Genome size variation in *Deschampsia cespitosa* sensu lato (Poaceae) in Eurasia

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Abstract

The grass *Deschampsia cespitosa* is a variable taxon out of which many varieties, subspecies and endemic species have been separated. In this paper, the variation in genome size (GS) and ploidy of this grass including several of its subspecies and two related species in Eurasia was investigated by flow cytometric (FCM) measurements. GS and ploidy data were also related to specific environments and reproduction mode. Ploidy levels found by FCM were confirmed by chromosome counts of diploid (2n = 26) and tetraploid (2n = 52) samples. Seminiferous (seed bearing) *D. cespitosa* was mainly diploid (GS between 3.754 and 5.438 pg/1C). GS variation in diploids showed a geographic pattern with a significant difference (H = 41,441, P < 0.001) between European (median = 4.377 pg) and Asian (median = 4.881 pg) accessions. Genome size (1C) in tetraploids ranged from 7.9426 to 9.0399 pg. Tetraploid seminiferous *D. cespitosa* was found mostly in disturbed habitats in western and southern Europe, while tetraploids in Asia were registered in wet Arctic habitats. Genome size (1C) between 8.3278 and 8.8603 pg of the pseudoviviparous plants (spikelets produce plantlets asexually) of wet habitats in central and northern Europe indicated tetraploidy. A putative triploid (GS 6.6817 pg) was detected in Iceland. Summing up, we found a high variation in GS on the geographic scale with significant regional differences in diploid *D. cespitosa*. Among the tetraploids, the asexually reproducing plants were bound to specific habitats, while the seminiferous plants showed a habitat preference similar to the diploids.

Keywords *Deschampsia* · Flow cytometry · Genome size · Geographical variation · Polyploidy · Pseudovivipary

Introduction

There is increasing evidence that all plant lineages have undergone several whole-genome doubling (WGD) events, i.e., cycles of polyploidization (e.g., Jiao et al. 2011; Soltis et al. 2014; Wendel et al. 2018) although the exact number and timing of these events are still disputed (Ruprecht et al. 2017). From analyzing the whole-genome data of major lineages, Jiao et al. (2011) infer at least two events of ancient whole-genome duplications to have occurred about 319 and 192 million years ago; this is long before the divergence of monocots and eudicots. Further duplications within various lineages provided raw material for plant evolution, which makes the study of its relationship with lineage diversification essential (Weiss-Schneeweiss and Schneeweiss 2013).

Grasses (Poaceae) are among those groups that have experienced ancient WGDs (Jiao et al. 2011) affecting the entire family, and further ancient or more recent polyploidizations in several lineages such as those in *Andropogon*...
gerardii Vitman (Keeler and Davis 1999), in the genera Hor-deum L. (Eilam et al. 2009), as well as in Zea L. and Triti-cum L. (Kellogg 2015), have played a major role in evolution. The genus Deschampsia is another group experiencing polyploidization followed by chromosomal rearrangements (Amosova et al. 2017), as evidenced also by different ploidy levels, chromosome numbers (Amosova et al. 2015, 2017) and a high variation in genome size (1C from 5 to 5.5 pg) for diploids (Bennett et al. 1982; Murray et al. 2005; Pascual-Díaz et al. 2020). However, we found only one value for a tetraploid plant (1C = 9 pg) reported by Bennett et al. (1982). The basic chromosome number of the genus (x = 13) differs from other members of the core Pooidae (mostly x = 7), suggesting polyploidy incidence predating the diversification of the genus. In the traditional narrow sense, with exclusion of taxa formerly ascribed to Deyeuxia Clarion ex P.Beauv. (Saarela et al. 2017) and Scribneria Hack (Soreng et al. 2015), Deschampsia comprises ca. 30 species of mostly perennials and a few South American annual species (Chiapella and Zuloaga 2010). Hybridization and reticulate evolution are common in this group of grasses (Wölk and Röser 2017) and might obscure phylogenetic relationships that are traditionally based on only a few selected molecular markers.

*Deschampsia cespitosa* (L.) P.Beauv., the tufted hair grass, is the most common species of the genus, found in all continents, with a disjunct common distribution among the northern hemisphere landmasses, southern South America, Australasia and South Africa. *Deschampsia cespitosa* is a tussock-forming, wind-pollinated, self-incompatible grass, occurring in habitats with moderate to high moisture in a cold-temperate climate. This grass has established populations in similar habitats in regions separated by thousands of kilometers, where appropriate ecological conditions are present.

### Taxonomic challenges

The high morphological variation of *Deschampsia cespitosa* has led to a confusing taxonomy in the northern hemisphere (e.g., Böcher et al. 1968; Porsild and Cody 1980) where several Central European populations in particular ecological settings have been treated as narrow endemic species (Conert 1987; Lauber and Wagner, 1998), or subspecies (Chiapella 2000). Intergradation and hybridization between geographic variants that may be considered subspecies were assumed by Clarke (1978). Taxa from Russia and eastern Asia were also treated either as different species (Probatova 1985; Czerepanov 1995; Tzvelev and Probatova 2012) or as infraspecific taxa by Chiapella and Probatova (2003). Several endemic taxa were described by Tzvelev et al. (2015) and Tzvelev and Probatova (2019). In contrast, the lumping treatment in the Flora of China (Wu and Phillips 2006) recognizes only two species (*D. koe lerioides* Regel and *D. cespitosa*) with four subspecies of *D. cespitosa*. The occurrence of pseudovivipary or facultative pseudovivipary has further complicated systematic approaches. With this form of clonal reproduction, the entire spikelet is transformed into a small plantlet. Because of the uncertain taxonomy, especially in the northern hemisphere, some authors (e.g., Kawano 1963; Rothera and Davy 1986) preferred to use the informal rank of *D. cespitosa* complex which includes all morphologically similar varieties, subspecies and regional species.

### Karyological variation

*Deschampsia cespitosa* has been studied in various regional samples (e.g., Kawano 1963; Albers 1980; Garcia-Suarez et al. 1997; Murray et al. 2005; Amosova et al. 2017). Eurasian accessions of *D. cespitosa* were found to be mostly diploid with 2n = 26 (Kawano 1963; Albers 1975, 1980; Garcia-Suarez et al. 1997; Dobes and Vitek 2000) with occasional reports of 2n = 28 (Kawano 1963) or 2n = 26 + 1B (Marhold et al. 2007). Aneuploidy was reported for both the diploids (Kawano 1963) and the polyploids (tetraploids or occasionally also triploids; Kawano 1963; Albers 1980; Hedberg 1958). Deviating numbers such as 2n = 41, 49, 50 (Löve and Löve 1975; Albers 1980) and 2n = 42 (Sokolovskaya and Probatova 1975) have also occasionally been reported for the polyploids. In a detailed survey of ploidy levels in populations of *D. cespitosa* across Great Britain, both diploids and tetraploids (2n = 26 and 2n = 52, respectively) were reported by Rothera and Davy (1986). This survey revealed that tetraploids are the predominant cytotype on the island. The presence of diploids and polyploids has also been documented in several of the taxa of different ranks related to *D. cespitosa* recognized from Russia (Chiapella and Probatova 2003; Tzvelev and Probatova 2019). It has been hypothesized that morphological variation may be correlated with ploidy levels (Chiapella and Probatova 2003).

