

Genetic analysis of Latvian *Salix alba* L. and hybrid populations using nuclear and chloroplast DNA markers

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Latvia is at the northern border of the species range of *S. alba* L. in Europe, and there has been some debate whether the Latvian populations of *S. alba* are autochthonous, as this species has long been planted in Latvia as an ornamental tree in gardens, parks and roadsides. In addition, there is increasing interest in the use of several *Salix* species (including *S. alba*) as bioenergy crops. Natural *S. alba* stands throughout Latvia, as well as stands of possibly hybrid origin were analysed using nuclear and chloroplast markers. Our results showed that *S. alba* populations are probably natural, and that the rate of vegetative reproduction is low, supporting the evidence that Latvia is within the natural range of *S. alba*. These results provide the basis for the identification of possibly introduced or artificially regenerated stands of *S. alba* in Latvia. In addition, our results confirm that *S. alba* hybridises with *S. fragilis*, and that natural stands including hybrid individuals can be established. The analysis of chloroplast markers indicated that the predominant hybridisation occurs by fertilisation of *S. fragilis* by *S. alba* pollen; however, the extent of haplotype sharing between these two species should be further investigated.

Keywords: Willow, Population Structure, Autochthonous, Provenance

Introduction

There is an increasing interest of using short rotation species such as white willow (*Salix alba* L.) as a source of biomass in Latvia (Pucka & Lazdina 2013). Currently, foreign clones of this species are mainly utilised in plantations, though there are plans to identify high-yield locally adapted clones. To this purpose, information about the structure, provenance and diversity of *Salix alba* populations in Latvia will assist in the identification of the most valuable genetic resources for further utilisation.

In Latvia, as well as Estonia and Lithuania, white willow has long been planted as an ornamental tree in gardens, parks and roadsides. White willow has intensively spread into surrounding areas due to the high reproductive capacity of this species, both sexual and vegetative. As a consequence, different opinions exist among researchers about whether this species is introduced or if it is native. The status of this species in Latvia is further complicated by two other factors. Firstly, white willow

has many sub-species and varieties, and also readily forms hybrids with other *Salix* species. Cinovskis et al. (1993) consider these sub-species and varieties as cultivated clones or decorative forms, which should not be considered at a taxonomic level. Secondly, various studies consider that Latvia is at the northern range boundary (or possibly outside) of the natural distribution area of this species, and therefore the provenance of particular white willow individuals or populations may be difficult to determine. Several studies (Meusel et al. 1965, Wagenitz 1981) place the distribution boundary of white willow to the north of Latvia, extending from lake Onega in Russia along the Svir River to lake Ladoga, then along the Neva River, the Gulf of Finland, and the east coast of the Baltic Sea, thus encompassing all of the Baltic States (Estonia, Latvia and Lithuania – Fig. 1). In contrast, other authors considered the boundary to be in northern Latvia, based on the extensive natural white willow stands found along the Gauja River (Rasiņš

1959). This northern boundary along the Latvian-Estonian border was also reported in the USSR Atlas of Woody Plants (Skvortsov & Svjaseva 1977), as well as in the European Flora Atlas, where *S. alba* stands in Latvia and Lithuania are reported as autochthonous, while Estonian stands are considered to be of unknown provenance (Jalas & Suominen 1976). The autochthonous status of white willow stands along the Gauja and Daugava rivers has been emphasised by Latvian dendrologists (Lange et al. 1978, Evarts-Bunders 2005). Evidence supporting this conclusion includes the presence of white willow in mixed species stands (including *Alnus incana*, *Alnus glutinosa*, *Ulmus glabra* and other species), as well as the mixed age structure of white willow stands (Dreimanis & Vulaks 1974, Laivins et al. 2009), while white willow is considered to form only even-aged stands at the northern distribution boundary (Lange et al. 1978, Korchagin 1957, Skvortsov & Svjaseva 1977). Other reports suggest that the northern distribution boundary might be in the southern part of the Baltic states, with mixed age natural stands of white willow found only along the Svete River in Latvia, and in particular along the Nemuna, Neri and Neveži rivers in Lithuania (Cinovskis 1996, Mauriņš & Zvirgzds 2006, Navasaitis et al. 2003).

Despite these uncertainties about the provenance and distribution area of white willow in the Baltic States, this species is found throughout this region, and was mentioned in the first description of the flora of the region (Fischer 1778), as well as in further descriptions of the flora of the

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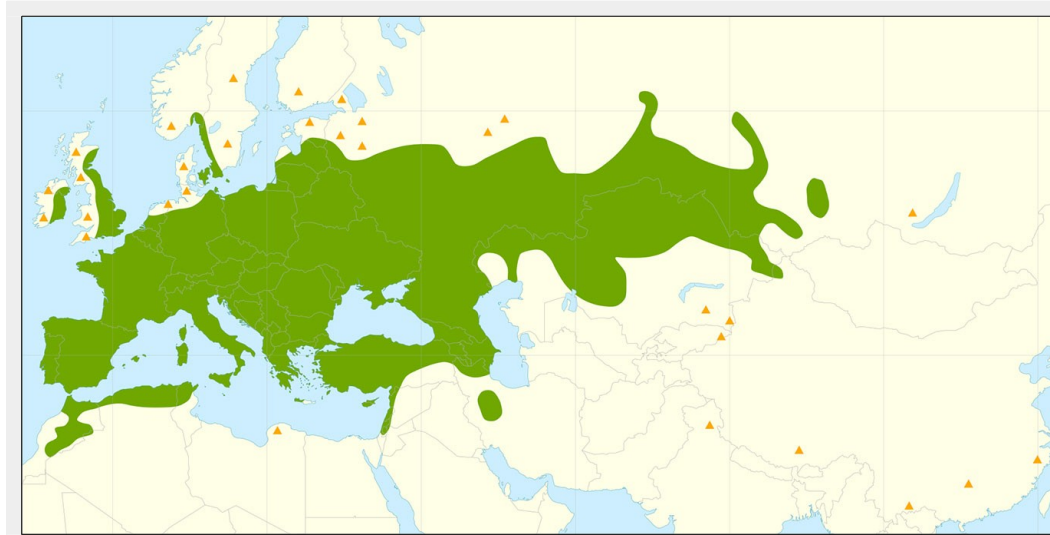


