant DNA ploidy levels having the following mean relative fluorescence intensity values (± SD): 2x = 0.30 ± 0.01 (range: 0.24–0.36, 50% variation, n = 849), 4x = 0.57 ± 0.01 (range: 0.54–0.64, 18.5% variation, n = 5975). The average coefficient of variance (CV) was 1.68% (particular CVs are given in S1 Table). The tetraploid cytotype strongly prevailed over the diploid one (2x = 13%, 4x = 87%). Diploids were found frequently in mixed populations with prevailing tetraploids. For the first time we managed to detect a few triploid (8) and pentaploid (4) individuals in a mixed-ploidy population of diploids and tetraploids (3x = 0.44 ± 0.01 (range: 0.42–0.46, 9.5% variation, n = 8), 5x = 0.73 ± 0.02 (range: 0.71–0.77, 8.5% variation, n = 4); Fig 2, S3 and S4 Figs).

To assess the potential for heteroploid hybridization, we analysed 70 achenes from the mixed population (i.e. population UP0466; S1 Table). From diploid maternal plants (33 seeds overall), 82% of the progeny (27 achenes) was diploid (with 3x endosperm) and 18% (6) triploid (4x endosperm). Tetraploid maternal individuals (37 seeds overall) produced 94% (35 achenes) of tetraploid seeds (with 6x endosperm) and one triploid (5x endosperm) and one pentaploid seed (9x endosperm, S5 Fig).

To calibrate the measurements and detect differences between particular diploid subspecies, we also estimated absolute genome size for a reduced set of accessions (78 plants from 60 populations—U. dioica; 17 plants from 10 populations—closely related species; S2 Table, S6 Fig). Core diploid subspecies (U. d. subsp. kurdistanica, subsp. pubescens, subsp. sondenii and subsp. subinermis) did not differ from each other in absolute genome size whereas the other closely related species (i.e. species closely related to U. dioica crown clade in recent phylogenies) U. bianorii, U. d. subsp. cypria, U. kioviensis and U. simensis had significantly greater DNA content. Only U. atrovirens, which is also ranked among the closely related species of U. dioica, was assigned to the group of core diploid subspecies. We, for the first time, estimated the absolute genome size of U. d. subsp. kurdistanica, U. d. subsp. cypria and the triploid cytotype of U. dioica (a plant morphologically identical with U. d. subsp. dioica). Absolute genome size was determined for all the mentioned species and subjected to ANOVA and Tukey’s HDS test (p <0.001; Fig 3, Table 1, S7 Fig).

Chromosome counts
All DNA ploidy levels were verified by subsequent chromosome counts. Ten plants were checked for their chromosome numbers using fluorescent karyology. The chromosome number of 2n = 26 was ascertained for diploids (three plants classified as U. d. subsp. pubescens, subsp. sondenii and subsp. subinermis, respectively), 2n = 39 was ascertained for triploids (one plant morphologically identical with U. d. subsp. dioica), 2n = 52 was ascertained for
tetraploids (five plants assigned to *U. d. subsp. dioica*) and 2n = 65 was ascertained for pentaploids (one plant also morphologically identical with *U. d. subsp. dioica*; Fig 4, Table 1).

**Ecological relations**

**Bioclimatic and geographic pattern.** To verify the habitat and ecological preferences of individual ploidy levels, we used basic modelling. Our analysis of bioclimatic data in relation to the ploidy levels of individual populations and their exactly recorded positions shows that the variability of individual ploidy levels is interdependent. This is also confirmed by field observations, as in most cases diploid individuals grow in mixed populations with the tetraploid cytotype, which evinces a high degree of plasticity (e.g. ecological or morphological) shared with diploid individuals. In the stepwise selection analysis, the following features were the most contributing to group separation: BIO3 (Isothermality = BIO2/BIO7 × 100; this quantifies how much day-to-night temperatures oscillate relative to the summer-to- winter (annual) oscillations), BIO5 (Max Temperature of Warmest Month) and BIO17 (Precipitation of Driest Quarter; Fig 5).

We found a positive correlation between relative fluorescence intensity and longitude (cor = 0.161, p < 0.001; Fig 6A) and latitude (cor = 0.133, p < 0.001; Fig 6B) in the complete dataset of European populations. In the reduced dataset of diploid accessions (the complete dataset of diploids excluding *U. d. subsp. kurdistanica*), we detected a positive correlation between relative fluorescence intensity and longitude (cor = 0.296, p < 0.001). Diploid taxa growing in relict habitats preferred lower elevations compared to the ubiquitous tetraploid cytotype. Correlations of relative fluorescence intensity with elevation were significant for each of the datasets fitted with a linear function (0–500 m above sea level: cor = −0.305, p < 0.001; >500: cor = 0.344, p < 0.001) and in all data fitted with a quadratic function (cor = 0.208, p < 0.001; Fig 6C).
Affinity to synanthropic habitats. To determine habitat preferences, especially of relict diploids, we used the data from assessment of individual locations. Using Pearson’s chi-squared test, we have determined with a high degree of confidence that the probability of occurrence of a diploid population depends on the type of environment ($p < 0.001$). The distribution of diploid and tetraploid populations with respect to the environment is presented as in a contingency table (Table 2, Fig 7), along with associated standard residuals. The diploid cytotype of *U. dioica* (*U. d. subsp. kurdistanica*, *subsp. pubescens*, *subsp. sondenii*, *subsp. subinermis*) tends to occur in less human-affected habitats (habitat type 3 and 4 on the four-level scale of synanthropy; Table 3). A special case is the diploid subspecies *U. d. subsp. pubescens* from the Po river basin, which occurs exclusively in highly synanthropic and strongly human-affected locations (habitat type 1 and 2 on the four-level scale of synanthropy; mode value for all diploid subspecies of *U. dioica*: 3; mode value for *U. d. subsp. pubescens*: 2). This stands in contrast to the tetraploid cytotype, which occurs in habitats of all types, although it prefers environments with an increased degree of synanthropy.

Discussion

During our large-scale screening of *Urtica dioica* across Europe and West Asia, we found two major ploidy levels: widely distributed tetraploids and less frequent diploids. We have not
proved any strong correlation supporting either the generally suggested hypothesis that polyploids are more abundant at higher elevations and latitudes or the idea that diploids are confined to Southern European glacial refugia whereas polyploids occur across broader geographic ranges [89], often shifted to harsh environments. Instead, our results suggest that diploid plants show some degree of affinity to habitats less affected by human activities, in contrast to tetraploids, which tend to grow in human-made or strongly influenced habitats. Furthermore, our study has revealed a significant difference in absolute genome size between *U. dioica* and its closely related species *U. bianorii, U. dioica* subsp. *cypria, U. kioviensis* and *U. simensis* [72, 76, 90–95].

We are aware that the frequency of diploids detected would be significantly lower had we chosen an entirely random sampling strategy instead of partly preferentially targeting relict and natural habitats. Additional occurrences of the alluvial diploid cytotype (~*U. d. subsp. subinermis*) can be expected in Western Europe (especially in France and the United Kingdom). On the other hand, we sampled numerous relict habitats in Spain and northern Iran, and detected only tetraploids there, so the occurrence of a diploid cytotype is, in concordance with previously published chromosome counts [94, 96, 97], less probable in these two countries.

**Major and minor cytotypes of *Urtica dioica***

The widely distributed tetraploids and the less frequent diploids possess the chromosome numbers of 2n = 2x = 26 and 2n = 4x = 52, respectively, ascertained here and also reported
However, for the first time we managed to capture a small percentage of very rare triploid (8 individuals) and pentaploid (4 individuals) cytotypes, both in mixed-ploidy populations of diploids and tetraploids. The origin of these minor cytotypes is discussed below.

For some plants, we detected abnormal values of relative genome size (8 tetraploid individuals, range of 2C pg: 1.66–1.74), which could be explained by aneuploidy. The values might correspond to a loss of four chromosomes, i.e. to the frequently reported chromosome number 2n = 48 [61–64, 98–102]. Unfortunately, we did not succeed in cultivating any of these aberrant individuals, so we cannot confirm any hypothetical aneuploid counts.