Thus, *Deschampsia cespitosa* provides an interesting system to study the role of polyploidy in the context of biogeographical and habitat features. Using flow cytometry (FCM), this paper reports the genome size and ploidy-level variation of multiple populations of *D. cespitosa* to (i) identify biogeographic patterns related to genome size and ploidy levels; (ii) examine to which extent ploidy is related to specific environments and asexual reproduction.

### Materials and methods

#### Plant material

Plant leaves of 129 populations of *Deschampsia cespitosa* and two related species for FCM were collected in the field and dried under standard herbarium conditions. Tillers were...
also transplanted to the Botanical Garden of the University of Vienna (HBV) to provide the source of fresh leaf tissue for genome size measurements and root tips for chromosome counts. Plantlets of pseudoviviparous origin were also used for genome size measurement and grown in petri dishes for collecting root tips. Finally, fresh material was used for only a few samples from Lake Constance and for a small number of Austrian samples. The majority of samples for genome size measurement, however, was taken from the dried vouchers collected during the field trips. In a pilot test, we have followed the performance of fresh and dried leaf samples by measuring after several intervals over 5 months (data in Online Resource 1). The vast majority of our samples was stored between 3 and 4 months before processing. Vouchers of the samples are deposited in the herbaria WU and CSH (for duplicates of Chinese samples).

With the broad concept of *Deschampsia cespitosa*, we follow here Chiapella and Probatova (2003) and further relevant sources (Clarke 1980; Chiapella 2000; Wu and Phillips 2006) as none covers the entire investigated area (see Online Resource 2). We were not able to collect the newly described endemic taxa by Tzvelev and Probatova (2019). Regarding Europe, we are aware of several variants given subspecific or specific rank (Clarke 1980; Conert 1987; Chiapella 2000) or regarded as evolutionary distinct regional units (Peintinger et al. 2012) or narrowly distributed neo-endemics (Heydel et al. 2017). However, for the purpose of this large-scale study we could not collect all those variants that in part may reflect ecotypical differentiation. Following Peintinger et al. (2012), we have separated the *D. cespitosa* subsp. *rhena*na (Gremli) Kerguélen and *D. cespitosa* subsp. *littoralis* (Gaudin) K.Richter.

Additionally, we have included in this investigation two rare species related to *D. cespitosa*, namely *D. koelerioides* Regel, which was also considered a subspecies of the former by Tzvelev (1976), and *D. argentea* Lowe from Macaronesia which may be an endemic derivative of *D. cespitosa*. These samples were not included in the statistical analyses.

**Genome size measurements using flow cytometry (FCM)**

Twenty-five milligrams of fresh or dried leaf tissue was co-chopped (Galbraith et al. 1983) along with fresh standard leaf material (*Solanum pseudocapsicum* L., 1C = 1.295 pg; Temsch et al. 2010) in Otto’s buffer I (Otto et al. 1981) using a sharp razor blade. The resulting nuclear isolate was filtered through a 30-µm nylon mesh. Subsequently, double-stranded RNA was removed by a half an hour treatment with RNase A (Sigma-Aldrich, USA) at 37 °C. Afterward, Otto’s buffer II (Otto et al. 1981) that contained 50 mg/L propidium iodide (PI; AppliChem GmbH, Germany) was added. The preparations were allowed to incubate before measurement for at least 1 h in the refrigerator or overnight. The samples were measured using a flow cytometer CyFlow ML or CyFlow space (both Sysmex Partec GmbH, Germany), equipped with a diode-pumped laser (532 nm, 100 mW, Cobolt AB, Sweden). The 1C values of each sample were calculated in respect of a linear relationship between the mean fluorescence intensity (FI) of the G1 nuclei population of the standard and the samples. 1C value sample = mean FI sample G1 / mean FI standard G1 * 1C-value standard. From 3,333 up to 10,000 particles were measured per preparation. Means and standard deviations for several runs are given in Table 1. For visualization of different genome sizes, squared decimal size classes were used with genome sizes < 4.1 pg combined in a single class. The DNA ploidy estimated from the FCM data was calibrated by chromosome counts in one tetraploid and two diploid samples.

**Chromosome number analyses and Feulgen densitometry (FDM)**

Feulgen densitometry (FDM) was used for chromosome number analyses in order to verify the ploidy level in two diploid and one tetraploid individuals (genome size data not given. Table 1 contains only FCM data of these samples). Root tips were harvested from healthy plantlets of pseudoviviparous plants grown in petri dishes and from samples of seminiferous (seed bearing) plants grown in HBV. For chromosome number analyses, selected root tip meristems were pre-treated with 0.002 M 8-hydroxyquinoline in darkness for 2.5 h at room temperature and 2.5 h at 4 °C, fixed in methanol/acetic acid (3:1) overnight and stored at − 20 °C until use. The fixed root tips were washed six times in distilled water together with fixed root tips from an internal standard and hydrolyzed in 5 N HCl in an ultra-thermostatic water bath (Model LTD6, Grant Instruments Ltd., Cambridge, Barrington, England) at 20.0 °C for 60 min. The hydrochloric acid was removed by three washings with distilled water, followed by the staining step with Schiff’s reagent (Merck, Darmstadt, Germany) for 1.5 h at room temperature under light protection. The samples were subsequently washed six times with SO₂ water (0.02 M potassium metabisulfite dissolved in 0.01 N HCl) over a time period of 45 min. Each root tip was squashed in 45% acetic acid on a slide under a cover slip. After removal of the cover slip, the slides were shortly fixed with ethanol (96%) and finally air-dried.