Fig. 1 - Natural distribution range of white willow (*Salix alba* L. – in green). Orange triangles represent putative introduced and/or naturalized (synanthropic) populations. Source: Hous-ton Durrant et al. (2016).

Baltic region. In the 19th century, *Salix alba* was reported to be a widespread species in the eastern Baltic region, commonly found in natural stands along sandy riversides and moist forests, as well as in artificial plantations (Klinge 1883). In the Atlas of Latvian Woody Plants, *Salix alba* is reported in 1349 sites as the dominant species (in both plantations and natural stands, grid size 5x5 km, frequency 48 %; Laivinš et al. 2009 – Fig. 2), and is the most common *Salix* species in Latvian parks and plantings (Cinovskis et al. 1974). Many *S. alba* cultivars have been cultivated for long time: cultivar ‘Calva’ was first described in 1700, ‘Chermesina’ and ‘Sericea’ approximately in 1840 (Krüsmann 1978). The variety or cultivar ‘Vitellina’ was described as a separate species by Linnaeus in 1671 (*S. vitellina* L. – Evarts-Bunders 2005), but the taxonomic status of these cultivars, sub-species or species is currently disputed (Cinovskis 1996). In this study, the samples collected from the National Botanic Garden (Salaspils, Latvia) are designated as cultivated clones or varieties. These cultivated *S. alba* varieties were compared with the putative natural populations of *S. alba* in Latvia to determine the genetic similarity between the cultivated and natural forms, and to possibly identify the escape of cultivated clones into natural habitats.

Various kinds of DNA markers have been

used to analyse *Salix* species, including dominant markers such as AFLPs, ISSR and RAPDs (Meneghetti et al. 2007), as well as co-dominant markers such as SSRs (Perdereau et al. 2014). *Salix* species have also been extensively studied with maternally-inherited chloroplast markers (Palmé et al. 2003, Perdereau et al. 2014), which enables the differentiation of lineages, as well as the direction of hybridisation. *S. alba* and *S. fragilis* are both polyploids, with *S. alba* reported as being stably tetraploid (Håkansson 1955), and *S. fragilis* being mainly tetraploid, but with other ploidy levels also identified (Barcaccia et al. 2003). SSR markers have been developed for a range of *Salix* species, and many SSR markers showed cross-species amplification in other *Salix* species and genera such as *Populus* (Barker et al. 2003, Hoshikawa et al. 2009). Analysis of polyploid species such as *S. alba* and *S. fragilis* with codominant markers such as SSRs is complicated by the difficulty of determining the allele dosage, and therefore they are often converted to binary, dominant marker genotypes for analysis. However, some *Salix* SSR markers have been reported to have disomic-mono-genic inheritance in tetraploid *S. alba* and *S. fragilis*, and therefore can be analysed as diploid, codominant markers (King et al. 2010). Identification of such SSR markers is time-consuming as the segregation of alle-

les must be determined in the progeny of controlled crosses.

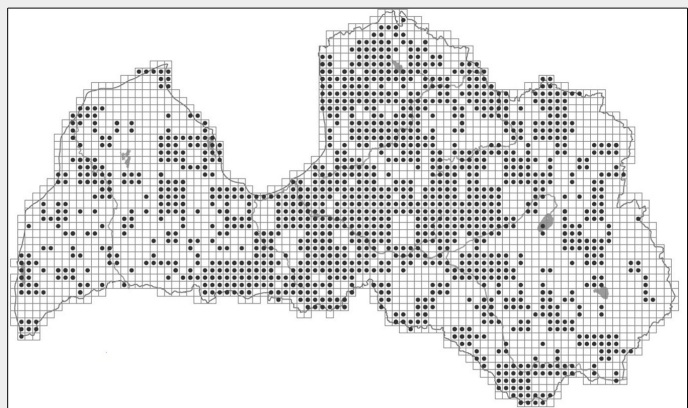
There is evidence that interspecific hybridisation is common between a range of *Salix* species (Palmé et al. 2003), and controlled crosses have shown that *S. alba* and *S. fragilis* can also readily hybridise (Triest et al. 2000). Additionally, continuous variation of morphological characteristics between these species has been reported in natural populations (Triest 2001). Despite this morphological continuum between the two species, DNA marker studies have indicated that the two species are clearly differentiated, suggesting that introgressive hybridisation is not widespread (Triest 2001, Oberprieler et al. 2013).

Given the increasing interest in Latvia of utilising *S. alba* as a short-rotation plantation species, we used nuclear and chloroplast markers to analyse the genetic diversity and population structure of natural *S. alba* stands throughout Latvia, with the aim of characterising the available genetic resources for this species in this region. In addition, stands of *S. fragilis* and putative hybrid individuals were analysed to investigate hybridisation between these two species. As the autochthony of Latvian *S. alba* populations is uncertain, individuals representing cultivated putative subspecies and other decorative forms were analysed, to determine their genetic similarity to natural *S. alba* stands in Latvia.