Data from our screening of seeds from both diploid and tetraploid maternal plants from a mixed-ploidy field population in southern Moravia (south-eastern Czech Republic—UP0466; S1 Table) suggest that gene flow between the two cytotypes can occur. Besides the ploidy level of the embryo, we also paid special attention to the ploidy of the endosperm in order to decipher the contribution of the paternal cytotype and thus to determine the seed formation pathway [80, 103]. The greater frequency of triploid embryos in seeds of diploid maternal plants might be in line with the greater frequency of tetraploid plants and thus the larger greater amount and pressure of diploid pollen grains (from tetraploid plants). Another explanation, not mutually exclusive with the previous, supposes that the spatial pattern of (often large) male and female clones of both cytotypes at the site may play a role. Although our data indicate the origin of triploid seeds via heteroploid crosses, we cannot fully exclude the ability of diploid
plants to produce triploid seeds via unreduced gametes (i.e. reduced gamete fusion with a male unreduced gamete from diploid plants), in general, the formation of unreduced gametes is not frequent [104, 105]. The frequency of triploid seeds (18% from diploid and 3% from tetraploid maternal plants) also contradicts the frequency of adult triploid plants in mixed diploid-

Table 2. Contingency table with standard residuals of diploid and tetraploid populations depending on habitat type.

| Ploidy level | Environment type 1 | Environment type 2 | Environment type 3 | Environment type 4 |
|--------------|---------------------|---------------------|---------------------|---------------------|
| Diploids (2x) | 1 (-3.90)           | 81 (-0.04)          | 182 (1.38)          | 4 (3.07)            |
| Tetraploids (4x) | 57 (3.90)          | 278 (0.04)          | 580 (-1.38)         | 1 (-3.07)           |

https://doi.org/10.1371/journal.pone.0218389.t002

Fig 6. Relations between relative fluorescence intensity and longitude, latitude and elevation fitted with a linear or quadratic function. (A) Correlation of relative fluorescence intensity with longitude fitted with a linear function (complete dataset of European populations); (B) Correlation of relative fluorescence intensity with latitude fitted with a linear function (complete dataset of European populations); (C) Correlation of relative fluorescence intensity with elevation fitted with a quadratic function.
tetraploid populations (8 individuals). The most plausible explanation seems to be a triploid block (lower fitness of or strong selection against triploid seedlings, or lower germination rates of triploid seeds, or their inability to germinate, compared to diploid and tetraploid ones \[106\]). The detection of a pentaploid individual (with 9x endosperm) in the offspring of a tetraploid plant indicates the formation of an unreduced gamete at the 4x level and its fusion with a reduced (x) gamete from a diploid plant. Alternatively, pentaploids might originate from crosses between tetraploids and hexaploids, but we detected neither adult hexaploid

tetraploid populations (8 individuals). The most plausible explanation seems to be a triploid block (lower fitness of or strong selection against triploid seedlings, or lower germination rates of triploid seeds, or their inability to germinate, compared to diploid and tetraploid ones \[106\]). The detection of a pentaploid individual (with 9x endosperm) in the offspring of a tetraploid plant indicates the formation of an unreduced gamete at the 4x level and its fusion with a reduced (x) gamete from a diploid plant. Alternatively, pentaploids might originate from crosses between tetraploids and hexaploids, but we detected neither adult hexaploid

Table 3. Evaluation of the affinity of particular _Urtica dioica_ cytotypes to human-affected habitats.

| Level | Vegetation types and habitats | Degree of influence by man |
|-------|-------------------------------|---------------------------|
| 1     | intensively managed habitats (agricultural, ruderal, etc.), road margins, urbanized areas | highly nitrophilous and intensively human-affected locations |
| 2     | extensively cultivated landscapes, agricultural marginal habitats, cultivated and plantation-like forests | partly synanthropic and extensively cultivated locations |
| 3     | semi-natural vegetation, recent vegetation +/- corresponding to the potential natural vegetation * | semi-natural habitats |
| 4     | tundra marches, Mediterranean alpine zones, natural alluvial associations, other relict habitats | least human-affected habitats (primary habitats) |

* sensu Tüxen 1956 \[87\]

https://doi.org/10.1371/journal.pone.0218389.t003
plants nor pentaploids with the embryo: endosperm ploidy ratio indicating this hybridization history (5x embryo: 7x endosperm). A combination of a more extensive seed screen (incl. experimental hybridization) and molecular analyses should be carried out to assess the rate of gene flow. Nevertheless, we have confirmed the possibility of heteroploid hybridization, which might cause genetic erosion and therefore pose a threat to the far less abundant diploid populations.

**Diploids as indicators of natural habitats versus synanthropic invasive tetraploids?**

We detected geographically stratified elevational and ecological segregation. In Central Europe, the Balkans and the Baltic region, diploids are likely confined to lowland alluvial, especially white willow, gallery forests. In addition, river banks and the surroundings of water bodies, together with forest-tundra stands and ravine forests, are the predominating habitats of diploids in Northern Europe. By contrast, diploids in Anatolia tend to occupy natural habitats at higher elevations (e.g. screes). The species assembly of ancient Central European semi-natural alluvial forests was formed in the Early Holocene. Since the Neolithic period, the floodplains of lowland rivers experienced vast changes caused by erosion, soil deposition and eutrophication. The human-driven decline of woodlands, especially in the Medieval period, and changes in species composition led to the fragmentation of semi-natural woodlands, which are currently confined to more or less small patches within agricultural landscapes [107, 108]. The diploid cytotype of *U. dioica* is restricted to well preserved alluvial forests in Central and Western Europe, so diploids may also indicate relict habitats of this type. The rather narrow ecological niche of diploids compared to tetraploids might indicate, besides other phenomena, ploidy-related drought tolerance and greater plasticity in polyploids allowing tetraploid to occupy a broader spectrum of habitats [109–111]. Similar ecological diploid-polyploid differentiation has been described in the grass species *Deschampsia cespitosa* (L.) P. Beauv. (tussock grass) in Britain [112] and *Dactylis glomerata* L. (cock’s-foot) in Spanish Galicia [22]. Diploids appeared to be restricted mainly to low-density forest-floor habitats in woodlands of mostly ancient, semi-natural origin whereas tetraploids were found in varied habitats, but they predominated in open places such as in meadows, pastures, plantations, their verges and waste grounds. Based on our observations, both cytotypes are ecologically differentiated, but tetraploids do not exhibit local adaptation. Instead, they have greater fitness across both diploid- and polyploid-occupied regions.

In contrast to diploids in Central Europe, Anatolian diploids occur at higher elevations compared to tetraploids and also in different habitats. Instead of European lowland and ravine forest they mostly occur in mountain screes. Analogously to Europe, however, the vast majority of Anatolian populations (from various habitats, including screes) are formed by tetraploids. Surprisingly, even in Anatolia a single diploid was found in an alluvial population (Cappadocia—population UP0038; S1 Table).

In general, however, we have not confirmed the frequently made assumption that polyploids are more abundant at higher elevations and latitudes because of their potentially greater ecological tolerance and colonization ability [1, 17, 113–116]. However, considering only the invasiveness of polyploids, our results are well in agreement with general suggestions. The widespread tetraploid cytotype of *U. dioica* is also often supposed to be an allopolyploid or a group of allopolyploids with different evolutionary histories (e.g. [117]). Polyploidyization and hybridization likely went hand in hand, resulting in rapid divergence of the neopolyploid. Tetraploids were possibly predisposed to spread into ranges thanks to their potential for subsequent adaptation due to greater genetic diversity, higher survival rates and better fitness...
ascribed to the heterosis effect, restoring sexual reproduction following hybridization [37]. A more or less stable occurrence of diploids in semi-natural habitats and tetraploids in mainly human-made habitats, together with a recent spread of tetraploids, has also been reported for Centaurea stoebe [32, 33, 118] and Seseli libanotis (L.) W.D.J.Koch [119]. Although a positive correlation between invasiveness and ploidy seems to be in conflict with a negative correlation between invasiveness and genome size [120], it is their interaction that underlies their actual effects on plant phenotypes and physiology, and, ultimately, on invasion success [36].