The preparations were analyzed using the AxioPlan light microscope (Carl Zeiss, Vienna, Austria) equipped with a CCD black–white camera. Images of the metaphase plates were captured using ZEN software (Carl Zeiss, Vienna, Austria). Genome size was estimated from telophases of the objects and the standard in order to unambiguously assign the ploidy level to the genome size estimates on the basis of single root tips. Therefore, the integrated optical density
## Table 1
The studied Deschampsia populations, their holoploid and monoploid genome size values (1C and 1Cx, pg), ploidy level, collection site including latitude, longitude and altitude

| Nr | Country | Prov/State/County | Latitude | Longitude | Altitude (m) | 1C (pg) | S.D. | n | runs | 1Cx (pg) DNA-Ploidy | Mat (d/f) | Rep | (s/v) | Hab class | Subsp. | Sample ID |
|----|---------|-------------------|----------|-----------|--------------|---------|------|---|------|----------------------|-----------|-----|------|-----------|-------|-----------|
| 1  | ARM     | Tawusch           | 40.83593 | 4.3282    | 1             | 2x       |       |   |      |                       |           |     |      | Wet       | cespitosa | 228       |
| 2  | AUT     | Steiermark        | 47.42556 | 4.3924    | 1             | 2x       |       |   |      |                       |           |     |      | Wet       | cespitosa | 1         |
| 3  | AUT     | Steiermark        | 47.37056 | 4.3924    | 1             | 2x       |       |   |      |                       |           |     |      | Wet       | cespitosa | 2         |
| 4  | AUT     | Niederösterreich  | 47.91556 | 4.3671    | 1             | 2x       |       |   |      |                       |           |     |      | Wet       | cespitosa | 12        |
| 5  | AUT     | Steiermark        | 47.63889 | 4.3732    | 1             | 2x       |       |   |      |                       |           |     |      | Wet       | cespitosa | 15        |
| 6  | AUT     | Niederösterreich  | 47.71889 | 4.3769    | 1             | 2x       |       |   |      |                       |           |     |      | Wet       | cespitosa | 24        |
| 7  | AUT     | Niederösterreich  | 47.54889 | 4.4079    | 1             | 2x       |       |   |      |                       |           |     |      | Wet       | cespitosa | 48        |
| 8  | AUT     | Wien              | 48.19289 | 4.4067    | 1             | 2x       |       |   |      |                       |           |     |      | Wet       | cespitosa | 15        |
| 9  | AUT     | Vorarlberg        | 47.50667 | 4.3710    | 1             | 4x       |       |   |      |                       |           |     |      | Wet       | rhenana   | 990       |
| 10 | AUT     | Niederösterreich  | 47.99417 | 4.6660    | 3             | 2x       |       |   |      |                       |           |     |      | Forest    | cespitosa | 994       |
| 11 | AUT     | Wien              | 48.19305 | 4.3741    | 3             | 4x       |       |   |      |                       |           |     |      | Wet       | cespitosa | 995a      |
| 12 | AUT     | Niederösterreich  | 47.85691 | 4.2522    | 3             | 2x       |       |   |      |                       |           |     |      | Ruderal   | cespitosa | 998b      |
| 13 | AUT     | Canton Vaud       | 46.67133 | 4.4629    | 1             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998c      |
| 14 | AUT     | Canton Thurgau    | 47.04633 | 4.4064    | 1             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998d      |
| 15 | AUT     | Canton Vaud       | 46.67228 | 4.3759    | 1             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998e      |
| 16 | AUT     | Canton Thurgau    | 47.65528 | 4.5192    | 4             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998f      |
| 17 | AUT     | Canton Vaud       | 47.65528 | 4.5192    | 4             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998g      |
| 18 | AUT     | Canton Thurgau    | 47.65528 | 4.5192    | 4             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998h      |
| 19 | AUT     | Canton Thurgau    | 47.65528 | 4.5192    | 4             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998i      |
| 20 | AUT     | Canton Thurgau    | 47.65528 | 4.5192    | 4             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998j      |
| 21 | AUT     | Canton Thurgau    | 47.65528 | 4.5192    | 4             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998k      |
| 22 | AUT     | Canton Thurgau    | 47.65528 | 4.5192    | 4             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998l      |
| 23 | AUT     | Canton Thurgau    | 47.65528 | 4.5192    | 4             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998m      |
| 24 | AUT     | Canton Thurgau    | 47.65528 | 4.5192    | 4             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998n      |
| 25 | AUT     | Canton Thurgau    | 47.65528 | 4.5192    | 4             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998o      |
| 26 | AUT     | Canton Thurgau    | 47.65528 | 4.5192    | 4             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998p      |
| 27 | AUT     | Canton Thurgau    | 47.65528 | 4.5192    | 4             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998q      |
| 28 | AUT     | Canton Thurgau    | 47.65528 | 4.5192    | 4             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998r      |
| 29 | AUT     | Canton Thurgau    | 47.65528 | 4.5192    | 4             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998s      |
| 30 | AUT     | Canton Thurgau    | 47.65528 | 4.5192    | 4             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998t      |
| 31 | AUT     | Canton Thurgau    | 47.65528 | 4.5192    | 4             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998u      |
| 32 | AUT     | Canton Thurgau    | 47.65528 | 4.5192    | 4             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998v      |
| 33 | AUT     | Canton Thurgau    | 47.65528 | 4.5192    | 4             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998w      |
| 34 | AUT     | Canton Thurgau    | 47.65528 | 4.5192    | 4             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998x      |
| 35 | AUT     | Canton Thurgau    | 47.65528 | 4.5192    | 4             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998y      |
| 36 | AUT     | Canton Thurgau    | 47.65528 | 4.5192    | 4             | 2x       |       |   |      |                       |           |     |      | Wet       | littoralis | 998z      |
### Table 1 (continued)

