

Characterization of newly developed expressed sequence tag-derived microsatellite markers revealed low genetic diversity within and low connectivity between European *Saccharina latissima* populations

Jaromir Guzinski^{1,3} · Stéphane Mauger¹ · J. Mark Cock² · Myriam Valero¹

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Abstract The kelp *Saccharina latissima* is a species of high ecological and economic importance. We developed a novel set of *S. latissima*-specific genetic markers that will find applications in conservation biology, biodiversity assessment, and commercial exploitation of this macroalga. Thirty-two expressed sequence tag (EST)-derived microsatellite markers (SSRs) were developed and characterized in this study using publically available EST sequences. Twenty-seven percent of the 7064 analyzed ESTs contained repeat motifs, and polymerase chain reaction (PCR) amplification primers were designed for 96 selected loci. Fifty-one (53 %) of the primer pairs amplified their target loci, of which 32 (33 %) were polymorphic within a sample of 96 *S. latissima* sporophytes collected from six localities distributed along the European Atlantic coast from Southern Brittany (France) to Spitzbergen (Norway). The 32 loci harbored moderate levels

of polymorphism with 2–13 alleles per locus (mean 5.4). The 25 loci that were retained for population genetic analyses revealed substantial genetic differentiation among the European populations (pairwise F_{ST} values ranging from 0.077 to 0.562) that did not follow any pattern of isolation by distance. In addition, within-population genetic diversity was generally low ($H_s < 0.323$). Two non-mutually exclusive hypotheses were proposed to explain this low diversity pattern: (1) lower variability of the EST-derived microsatellites compared to the random distribution of SSRs developed from genomic DNA since the former are frequently located in coding regions, which are generally less variable, or (2) reduced effective population size of *S. latissima*. The particularly high genetic differentiation between the French and Scandinavian *S. latissima* populations is in agreement with the reported ecotypic differentiation, which may reflect an important resource for genetic improvement. The pattern of genetic diversity revealed in this study thus suggests that care should be taken to avoid the transfer of strains between different geographic regions.

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✉ Jaromir Guzinski
jaromir.guzinski@sb-roscoff.fr

¹ CNRS, UPMC, Sorbonne Universités, PUCCH, UACH, UMI 3614, Evolutionary Biology and Evolution of Algae, Station Biologique de Roscoff, Place Georges Teissier, CS 90074 Roscoff, France

² CNRS, UPMC, Sorbonne Universités, UMR8227, Laboratoire Végétaux Marins et Biomolécules, Équipe Génétique des Algues, Station Biologique de Roscoff, Place Georges Teissier, CS 90074 Roscoff, France

³ CNRS, UPMC, Sorbonne Universités, UMR7144, Adaptation et diversité en Milieu Marin, Équipe Diversité et Connectivité dans le Paysage Marin Côtier, Station Biologique de Roscoff, Place Georges Teissier, CS 90074 Roscoff, France

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Introduction

Saccharina latissima (Linnaeus) C.E. Lane, C. Mayes, Druehl & G.W. Saunders, known by the common names of sea belt or sugar kelp, is a perennial brown macroalga (class Phaeophyceae) of the family Laminariaceae. This species exhibits circumpolar distribution in the northern hemisphere, occurring in polar and temperate coastal waters from intertidal to a lower depth limit of about 30 m (Bolton et al. 1983;

Gerard 1988). *Saccharina latissima* forms continuous dense stands (“kelp forests”) on shallow rocky coasts, occurring in clear and turbid waters, preferentially in areas sheltered from strong wave action (Bekkby and Moy 2011). In Europe, its range extends from Spitsbergen, Norway as far south as Portugal. *Saccharina latissima* requires rocky substrate as an attachment point and does not occur where this surface is replaced by long stretches of sandy coast (Bartsch et al. 2008). Both the wide latitudinal and vertical distribution of this species suggest the occurrence of ecotypic differentiation between populations with respect to light and temperature (Gerard and Du Bois 1988; Müller et al. 2008). However, knowledge about genetic diversity and connectivity is still lacking, and this information will be important not only for elucidating population structure across the range of *S. latissima* but also for improved exploration of genetic resources in the wild.

In general, the dispersal capacities of macroalgae are limited, on the order of few meters up to several kilometers, compared with distances of hundreds of meters to several hundred kilometers for other marine organisms such as invertebrates or fish characterized by a pelagic larval dispersal phase (Kinlan and Gaines 2003). In kelps, dispersal is primarily mediated by short-lived propagules, and the extent of dispersal depends on factors such as the morphology of the blade (i.e., location of the sporophylls on the thallus and whether they possess swimming aids such as air bladders; see Valero et al. 2011 for a review), as well as on the orientation of currents and habitat continuity (Billot et al. 2003; Alberto et al. 2010; Coleman et al. 2011). In a recent meta-analysis of genetic differentiation in macroalgae, Durrant et al. (2014) found that most species exhibited a clear pattern of isolation by distance with high-level genetic differentiation at distances greater than 50–100 km. However, occasional long-distance dispersal, even on the scale of tens to hundreds of kilometers, can occur due to pieces of fertile thalli breaking off and being carried away from the point of origin by currents (Thiel and Gutow 2005; Alberto et al. 2011; Coleman et al. 2011). Additionally, Cie and Edwards (2011) have uncovered a mechanism facilitating vertical transport of kelp zoospores into the shallower portions of the water column, which has been proposed to enable long-distance dispersal of zoospores via current action. Indeed, it has recently been suggested that spore dispersal is not likely to be a limiting factor in maintaining connectivity between Norwegian *S. latissima* populations (Andersen 2013). Therefore, it remains an open question as to whether *S. latissima* populations are genetically distinct (limited dispersal—spores recruiting within their population of origin) or whether there is some degree of genetic homogenization and thus lack of pronounced spatial or geographic structure (long-distance dispersal, even if only occasional, resulting in a certain number of spores recruiting to distant populations). Until now, this question has not been investigated, largely because the necessary genetic tools have been lacking.