Materials and methods

Leaf samples of *S. alba* and putative hybrids with *S. fragilis* (based on phenotypic traits) were collected from 14 stands in Latvia, and one in Lithuania (Fig. 3, Tab. 1). Leaf morphological traits were analysed to differentiate the two species and to identify putative hybrids, particularly leaf colour/pubescence and serration. The populations were designated as *S. alba*, *S. fragilis* or hybrid populations, according to the dominant species, and the sampled individuals were morphologically selected according to the dominant species in each stand (Tab. 1). Individuals with hybrid morphol-

Fig. 2 - Distribution of white willow (*Salix alba* L.) in Latvia. Source: Laivinš et al. (2009).



ogy were dominant in only one population (Dviete). In addition, leaf samples from one individual of each of the *S. alba* cultivars 'Chermesina', 'Britzenis', 'Calva', 'Coccinea', 'Vitellina', 'Sericea', one individual of *S. alba* and two individuals of *Salix × rubens* Schrank (hybrid between *S. alba* and *S. fragilis*) were collected from the National Botanic Garden in Salaspils, Latvia (Tab. 1).

DNA from leaves was isolated using a CTAB-based method (Porebski et al. 1997). A total of 331 individuals were genotyped using eight nuclear SSR markers and six angiosperm chloroplast markers. Genotyping was carried out using nuclear SSR markers SB243, SB93, SB38, SB80, SB201, SB24, SB194 and SB199, which were reported to be informative in a range of *Salix* species (Barker et al. 2003). Six angiosperm chloroplast markers ccmp2, ccmp4, ccmp6, ccmp10, ccmp7 and ccmp3 were also utilised (Weising & Gardner 1999), which had previously been used to analyse *S. caprea* (Palmé et al. 2003, Perdereau et al. 2014). The PCR reactions were multiplexed as follows: set A (SB243, SB93, SB38); set B (SB80, SB201); set C (SB24, SB194, SB199); set D (ccmp2, ccmp3, ccmp10); set E (ccmp 4, ccmp6, ccmp7). Each forward primer in the multiplex sets was labelled with a different fluorophore (6-FAM, HEX or TMR) to facilitate visualisation using capillary electrophoresis. PCR reactions for the nuclear SSR markers were carried out in a 20 µl solution comprising: 4 µl of 5× HOT FIRE-Pol® Blend Master Mix (containing 10 mM MgCl₂, 2 mM dNTPs and BSA – Solis BioDyne, Estonia), 0.4 µM final concentration of each primer and 1 µl genomic DNA. The PCR profile consisted of an initial denaturation at 95 °C for 15 min, followed by 35 cycles of denaturation at 94 °C for 40 s, annealing at 48 °C for 60 s, and extension at 72 °C for 120 s. Final extension of amplified DNA occurred at 72 °C for 10 min. PCR reactions for the chloroplast SSR were carried out in a 10 µl solution comprising: 2 µl of 5× HOT FIREPol® Blend Master Mix containing 10 mM MgCl₂, 2 mM dNTPs and BSA (Solis BioDyne, Estonia), 0.2 µM final concentration of each primer and 1 µl genomic DNA. The PCR profile consisted of an initial denaturation at 95 °C for 15 min followed by 25 cycles of denaturation at 94 °C for 60 s, annealing at 53 °C (set D) or 50 °C (set E) for 60 s, and extension at 72 °C for 60 s. Final extension of amplified DNA occurred at 72 °C for 10 min. All PCR reactions were carried out in a Mastercycler® ep Gradient thermal cycler (Eppendorf, Hamburg, Germany). Amplification fragments were separated on an ABI Prism 3130xl® Genetic Analyzer (Applied Biosystems, Foster City, CA, USA) and visualized with the software GeneMapper® ver. 3.5 (Applied Biosystems).

The nuclear SSR genotypes were encoded as binary data to facilitate analysis of the polyploid species *S. alba* and *S. fragilis*. Two of the utilised markers (SB38 and SB194) have been reported to have disomic-monogenic inheritance in tetraploid

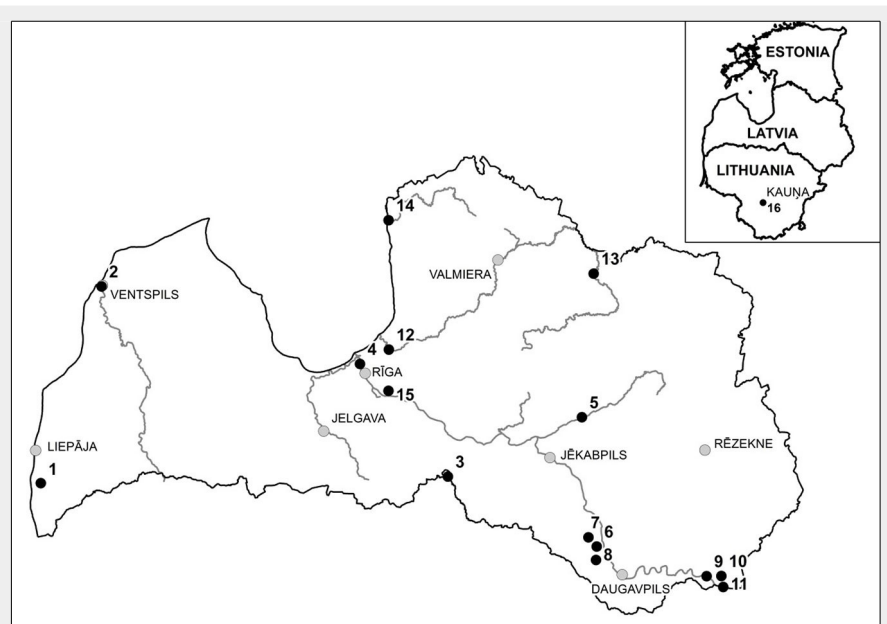


Fig. 3 - Location of sampled stands (black circles) in Latvia and Lithuania. Population numbers are the same as in Tab. 1 (see below).