| Nr | Country      | Prov/State/County | Latitude       | Longitude      | Altitude (m) | 1C (pg) | S.D. | n runs | 1Cx (pg) | DNA Ploidy | Mat (d/f) | Repr (s/v) | Hab class       | Subsp.        | Sample ID |
|----|--------------|-------------------|----------------|----------------|--------------|----------|------|--------|----------|------------|-----------|------------|----------------|---------------|-----------|
| 37 | CHN          | Sichuan           | 31.731949      | 100.735661     | 3908         | 5.3232   |      | 1      | 5.3232   | 2x         | d         | s          | Ruderal orientalis** | 209           |           |
| 38 | CHN          | Sichuan           | 31.708748      | 102.312484     | 4010         | 5.0215   |      | 1      | 5.0215   | 2x         | d         | s          | Ruderal orientalis** | 210           |           |
| 39 | CHN          | Sichuan           | 30.279167      | 99.553056      | 4642         | 5.1593   |      | 1      | 5.1593   | 2x         | d         | s          | Wet cespitosa s.l. | 214           |           |
| 40 | CZE          | Severomoravsky    | 49.819153      | 17.666156      | 440          | 4.2363   |      | 1      | 4.2363   | 2x         | d         | s          | Wet cespitosa      | 29            |           |
| 41 | DEU          | Bayern            | 48.872083      | 13.047889      | 753          | 4.3616   |      | 1      | 4.3616   | 2x         | f         | s          | Wet cespitosa      | 3             |           |
| 42 | DEU          | Bayern            | 48.018300      | 11.141825      | 533          | 4.5704   | 0.1649| 6      | 4.5704   | 2x         | f         | s          | Forest cespitosa   | 4             |           |
| 43 | DEU          | Baden-Württemberg | 47.686817      | 9.086317       | 395          | 4.3220   | 0.0760| 6      | 4.3220   | 2x         | d         | s          | Wet cespitosa      | 5             |           |
| 44 | DEU          | Schleswig-Holstein| 53.618611      | 13.616944      | 60           | 4.3278   |      | 1      | 4.3278   | 2x         | d         | s          | Forest cespitosa   | 18            |           |
| 45 | DEU          | Thüringen         | 50.809444      | 10.407778      | 540          | 3.9323   |      | 1      | 3.9323   | 2x         | d         | s          | Forest cespitosa   | 19            |           |
| 46 | DEU          | Nordrhein-Westfalen| 50.514722      | 6.285833       | 540          | 4.0363   |      | 1      | 4.0363   | 2x         | d         | s          | Ruderal cespitosa  | 20            |           |
| 47 | DEU          | Sachsen-Anhalt    | 51.748333      | 10.935000      | 420          | 3.7543   |      | 1      | 3.7543   | 2x         | d         | s          | Forest cespitosa   | 21            |           |
| 48 | DEU          | Schleswig-Holstein| 54.093056      | 10.094167      | 30           | 4.5338   |      | 1      | 4.5338   | 2x         | d         | s          | Forest cespitosa   | 22            |           |
| 49 | DEU          | Baden-Württemberg | 47.679169      | 9.322386       | 450          | 4.5940   | 0.0591| 7      | 4.5940   | 2x*        | f         | s          | Ruderal cespitosa  | 37            |           |
| 50 | DEU          | Baden-Württemberg | 47.670556      | 9.213889       | 400          | 8.7546   | 0.1823| 10     | 4.3773   | 4x         | d         | v          | Wet rhenana      | 152           |           |
| 51 | DEU          | Baden-Württemberg | 47.747500      | 9.143056       | 400          | 8.4793   | 0.3500| 9      | 4.2397   | 4x         | d         | v          | Wet rhenana      | 153           |           |
| 52 | DEU          | Bayern            | 47.561111      | 9.644167       | 400          | 8.6444   |      | 1      | 4.3222   | 4x         | f         | v          | Wet rhenana      | 996           |           |
| 53 | DEU          | Baden-Württemberg | 47.669722      | 9.329167       | 400          | 8.8603   | 0.0459| 3      | 4.4302   | 4x         | f         | v          | Wet rhenana      | 997           |           |
| 54 | DEU          | Baden-Württemberg | 47.703056      | 9.043611       | 400          | 8.7967   | 0.1694| 5      | 4.3984   | 4x         | f         | v          | Wet rhenana      | 999           |           |
| 55 | ESP          | Castilla y León  | 42.855970      | −6.817722      | 1700         | 7.9426   | 0.0633| 3      | 3.9713   | 4x         | d         | s          | Wet cespitosa     | 94            |           |
| 56 | ESP          | Ávila. Piedrahita | 40.423170      | −5.293222      | ?            | 4.2178   | 0.0065| 3      | 4.2178   | 2x         | d         | s          | Wet cespitosa     | 95            |           |
| 57 | EST          | Ida-Viru County   | 59.004167      | 27.360556      | 40           | 4.3729   | 0.0079| 3      | 4.3729   | 2x         | d         | s          | Forest cespitosa  | 50            |           |
| 58 | EST          | Ida-Viru County   | 58.981490      | 27.183890      | 40           | 4.3863   | 0.0551| 3      | 4.3863   | 2x         | d         | s          | Forest cespitosa  | 142           |           |
| 59 | FIN          | Etelä-Suomen      | 60.343889      | 25.603528      | 8            | 4.3677   |      | 1      | 4.3677   | 2x         | d         | s          | Forest cespitosa  | 39            |           |
| 60 | FIN          | Etelä-Savo        | 61.672528      | 27.276694      | 88           | 4.2418   |      | 1      | 4.2418   | 2x         | d         | s          | Ruderal cespitosa | 40            |           |
| 61 | FIN          | Lapland           | 65.616306      | 26.718333      | 134          | 4.5808   |      | 1      | 4.5808   | 2x         | d         | s          | Forest cespitosa  | 41            |           |
| 62 | FIN          | Lapland           | 67.973283      | 23.681267      | 256          | 4.3201   |      | 1      | 4.3201   | 2x         | d         | s          | Forest cespitosa  | 42            |           |
| 63 | FIN          | Enotekió          | 69.060944      | 20.770000      | 494          | 4.4937   | 0.0263| 4      | 4.4937   | 2x         | d         | s          | Ruderal cespitosa | 43            |           |
| 64 | FIN          | Enotekió          | 69.089222      | 20.782000      | 595          | 4.0615   | 0.0748| 5      | 4.0615   | 2x         | d         | s          | Wet cespitosa     | 44            |           |
| 65 | FIN          | Northern Ostrobothnia | 65.028944   | 25.446111      | 2            | 4.5265   |      | 1      | 4.5265   | 2x         | d         | s          | Wet cespitosa     | 45            |           |
| Nr | Country | Prov/State/County | Latitude  | Longitude  | Altitude (m) | Nr Cx (pg) | S.D. | Mat (d/f) | Repr (s/v) | Hab class | Subsp. | Sample ID |
|----|---------|------------------|-----------|------------|--------------|-----------|------|-----------|------------|-----------|-------|-----------|
| 66 | FIN     | Central Ostrobothnia | 63.817194 | 22.978056 | 2            | 4.0070    |       | 2x        | d          | s        | Forest | cespitosa | 46    |
| 67 | FRA     | Auvergne. Massif Central | 45.066486 | 4.079469 | 1100         | 8.4467    |       | 4x        | d          | s        | Ruderal | cespitosa | 31    |
| 68 | FRA     | Auvergne. Massif Central | 45.070831 | 4.126953 | 1200         | 8.0231    | 0.0651 | 4x        | d          | s        | Ruderal | cespitosa | 32    |
| 69 | FRA     | Rhone Alpes. Vercors | 45.071469 | 5.494672 | 730          | 4.2972    |       | 2x        | d          | s        | Wet     | cespitosa | 33    |
| 70 | FRA     | Rhone Alpes. M. Vanoise | 45.388144 | 6.567428 | 1400         | 4.3943    |       | 2x        | d          | s        | Wet     | cespitosa | 34    |
| 71 | FRA     | Haut-Jura | 46.496972 | 6.083319 | 1010         | 4.5695    | 0.1122 | 7         | d          | s        | Forest | cespitosa | 35    |
| 72 | GBR     | England. North Yorkshire | 54.245978 | −0.682961 | 140          | 8.5221    | 0.2794 | 4         | d          | s        | Ruderal | cespitosa | 14    |
| 73 | GBR     | Scotland. Ross & Cromarty | 57.600000 | −4.951189 | 125          | 4.2978    | 0.0831 | 4         | d          | s        | Ruderal | cespitosa | 13a   |
| 74 | GBR     | Scotland. Ross & Cromarty | 54.245978 | −0.682961 | 140          | 8.2434    | 0.0425 | 4         | d          | s        | Ruderal | cespitosa | 13b   |
| 75 | HRV     | Zagorje | 45.947222 | 15.825000 | 4.3362       | 1         | 4.3362 | 2x        | d          | s        | Wet     | cespitosa | 141   |
| 76 | ISL     | Western Region | 64.600556 | −22.019444 | 35          | 4.5419    | 0.1443 | 11        | d          | s        | Grass   | cespitosa | 96    |
| 77 | ISL     | Western Region | 64.877222 | −23.685833 | 200         | 8.5727    | 0.0112 | 3         | d          | v        | Grass   | alpina    | 97    |
| 78 | ISL     | Western Region | 64.877222 | −23.685833 | 150         | 4.4866    | 0.0153 | 3         | d          | s        | Ruderal | cespitosa | 98    |
| 79 | ISL     | Western Region | 65.550278 | −24.352778 | 5           | 4.4427    | 0.0070 | 3         | d          | s        | Ruderal | cespitosa | 100   |
| 80 | ISL     | Northeastern Region | 65.595833 | −17.177500 | 260         | 4.4680    | 0.0071 | 3         | d          | s        | Forest  | cespitosa | 105   |
| 81 | ISL     | Northeastern Region | 66.048333 | −17.345000 | 20          | 4.5668    | 0.0064 | 3         | d          | s        | Ruderal | cespitosa | 106   |
| 82 | ISL     | Southern Region | 63.891944 | −21.364444 | 10          | 4.2975    | 0.0646 | 9         | d          | s        | Ruderal | cespitosa | 109   |
| 83 | ISL     | Northeastern Region | 65.669390 | −15.095570 | 155         | 4.5180    | 0.0141 | 3         | d          | s        | Grass   | cespitosa | 115   |
| 84 | ISL     | Northeastern Region | 66.361740 | −14.947700 | 11          | 4.4531    | 0.0084 | 3         | d          | s        | Grass   | cespitosa | 116   |
| 85 | ISL     | Northeastern Region | 66.361740 | −14.947700 | 11          | 8.3636    | 0.0201 | 3         | d          | v        | Wet     | alpina    | 117   |
| 86 | ISL     | Northeastern Region | 65.512222 | −18.602778 | 290         | 4.4406    | 0.0585 | 4         | d          | s        | Grass   | cespitosa | 108a  |
| 87 | ISL     | Northeastern Region | 65.512222 | −18.602778 | 290         | 8.3278    | 0.0202 | 3         | d          | v        | Wet     | alpina    | 108b  |
Table 1 (continued)