**Taxonomic consequences**

In the two most recent phylogenies [67, 68], the crown clade of Urtica (predominantly formed by U. dioica) consist of different additional related taxa, depending on the molecular markers used. Based on a concatenated tree (combining nuclear and plastid markers [68]), U. dioica in the strict sense, an exclusively Eurasian group including U. dioica (except for subsp. cypria), U. kioviensis from western Eurasia and U. platyphylla Wedd. from Northeastern Eurasia is a sister group to Mediterranean endemics (U. atrovirens, U. bianorii) and two African species (U. massaica Mildbr., U. simensis). Together these taxa form a well-supported cluster. Our genome size data partly support this concept. Urtica bianorii and U. d. subsp. cypria clearly fall outside of the U. dioica s.str. group in published phylogenies, which is in concordance with our genome size data. Urtica kioviensis, which could not be separated from U. dioica s.str. in previous phylogenies [68], could be reliably separated from the Eurasian U. dioica s.str. clade using genotyping-by-sequencing data [121], and this separation is well supported and justified by our genome size estimations. Only U. atrovirens, which is ranked close to U. dioica s.str., did not significantly differ from diploid subspecies of U. dioica even though it is distinctive morphologically [93, 94]. We have thus confirmed that genome size can significantly contribute to the delineation and detection of taxa, and that differences between genome size values may be indicative of genetic distance (see e.g. [103, 122, 123]).

In addition to other already discussed reasons to recognize several intraspecific taxa of U. dioica at the subspecies level (i.e. extreme morphologic forms and sexual morphs), polyploidy evidently shapes the structured pattern confining diploid cytotypes to relict habitats (e.g. alluvial forests, tundra marches or Mediterranean alpine zones). The diploid subspecies (subsp. kurdistanica, subsp. pubescens, subsp. sondenii and subsp. subinermis) are more or less morphologically, ecologically and geographically defined and capture a considerable part of the morphological diversity present in Western Eurasian Urtica dioica. However, any clear delineation of some of them is anything but straightforward and even molecular approaches have failed to resolve infraspecific relationships [67, 68]. Although published chromosome counts/ploidy levels are very scarce, ploidy is widely accepted as a trait in the delineation of Urtica dioica subsp. dioica (tetraploid) and the rest of the subspecies [56]. Here we generally confirm that the diploid level (with the chromosome number of 2n = 26) is associated with plants morphologically assigned to U. d. subsp. subinermis, U. d. subsp. sondenii, U. d. subsp. pubescens and U. d. subsp. kurdistanica. We did not find any significant differences in genome size between the subspecies, so genome size cannot serve as a supportive character in the delineation of homoploid taxa as in some another plant groups [123–126].

One particular matter for debate is the delimitation and geographic distribution of U. d. subsp. pubescens. Geltman [127, 128] regards it as an endemic of wetland territories in the Volga delta and its surroundings and in the lower Dnieper region whereas in its wide circumscription the species occupies a geographic area spanning Southern and Eastern Europe, western Turkey [72], Georgia and Azerbaijan [121]. According to Weigend [71] it can be identified by its green-grey leaf colour, a distinctly hairy stem and leaves on both sides, and based on the
ratio of the width to the length of the lamina. However, minor morphological differences between populations (unpublished data) and, in addition, genetic differences between European and West Asian populations [121] may indicate a mosaic-like structure and different evolutionary histories within subsp. *pubescens* in its broad circumscription. We carried out an extensive screening of ploidy levels in populations of 'hairy' nettles from the Po river basin (northern Italy), tentatively assigned to *U. d.* subsp. *pubescens.* Across the basin and in adjacent mountain valleys, we found mostly diploid plants, even though this area is surrounded by expanses dominated by tetraploids (even from the south, i.e. on the slopes of the Apennines). Worth mentioning are two aspects: First, this is the only large area in our study that is most likely occupied nearly exclusively by diploid plants (S1 Fig); otherwise, diploids occur as a rule in mixed-ploidy populations, accompanied by tetraploids. Second, the Po river diploids regularly occur both in a wide range of highly synanthropic types of habitats and in semi-natural alluvial vegetation. Diploid populations might have survived the last glaciation in an refugium extending along the lower elevations of the southern Alps and in adjoining areas, as demonstrated for many alpine plants as well as for beech (*Fagus*) and some insect species [129–132]. Alternatively, diploids might have survived in more southerly located refuges in the Apennine Peninsula [133, 134]. In any case, the Po river diploids definitely deserve a further biosystematic/taxonomic evaluation.

The genome size of *U. d.* subsp. *kurdistanica* corresponds to 2n = 26 (diploid level)—the same as in the other diploid subspecies. However, chromosome counts are not available for this subspecies, so certain deviations from this number cannot be fully excluded. Nevertheless, ours is the first DNA ploidy level estimation for this subspecies. Both localities visited over the course of our study (Mt. Erciyes Dağı (Argaeus) in Cappadocia the Gusguta valley in the Bolkar Dağları Mts. in southern Anatolia) are also mentioned by Weigend [72], which confirms the taxonomical identity of the plants under study. They occur on high-mountain screes that are only marginally influenced by human activities (pastures) and therefore fall within the broad concept that diploids tend to inhabit natural or semi-natural habitats. Long-term survival of these diploid populations seems to be a plausible explanation at least for two reasons: In Anatolia there was no major Pleistocene ice-sheet similar to those covering the European Alps or Scandinavia and only mountain peaks exceeding the height of ca 2200 m were glaciated [135, 136]. Furthermore, higher elevations provided moist conditions contrasting with the drier climate that prevailed in lower elevations of Anatolia during glacial periods [137].

We did not find diploids among a total of 80 plants from Iran. Weigend [72] reported two subspecies of *U. dioica* from this country, namely subsp. *dioica* and subsp. *kurdistanica.* Our plants can be more or less identified as subsp. *dioica,* and their ploidy is thus in line with the general picture of diploid subsp. *kurdistanica* and tetraploid subsp. *dioica.* Still, several populations (north of Tehran—Mt. Damavand and its surroundings) formed a unique monoecious population and their inflorescences consisted of equal numbers of male and female flowers (male in the upper part of the inflorescence), which does not correspond to the morphological description of either subsp. *dioica* or subsp. *kurdistanica.*

Finally, despite our extensive screening, we failed to find diploid plants at several localities of the morphologically defined taxon *U. d.* subsp. *subinermis.* This subspecies therefore has to be considered only supposedly diploid, as no chromosome counts are presented in the respective papers. This applies, for example, to the Neusiedler See lake (northeastern Austria), where precisely defined localities of plants morphologically assigned to *U. d.* subsp. *subinermis* are mentioned by Geltman [127] and Weigend [71]. Tetraploid plants found over the course of our study (5975 plants sampled) morphologically resemble *U. d.* subsp. *subinermis,* which raises the question as to whether (auto)polyploidization has taken place in this subspecies,
which would make the pattern of genomic evolution within the diploid-tetraploid complex of *Urtica dioica* considerably more complicated.