S. alba and *S. fragilis* (King et al. 2010), however this has not been demonstrated for the other SSR markers utilised in this study. The chloroplast marker data was combined into haplotypes. The pairwise Nei's genetic distances between populations and varieties were calculated from the nuclear SSR data (encoded as binary data), and used to construct a neighbour-joining dendrogram. Identification of vegetatively propagated clones was done by simple matching of nuclear SSR fragments. Principal coordinates analysis (PCoA) was done based on the

pairwise Nei's genetic distance matrix. Genetic analyses and Mantel's tests were done using the software package GenAlEx ver. 6.5 (Peakall & Smouse 2012) and Darwin (Perrier & Jacquemoud-Collet 2006). The robustness of the dendrograms was examined by bootstrapping analysis (1000 bootstraps). Polymorphism information content (PIC) values were calculated as $PIC = 1 - (p^2 + q^2)$, where p is the fragment frequency and q is the no-fragment frequency (Nei 1973). Dendrograms were constructed using the software MEGA 5.2 (Tamura et al.

Tab. 1 - Locations and description of sampled stands in Latvia and Lithuania.

Pop. no.	Location	River catchment	Coordinates	Species/Cultivar	Sample size
1	Nica	Barta	56° 21'; 21° 04'	<i>S. alba</i>	24
2	Ventspils	Venta	57° 23'; 21° 34'	<i>S. alba</i>	26
3	Vecmemele	Memele/Lielupe	56° 24'; 24° 54'	<i>S. alba</i>	23
4	Voleri/Riga	Daugava	57° 00'; 27° 04'	<i>S. alba</i>	24
5	Laudona	Aiviekste/Daugava	56° 42'; 26° 11'	<i>S. alba</i>	24
6	Dviete	Dviete/Daugava	56° 03'; 26° 16'	<i>S. alba</i> × <i>S. fragilis</i>	15
7	Bebrene	Dviete/Daugava	56° 04'; 26° 12'	<i>S. alba</i>	7
8	Ilukste	Daugava	55° 57'; 26° 16'	<i>S. fragilis</i>	24
9	Lielindrica	Daugava	55° 51'; 27° 20'	<i>S. fragilis</i>	24
10	Lielindrica	Daugava	55° 50'; 27° 19'	<i>S. fragilis</i>	24
11	Piedruja	Daugava	55° 47'; 27° 26'	<i>S. fragilis</i>	24
12	Adaži	Gauja	57° 04'; 24° 21'	<i>S. alba</i>	18
13	Vireši	Gauja	57° 27'; 26° 20'	<i>S. alba</i>	27
14	Salaca	Salaca	57° 45'; 24° 21'	<i>S. alba</i>	21
15	Salaspils/ National Botanic Garden	Daugava	56° 51'; 24° 21'	'Chermesina' 'Britzenis' 'Calva' 'Coccinea' 'Vitellina' 'Sericea' <i>S. alba</i> <i>Salix × rubens</i>	1 1 1 1 1 1 1 2
16	Kaunas	Nemuna	54° 54'; 23° 53'	<i>S. alba</i>	17

Tab. 2 - Parameters of genetic diversity for the analysed populations.

Population	N	No. of fragments	No. of fragments (freq. \geq 5%)	Mean PIC	No. of unique fragments
Nica (<i>S. alba</i>)	24	54	46	0.231	2
Ventspils (<i>S. alba</i>)	26	25	21	0.093	0
Vecmemele (<i>S. alba</i>)	23	40	31	0.146	0
Voleri/Riga (<i>S. alba</i>)	24	38	35	0.163	0
Laudona (<i>S. alba</i>)	24	37	29	0.127	1
Dviete (<i>S. alba</i> \times <i>S. fragilis</i>)	15	28	28	0.104	0
Bebrene (<i>S. alba</i>)	7	27	27	0.114	1
Ilukste (<i>S. fragilis</i>)	24	30	20	0.089	0
Lielindrica (<i>S. alba</i>)	24	35	26	0.115	1
Lielindrica (<i>S. fragilis</i>)	24	29	28	0.130	0
Piedruja (<i>S. fragilis</i>)	24	27	22	0.095	0
Adaži (<i>S. alba</i>)	18	38	38	0.156	0
Vireši (<i>S. alba</i>)	27	35	32	0.135	0
Salaca (<i>S. alba</i>)	21	33	24	0.124	1
Kaunas (<i>S. alba</i>)	17	42	42	0.178	1
Mean	21.786	34.533	29.071	0.130	-

2011) and FigTree v 1.4.2 (<http://tree.bio.ed.ac.uk/software/figtree/>).