| Nr | Country | Prov/State/County     | Latitude   | Longitude | Altitude (m) | 1C (pg) S.D. n runs | 1Cx (pg) DNA Ploidy Mat (d/f) Repr (s/v) Hab class Subsp. Sample ID |
|----|---------|-----------------------|------------|-----------|---------------|---------------------|-----------------------------|---------------------------------------------------------------------|
| 88 | ISL     | Northeastern Region   | 65.512222  | −18.602778| 290           | 6.6817 0.0727 4     | 4.4545 3x d s Wet cesp X alp 108c                                   |
| 89 | ITA     | Bolzano               | 46.770067  | 11.966483 | 1079          | 4.4418              | 1 4.4418 2x f s Forest cespitosa 10                                |
| 90 | ITA     | Udine                 | 46.496567  | 13.690333 | 857           | 4.3915              | 1 4.3915 2x f s Forest cespitosa 11                                |
| 91 | ITA     | Toscana, Pistoia      | 44.126389  | 10.63611  | 1560          | 8.6328              | 1 4.3164 4x d s Grass cespitosa 23                                |
| 92 | KGZ     | Osh Region            | 39.490520  | 72.911420 | 3560          | 4.9631              | 1 4.9631 2x d s Wet cespitosa s.l. 227                             |
| 93 | KOR     | Cheju Island          | 33.361872  | 126.51726 | 1680          | 4.2285 0.2336 6     | 4.2285 2x d s Forest cespitosa 36                                |
| 94 | MKD     | Municip Struga        | 41.260000  | 20.531944 | 8.4416        | 0.0801              | 3 4.2208 4x d s Wet cespitosa 144                                 |
| 95 | NOR     | Viken                 | 60.123611  | 10.374722 | 200           | 4.4086 0.1026 4     | 4.4086 2x d s Grass cespitosa 158                                 |
| 96 | POL     | Sulecyn County        | 52.292317  | 15.058283 | 90            | 4.3018              | 1 4.3018 2x d s Ruderal cespitosa 26                               |
| 97 | POL     | Magura Witowska       | 49.315811  | 19.831108 | 830           | 4.0800              | 1 4.0800 2x d s Forest cespitosa 28                               |
| 98 | POL     | Lublin, Chelm County  | 51.143056  | 23.311389 | 210           | 3.8728              | 1 3.8728 2x d s Ruderal cespitosa 76                               |
| 99 | RUS     | Tyumen Region         | 57.105000  | 66.106389 | 70            | 3.9999 0.0104 2     | 3.9999 2x d s Forest cespitosa 78                                 |
| 100| RUS     | Sverdlkov Region      | 57.146111  | 60.215556 | 290           | 4.3737 0.0793 3     | 4.3737 2x d s Grass cespitosa 82                                  |
| 101| RUS     | Sverdlkov Region      | 59.583333  | 69.300000 | 30            | 4.3794 0.0106 3     | 4.3794 2x d s Forest cespitosa 84                                 |
| 102| RUS     | Leningrad Region      | 60.649710  | 33.11631  | 10            | 4.3966 0.0376 3     | 4.3966 2x d s Forest cespitosa 87                                 |
| 103| RUS     | Sakhalin Region       | 48.225640  | 142.54124 | 30            | 4.2112 0.0327 3     | 4.2112 2x d s Ruderal cespitosa s.l. 88                            |
| 104| RUS     | Tver Region           | 57.777660  | 35.223740 | 160           | 4.3066              | 1 4.3066 2x d s Grass cespitosa 89                                 |
| 105| RUS     | Moscow Region         | 55.091185  | 37.501470 | 180           | 4.5834              | 1 4.5834 2x d s Ruderal cespitosa 121                               |
| 106| RUS     | Murmansk Region       | 67.860000  | 34.420000 | 380           | 4.4595              | 1 4.4595 2x d s Ruderal cespitosa 122                               |
| 107| RUS     | Karelia               | 66.301960  | 33.286260 | 0             | 4.7046              | 1 4.7046 2x d s Grass cespitosa 131                               |
| 108| RUS     | Krasnodar Region      | 44.070500  | 39.703830 | 1500          | 4.5176              | 1 4.5176 2x d s Ruderal cespitosa 134                               |
| 109| RUS     | Taymir Region         | 69.003944  | 91.008389 | 48            | 8.4955              | 1 4.2478 4x d s Wet cespitosa 149                                  |
| 110| RUS     | Yakutia               | 70.714660  | 127.417820| 8             | 4.5432              | 1 4.5432 2x d s Wet obensis 217                                   |
| 111| RUS     | Yakutia               | 71.847340  | 126.909180| 220           | 4.8735              | 1 4.2368 4x d s Wet obensis 218                                   |
| 112| RUS     | Yakutia               | 72.001590  | 129.10933 | 10            | 4.6751 0.2681 3     | 4.6751 2x d s Ruderal submutica 219                                |
| 113| RUS     | Yakutia               | 71.563930  | 128.760910| 30            | 8.4726              | 1 4.2363 4x d s Wet submutica 220                                 |
| 114| RUS     | Yakutia               | 66.775500  | 123.371460| 30            | 4.5629              | 1 4.5629 2x d s Wet submutica 221                                 |
| 115| RUS     | Burjatja              | 52.909170  | 108.160600| 450           | 4.9784              | 1 4.9784 2x d s Wet turczaninowii 222                              |
| 116| RUS     | Irkutskaja Region     | 53.398330  | 107.438300| 450           | 4.7819              | 1 4.7819 2x d s Wet turczaninowii 223                              |
| 117| RUS     | Chukotka              | 66.115440  | −170.52000 | 30            | 4.7392              | 1 4.7392 2x d s Wet obensis 235                                   |
| 118| RUS     | Chukotka              | 64.714900  | −174.09050 | 30            | 4.8121              | 1 4.8121 2x d s Wet beringensis 236                                |
| 119| RUS     | Novgorod Region       | 57.082310  | 30.751470 | 90            | 4.4866              | 1 4.4866 2x d s Wet cespitosa s.l. 237                             |
| 120| RUS     | Kostroma Region       | 58.349722  | 42.216111 | 130           | 4.2465 0.0184 2     | 4.2465 2x d s Wet cespitosa s.l. 239                             |
Table 1 (continued)