As a primary producer, *S. latissima* plays a crucial role in the functioning of the subtidal ecosystem and dense *S. latissima* stands provide a habitat for a plethora of marine organisms (Bekkby and Moy 2011). Additionally, this is a species of high economic importance, with applications in the food, bioenergy, cosmetic, and pharmaceutical industries. *Saccharina latissima* is considered to be a species of choice for the production of brown seaweed biomass in Europe (Peteiro and Freire 2013) because it has a shorter life cycle than those of *Laminaria digitata* and *Laminaria hyperborea*, the other two most abundant kelps along the European coast. Furthermore, *S. latissima* is closely related to *Saccharina japonica*, a domesticated species that occurs along the northwestern coasts of the Pacific Ocean. Indeed, *S. japonica* is one of the most economically important types of seaweed in the seaweed farming industry with a long history of breeding programs and artificial seedling-rearing techniques (Tseng 2001). The complete genome of *S. japonica* has been recently published providing a vital resource for algal genome studies and improvement of traits of economic importance (Ye et al. 2015). For these reasons, *S. latissima* is a good model species for studying algal domestication in Europe, and, as a likely commercially important species, it is about to become a subject of studies aimed at identifying strains exhibiting traits of interest for cultivation.

Given the significant ecological and commercial importance of *S. latissima*, it is of particular concern that over the past 5 to 10 years, this species has experienced large-scale population declines at sites in Norway, where 50 % of Europe’s *S. latissima* forests are found, as well as in Sweden and Germany (Bekkby and Moy 2011). It is therefore urgent to develop a set of species-specific genetic markers that could be employed to elucidate and monitor *S. latissima* population structure and hence provide guidance on how to best implement conservation and management strategies.

Microsatellites, also known as simple sequence repeats (SSRs), are among the most commonly used marker types in evolutionary and ecological studies. SSRs are appropriate for the determination of genetic diversity because they are abundant in the genome, are highly polymorphic, are highly reliable, and usually have a codominant mode of inheritance. The development of microsatellites has traditionally involved a large input of time and labor for the construction, enrichment, and sequencing of genomic libraries (Edwards et al. 1996). In contrast, identification of SSRs from expressed sequences (EST-SSRs) is a fast, efficient, and relatively low-cost alternative (Bouck and Vision 2007). EST-SSRs facilitate physical mapping and tend to be widely transferable between species and even between genera (Liu et al. 2012). Here, we describe the development of a suite of EST-based (Heinrich et al. 2012), *S. latissima*-specific microsatellite markers. Additionally, we test the ability of these newly developed

SSRs to differentiate between individuals from six natural populations of *S. latissima* distributed from Brittany to Norway and investigate the importance of inter-population connectivity. We also discuss the potential uses of these markers for applied research (conservation, strain selection) on this kelp species.

Materials and methods

Sample collection and DNA extraction

Sixteen mature *S. latissima* sporophytes were collected from each of six sampling locations. The six sites, SB—Mousterlin, France, NB—Roscoff, France, PDC—Audresselles, France, DEN—Ebeltoft, Denmark, SWE—Tjarno, Sweden, and NOR—Ny-Alesund, Spitsbergen, Norway, followed a latitudinal gradient (Fig. 1). Upon collection, a disk of tissue (diameter 2 cm) was cut out from a spot that was free of epiphytes and stored with silica gel. If possible, the sample was

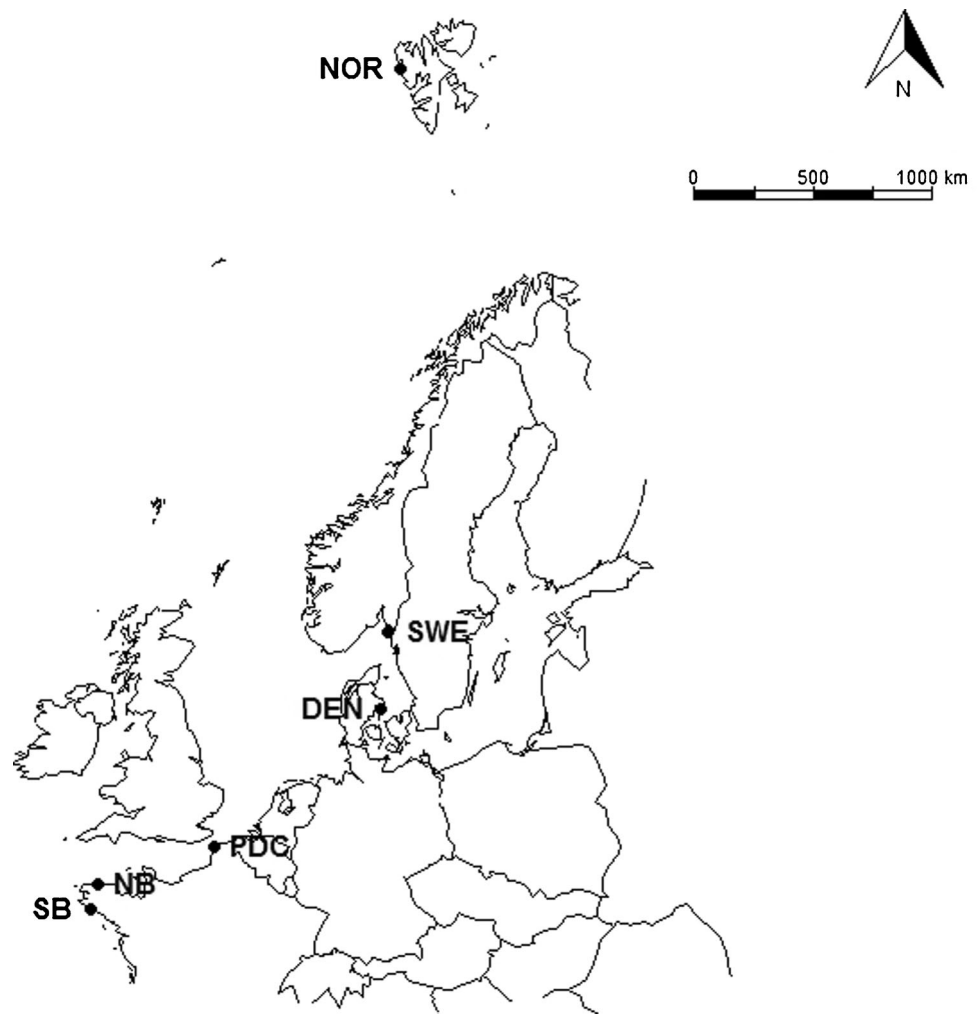
taken from close to the junction of the stipe and the blade as the aim was to sample the youngest tissue.