**Conclusion**

Our large-scale cytogeographic screening of *Urtica dioica* has revealed a complex pattern across a major part of the species’ distribution range, consisting of a widespread tetraploid cytotype, low-abundant scattered diploids and sporadically occurring triploid and pentaploid plants. We have not found any differences in genome size (Cx-values) between most subspecies of *U. dioica* (*U. d. subsp. dioica*, subsp. *kurdistanica*, subsp. *pubescens*, subsp. *sondenii* and subsp. *subinermis*). On the other hand, *U. d. subsp. cypria* does differ in genome size from the rest of *U. dioica*. Moreover, Cx-values of closely related species (*U. bianorii*, *U. kioviensis* and *U. simensis*) clearly differ from those of *U. dioica*, and genome size can thus serve as a valuable supportive character in the delimitation of *U. dioica*. We have also found positive correlations between genome size and longitude and latitude in our complete dataset of European populations and a positive correlation of genome size with longitude in a reduced dataset of diploid accessions (the complete dataset of diploids excluding *U. d. subsp. kurdistanica*). Diploid taxa growing in relict habitats are more frequent at lower elevations. In addition, our study has revealed a significant affinity of diploids to less human-influenced semi-natural habitats (this does not apply diploids from the Po river basin, assigned to *U. d. subsp. pubescens*) and (in the European range) to lower elevations. The tetraploid cytotype, by contrast, tends to thrive even in highly synanthropic sites and is able to expand to higher elevations.

**Supporting Information**

**S1 Table.** List of analyses of *Urtica dioica* (sorted by population identification number).

For each population, the following information is provided: geographic coordinates in the WGS-84 system, elevation, country abbreviation, collector’s initials, number of analysed plants in simultaneous analyses, relative fluorescence intensity, DNA-ploidy level, and coefficient of variance of the standard and sample peaks.

(PDF)

**S2 Table.** List of analyses (absolute genome size) of *Urtica dioica* (sorted by taxon and population identification number).

For each population, the following information is provide: geographic coordinates in the WGS-84 system, elevation, country abbreviation, collector’s initials, absolute genome size—2C-value (pg), ploidy level, and coefficient of variance of standard and sample peaks.

(PDF)

**S1 Fig.** Map of locations of *Urtica dioica* samples collected in the Po river basin (northern Italy). The size of the circles reflects the number of populations. The blue line indicates the outline of the Po river basin.

(TIF)

**S2 Fig.** Map of locations of *Urtica dioica* samples collected in the Czech Republic and Slovakia. The size of the circles reflects the number of populations.

(TIF)

**S3 Fig.** Relative fluorescence intensity variation in *Urtica dioica*. Two dominant ploidy levels were detected (red—2x and yellow—4x).

(TIF)
S4 Fig. Flow cytometric histogram of all detected cytotypes of *Urtica dioica*. Simultaneous analysis—from the left: 2x—diploid cytotype, 3x—triploid, 4x—tetraploid, 5x—pentaploid, *Bellis perennis*—the internal standard. (TIF)

S5 Fig. Proportions of cytotypes of *Urtica dioica* seeds. (A) Ratio of diploid and triploid seeds from a 2x maternal plant (from the mixed-ploidy population); (B) Ratio of triploid, tetraploid and pentaploid seeds from a 4x maternal plant (from a mixed-ploidy population). (TIF)

S6 Fig. Map of locations of closely related species. Species closely related to the *U. dioica* clade in recent phylogenies, namely: *U. atrovirens*, *U. bianorii*, *U. kioviensis*. The top-left section shows the one population of *U. simensis* in Ethiopia. The size of the circles reflects the number of populations. For more details see S2 Table. (TIF)

S7 Fig. Absolute genome size variation in *Urtica dioica* and closely related species. Diploid cytotype—*U. d*. subsp. *kurdistanica*, subsp. *pubescens*, subsp. *sondenii* and subsp. *subinermis*; tetraploid cytotype—*U. d*. subsp. *dioica*; closely related species—*U. atrovirens*, *U. bianorii*, *U. d*. subsp. *cypria*, *U. kioviensis* and *U. simensis*). Numbers of analysed individuals are presented in parentheses. (TIF)

**Acknowledgments**

We are grateful to R. Bartošová, M. Dudáš, Š. Hořčicová, A. Hrabalík, Z. Chumová, F. Kolář, C. Pachschwöll, H. Rydlo, B. Senius, K. Senius, K. Šemberová and many others for their help during fieldwork or for providing plant samples, P. Neumann (UMBR) for allowing us to a fluorescent microscope, and F. Rooks for helping out with the language of our raw manuscript.

Funding: The research was supported by the Czech Science Foundation (project no. GA 17-20201S), Charles University (GAUK project no. 1206617 and the Research Centre programme no. 204069) and the Czech Academy of Sciences (long-term research development project no. RVO 67985939). The funders had no role in the design of the study, collection and analysis of data, the decision to publish this paper or the preparation of the manuscript.

**Author Contributions**

**Conceptualization:** Tomáš Urfus.

**Data curation:** Ludmila Rejlová, Jindřich Chrtek, Pavel Trávníček, Magdalena Lučanová, Petr Vít, Tomáš Urfus.

**Formal analysis:** Pavel Trávníček, Magdalena Lučanová, Petr Vít.

**Funding acquisition:** Ludmila Rejlová, Jindřich Chrtek.

**Investigation:** Ludmila Rejlová, Magdalena Lučanová, Tomáš Urfus.

**Methodology:** Ludmila Rejlová, Jindřich Chrtek, Pavel Trávníček, Magdalena Lučanová, Petr Vít, Tomáš Urfus.

**Supervision:** Jindřich Chrtek, Tomáš Urfus.

**Writing – original draft:** Ludmila Rejlová, Jindřich Chrtek, Tomáš Urfus.
Writing – review & editing: Ludmila Rejlova, Pavel Trávníček.