| Nr | Country Prov/State/County | Latitude | Longitude | Altitude (m) | ICx (pg) | DNA runs | DNA Ploidy | Mat (d/f) | Repro | Hab class | Subsp. | Sample ID |
|----|---------------------------|----------|-----------|--------------|----------|----------|-----------|-----------|-------|-----------|--------|-----------|
| 121 | RUS Voronezh Region | 51.195250 | 40.308080 | 100 | 4.0687 | 1 | 4.0687 | 2x | d | Forest | cespitosa | 241 |
| 122 | SVK Mala Fatra | 49.253836 | 19.045972 | 55 | 0.0244 | 3 | 4.0507 | 2x | d | Forest | cespitosa | 27 |
| 123 | SWE Skane | 55.713330 | 13.306110 | 50 | 4.2346 | 1 | 4.2346 | 2x | d | Grass | koelerioides | 25 |
| 124 | UKR Chernivtsi Region | 48.291840 | 25.876510 | 4.2715 | 1 | 4.2715 | 2x | d | Wet | koelerioides | 111 |
| 125 | UKR Zarkapattia Region | 48.150800 | 24.357580 | 1680 | 4.2715 | 1 | 4.2715 | 2x | d | Wet | koelerioides | 111 |

If more than one run per population was performed, mean values along with standard deviation (S.D.) are indicated. Country codes according to ISO 3166 standard: AUT Austria, ARM Armenia, CHE Switzerland, CZE Czech Republic, DEU Germany, ESP Spain, EST Estonia, FIN Finland, GRE Greece, HUN Hungary, IRL Ireland, ITA Italy, JAP Japan, KOR South Korea, LIT Lithuania, LTU Latvia, MKD North Macedonia, NLD Netherlands, POL Poland, POR Portugal, RUS Russian Federation, SWE Sweden, SVK Slovakia, UKR The Ukraine; sample (Mat): dry (d), fresh (f); reproduction (Repr): seminiferous (s), pseudoviviparous (v); habitat class (Habclass): forest/ruderal/wetland/grassland.

*DNA ploidy confirmed by chromosome count
**Approaching D. koelerioides in panicle shape (more contracted) and color (due to mostly golden glumes)

BIogeography, habitat and reproductive system

We have divided the entire Deschampsia cespitosa samples into two regions (Europe and Asia) separated by the 60th degree longitude (Urals). Longitudinal classes covering 10 degrees were applied for visualization of genome size variation related to geography. Four habitat classes (Habclass) based on descriptive sampling data were distinguished: forest (including clearings); grass (meadows, pastures); ruderal (wasteland, roadsides); and wet (wet meadows, lake shores, riverbanks, riparian gravel). Plants were collected in reproductive phase (flowering or with proliferous spikelets), or the reproductive system was determined from monitoring records (observed by MP).