Total genomic DNA was extracted from 5 to 10 mg of dry tissue using the Nucleospin 96 plant kit (Macherey-Nagel, Germany). The extractions were performed according to the manufacturer's instructions except that samples were left in the lysis buffer at room temperature for 1 h rather than being heated to 65 °C for 30 min. The extracted DNA was eluted in 200 µL of the supplied elution buffer.

EST-SSR screening and primer design

A total of 400,503 *S. latissima* ESTs obtained from the Sequence Read Archive (SRA) database (accession number SRR305166; Heinrich et al. 2012) were screened for di-, tri-, tetra-, and pentanucleotide motif SSRs using the Sputnik program (<http://espressoftware.com/sputnik/>). Seven thousand sixty-four ESTs were found to contain at least one SSR, with 438 (6 %) containing a dinucleotide repeat microsatellite, 2330 (33 %) containing a trinucleotide repeat microsatellite, 1765 (25 %) containing a tetranucleotide repeat microsatellite,

Fig. 1 Map of the six sampled locations: SB - Mousterlin, France (47°50'27.4"N, 4°02'18.1"W), NB - Roscoff, France (48°43'41.5"N, 4°00'20.6"W), PDC - Audresselles, France (50°49'27.8"N, 1°35'40.9"E), DEN - Ebeltoft, Denmark (56°10'04.5"N, 10°43'51.2"E), SWE - Tjarno, Sweden (58°52'31.9"N, 11°06'12.6"E), NOR - Ny-Alesund, Spitsbergen, Norway (78°55'30"N, 11°55'20"E).



and 2531 (36 %) containing a pentanucleotide repeat microsatellite (Table 1). For loci where the flanking sequence on either side of the repeat was of sufficient length, PCR amplification primers were designed in Primer3 (Rozen and Skaletsky 1999). In-house scripts were used to specify the primer selection conditions, which used default values for all parameters except the optimum melting temperature, which was set to 60 °C (range 57–62 °C), and the expected product size, which was set at 350 bp.

This resulted in the design of 1878 primer pairs corresponding to 270 (14.4 %) di-, 849 (45.2 %) tri-, 428 (22.8 %) tetra-, and 331 (17.6 %) pentanucleotide repeat loci (Table 1). In order to cost-effectively test these primers for amplification, and then check the amplified loci for levels of polymorphism within the sampled individuals, we reduced the number of loci used in the trials to 96 (i.e., capacity of a microtiter plate), employing the following criteria (see Table 1 for details). First, we eliminated all loci which contained an unknown base (“N”) within the repeat region. Next, for the di- and trinucleotide motif SSRs, only those loci which comprised at least eight repeat units were retained, and for the tetra- and pentanucleotide motif SSRs, only those loci which comprised at least five repeat units were retained (184 loci remaining—Table 1). Finally, the number of loci was reduced to 96 such that loci with the highest number of repeats were selected for each of the four motif categories.

Amplification trials using the 96 primer pairs

Ninety-six primer pairs were tested for amplification using DNA from six randomly chosen *S. latissima* individuals, one individual from each locality. Amplifications were carried out in 10 µL reaction volumes with each reaction comprising 2 µL of DNA template diluted 1:50, 1× GoTaq Flexi buffer (Promega Corporation), 2 mM MgCl₂, 150 µM of each dNTP

Table 1 Steps involved in the selection of 96 *Saccharina latissima* SSR loci from 7064 initial candidates obtained by analysis of cDNA library sequence data

Reduction step	Di	Tri	Tetra	Penta	Number of retained loci
Sputnik output	6%	33%	25%	36%	7064
Primer3 output	14%	45%	23%	18%	1878
Loci with “N” base removed	14%	45%	23%	18%	1871
Repeat number >8 (di and tri) and >5 (tetra and penta)	20%	30%	32%	19%	184
Final selection	17%	33%	33%	17%	96

The percentage of loci with specific motif types is also presented for each step of the selection process

Di SSR loci with dinucleotide repeat motifs, *Tri* SSR loci with trinucleotide repeat motifs, *Tetra* SSR loci with tetranucleotide repeat motifs, *Penta* to SSR loci with pentanucleotide repeat motifs

(Thermo Fisher Scientific Inc., USA), 30 pmol forward primer, 30 pmol reverse primer, 30 pmol fluorescent-labeled M13 (-21) primer (Eurofins MWG Operon, Germany), and 0.35 U GoTaq Polymerase (Promega). The 5' end of each forward primer included an M13 (-21) universal sequence tag (5'-TGTTAAACGACGGCCAGT-3') to enable the incorporation of the universal fluorescent-labeled M13 (-21) primer for detection of the amplified products on an ABI3130 XL DNA Analyzer (Applied Biosystems, USA). The 96 loci were divided into four randomly selected groups (24 loci per group), and each of the four groups of loci was labeled with a different color. Amplifications were carried out in a BioRad DNA Engine Peltier Thermal Cycler under the following conditions: initial denaturation at 95 °C for 5 min, 12 cycles of denaturation at 95 °C for 30 s, annealing at 65 °C for 30 s (-1 °C per cycle - touchdown to 53 °C), and extension at 72 °C for 30 s, and 30 cycles of denaturation at 95 °C for 30 s, annealing at 53 °C (optimum M13 (-21) annealing temperature) for 30 s, extension at 72 °C for 30 s, followed by a final extension of 72 °C for 10 min.

The PCR products labeled with the four different colors were pooled (24 pools in total) and diluted 1:10 with water. Next, 2 µL of the diluted PCR product pool was added to 10 µL of loading buffer made up of 0.5 µL of the SM594 size standard (Mauger et al. 2012) and 9.5 µL of Hi-Di formamide, denatured at 95 °C for 3 min, and run in an ABI 3130 XL capillary sequencer (Applied Biosystems, USA). Genotypes were scored manually in Genemapper version 4.0 (Applied Biosystems).

Forty-five of 96 primer pairs did not amplify a product from any of the six individuals despite several re-amplifications and were not further analyzed.

Investigating levels of polymorphism for 51 EST-SSR loci

The 51 loci retained after the amplification tests were subsequently checked for levels of polymorphism within the sample of 96 *S. latissima* individuals. Amplifications were carried out in the same manner as described for the amplification trials.