References

1. Grant V. Plant speciation. 1st ed. New York: Columbia University Press; 1981.
2. Rieseberg LH, Willis JH. Plant speciation. Science. 2007; 317(5840):1–9. https://doi.org/10.1126/science.1137729
3. Symonds VV, Soltis PS, Soltis DE. Dynamics of polyploid formation in *Tragopogon* (Asteraceae): Recurrent formation, gene flow, and population structure. Evolution. 2010; 64(7):1964–2003. https://doi.org/10.1111/j.1558-5646.2010.00978.x PMID: 20199558
4. Renny-Byfield S, Wendel JF. Doubling down on genomes: Polyploidy and crop plants. Am J Bot. 2014; 101(10):1711–1725. https://doi.org/10.3732/ajb.1400119 PMID: 25090999
5. Jiao Y, Wickett NJ, Ayyampalayam S, Chanderbali AS, Landherr L, Ralph PE, et al. Ancestral polyploidy in seed plants and angiosperms. Nature. 2011; 473(7370):67–70. https://doi.org/10.1038/nature09916 PMID: 21478875
6. Otto SP, Whitton J. Polyploid incidence and evolution. Annu Rev Genet. 2000; 34(1):401–437. https://doi.org/10.1146/annurev.genet.34.1.401 PMID: 11092833
7. Wood TE, Takebayashi N, Barker MS, Mayrose I, Greenspoon PB, Rieseberg LH. The frequency of polyploid speciation in vascular plants. Proc Natl Acad Sci. 2009; 106(33):13875–13879. https://doi.org/10.1073/pnas.0811575106 PMID: 19667210
8. Ohri D. Genome size variation and plant systematics. Ann Bot. 1998; 82(2):75–83. https://doi.org/10.1006/anbo.1998.0765
9. Thompson JD, Lumaret R. The evolutionary dynamics of polyploid plants: origins, establishment and persistence. Trends Ecol Evol. 1992; 7(9):302–307. https://doi.org/10.1016/0169-5347(92)90228-4 PMID: 21236040
10. Wolfe KH. Yesterday’s polyploids and the mystery of diploidization. Nat Rev Genet. 2001; 2:333–341. https://doi.org/10.1038/35072009 PMID: 11331899
11. Mandáková T, Joly S, Krzywinski M, Mummenhoff K, Lysak MA. Fast diploidization in close mesopolyploid relatives of *Arabidopsis*. Plant Cell. 2010; 22(7):2277–2290. https://doi.org/10.1105/tpc.110.074526 PMID: 20639445
12. Renny-Byfield S, Kovarik A, Kelly LJ, Macas J, Novak P, Chase MW, et al. Diploidization and genome size change in allopolyploids is associated with differential dynamics of low- and high-copy sequences. Plant J. 2013; 74(5):829–839. https://doi.org/10.1111/tpj.12168 PMID: 23517128
13. Soltis DE, Soltis PS, Tate JA. Advances in the study of polyploidy since Plant speciation. New Phytol. 2003; 161(1):173–191. https://doi.org/10.1046/j.1469-8137.2003.00948.x
14. Leitch IJ, Bennett MD. Genome downsizing in polyploid plants. Biol J Linn Soc. 2004; 82(4):651–663. https://doi.org/10.1111/j.1095-8312.2004.00349.x
15. Leitch I, Soltis DE, Soltis PS, Bennett MD. Evolution of DNA amounts across land plants (Embryophyta). Ann Bot. 2005; 95(1):207–217. https://doi.org/10.1093/aob/mci014 PMID: 15596468
16. Rice A, Glick L, Abadi S, Einhorn M, Kopelman NM, Salman-Minkov A, et al. The Chromosome Counts Database (CCDB)—a community resource of plant chromosome numbers. New Phytol. 2015; 206(1):19–26. https://doi.org/10.1111/nph.13191 PMID: 25423910
17. Levin DA. The role of chromosomal change in plant evolution. New York: Oxford University Press; 2002.
18. Francis D, Davies MS, Barlow PW. A strong nucleotypic effect on the cell cycle regardless of ploidy level. Ann Bot. 2008; 101(6):747–757. https://doi.org/10.1093/aob/mcn038 PMID: 18339642
19. Maherali H, Walden AE, Husband BC. Genome duplication and the evolution of physiological responses to water stress. New Phytol. 2009; 184(3):721–731. https://doi.org/10.1111/j.1469-8137.2009.02997.x PMID: 19703115
20. Ramsey J. Polyploidy and ecological adaptation in wild yarrow. Proc Natl Acad Sci. 2011; 108(17):7096–7101. https://doi.org/10.1073/pnas.1016631108 PMID: 21402904
21. Madlung A. Polyploidy and its effect on evolutionary success: old questions revisited with new tools. Heredity. 2013; 110(2):99–104. https://doi.org/10.1038/hdy.2012.79 PMID: 23149549
22. Lumaret R, Guillerm JL, Delay J, Ait Lhaj Loutfi A, Izzo J, Jay M. Polyploidy and habitat differentiation in *Dactylis glomerata* L. from Galicia (Spain). Oecologia. 1987; 73(3):436–446. https://doi.org/10.1007/BF00385262 PMID: 28311527
23. Burnier J, Arrigo N, Küpfer P, Alvarez N. Genetic structure and evolution of Alpine polyploid complexes: Ranunculus kuepferi (Ranunculaceae) as a case study. Mol Ecol. 2009; 18(17):3730–3744. https://doi.org/10.1111/j.1365-294X.2009.04281.x PMID: 19674303

24. Treier UA, Breonnimann O, Normand S, Guisan A, Schaffner U, Steinger T, et al. Shift in cytotype frequency and niche space in the invasive plant Centaurea maculosa. Ecology. 2009; 90(5):1366–1377. https://doi.org/10.1890/08-0420.1 PMID: 19537556

25. Salmon A, Ainouche ML, Wendel JF. Genetic and epigenetic consequences of recent hybridization and polyploidy in Spartina (Poaceae). Mol Ecol. 2005; 14(4):1163–1175. https://doi.org/10.1111/j.1365-294X.2005.02488.x PMID: 15773943

26. Cavanagh CR, Chao S, Wang S, Huang BE, Stephen S, Kiani S, et al. Genome-wide comparative diversity uncovers multiple targets of selection for improvement in hexaploid wheat landraces and cultivars. Proc Natl Acad Sci. 2013; 110(20):8057–8062. https://doi.org/10.1073/pnas.1217133110 PMID: 23630259

27. Těšitelová T, Jersáková J, Roy M, Kubátová B, Těšítel J, Urfus T, et al. Ploidy-specific symbiotic interactions: divergence of mycorrhizal fungi between cytotypes of the Gymnadenia conopsea group (Orchidaceae). New Phytol. 2013; 199(4):1022–1033. https://doi.org/10.1111/nph.12348 PMID: 23731358

28. Linder HP, Barker NP. Does polyploidy facilitate long-distance dispersal? Ann Bot. 2014; 113(7):1175–1183. https://doi.org/10.1093/aob/mcu047 PMID: 24694830

29. Rosche C, Hensen I, Lachmuth S. Local pre-adaptation to disturbance and inbreeding-environment interactions affect colonisation abilities of diploid and tetraploid Centaurea stoebe. Plant Biol. 2018; 20(1):75–84. https://doi.org/10.1111/pib.12628 PMID: 28921779

30. Rosche C, Hensen I, Mráz P, Durka W, Hartmann M, Lachmuth S. Invasion success in polyploids: the role of inbreeding in the contrasting colonization abilities of diploid versus tetraploid populations of Centaurea stoebe s.l. J Ecol. 2017; 105(2):425–435. https://doi.org/10.1111/1365-2745.12670

31. Schlaepfer DR, Edwards PJ, Semple JC, Billeret R. Cytogeography of Solidago gigantea (Asteraceae) and its invasive ploidy level. J Biogeogr. 2008; 35(11):2119–2127. https://doi.org/10.1111/j.1365-2699.2008.01937.x

32. Španiel S, Marhold K, Hodálova I, Lihová J. Diploid and tetraploid cytotypes of Centaurea stoebe (Asteraceae) in Central Europe: Morphological differentiation and cytotype distribution patterns. Folia Geobot. 2008; 43(2):131–158. https://doi.org/10.1007/s12224-008-9008-7

33. Mráz P, Bourchier RS, Treier UA, Schaffner U, Müller-Schärer H. Polyploidy in phenotypic space and polyploidy in the nettle Polyploidy evolution in the nettle.Centaurea stoebe s.l. Int J Plant Sci. 2011; 172(3):386–402. https://doi.org/10.1086/658151

34. Zožomová-Lihová J, Malánová-Krásná I, Vl P, Urfus T, Senko D, Svitok M, et al. Cytotype distribution patterns, ecological differentiation, and genetic structure in a diploid-tetraploid contact zone of Cardamine amara. Am J Bot. 2015; 102(8):1380–1395. https://doi.org/10.3732/ajb.1500052 PMID: 26290560

35. Baduel P, Arnold B, Weisman CM, Hunter B, Bombikes K. Habitat-associated life history and stress-tolerance variation in Arabidopsis arenosa. Plant Physiol. 2016; 171(1):437–451. https://doi.org/10.1104/pp.15.01875 PMID: 26941193

36. Pandit MK, Pocock MJO, Kunin WE. Ploidy influences rarity and invasiveness in plants. J Ecol. 2011; 99(5):1108–1115. https://doi.org/10.1111/j.1365-2745.2011.01838.x

37. te Beest M, Le Roux JJ, Richardson DM, Brysting AK, Suda J, Kubesová M, et al. The more the better? The role of polyploidy in facilitating plant invasions. Ann Bot. 2012; 109(1):19–45. https://doi.org/10.1093/aob/mcu047 PMID: 22040744

38. Husband BC, Sabara HA. Reproductive isolation between autotetraploids and their diploid progenitors in fireweed, Chamerion angustifolium (Onagraceae). New Phytol. 2003; 161(3):703–713. https://doi.org/10.1046/j.1469-8137.2004.00998.x

39. Kao RH. Asexuality and the coexistence of cytotypes. New Phytol. 2007; 175(4):764–772. https://doi.org/10.1111/j.1469-8137.2007.02145.x PMID: 17685991

40. Sonnleitner M, Flatscher R, Escobar Garcia P, Rauchová J, Suda J, Schneeweis GM, et al. Distribution and habitat segregation on different spatial scales among diploid, tetraploid and hexaploid cytotypes of Senecio carniolicus (Asteraceae) in the Eastern Alps. Ann Bot. 2010; 106(6):967–977. https://doi.org/10.1093/aob/mcq192 PMID: 20980930