Results

Of the six analysed chloroplast markers, only two were polymorphic, which detected two polymorphic alleles each (CCMP3: 113bp and 114bp; CCMP10: 110bp and 111bp). These were combined into haplotypes, and only two haplotypes were identified within the analysed samples (designated as HapA and HapB). HapA was the most commonly identified haplotype, with 236 individuals (71%), while HapB was found in 95 individuals (29%). Most of the analysed stands contained individuals of only one haplotype, with the exception of the Vireši stand (HapA frequency 0.81) and the Vecmemele

stand (HapA frequency 0.96). HapB was found in all individuals collected from *S. fragilis* and hybrid stands (Dviete, Piedruja, Lielindrica – *S. fragilis*, Ilukste). In addition, two individuals with HapB were identified from samples collected in the National Botanic Garden ('Vitellina', *S. x rubens*).

Three of the analysed SSR markers amplified a maximum of two alleles per individual (SB38, SB194 and SB201), two of which (SB38 and SB194) were previously reported to have disomic-monogenic inheritance in tetraploid *S. alba* and *S. fragilis* (King et al. 2010). The remaining markers amplified more than two alleles in some of the analysed individuals, therefore the genotypes were analysed as dominant binary markers. The eight SSR markers identified a total of

75 fragments. Of these, 33 (44%) had a minor allele frequency of less than 0.05. The number of fragments detected in each population ranged from 25 (Ventspils) to 54 (Nica). The average number of fragments detected over all the analysed populations was 34.5. The average PIC values for the loci within each population ranged from 0.088 (Ilukste) to 0.231 (Nica). One unique allele was found in each of five populations (Laudona, Bebrene, Lielindrica – *S. alba*, Salaca, Kaunas), and two unique alleles were identified in the Nica population (Tab. 2). The Ventspils population was the least genetically diverse, and 14 of the 26 analysed individuals belonged to just three distinct clones. Evidence of vegetative propagation to a lesser extent was also found in Vireši (2 individuals, 1 clone), Ilukste (5 individuals, 2 clones) and Dviete (4 individuals, 2 clones).

The calculated Nei's genetic distance based on nuclear SSR marker genotypes, ranged from 0.006 (Piedruja-Ilukste) to 0.113 (Bebrene-Ilukste). The average pairwise Nei's genetic distance was 0.051. Dendrogram analysis grouped the *S. fragilis* and hybrid populations into one cluster, and the *S. alba* populations into another (Fig. 4). However, only a few clusters had over 50% bootstrap support, and there was also no evidence of isolation by distance, with a Mantel test indicating no significant correlation between genetic distance and geographic distance, both for all populations and for the *S. alba* populations only. AMOVA results revealed some differentiation between the analysed populations, with 37% of the total genetic variation found between all populations (PhiPT = 0.371, $p < 0.001$, 999 permutations). The separate analysis of *S. alba* populations

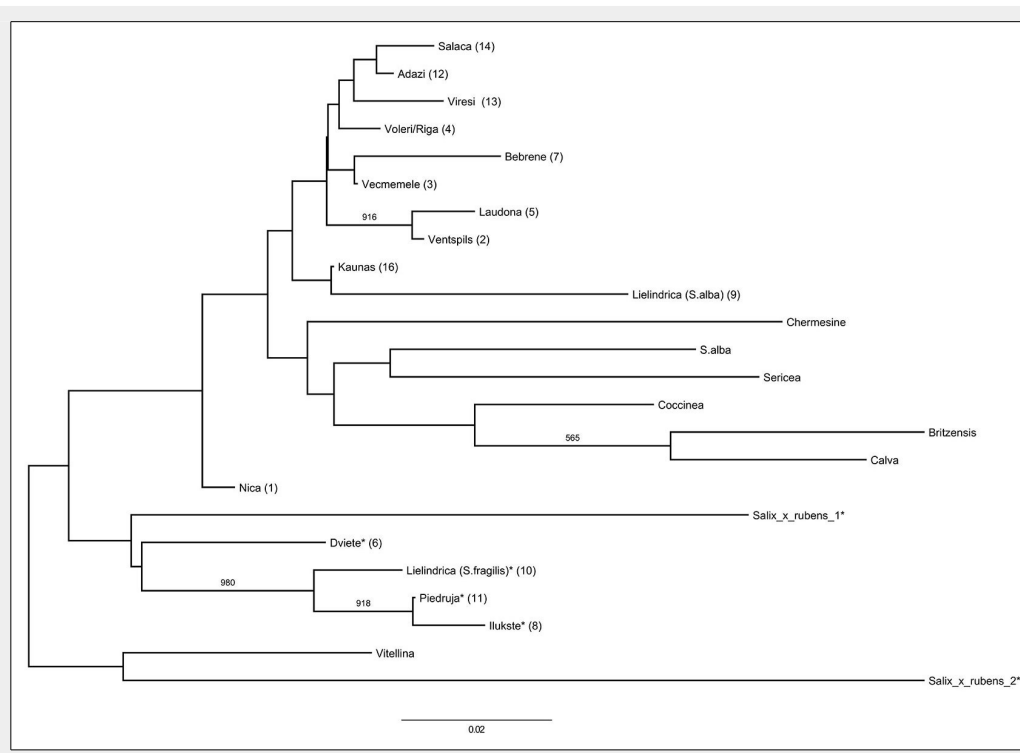


Fig. 4 - Neighbour-joining dendrogram of Nei's genetic distances between populations and varieties. (*): *S. fragilis* populations and hybrid varieties/individuals; (NBD): samples collected from the National Botanic garden. Numbers in brackets refer to populations labels reported in Tab. 1. Numbers on dendrogram branches indicate those having more than 50% bootstrap support (out of 1000 replications).

revealed that between-population variability accounted for 25% of the total genetic variation ($\Phi_{PT}=0.254$, $p<0.001$, 999 permutations). The populations in the *S. alba* cluster show some geographical grouping, with the Salaca and Vireši populations found in northern Vidzeme, the Adaži and Voleri populations in the vicinity of Riga, and the Bebrene and Vecmemele populations found in the eastern region of Latvia (Fig. 3). One exception was the close genetic relationship of the Ventspils and Laudona population. These populations are located over 300 km distant from each other, but were among the most closely related pair of populations analysed, with a pairwise Nei's genetic distance of 0.010. The Nica population was at an intermediate position between the two clusters.