Within the sample of 96 *S. latissima* individuals, 36 of the 51 loci were polymorphic. However, two of the polymorphic loci failed to amplify in many of the individuals originating from NB, PDC, DEN, and SWE (Sacl-70 amplified in 76 out of 96 samples, and Sacl-77 in 72 samples). As re-amplifications did not resolve this issue, these two loci were not retained for further analyses (see Supplementary Table 1 for the primer sequences and repeat motif details for these two loci, and Supplementary Table 2 for information on the 15 monomorphic loci). Two pairs of linked loci were detected among the remaining 34 loci (see “Statistical analyses” section) and, after removal of one locus from each pair, the remaining 32 loci were used for population genetics analyses.

Multiplex development

In order to optimize the usage of the 32 loci in future population genetics studies, we used Multiplex Manager v 1.2 (Holleley and Geerts 2009) with default parameters to organize the amplification of the loci in as few multiplex reactions as possible. Three multiplexes were created: multiplexes 1 and 2 comprised 10 loci each, and multiplex 3 comprised 12 loci (Table 2). For each multiplex, two disparate sets of loci were amplified separately to minimize primer dimer formation, but the two sets were pooled before genotyping. For multiplex 1, these were set M1i and set M1ii, whereas for multiplex 2, these were set M2i and set M2ii, and for multiplex 3, these were set M3i and set M3ii (Table 2). Each of the six multiplex sets was amplified in the same manner as described for the amplification trials, except that we added 40 pmol of each forward fluorescently labeled primer (Eurofins MWG Operon, Ebersberg, Germany), 40 pmol of each reverse primer, and 0.5 U GoTaq Polymerase. Amplifications were carried out in a BioRad DNA Engine Peltier Thermal Cycler under the following conditions: initial denaturation at 95 °C for 5 min, 10 cycles of denaturation at 95 °C for 30 s, annealing at 65 °C for 30 s (-1 °C per cycle - touchdown to 55 °C), and extension at 72 °C for 30 s, and 30 cycles of denaturation at 95 °C for 30 s, annealing at 55 °C for 30 s, extension at 72 °C for 30 s, followed by a final extension of 72 °C for 10 min.

Statistical analyses

We used Genepop v 4.3 (Rousset 2008) to test for linkage disequilibrium between all pairs of the 34 loci within each of the six sampling sites as well as for all localities combined (global test employing Fisher's method). This analysis used the following Markov chain parameters: dememorization, 100,000; batches, 1000; and iterations per batch, 50,000. All significance levels were adjusted from an alpha value of 0.01 using sequential Bonferroni correction to allow for multiple tests on the same dataset (Rice 1989). We omitted locus Sacl-14 from further analyses since it exhibited significant linkage disequilibrium with locus Sacl-66 in three localities (NB, SB, DEN) and across all populations. We also omitted locus Sacl-72 since its allele patterns were identical to those of locus Sacl-08 (i.e., Sacl-72 alleles were always one base larger than Sacl-08 alleles) for all individuals with the exception of a genotype for a single NOR individual.

For each of the remaining 32 loci, we calculated the following statistics using the combined sample of all 96 amplified individuals with the indicated R (R Development Core Team 2015) package. Number of alleles (N_A) and the polymorphic information content (PIC) were estimated in *PopGenKit* (Paquette 2012), whereas *hierfstat* (Goudet 2014) was used to obtain the observed heterozygosity (H_O) and the overall gene diversity (H_T) (Nei 1987). All

significance levels were adjusted using sequential Bonferroni corrections (Rice 1989). Next, we computed for each of the 32 loci within each locality, allelic richness (Ar), H_O , and within-population gene diversity (H_S) (Nei 1987) using *hierfstat*, with the per locus “All Pops” Ar and H_S obtained by averaging these two parameters over the six localities. F_{IS} values for each locus within each locality were calculated in *Demerelate* (Kraemer and Gerlach 2013). Significance for each estimate was obtained by generating 10,000 bootstraps. The “All Pops” F_{IS} and F_{ST} parameters were calculated in *hierfstat*.

The frequencies of null alleles of the 32 loci were estimated using the Brookfield 1 equation (Brookfield 1996). The equation was run for each of the genotyped loci, separately for each of the six sampling locations, using MicroChecker 2.2.3 (Van Oosterhout et al. 2004). The confidence interval for the Monte Carlo simulations of the homozygote frequencies was set to 95 %.

Subsequent analyses were performed using a reduced set of 25 loci (after removing loci with null alleles and/or exhibiting odd allelic patterns, see “Results” section and Supplementary Material 1). For each of the six localities, we estimated the mean Ar and mean H_S by averaging over the 25 locus-specific values presented in Supplementary Table 3 (for underlined loci only). The H_S means were compared between the six populations using the Kruskal-Wallis test. F_{IS} for each of the six populations was estimated as the multilocus estimate of Wright's inbreeding coefficient using *diveRsity* (Keenan et al. 2013). Significance was tested with 10,000 bootstrap iterations, and the values were considered to be significant if the bias-corrected 95 % confidence intervals did not overlap zero. The global (i.e., over all loci and all populations) values for H_S , F_{IS} , and F_{ST} were estimated by calculating the average of the over-population means (obtained from *hierfstat*) for each of these parameters over the 25 loci.

To estimate the level of genetic differentiation between the six sampling localities, pair-wise F_{ST} (Weir and Cockerham 1984) values were computed in *hierfstat*. The significance of the comparisons was estimated by performing 10,000 bootstraps over loci, with the comparisons judged significant if the bootstrap generated confidence intervals did not overlap zero.