41. Kolář F, Čertner M, Suda J, Schönsweeter P, Husband BC. Mixed-ploidy species: progress and opportunities in polyploid research. Trends Plant Sci. 2017; 22(12):1041–1055. https://doi.org/10.1016/j.tplants.2017.09.011 PMID: 289054346
42. Lafuma L, Balkwill K, Imbert E, Verlaque R, Maurice S. Ploidy level and origin of the European invasive weed Senecio inaequidens (Asteraceae). Plant Syst Evol. 2003; 243(1-2):59–72. https://doi.org/10.1007/s00606-003-0075-0

43. Buggs RJA, Pannell JR. Ecological differentiation and diploid superiority across a moving ploidy contact zone. Evolution. 2007; 61(1):125–140. https://doi.org/10.1111/j.1558-5646.2007.00010.x PMID: 17300432

44. Obbard DJ, Harris SA, Buggs RJA, Pannell JR. Hybridization, polyploidy, and the evolution of sexual systems in Mercurialis (Euphorbiaceae). Evolution. 2006; 60(9):1801. https://doi.org/10.1554/06-104.1 PMID: 17089965

45. Čertner M, Fenclová E, Kúr P, Kolář F, Koutecký P, Krahulcová A, et al. Evolutionary dynamics of mixed-ploidy populations in an annual herb: dispersal, local persistence and recurrent origins of polyploids. Ann Bot. 2017; 120(2):303–315. https://doi.org/10.1093/ab/mcx032 PMID: 28398545

46. Bredemann G, Garber K. Die grosse Brennessel Urtica dioica L: Forschungen über ihren Anbau zur Fasergewinnung. Berlin: Akademie-Verlag; 1959.

47. Bacci L, Baronti S, Predieri S, di Virgilio N. Fiber yield and quality of fiber nettle (Urtica dioica L.) cultivated in Italy. Ind Crops Prod. 2009; 29(2-3):480–484. https://doi.org/10.1016/j.indcrop.2008.09.005

48. Di Virgilio N, Papazoglou EG, Jankauskienė Z, Di Lonardo S, Praczuk M, Wielgusz K. The potential of stinging nettle (Urtica dioica L.) as a crop with multiple uses. Ind Crops Prod. 2015; 68:42–49. https://doi.org/10.1016/j.indcrop.2014.08.012

49. Farzami B, Ahmadvand D, Vardasi S, Majinfar FJ, Khaghani S. Induction of insulin secretion by a component of Urtica dioica leave extract in perifused Islets of Langerhans and its in vivo effects in normal and streptozotocin diabetic rats. J Ethnopharmacol. 2003; 89(1):47–53. https://doi.org/10.1016/S0378-8741(03)00220-4 PMID: 14522431

50. Mohammadi A, Mansoori B, Aghapour M, Baradaran B. Urtica dioica dichloromethane extract induce apoptosis from intrinsic pathway on human prostate cancer cells (PC3). Cell Mol Biol. 2016; 62(3):78–83. PMID: 27064877

51. Olsen C. The Ecology of Urtica dioica. J Ecol. 1921; 9(1):1. https://doi.org/10.2307/2255757

52. Ivins J. Concerning the Ecology of Urtica dioica L. J Ecol. 1952; 40(2):380. https://doi.org/10.2307/2256806

53. Pagad S, Genovési P, Carnevali L, Schigel D, McGeoch MA. Introducing the Global Register of Introduced and Invasive Species. Sci Data. 2018; 5:170202. https://doi.org/10.1038/sdata.2017.202 PMID: 29360103

54. Domin K. Monografická studie o kopřivě dvoudomé (Urtica dioica L.)—1. pokračování. Časopis čes lékařstva. 1944; 57:59–75.

55. Domin K. Monografická studie o kopřivě dvoudomé (Urtica dioica L.). Časopis čes lékařstva. 1944; 3:92–94.

56. Grosse-Veldmann B, Weigend M. Weeding the nettles III: Named nonsense versus named morphotypes in European Urtica dioica L. (Urticaceae). Phytotaxa. 2015; 208(4):239. https://doi.org/10.11646/phytotaxa.208.4.1

57. Grosse-Veldmann B, Weigend M. The geometry of gender: hyper-diversification of sexual systems in Urtica L. (Urticaceae). Cladistics. 2018; 34(2):131–150. https://doi.org/10.1111/cla.12193

58. Geltman D. On the chromosome numbers of Urtica dioica s. l. (Urticaceae)—(On the basis of the material from the middle zone of the European part of the SSSR). Bot Zhurn. 1981; 67:78–89.

59. Jonsell B, editor. Flora Nordica. Volume 1: Lycopodiaceae—Polygonaceae. vol. 1. Stockholm: Ber- gius Foundation; 2000.

60. Lippert W. Chromosomenzahlen von Pflanzen aus Bayern und anderen Gebieten. Berichte Bayer Bot Ges. 2006; 76:85–110.

61. Löve A, Löve D. Chromosome numbers of central and northwest European plant species. Stockholm: Almqvist & Wiksell; 1961.

62. Löve A, Löve D. Chromosome numbers of northern plant species. No. No. 3 in Reports, Ser. B. Reykjavik: University Institute of Applied Science, Dept. of Agriculture; 1948.

63. Mráz P. Chromosome number and DNA ploidy level reports from Central Europe—2. Biologia (Bratisl). 2006; 61(1):115–120.

64. Tropicos.org. Missouri Botanical Garden.; 2018 [cited 2018 Dec 4]. Database: [Internet]. Available from: http://www.tropicos.org/Name/33400020?tab=chromosomcounts.

65. Farag MA, Weigend M, Luebert F, Brokamp G, Wessjohann LA. Phytochemical, phylogenetic, and anti-inflammatory evaluation of 43 Urtica accessions (stinging nettle) based on UPLC–Q-TOF-MS
metabolomic profiles. Phytochemistry. 2013; 96:170–183. https://doi.org/10.1016/j.phytochem.2013.09.016 PMID: 24169378

66. Wu ZY, Monro AK, Milne RI, Wang H, Yi TS, Liu J, et al. Molecular phylogeny of the nettle family (Urticaceae) inferred from multiple loci of three genomes and extensive generic sampling. Mol Phylogenet Evol. 2013; 69(3):814–827. https://doi.org/10.1016/j.ympev.2013.06.022 PMID: 23850510

67. Henning T, Quandt D, Grosse-Veldmann B, Monro A, Weigend M. Weeding the Nettles II: A delimitation of *Urtica dioica* L. (Urticaceae) based on morphological and molecular data, including a rehabilitation of *Urtica gracilis* Ait. Phyto taxa. 2014; 162(2):61. https://doi.org/10.11646/phytotaxa.162.2.1

68. Grosse-Veldmann B, Nürk NM, Smissen R, Breitwieser I, Quandt D, Weigend M. Pulling the sting out of nettle systematics—A comprehensive phylogeny of the genus *Urtica* L. (Urticaceae). Mol Phylogenet Evol. 2016; 102:9–19. https://doi.org/10.1016/j.ympev.2016.05.019 PMID: 27211697

69. Chrtek J. Urticaceae. In: Rechinger KH, editor. Flora Iranica: Flora des iranischen Hochlandes und der umrahmenden Gebirge: Persien, Afganistan, Teile von West-Pakistan, Nord-Iraq, Azerbaidjan, Turkmenistan. Ht. 105: Urticaceae. vol. 105. Graz: Akademische Druck- u. Verlaganstalt; 1974. p. 1–20.

70. Townsend CC, Guest E, editors. Flora of Iraq. Vol. III: Leguminosae. vol. 3. Baghdad: Ministry of Agriculture of the Republic of Iraq; 1974.