Principal coordinate analysis (PCoA) using the Nei's genetic distance matrix calculated from nuclear marker data showed that the analysed individuals from the *S. fragilis* populations and putative hybrid individuals were mostly distinguished from the *S. alba* populations (Fig. 5). The putative hybrid individuals were intermediate between the two pure species clusters, but were closer to *S. fragilis*, corresponding to the neighbour-joining dendrogram. Twelve individuals from the Nica population were clustered with the *S. alba* individuals, while 12 individuals were clustered with the other *S. fragilis*/hybrid individuals according to the nuclear SSR marker data (Fig. 5), which is also reflected in the intermediate position of the Nica population in the neighbour-joining dendrogram. In addition, six individuals from the Vireši population, one from the Lielindrica (*S. alba*) and three individuals from the Vecmemele population were clustered with the *S. fragilis*/hybrid individuals. Of these 21 individuals (morphologically identified as *S. alba*, though clustering with *S. fragilis*/hybrid individuals based on nuclear SSR data), six had the chloroplast haplotype B, characteristic of *S. fragilis* (5 individuals from Vireši and one from Vecmemele).

The individuals labelled as either HapA or HapB, as determined by the chloroplast markers, were clustered by PCoA using the nuclear SSR marker data (Fig. 6). The majority of the individuals distinctly cluster into two groups, reflecting their designation as *S. alba* or *S. fragilis*/hybrids. The majority of the hybrid individuals (identified phenotypically or by nuclear SSR marker data) as well as the *S. fragilis* populations were designated as HapB. However, there were 16 individuals which were grouped with the *S. fragilis*/hybrids individual that were designated as HapA. The majority of these (12 of the 16 individuals) were from the Nica population, three were from Vecmemele, one from Lielindrica (*S. alba*) population, and one sample from the botanic garden (*S. × rubens*). In addition, two individuals designated as HapB were clearly grouped with the *S. alba* individuals according to the nuclear SSR marker data

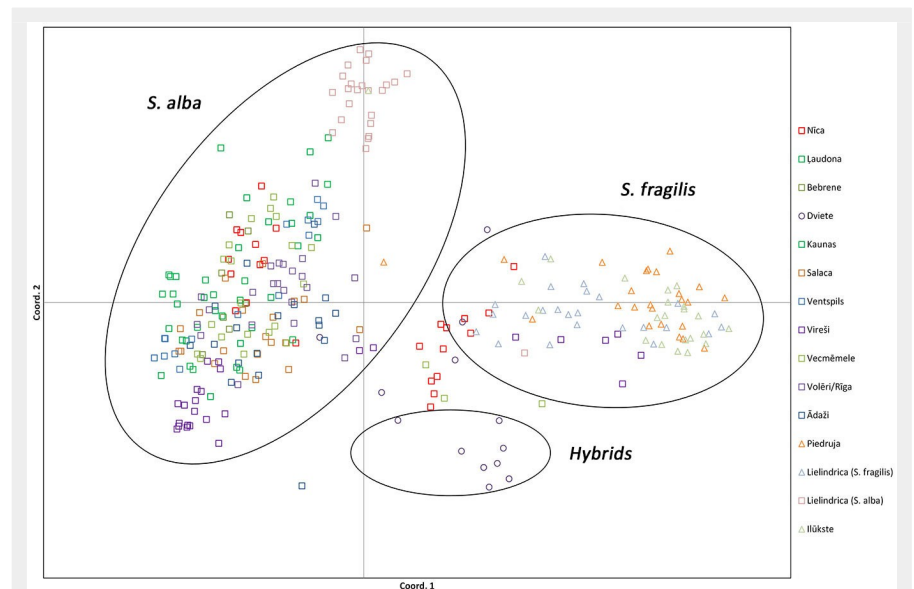


Fig. 5 - Principal Coordinates Analysis of nuclear SSR genotype data. Individuals are labelled according to the population phenotype listed in Tab. 1 (*S. alba*, *S. fragilis* or hybrids).

(one individual from each of the populations Ilukste and Piedruja).

The individuals collected from the National Botanic Garden were split between the *S. alba* and *S. fragilis*/hybrid clusters. Most samples were clustered with the *S. alba* populations, however, the two individuals identified as *Salix × rubens* Shrank (*S. alba* × *S. fragilis*) clustered with the *S. fragilis*/hybrids, as well as the variety 'Vitellina'.

Discussion

The nuclear SSR data did not identify any highly differentiated *S. alba* populations in Latvia. The levels of diversity (number of fragments identified and average PIC) within each population were similar, even in the two populations with a smaller number of sampled individuals (Bebrene: 7 individuals; and Ventspils: 26 individuals), but

with 14 analysed individuals belonging to just three clones, i.e., 15 genetically distinct individuals. This indicates a high level of genetic diversity within and between individuals in a single population, as expected from a dioecious species with wind dispersed seeds (Triest et al. 2000).