To gain further insights into *S. latissima* population subdivisions across our sampling range, and in particular to determine if the six clusters defined based on the sampling localities reflect the biological reality, we used the Discriminant Analysis of Principal Components (DAPC) method implemented in the *adegenet* R package (Jombart 2008; Jombart et al. 2010). This multivariate statistical approach seeks to provide an efficient description of genetic substructure through the use of a limited number of synthetic variables called the discriminant functions. DAPC infers the number of clusters of genetically related individuals (i.e., the number of distinct populations) by partitioning the original variables (allele frequencies) into within-group and

Table 2 Technical information (primer sequence, dye, and multiplex), characteristics (repeat motif, expected product size, and observed size range), and polymorphism estimates (N_A , H_O , H_T , PIC) for 32 EST-SSR loci based on amplification from genomic DNA of 96 *Saccharina latissima* sporophytes

Locus	Primer sequence (5'–3')	Repeat motif in the EST	Dye	Contig number on GenBank BioProject PRJNA80101	Expected product size (bp)	Observed size range (bp)	N_A	H_O	H_T	PIC	Multiplex
Sacl-03	F: GTTGTACGTTGGCGTTG R: TTCAATAATCGCAGGAAGCAC	(CA) ₉	PET	10167	212	204–214	6	0.479	0.679	0.625	M1i
Sacl-06	F: TTTGTAGATAGTCCGCTGGT R: GTAGCCTGCCGAAGAAATAAA	(AG) ₁₅	PET	13119	158	141–165	9	0.271	0.565	0.519	M1ii
Sacl-08	F: CAGACCTCACCAAGGCAAAG R: TCCACGCACATACAGCAAA	(TA) ₁₁	PET	14056	130	118–144	4	0.073	0.071	0.069	M3ii
Sacl-09	F: ACCGTTCCCAATAACTACCC R: ATGGACCTATCGCCACA	(AGC) ₁₃	PET	7028	332	320–332	5	0.531	0.425	0.385	M1i
Sacl-11	F: AAGAGGTGGTTGCTCGGTT R: CACTGATGGATGCCCTGTC	(CTG) ₁₂	FAM	8341	302	294–306	5	0.281	0.356	0.322	M1i
Sacl-13	F: CTCAGGTGTTCCGGTGCCT R: GCTCGGTAGATGGTTAGTC	(GCA) ₉	FAM	13741	268	235–244	3	0.083	0.147	0.139	M3i
Sacl-19	F: GCGTATTCCTAAACACCTCCC R: CGATGACTGCCACCACAG	(ACC) ₉	PET	2896	175	163–184	5	0.156	0.533	0.428	M2i
Sacl-21	F: TCTCAACTCCAACAGAGCGT R: AGCAGCAGAAGCAGGCGA	(GCT) ₉	FAM	17486	184	151–166	5	0.177	0.203	0.194	M3i
Sacl-24	F: GCCAACCTATCATTCAAAGCA R: TGTGGGAGCAACATCCTCA	(TGA) ₁₁	PET	16267	103	87–132	13	0.375	0.515	0.474	M2i
Sacl-25	F: TTCGTTTCAGTTGGTGGGTT R: TGCTGTAGTAGTATTCTTCGCC	(TTGA) ₂₃	NED	7352	339	323–343	4	0.156	0.472	0.373	M1i
Sacl-27	F: GCGTTTGGTTAGCAGTGTGA R: GTGCCCTCCCTACTCCCGT	(AT) ₈	NED	17361	314	276–312	5	0.000	0.082	0.081	M3i
Sacl-32	F: CTCGTTGTCTGCTTGCTC R: CTTTCGCCGTCACCTACTACA	(GCTT) ₅	PET	2655	218	192–200	3	0.031	0.031	0.031	M3i
Sacl-33	F: TTTCGCTCTGTCTCTCCT R: AAATCACAACACAAGGCTGCT	(CTTT) ₆	FAM	20594	204	182–186	2	0.042	0.118	0.110	M3ii
Sacl-37	F: AACCGCTCTTGTGTTGATG R: CTCCTTCCCTCCCTCTCC	(GGGA) ₆	NED	12415	149	146–154	3	0.292	0.311	0.271	M1i
Sacl-38	F: TACGATTGCGTGGTTGT R: ACGAAGAGATTGCGACAAA	(TA) ₇	NED	15061	136	122–152	8	0.635	0.846	0.823	M2i
Sacl-41	F: GTGGCGTTAGATGCTGTATGG R: CGTGGACAAAAGTAGGAAAGGG	(GCTTA) ₁₁	VIC	136	340	262–352	6	0.104	0.139	0.134	M3ii
Sacl-47	F: CGACACAATCGCAGTCAG R: GCCCGACACACTCAAGAC	(CTTGA) ₆	FAM	11611	165	154–174	3	0.365	0.411	0.353	M1i
Sacl-49	F: GCTCTCCACCTCGACTAA R: TACCCTCAGCCTCCAGAG	(AT) ₁₃	NED	7113	333	315–335	11	0.875	0.796	0.765	M2ii
Sacl-54	F: CGACGCTGACCTGACACA R: CCTTCCAATCTCTTCCCTTCCA	(GA) ₈	NED	16526	225	221–227	4	0.323	0.625	0.552	M1ii
Sacl-56	F: GGAGAGCGTCGGATAGACC R: GAGAATAGCACAGCAGCGAAC	(AT) ₁₀	VIC	1453	132	112–118	2	0.021	0.041	0.040	M3i
Sacl-60	F: GTCCAGGTGCGTGGTTAG R: GCGAGCAGTTGAAAGGTGG	(TGC) ₈	VIC	7664	200	179–182	2	0.010	0.010	0.010	M3ii
Sacl-65	F: ATCTCCAAACCACACAAG R: CATCATCGTCAAGAACTCGG	(CCA) ₁₀	PET	6095	344	339–369	6	0.115	0.643	0.580	M2i
Sacl-66	F: TATGTATGTCGGGAGACGGG R: GGGATTAGCAACTGAAACCA	(TTG) ₉	PET	20551	264	202–238	12	0.521	0.835	0.813	M2ii
Sacl-68	F: GGTGGATTCTTTGGACGA R: AAATGTGCTTGGGTCGGG	(GGT) ₈	FAM	19139	162	155–158	2	0.073	0.501	0.375	M2i
Sacl-74	F: CCTAAAGTTCTACCTGGGCAA R: TCACAAGGACCACATTCCAAC	(CCTT) ₆	PET	19201	279	256–284	7	0.260	0.291	0.279	M1i
Sacl-75	F: CTCGTGTCGCCCTTCATC R: CTGTCTCCAGAACTCGCC	(TC) ₅	NED	13067	256	236–238	2	0.010	0.010	0.010	M3ii
Sacl-78	F: GTTGTCTGCTGCTTCTAATCGG R: GTCCATTTCTTGCTGCTG	(GCTT) ₅	FAM	17312	291	265–289	3	0.042	0.041	0.041	M3i
Sacl-81	F: ACTTTGGCTCGGTCTGCTT R: CCTCTCCCTTACCTACCTCC	(CG) ₆	FAM	9492	329	321–337	6	0.469	0.694	0.642	M2ii
Sacl-88	F: GAAACGGTGAAGTACTGATGAC R: ACAAGGACGAACAGAGAACGA	(AC) ₅	PET	1052	189	171–175	3	0.021	0.062	0.059	M3ii