Statistical analysis

SPSS statistics 25.0 (IBM, Armonk, New York, USA) was used for the analyses and visualization (box plots, bar charts and scatter diagrams) of the overall genome size variation, the variation within the ploidy levels, and its association with habitat classes. Nonparametric options were used to test for differences in sample parameters (Kruskal–Wallis tests). A linear multivariate regression was used to evaluate the effects of altitude, habitat and bioclimatic factors on the genome size of diploid plants. We extracted 19 bioclimatic variables of climate data for each sample location using ArcGIS 10.0.4 from the WorldClim database (Fick and Hijmans 2017; https://worldclim.org/data/worldclim21.html), which are the average for the years 1970–2000 at a resolution of 5 min. We ran pairwise correlation analyses among the 19 bioclimatic variables (BIO1–BIO19) and choose nine with low correlation ($R < 0.8$) which are BIO1 (annual mean temperature), BIO2 (mean diurnal range), BIO3 (isothermality), BIO4 (temperature seasonality), BIO5 (max temperature of warmest month), BIO8 (mean temperature of wettest quarter), BIO9 (mean temperature of driest quarter), BIO10 (annual precipitation) and BIO12 (precipitation of warmest quarter). The AIC was employed to determine the best model in stepwise backward multiple regression. The partial effects of each dependent variable in the best model are shown in partial plots according to Grace et al. (2016).
Genome size variation and ploidy

The geographical distribution of the entire sample is shown on the map in Fig. 1. The genome size (1C) ranged from 3.754 to 5.438 pg in diploid and from 7.943 to 8.973 pg in tetraploid *Deschampsia cespitosa* (Fig. 2). We report here also the genome size of 4.864–5.235 pg found in *D. koelerioides* and 4.753 pg for *D. argentea*. Genome size data and DNA ploidy of the investigated population samples as revealed by the FCM measurements are provided in Table 1. Mitotic metaphase chromosomes for a diploid plant (2n = 26, 1C = 4.374 pg) of population 13 and a tetraploid plant (2n = 52, 1C = 8.973 pg) of population 20 are shown in Online Resource 3. The genome size of one individual of 6.682 pg suggested a putative DNA triploid, sampled in a mixed population (therefore, different numbers 86–88 in Table 1) of seminiferous and pseudoviviparous plants. Genome size (1C, pg) clearly varied between diploids, tetraploids and the putative triploid (Fig. 2). The monoploid genome sizes (1Cx, pg) showed geographically structured variation within the diploids, and between some diploids and all tetraploids, but not within tetraploids (Table 1, Fig. 3).
Biogeographic patterns and environmental determinants of genome sizes

Monoploid genome sizes (Cx, pg) of Asian (median = 4.881) and European (median = 4.377) diploid populations (Fig. 3) were significantly different ($H = 41.441, P < 0.001$). Similarly, Asian diploids and all European tetraploids (median = 4.314) were significantly ($H = 58.007, P < 0.001$) different. There was a weak although nonsignificant tendency toward smaller genome sizes (i.e., down-sizing) in the European tetraploids compared to the diploids of the same region. The sample of 4x Asia was too small ($n = 3$) for any meaningful statistics. Variation in monoploid genome size from Western Europe to eastern Asia along longitude classes of 10 degree is shown in Online Resource 4.

While the tetraploid pseudoviviparous variants are clustered in Iceland and around Lake Constance (insets in Fig. 1), the distribution of the tetraploid seminiferous plants did not show a distinct geographic pattern. We found them in southern and Western Europe as well as in Arctic Siberia. No tetraploid seminiferous plants were found in northern Europe and in remaining Asia.

The genome size variation of the entire European and Asian sample of diploid plants was best explained by a model containing altitude, BIO5 (max temperature of warmest month) and BIO12 (annual precipitation). The altitude had a positive effect ($R_\partial = 0.53, P < 0.001$), and BIO5 ($R_\partial = -0.26, P < 0.001$) as well as BIO12 ($R_\partial = -0.25, P = 0.011$) had a negative effect on genome size (GS increment/decrement) shown in the partial plots (Fig. 4). The same analysis did not find any significant effect on genome size within the European diploids. European plants with small genomes ($1C < 4.1 \text{ pg}$) were confined to low and moderate altitudes (< 830 m) mainly in temperate latitudes (49–52° N) of Central Europe, while plants with large genomes were found in the whole range of altitudes being scattered across all the studied area in Europe (Fig. 5a, b).

### Ploidy in relation to ecology and reproduction

Frequencies of diploids and tetraploids considering their reproduction mode were analyzed in the four habitat types (Fig. 6). Both diploids and tetraploids showed a preference for wet habitats. Diploids were found in all environment types, whereas tetraploids were not found in forests. Seminiferous plants were either diploid or tetraploid, while pseudoviviparous plants were only tetraploid. Separating the tetraploids by habitat class, reproduction mode and region indicated the clear preference (91%) of pseudoviviparous plants for wet habitats, while a high proportion (50%) of the nonviviparous western tetraploid plants was found in ruderal habitats (Online Resource 5).

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| Region | Euro 4x | Euro 2x | Asia 4x | Asia 2x |
|--------|---------|---------|---------|---------|
| 1Cx (pg) | 5.5 | 5.0 | 4.5 | 4.0 | 3.5 |

*Fig. 3* Variation in monoploid genome size of *Deschampsia cespitosa* (Cx, pg) between diploids ($n$(Asia) = 29; $n$(Europe) = 71) and tetraploids ($n$(Asia) = 3; $n$(Europe) = 21)

*Fig. 4* Plots showing partial effects of predictors on genome size of diploid *Deschampsia cespitosa* revealed by a multivariate linear regression of a altitude, b BIO5 (max. temperature of warmest month), and c BIO12 (annual precipitation). Shaded areas represent 95% confidence intervals.
Discussion

Distribution patterns and ecology of Deschampsia cespitosa in relation to ploidy levels and genome size

This study confirmed that seminiferous Deschampsia cespitosa is predominantly diploid in Eurasia. In addition, it revealed a pattern of variation in genome size related to biogeography between diploid European and Asian plants. Several tetraploid populations were found, but their number was too low to draw more general conclusions on their distribution apart from the fact that the tetraploid seminiferous plants tended to occur more often in disturbed habitats in Southern and Western Europe and in wet habitats in East Asia, whereas tetraploid pseudoviviparous plants were restricted to periglacial or high-latitude regions and special environments. Diploid D. cespitosa was found in all environments but exhibited a preference for wet habitats. Polyploids were reported to be more common than diploids on the British Isles (Rothera and Davy 1986), and although there was no evidence that they were better adapted to cold environments, they were putatively associated with more disturbed habitats. Given the wide distribution of Subarctic and Arctic variants of diploid D. cespitosa (this paper; Kawano 1963; Tzvelev and Probatova 2019) and
the predominance of diploid *D. antarctica* E.Desv on the Antarctic Peninsula (González et al. 2016; Pascual-Díaz et al. 2020), it is unlikely that temperature alone affected the ploidy levels. Polyploids of various plant groups in arctic regions were hypothesized to be more successful than diploids in post-glacial (re-)colonization (Brochmann et al. 2004). Such polyploids were, however, often found to be of allopolyploid origin, and their success was hypothesized to result from effects of fixed heterozygosity. We do not know whether *D. cespitosa* tetraploids are of auto- or allopolyploid origin. Further genetic and molecular phylogenetic analyses are needed to infer their origin.