71. Weigend M. Die Erben Pokornys—Ein Beitrag zur Abgrenzung der Sippen *Urtica galeopsophila* und *Urtica pubescens* in Mittel- und Osteuropa. Hoppea, Denkschr Regensb Ges. 2005; 66:101–118.

72. Weigend M. *Urtica dioica* subsp. *cypris*, with a re-evaluation of the *U. dioica* group (Urticaceae) in western Asia. Wildenowia. 2006; 36(2):811–822. https://doi.org/10.3372/wi.36.36212

73. Geltman D. New sections and subsections of the genus *Urtica* (Urticaceae). Bot Zhurn. 1982; 67:1413–1416.

74. Simmons HG. Floran och vegetationen i Kiruna: en växtgeografisk studie med särskild hänsyn till kulturens inflytande. Vetenskapliga och praktiska undersökningar i Lappland. Stockholm: Nordiska bokhandeln in distribution; 1910.

75. Pollard AJ, Briggs D. Genealogical studies of *Urtica dioica* L. I. The nature of intraspecific variation in *U. dioica*. New Phytol. 1982; 92(3):453–470. https://doi.org/10.1111/j.1469-8137.1982.tb03403.x

76. Meikle RD. Flora of Cyprus. Vol. II. Kew: Bentham Moxon Trust, Royal Botanic Gardens; 1985.

77. Suda J, Trávníček P. Reliable DNA ploidy determination in dehydrated tissues of vascular plants by DAPI flow cytometry—new prospects for plant research. Cytom Part A. 2006; 69A(4):273–280. https://doi.org/10.1002/cyto.a.20253

78. Doležel J, Greilhuber J, Suda J. Estimation of nuclear DNA content in plants using flow cytometry. Nat Protoc. 2007; 2(9):2233–2244. https://doi.org/10.1038/nprot.2007.310 PMID: 17853881

79. Schönswetter P, Suda J, Popp M, Weiss-Schneweiss H, Brochmann C. Circumpolar phylogeography of *Juncus biglumis* (Juncaceae) inferred from AFLP fingerprints, cpDNA sequences, nuclear DNA content and chromosome numbers. Mol Phylogen et Evol. 2007; 42(1):92–103. https://doi.org/10.1016/j.ympev.2006.06.016 PMID: 16905337

80. Doležel J, Binarová P, Lucretti S. Analysis of Nuclear DNA content in plant cells by Flow cytometry. New Phytol. 1982; 92(3):453–470. https://doi.org/10.1111/j.1469-8137.1982.tb03403.x

81. A language and environment for statistical computing R Foundation for statistical computing, Vienna, Austria [Internet]. R Core Team; 2018. Available from: https://www.R-project.org/.

82. Mandáková T, Lysak MA. Chromosome preparation for cytogenetic analyses in *Arabidopsis*. In: Stacey G, Bircher J, Ecker J, Martin CR, Stitt M, Zhou JM, editors. Current protocols in plant biology. Hoboken, NJ, USA: John Wiley & Sons, Inc.; 2016. p. 43–51. Available from: http://doi.wiley.com/10.1002/cpb.20009.

83. Chumová Z, Záveská E, Mandáková T, Krak K, Trávníček P. The Mediterranean: the cradle of *Anthoxanthum* (Poaceae) diploid diversity. Ann Bot. 2017; 120(2):285–302. https://doi.org/10.1093/aob/mcx021 PMID: 28444200

84. Aiello-Lammens ME, Borja RA, Radosavljevic A, Vilela B, Anderson RP. spThin: an R package for spatial thinning of species occurrence records for use in ecological niche models. Ecography. 2015; 38(5):541–545. https://doi.org/10.1111/ecog.01132

85. Hijmans RJ, Graham CH. The ability of climate envelope models to predict the effect of climate change on species distributions. Global change biol. 2006; 12(12):2272–2281. https://doi.org/10.1111/j.1365-2486.2006.01256.x

86. Koutecký P. MorphoTools: a set of R functions for morphometric analysis. Plant Syst Evol. 2015; 301 (4):1115–1121. https://doi.org/10.1007/s00606-014-1153-2
87. Tüxen R, Preising E. Die heutige potentielle natürliche Vegetation als Gegenstand der Vegetationskartierung. Angewandte Pflanzensoziologie. Stolzenau (Weser): Selbstverl. d. Bundesanst. für Vegetationskartierung; 1956.

88. Agresti A. Introduction to categorical data analysis. Second edition ed. Wiley series in probability and statistics. Department of Statistics, University of Florida Gainesville, Florida: John Wiley & Sons, Inc.; 2007.

89. Ramsey J, Ramsey TS. Ecological studies of polyploidy in the 100 years following its discovery. Philos Trans R Soc B Biol Sci. 2014; 369(1648):20130352. https://doi.org/10.1098/rstb.2013.0352

90. Hegi G. Illustrierte Flora von Mittel-Europa: mit besonderer Berücksichtigung von Deutschland, Österreich und der Schweiz: zum Gebrauche in den Schulen und zum Selbstunterricht. Ill. Band. I. Teil: Dicotyledones. 1st ed. München: J. F. Lehmann; 1912.

91. Zólyomi B. Urtica kioviensis Rogovitsch neu für die Deutsche Flora. Verh Bot Ver Prov Brandenburg. 1936; 76:152–156.

92. Tutin TG, editor. Flora Europaea. Vol. 1: Lycopodiaceae to Platanaceae. vol. 1. Cambridge: Cambridge University Press; 1964.

93. Pignatti S. Flora d'Italia. vol. 1. Bologna: Edagricole; 1982.

94. Paiva J, LXIV. Urticaceae. In: Paiva J, Castroviejo S, editors. Flora Iberica: plantas vasculares de la Península Ibérica e Islas Baleares. Vol. III: Plumbaginaceae (partim)—Capparaceae. vol. 3. Madrid: Real Jardín Botánico de Madrid; 1993. p. 262–268.

95. Danhelka J, Lepší M. Kopřiv a luznina, Urtica kioviensis, na soutoku Moravy a Dyje. Zprávy Čes Bot Spol. 2004; 39:25–35.

96. Moreno M. Urticaceae. In: Valdés B, Talavera S, Galiamo EF, editors. Flora Vascular de Andalucia Occidental. vol. 1. Barcelona: Ketres Editora; 1987. p. 154–156.

97. Cabezudo B. Urticaceae. In: Torres C, Salazar C, editors. Flora Vascular de Andalucía Oriental. 2nd ed. Granada: Universidades de Almería; 2011. p. 913–914.

98. Darlington CD, Wylie AP. Chromosome atlas of flowering plants. 2nd ed. London: G. Allen & Unwin; 1955.

99. Bolchovskich ZV. Chromosomnye čisla cvetkovych rastenij. Leningrad: Akademija nauk SSSR, Nauka; 1969.

100. Marhold K, editor. Chromosome number survey of the ferns and flowering plants of Slovakia. Bratislava: Veda; 2007.

101. Tischler G. Die Chromosomenzahlen der Gefäßpflanzen Mitteleuropas. Den Haag: W. Junk; 1950.

102. Májovský J, Murín A. Karyotaxonomicky prehľad flóry Slovenska. Bratislava: Veda; 1987.

103. Loureiro J, Travníček P, Rauchová J, Urfus T, Vítech M, et al. The use of flow cytometry in the biosystematic, ecology and population biology of homoploid plants. Preslia. 2010; 82:3–21.