In previous studies of *S. alba* and *S. fragilis*, a higher differentiation of these two species was reported by using DNA markers as compared to morphological traits (Triest 2001). It has been suggested that this might be a consequence of the allotetraploid nature of the genomes of *S. alba* and *S. fragilis*, with the two species sharing one common or two highly related diploid ancestor species, but differentiated from each other by the genomic contributions of divergent second diploid ancestor species (Barcaccia et al. 2014). The allotetraploid

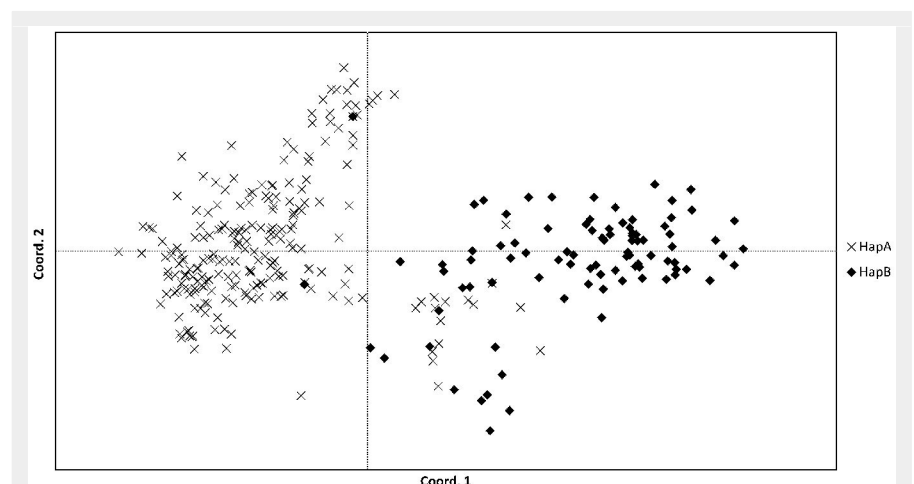


Fig. 6 - Principal Coordinates Analysis of nuclear SSR genotype data. Individuals are labelled according to chloroplast haplotypes HapA (predominantly *S. alba*) or HapB (predominantly *S. fragilis*/hybrids).

genomes of both species, may suggest disomic segregation of genetic loci. However, the SSR markers utilised in this study were developed from *Salix burjatica*, and have been shown to amplify in a wide range of other *Salix* species (Barker et al. 2003), thus explaining why more than two alleles were amplified at some of the analysed SSR loci.

Only two chloroplast haplotypes were identified within the analysed individuals, and the same two haplotypes were also identified in the cultivated clones and varieties collected from the National Botanic Garden, which are presumably of different provenance (not from the Baltic region). Analysis with additional chloroplast markers, which reveal further polymorphism/haplotypes, may allow a more fine-scale elucidation of the provenance of the Baltic *S. alba* populations. Analysis of additional reference material sourced from non-Baltic countries will allow a better understanding of the extent of chloroplast polymorphism revealed by the utilised markers and the levels of diversity within *S. alba* and *S. fragilis* at the European scale. The chloroplast haplotypes were able to differentiate the majority of *S. alba* (HapA) and *S. fragilis* (HapB) individuals. The majority of putative hybrid individuals had the *S. fragilis* haplotype (HapB), suggesting that the in general hybridisation events occur by fertilisation of *S. fragilis* mothers by *S. alba* pollen; however, there may be a lower frequency of reciprocal fertilisation of *S. alba* by *S. fragilis* pollen. Alternatively, there may be a low frequency of shared chloroplast haplotypes between *S. alba* and *S. fragilis* due to natural hybridisation events. A mixture of haplotypes was found only in two populations, Vireši and Vecmemele. The majority of individuals in these populations with the *S. fragilis* haplotype (HapB) also clustered with the *S. fragilis*/putative hybrid individuals according to the nuclear SSR marker data, while the individuals from these populations that clustered with the *S. alba* individuals based on nuclear SSR data also had the *S. alba* chloroplast haplotype (HapA). This suggests that hybridisation can occur in naturally established and regenerated stands or alternatively, that *S. alba* and *S. fragilis* individuals are growing in mixed stands, and that the morphological parameters that distinguish these species are homogenised by environmental and other growing site characteristics. However, the Nica population was homogeneous for the *S. alba* chloroplast haplotype (HapA), but was split into two clusters according to the nuclear SSR data. This suggests that this population may contain some individuals generated by hybridisation in the opposite direction (i.e., fertilisation of *S. alba* mother trees by *S. fragilis* pollen), as compared to the majority of the hybrids identified in this study (fertilisation of *S. fragilis* mother trees by *S. alba* pollen). This population is located in the south-western region of Latvia, which has a relatively higher density

of *S. fragilis* and also has a less continental climate than eastern Latvian regions. Such factors may influence the hybridisation pattern, including effects on the phenology of these species, for instance alteration of flowering times. These putative hybridisation occurrences should be analysed in more detail at a molecular and phenotypic level by analysing additional putative hybrid individuals, particularly in stands where a mixture of chloroplast haplotypes and morphological characteristics are found.