Table 2 (continued)

Locus	Primer sequence (5′–3′)	Repeat motif in the EST	Dye	Contig number on GenBank BioProject PRJNA80101	Expected product size (bp)	Observed size range (bp)	N_A	H_O	H_T	PIC	Multiplex
Sacl-90	F: ATTGTGTTGCTGGATGAGGAC R: ACCTTCCGCTCTCTCGCT	(GCAA) ₉	PET	4328	292	274–314	8	0.240	0.577	0.532	M2ii
Sacl-94	F: TGCCAAATAACACATTCCAGAG R: TGGTGCGAAGTCACGAAATAG	(CGGTA) ₅	VIC	11651	256	243–298	12	0.573	0.779	0.758	M1ii
Sacl-95	F: GGGAAGGAGGAAGAAGGTG R: TAGCGGAAAGAACGGGTAGT	(GTTGC) ₆	VIC	19139	201	188–203	4	0.479	0.505	0.400	M2i

Primer sequences exclude the M13(–21) universal sequence tag. 100% amplification was achieved for all loci

F forward primer (labeled with fluorescent dye), *R* reverse primer, *Expected product size* size in base pairs (bp) of the expressed sequence tag (EST) that possessed the microsatellite locus, N_A number of alleles, H_O observed heterozygosity, H_T overall gene diversity, PIC polymorphic information content

between-group components such that between-group variance is maximized. We used the six predefined clusters as the input for DAPC as the F_{ST} analysis showed all pairwise comparisons to be significant (see “Results” section).

Finally, we tested for isolation by distance (IBD) between the six *S. latissima* populations. The geographic distance was the “as crow flies” direct distance in kilometers between each of the localities, whereas the genetic distance was Rousset’s $F_{ST}/(1 - F_{ST})$ estimate for each pair of populations (Rousset 1997). The significance of the IBD relationship was tested statistically with a Mantel test in R package *vegan* (Oksanen et al. 2015) with 999 permutations.

Results

Of the 96 primer pairs selected for the amplification trials, amplification was successful for 51 loci (53 %) (Supplementary Fig. 1). Of these, two loci (Sacl-70 and Sacl-77) were removed due to inconsistent amplification in a larger sample of individuals, and two were dropped because they were found to be in linkage with other loci (Sacl-08 and Sacl-14). Of the 47 remaining EST-SSR loci, 32 loci (68 %) were polymorphic within the analyzed sample of 96 *S. latissima* sporophytes originating from six disparate geographic locations (Fig. 1). These 32 loci displayed variable levels of polymorphism, with PIC ranging from 0.010 to 0.823 with a mean of 0.350, and 11 (34 %) of the loci displaying values above 0.450. The number of alleles per locus ranged from 2 to 13 while the average over 32 loci was 5.41 alleles per locus (Table 2). Observed heterozygosity H_O was the highest at 0.875 for locus Sacl-49, whereas locus Sacl-27 was homozygous within all the scored individuals (Table 2). Mean H_O was 0.253 and the average overall gene diversity (H_T) was 0.385. H_O was lower than H_T at 24 of the 32 loci, probably

because of a Wahlund effect due to pooling of individuals from multiple genetic clusters, but other contributing causes (inbreeding or the presence of null alleles) might also be involved. To minimize the possibility of the Wahlund effect, the per-locus statistics were estimated within each of the six geographic localities (Supplementary Table 3).

Based on these data, seven loci were considered unusable for population genetics analyses within the context of our dataset and were thus removed from all subsequent analyses (details are given in Supplementary Material 1 and in Supplementary Table 3). Briefly, these seven loci (Sacl-06, Sacl-09, Sacl-27, Sacl-47, Sacl-49, Sacl-88, Sacl-90) were removed because they exhibited significant null allele frequency at multiple geographic localities, displayed low levels of polymorphism, or they exhibited odd allelic patterns with for instance fixed heterozygosities in some populations (highly negative F_{IS} ; Supplementary Table 3). When significant null allele frequency was detected for highly polymorphic loci, but only for a single population, such loci were retained for subsequent analyses (i.e., Sacl-24, Sacl-54, or Sacl-65; Supplementary Table 3). Note that although the seven loci that were removed were deemed unsuitable for our analyses, they could be of use for other types of studies or studies on other populations of *S. latissima*, thus they are also reported in this manuscript and were optimized as part of the multiplexes specified in Table 2.

Of the 25 loci retained, 18 were monomorphic within at least one of the six localities (Supplementary Table 3). Thus, the within-population observed allelic richness averaged over the 25 loci was low, ranging from 2.080 in SWE to 2.840 in SB (Fig. 2, Supplementary Table 3). The highest observed heterozygosity value was 0.938 for locus Sacl-38 within the NOR samples, while the lowest value (0.063) was observed within NB for five loci and within DEN for three loci