104. Ramsey J, Schemske DW. Pathways, Mechanisms, and rates of polyploid formation in flowering plants. Ann Rev of Ecol and Syst. 1998; 29(1):467–501. https://doi.org/10.1146/annurev.ecolsys.29.1.467

105. Kreiner JM, Kron P, Husband BC. Frequency and maintenance of unreduced gametes in natural plant populations: associations with reproductive mode, life history and genome size. New Phytol. 2017; 214(2):879–889. https://doi.org/10.1111/nph.14423 PMID: 28134436

106. Burton TL, Husband B. Fitness differences among diploids, tetraploids, and their triploid progeny in Chamerion angustifolium: mechanisms of inviability and implications for polyploid evolution. Evolution. 2000; 54(4):1182–1191. https://doi.org/10.1111/0014-3820.2000.tb00553.x PMID: 11005287

107. Douba J, Boublík K, Sležák M, Biurrun I, Nociar J, Havrdová A, et al. Vegetation classification and biogeography of European floodplain forests and alder carrs. Appl Veg Sci. 2016; 19(1):147–163. https://doi.org/10.1111/avsc.12201

108. Roberts N, Fyfe RM, Woodbridge J, Gaillard MJ, Davis BaS, Kaplan JO, et al. Europe’s lost forests: a pollen-based synthesis for the last 11,000 years. Sci Rep. 2018; 8(1):716. https://doi.org/10.1038/s41598-017-18646-7 PMID: 29335417

109. Nassar MA. Production of triploid cassava, Manihot esculenta Crantz by hybrid diploid gametes. Field Crops Res. 1992; 30(1-2):173–182. https://doi.org/10.1016/0378-4290(92)90066-I

110. Nassar MA. The synthesis of a new cassava-derived species, Manihot vieiri Nassar. Genet Mol Res. 2006; 5(3):536–541. PMID: 17117369

111. Pustovoitova T, Erermin G, Rassvetaeva E, Zhdanova N, Zholkевич V. Drought resistance, recovery capacity, and phytohormone content in polyploid plum leaves. Russ J Plant Physiol. 1996; 43(2):232–235.
112. Rothera SL, Davy AJ. Polyploidy and habitat differentiation in Deschampsia cespitosa. New Phytol. 1986; 102(3):449–467. https://doi.org/10.1111/j.1469-8137.1986.tb00822.x

113. Stebbins GL, Singh R. Artificial and natural hybrids in the Gramineae, tribe Hordeae. IV. Two triploid hybrids of Agropyron and Elymus. 1950; p. 7.

114. Brochmann C, Brysting AK, Alsos IG, Borgen L, Grundt HH, Scheen AC, et al. Polyploidy in arctic plants. Biol J Linn Soc. 2004; 82(4):521–536. https://doi.org/10.1111/j.1095-8312.2004.00337.x

115. Weiss-Schneeweiss H, Emadzade K, Jang TS, Schneeweiss GM. Evolutionary consequences, constraints and potential of polyploidy in plants. Cytogenet Genome Res. 2013; 140(2-4):137–150. https://doi.org/10.1159/000351727 PMID: 23796571

116. Chumova Z, Krejčíková J, Mandáková T, Suda J, Trávníček P. Evolutionary and taxonomic implications of variation in nuclear genome size: lesson from the grass genus Anthoxanthum (Poaceae). Plos One. 2015; 10(7):e0133748. https://doi.org/10.1371/journal.pone.0133748 PMID: 26207824

117. Geltman D. Some problems of phylogeny of the species of the subsection Urtica of the genus Urtica (Urticaceae). Bot Zhurn. 1990; 75:840–845.

118. Otisková V, Koutecký T, Kolář F, Koutecký P. Occurrence and habitat preferences of diploid and tetraploid cytotypes of Centaurea stoebe in the Czech Republic. Preslia. 2014; 86:67–80.

119. Němečková H, Krak K, Chrtk J. Complex pattern of ploidal and genetic variation in Seselis libanotis (Apiaceae). Ann Bot Fenn. 2019; 56:57–77. https://doi.org/10.5735/085.065.0111

120. Suda J, Meyerson LA, Leitch IJ, Pysˇek P. The hidden side of plant invasions: the role of genome size. Mol Ecol. 2009; 18(9):2055–2066. https://doi.org/10.1111/j.1365-294x.2008.03971.x PMID: 18959400

121. Yen H, Martin SL, Bekele WA, Latta RG, Diederichsen A, Peng Y, et al. Genome size variation in the genus Anthoxanthum (Poaceae). Cytogenet Genome Res. 2013; 140(2-4):137–150. https://doi.org/10.1159/000351727 PMID: 23796571

122. Vít P, Krak K, Trávníček P, Douda J, Lomonosova MN, Mandák B. Genome size stability across Eurasian Chenopodium species (Amaranthaceae). Bot J Linn Soc. 2016; 182(3):637–649. https://doi.org/10.1111/boj.12474

123. Baack EJ, Whitney KD, Rieseberg LH. Hybridization and genome size evolution: timing and magnitude of nuclear DNA content increases in Helianthus homopoloid hybrid species. New Phytol. 2005; 167(2):623–630. PMID: 15986412

124. James JK, Abbott RJ. Recent, allopatric, homoploid hybrid speciation: The origin of Senecio squalidus (Asteraceae) in the British Isles from a hybrid zone on Mount Etna, Sicily. Evolution. 2005; 59(12):2533–2547. https://doi.org/10.1554/05-306.1 PMID: 16526502

125. Suda J, Meyerson LA, Leitch IJ, Pysˇek P. The hidden side of plant invasions: the role of genome size. Mol Ecol. 2009; 18(9):2055–2066. https://doi.org/10.1111/j.1365-294x.2008.03971.x PMID: 18959400

126. Frajman B, Eggen F, Oxelman B. Hybrid origins and homoploid reticulate evolution within Heliosperma (Sileneae, Caryophyllaceae)—A multigene phylogenetic approach with relative dating. Syst Biol. 2009; 58(3):328–345. https://doi.org/10.1093/sysbio/syp030 PMID: 20525587

127. Geltman D. Cytotaxonomic studies of the species of the genus Urtica (Urticaceae) in Flora of the USSR. Bot Zhurn. 1984; 69:1524–1530.

128. Geltman D. Genus Urtica L. (Urticaceae) in the URSS. Nov System Plant Vasc. 1988; 25:68–80.

129. Schönswetter P, Stehlik I, Holderegger R, Tribsch A. Molecular evidence for glacial refugia of mountain plants in the European Alps. Mol Ecol. 2005; 14(11):3547–3555. https://doi.org/10.1111/j.1365-294X.2005.02683.x PMID: 16156822

130. Magri D. Patterns of post-glacial spread and the extent of glacial refugia of European beech (Fagus sylvatica). J Biogeogr. 2008; 35(3):450–463. https://doi.org/10.1111/j.1365-294X.2007.01803.x

131. Schmitt T. Biogeographical and evolutionary importance of the European high mountain systems. Front Zool. 2009; 6(1):9. https://doi.org/10.1186/1742-9994-6-9 PMID: 19480666

132. Kühne G, Kosuch J, Hochkirch A, Schmitt T. Extra-Mediterranean glacial refugia in a Mediterranean faunal element: the phylogeography of the chalk-hill blue Polyommatus coridon (Lepidoptera, Lycaenidae). Sci Rep. 2017; 7:43533. https://doi.org/10.1038/srep43533

133. Taberlet P, Fumagalli L, Wust-Saury AG, Cosson JF. Comparative phylogeography and postglacial colonization routes in Europe. Mol Ecol. 1998; 7(4):453–464. https://doi.org/10.1046/j.1365-294X.1998.00289.x PMID: 9628000

134. Hewitt GM. Speciation, hybrid zones and phylogeography: seeing genes in space and time. Mol Ecol. 2008; 10(3):537–549. https://doi.org/10.1046/j.1365-294X.2001.01202.x

135. Eric S. Changes in the physical environment in Turkey since the end of the Last Glacial. In: Brice WC, editor. The Environmental history of the Near and Middle East since the last ice age; 1978. p. 67–81.
136. Atalay I. Palaeosols as indicators of the climatic changes during Quaternary period in S. Anatolia. J Arid Environ. 1996; 32(1):23–35. https://doi.org/10.1006/jare.1996.0003

137. Weeb T, Bartlein PJ. Global changes during the last 3 million years: climatic controls and biotic responses. Ann Rev Ecol Syst. 1992; 23:141–173. https://doi.org/10.1146/annurev.es.23.110192.001041