In general, the nuclear SSR genotypic data, the chloroplast haplotypes and the phenotypic assessments of the analysed populations were well correlated. The individuals that were phenotypically identified as putative hybrids were clustered between the *S. alba* and *S. fragilis* individuals using the nuclear SSR markers, while the majority of these individuals had the chloroplast haplotype B (HapB) predominant in *S. fragilis*. The individuals sampled from the National Botanic Garden were genetically similar to the *S. alba* populations, with the exception of the individuals previously identified as *Salix × rubens* Shrank (hybrid of *S. alba* and *S. fragilis*) as well as the cultivar 'Vitellina'. The two *Salix × rubens* Shrank individuals were designated as having different haplotypes (HapA and HapB), indicating that the hybridisation direction may differ for each individual. Alternatively, the haplotype identified primarily in *S. fragilis* and hybrid individuals (HapB), also found at a lower frequency in *S. alba* populations, could be the result of natural hybridisation, as reported previously for other *Salix* species (Palmé et al. 2003). Previous investigations of the willow complex *S. alba* – *S. × rubens* – *S. fragilis* using DNA markers (De Cock et al. 2003) revealed that *S. × rubens* can be separated into two groups (*S. × rubens* and *S. × rubens* var. *basfordiana*), which clustered with *S. alba* and *S. fragilis*, respectively. A more precise taxonomic classification for the two *S. × rubens* individuals analysed in this study was not available, but the alternating chloroplast haplotypes in these two individuals may also support the results reported by De Cock et al. (2003). The individual identified as 'Vitellina' which was collected from the National Botanic Garden, clustered with the *S. fragilis* and hybrid individuals according to the nuclear SSR data and this individual also had the chloroplast haplotype predominantly found in *S. fragilis* and hybrid individuals (HapB). This was surprising, as 'Britzensis' is reported as a cultivar of the *S. alba* sub-species *vitellina* or variety 'Vitellina'. However, the 'Britzensis' individual analysed was genetically similar to the *S. alba* individuals, and also had the *S. alba* chloroplast haplotype (HapA), in contrast to the 'Vitellina' individual analysed in this study. A more detailed morphological assessment of the sampled 'Vitellina' individual and further genetic analysis of additional 'Vitellina' individuals could

help determine if this particular individual was mislabelled, or if other representatives of the variety 'Vitellina' also are genetically distinct from *S. alba*.

This is the first study which examined the population structure of *S. alba* and its hybrids in Latvia. Analysis of the *S. alba* individuals collected from the National Botanic Garden, which represent a range of cultivated forms and varieties, as well as hybrids, indicated that the individuals designated as *S. alba* sub-species, were genetically similar to the analysed *S. alba* natural populations, and that there is little genetic differentiation between the *S. alba* populations and sub-species, supporting the hypothesis that the classification of cultivated clones or decorative forms as sub-species and varieties is not taxonomically significant. Further analysis of natural *S. alba* populations from beyond the Baltic region will indicate if the level of genetic differentiation in *S. alba* is low at the European scale, as found for other broadleaf tree species, e.g., *Fraxinus excelsior* L. (Heuertz et al. 2004). The analysis of additional nuclear and chloroplast markers may also allow better differentiation of *S. alba* populations.

Webb (1985) proposed a set of criteria for determining the native status of a species: fossil evidence, historical evidence, habitat, geographical distribution, frequency of known naturalization, genetic diversity, reproductive pattern, possible means of introduction. Consideration of these criteria suggests that the majority of the examined *S. alba* populations are probably natural. There are no fossil records available of this species from Latvia, but historical records report that this species has been described as growing in natural stands in the territory of Latvia for over 200 years (Fischer 1778, Klinge 1883). The habitats where the majority of the analysed *S. alba* populations were collected were assessed as natural (with the exception of the Ventšpils population), and this species is distributed throughout the territory of Latvia (Laiviņš et al. 2009 – Fig. 2). The genetic diversity analyses did not identify any highly differentiated populations; however, the discriminative power of the used nuclear and chloroplast markers may not be sufficient to detect possible introductions, as the presumably foreign clones in the National Botanic Garden collection were not differentiated from the analysed populations. In this regard, our results are not conclusive, though it seems unlikely that all the analysed populations are introduced, particularly as Latvia is already on the border of the previously described distribution area for this species. There was only a small number of genetically identical individuals identified, indicating a low level of vegetative reproduction (again with the exception of the Ventšpils population). Although there is a long history of cultivation and planting of *S. alba* in parks and gardens in Latvia (Cinovskis et al. 1974), which

would provide a plausible means of introduction, the majority of the criteria suggest a natural origin of *S. alba* in Latvia. This supports the hypothesis that Latvia is within the natural distribution range of *S. alba*, thus many of the natural stands found in Latvia could be considered to be autochthonous.

Our results provide the basis for further, more detailed investigations of *S. alba* populations in Latvia, allowing the identification of possibly introduced or artificially regenerated stands, and the determination of potential *S. alba* genetic resource stands. Within this study, the Ventspils population, which was suspected to be at least partially artificially established from *in situ* observations, was confirmed to contain a high proportion of vegetatively propagated individuals, which could be indicative of reproduction *via* cuttings. In addition, these results confirm that *S. alba* hybridises with *S. fragilis*, and that hybrid individuals can be found in natural stands. The results of the chloroplast marker analysis indicate that the predominant hybridisation occurs by fertilisation of *S. alba* mother trees by *S. fragilis* pollen, however, the extent of haplotype sharing between these two species should be further investigated. In addition, stands with varying proportions of chloroplast haplotypes were identified in this study, and these stands should be examined in more detail using both molecular and phenotypic methods.

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