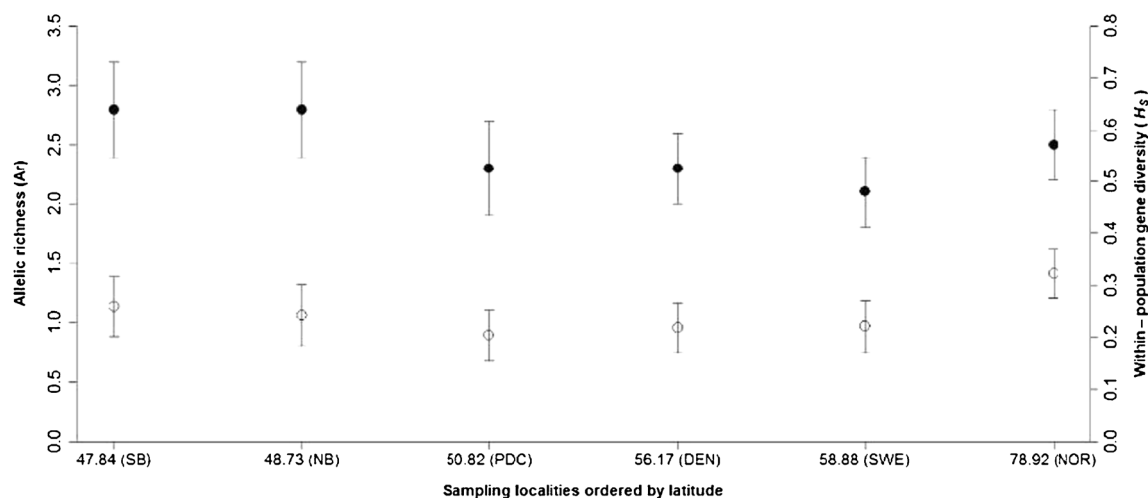


Fig. 2 Allelic richness (Ar) (filled symbol – left hand side y-axis) and within-population gene diversity (H_s) (open symbol – right hand side y-axis) averaged over 25 EST-SSR loci for each of the six sampling

localities (16 individuals per population) ordered by latitude. Error bars indicate standard error

(Supplementary Table 3). Similarly, the locus-specific, within-population gene diversity (H_s) values were generally low. The multilocus within-population H_s means ranged from 0.204 for PDC to 0.323 for NOR (Fig. 2, Supplementary Table 3). The H_s means were not significantly different between the six geographic localities (Kruskal-Wallis test $\chi^2(5)=4.549$, p value=0.473) and thus did not vary with latitude. This was also the case for the Ar means (Fig. 2). The within-population multilocus F_{IS} values were not significantly different from zero for any of the six geographic localities, which indicates that random mating commonly occurs within the studied populations.

Despite the generally low level of within-population polymorphisms displayed by the 25 loci (grand H_s mean of 0.245—Supplementary Table 3), there were 47 private alleles at 23 of 25 loci. The pattern of genetic differentiation among populations was highly significant with an overall F_{ST} mean of 0.259 and 10 loci exhibiting F_{ST} values above 0.300 (Supplementary Table 3).

All of the pairwise multilocus F_{ST} comparisons were significant. The least differentiated populations were SB and NB, with F_{ST} of 0.077 (Table 3). These populations exhibited approximately equal levels of divergence from PDC, with the geographically more distant SB being also slightly more distant genetically. The F_{ST} between DEN and SWE was the second lowest at 0.120, with both these populations being highly genetically differentiated from NOR (F_{ST} above 0.250 in both comparisons) (Table 3). The highest levels of differentiation were between the three north European populations and the three French populations, with F_{ST} ranging from 0.358 to 0.562 (Table 3). The IBD model was rejected (Mantel statistic $r=0.275$, significance=0.112) (Fig. 3) since the more geographically distant NOR population was less differentiated

from the three French localities than the less distant DEN and SWE populations.

The DAPC results correlate well with the F_{ST} estimates (Fig. 4). Axis 1 (which explains 45.12 % of the variance) separates the three French sampling localities from the three Scandinavian sampling sites (Fig. 4a, b), while axis 2 (which explains 10.83 % of the variance) separates NOR from the other five sampling localities (Fig. 4a) and axis 3 (which explains 6.03 % of the variance) partitions the French sampling site into a PDC and an SB/NB cluster (Fig. 4b). Indeed, the two pairs of localities displaying the lowest pairwise F_{ST} values (SB and NB, and DEN and SWE) were the least well defined by DAPC. There was no clear separation of these two pairs of sites within any of the first three principal components of the DAPC, as there was always some overlap between individuals from these two localities (Fig. 4a, b). The admixture plot (Fig. 4c), however, revealed that most of the SB/NB individuals were assigned to their population of origin, with only 9 out of 32 individuals originating from the SB/NB localities exhibiting some level of admixture. The level of

Table 3 Pairwise F_{ST} estimates for the six sampling localities

	SB	NB	PDC	DEN	SWE	NOR
SB	0.000					
NB	0.077	0.000				
PDC	0.179	0.157	0.000			
DEN	0.481	0.517	0.562	0.000		
SWE	0.472	0.509	0.559	0.120	0.000	
NOR	0.358	0.390	0.431	0.291	0.268	0.000

All comparisons are significant

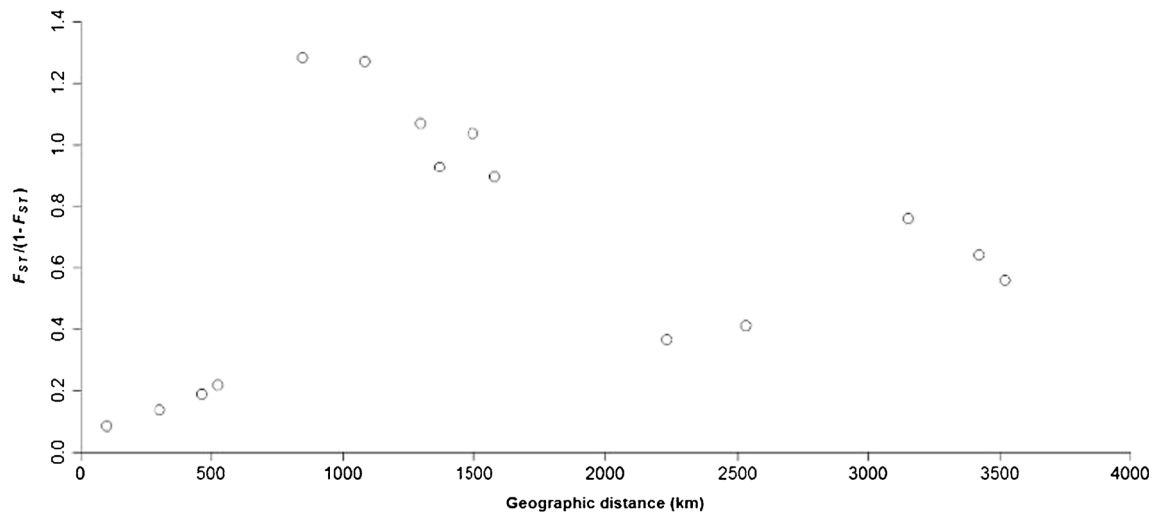


Fig. 3 Genetic estimates of pairwise differentiation ($F_{ST}/(1-F_{ST})$) plotted against geographic distance (as the crow flies) for the six sampling localities

admixture between PDC and the SB/NB localities was even lower (Fig. 4c). The admixture situation within the Scandinavia sampling localities mirrored that within the French sites, although the DEN/SWE admixture was less substantial in comparison to the SB/NB admixture (Fig. 4c). Finally, there was no evidence of admixture between the two sampling regions (i.e., France and Scandinavia; Fig. 4c).