The tetraploid and pseudoviviparous variants/subspecies of our sample were found in lake shore habitats or close to water courses in cold environments. Peintinger et al. (2012) considered the subsp. *rhenana* around Lake Constance a periglacial relict endemic with special adaptations in its reproduction (facultative pseudovivipary) to flooding and/or very harsh conditions in glacial periods. The other tetraploid and pseudoviviparous subsp. *alpina* (L.) Tzvelev occurring in the mountains of northern Europe (Clarke 1980) was thought to result from several independent polyploidization events of diploid lineages of *D. cespitosa* with the potential of reproducing by pseudovivipary (Hedberg 1958).

The C values found for diploid *Deschampsia cespitosa* were mostly lower than the genome sizes reported for the closely related *D. antarctica* (1C(pg) between 5.30 and 5.36; Pascual-Díaz et al. 2020). The trend toward higher genome size in lower (southern) latitudes observed for the diploid Asian *D. cespitosa* samples resulted from the higher genome size values of the Chinese samples collected mainly in high altitudes between 3500 and 4000 m. Correlations of genome size and altitude in other plant groups were found to be group-specific: They were negative in wild relatives of *Zea mays* L. (Poaceae; Laurie and Bennet 1985; Bilinski et al. 2018) and in *Arachis duranensis* Krapov & W.C.Greg (Fabaceae; Temsch and Greilhuber 2001) and positive in *Lagenaria siceraria* (Molina) Standl (Cucurbitaceae: Achigan-Dako 2008), and no correlation was inferred for *Sesleria albicans* Kit. ex Schult (Poaceae; Lysak et al. 2000). The association between high genome size, low maximum temperature of the warmest month and low annual precipitation is plausible concerning the moderate temperature. Difficult to interpret is the effect of low precipitation on the genome size; however, Fig. 4c shows a high variation between < 500 and 1000 mm that is obviously not well processed by the statistical model. A general caveat is that the coarse grid climate data may not well represent the specific conditions at the collection sites. This concerns especially the Chinese samples (considered different subspecies) as we found *Deschampsia* mostly close to water courses in the high Chinese mountain ranges (Z.X., H.S., J.G.: pers. obs.). Other investigations found a negative correlation in Liliaceae between genome size and precipitation seasonality (Carta and Peruzzi 2016). Jakob et al. (2004) found disparate genome size patterns in different lineages of *Hordeum marinum* L. (Poaceae) in association with climatic variables. They concluded that phylogenetic constraints might be more important than ecological determinants.

![Fig. 6 Numbers of diploid and polyploid Deschampsia cespitosa populations separated by reproduction mode (sem = seminiferous; vivip = pseudoviviparous) in four habitat classes.](image)
presence of cryptic evolutionary lineages connected to different vegetation history across Europe was inferred to be responsible for the complex patterns of genome size variation within morphologically similar groups of *Picris hieracioides* L. (Asteraceae; Slovák et al. 2009). This can likely be also the case in *D. cespitosa* in Europe and on a large continental scale, especially that many of the regional variants have been classified as distinct subspecies (e. g., Conert 1987; Chiapella and Probatova 2003).

No evidence for genome downsizing was found in Europe by comparing monoploid genome sizes of diploids and tetraploids. In Asia, the tetraploid sample was too small for any conclusions. The only significant difference in monoploid genome size was found between the Asian diploids and all other European samples. It may likely represent two different geographically and thus genetically distinct lineages. Further genetic analyses will allow for more insight into this observed pattern.

**Ploidy and reproductive mode (seed producing versus pseudovivipary)**

All pseudoviviparous plants investigated in our study had a genome size indicative of DNA tetraploidy. A reproductive switch to pseudovivipary in wild populations was often inferred to be associated with polyploidy and specific environmental conditions (Hedberg 1958; Sarapul'tsev 2001). Experimentally induced environmental stress (short-time daylight) was reported to trigger a switch from seminiferous to pseudoviviparous reproduction in *D. cespitosa* (Nygren 1949); however, the association with polyploidy was not tested. There is only a scarce record of diploid and pseudoviviparous variants of *Deschampsia*. An experimentally induced switch to this reproductive mode was observed in diploid European accessions transplanted to transects in California (Lawrence 1945). Another diploid plant collected in the wild with pseudoviviparous reproduction was reported by Hedberg (1958). A few diploid chromosome numbers were also reported for samples assigned to the usually tetraploid and pseudoviviparous northern variant (subsp. *alpina*), however, without reference to the reproductive mode (Kawano 1963). One of these records refers to a diploid chromosome number in a seminiferous accession under the name *D. alpina* Roem. ex Schult which was hypothesized to be a potentially slender variety of *D. cespitosa* s. str. (Nygren 1949).

**Conclusions**

Seminiferous *Deschampsia cespitosa* s.l. is mostly diploid throughout Eurasia. However, a high variation (CV% = 7.24) in genome size was found among the diploid populations across the geographic west–east gradient in Eurasia. Genome size was significantly lower in European accessions than in the Asian ones. Tetraploids of seminiferous plants were often found on disturbed habitats, albeit without any geographically or ecologically consistent pattern in their distribution. Pseudovivipary was found to be specific only to polyploids and associated with habitats of high moisture.

**Information on Electronic Supplementary Material**

Online Resource 1. Performance test of two *Deschampsia cespitosa* samples in FCM analysis during five months.

Online Resource 2. Classification of *Deschampsia* in Europe and Asia.

Online Resource 3. Mitotic metaphase chromosomes of *Deschampsia cespitosa*.

Online Resource 4. Regional variation in monoploid genome size of *Deschampsia cespitosa*.

Online Resource 5. Polyploids of *Deschampsia cespitosa* in relation to reproduction mode, habitat class, and region.

**Supplementary Information** The online version contains supplementary material available at https://doi.org/10.1007/s00606-021-01796-7.

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**Author contributions** All authors contributed to the study conception and design. Material collection and preparation were performed by JG, ZX, PV, MP, PW, IS, HS and JOC. Analyses in the laboratory were performed by EMT and HWS, and data analyses were performed by EM, JG, ZX, HS and PV. The first draft of the manuscript was written by JG and EMT, and all authors commented on previous versions of the manuscript. All authors read and approved the final manuscript.

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**Declarations**

**Conflict of interest** The authors declare that they have no conflict of interest.
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