Discussion

Of the 96 primer pairs that were tested for amplification with six *S. latissima* sporophytes, 51 (53 %) were deemed successful (though subsequently two of these loci were dropped due to inconsistency of amplification within a larger sample of individuals) (Supplementary Fig. 1). Our markers therefore exhibited a higher positive amplification rate than was reported by Wang et al. (2011) (36.5 %) for *Laminaria*-specific EST-SSRs. For the loci presented herein, the success of amplification was poorest for loci comprising tetranucleotide motifs (31.3 %) and best for those comprising dinucleotide motifs (81.3 %), whereas it was just under 60 % for loci comprising the trinucleotide and pentanucleotide motif types (Supplementary Fig. 1).

Utilization of expressed sequence tags to identify so-called genic SSRs is a relatively recent approach, but one that has been successfully employed in a vast array of taxa (Ellis and Burke 2007). The main advantage over the traditional method of isolation of microsatellite markers (in this case termed genomic SSRs) is that obtaining SSRs from ESTs is considerably less labor intensive and also cheaper (Zane et al. 2002; Ellis and Burke 2007). In this study, there was a high attrition rate among the newly developed markers, with only around 50 % of the amplification trials being successful, 33 % of

markers being polymorphic, and just over a quarter of the markers (25 out of 96—26 %) being retained for the population genetics analyses. However, the time and effort spent on acquiring these markers was not excessive, and a reasonably high number of markers were obtained. The traditional method of isolating SSRs, partly due to the abovementioned difficulties, rarely results in more than 20 markers being produced. Indeed, 16 genomic SSRs have recently been developed for *S. latissima* (Møller Nielsen M, Paulino C, Neiva J, Krause-Jensen D, Bruhn A, Serrão EA: Genetic variability of *Saccharina latissima* along a salinity boundary in the North Sea-Baltic transition, submitted). Note that as 184 loci were retained after the reduction step “Repeat no. > 8 (di & tri) & > 5 (tetra & penta)” (Table 1) and only 96 of these have been tested for amplification and polymorphism so far, there is potential to develop additional *S. latissima*-specific EST-SSRs from this analysis if needed.

A potential disadvantage of EST-SSRs as genetic markers is that they are commonly considered to harbor lower levels of polymorphism than genomic SSRs because they are derived from expressed, and thus conserved, genes (Gupta et al. 2003; Ellis and Burke 2007; Guichoux et al. 2011). However, various studies have demonstrated that the EST-derived SSRs were sufficiently diverse to be useful markers for population genetics analyses (Dong et al. 2009; Cubry et al. 2014; Teshome et al. 2015), and that conclusions regarding population structure and connectivity were highly congruent whether elucidated with genomic or EST-derived SSRs (Woodhead et al. 2005; Kim et al. 2008; Hu et al. 2011).

The number of alleles per locus we detected in our study (5.41 alleles per locus averaged over 32 loci) was higher than or comparable to the genomic or EST-SSR sets described in the literature. For example, in a closely related congener species *S. japonica*, the average number of alleles over 18 genomic SSR loci was 4.7 alleles per locus (Shi et al. 2007),

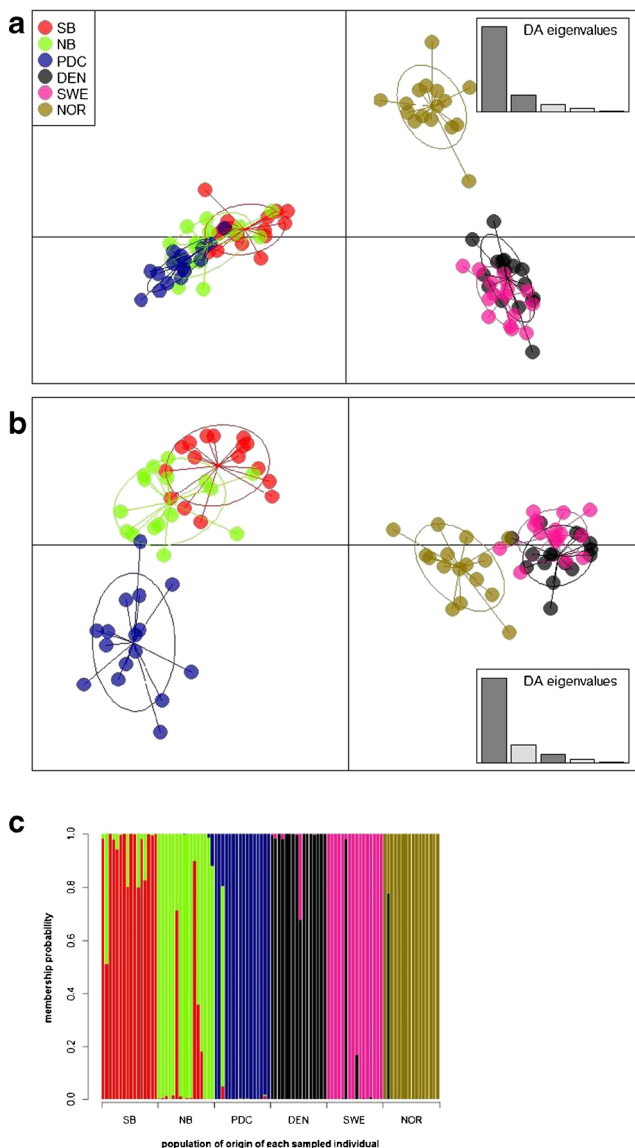


Fig. 4 Scatterplot of the **a**) first two components (axes 1 and 2), and **b**) of the first and third components (axes 1 and 3), of the discriminant analysis of principal components (DAPC) performed on 96 *S. latissima* sporophytes amplified at 25 microsatellite loci, using sampling location as prior clusters. Sampling groups are shown by different colours and inertia ellipses, and each point corresponds to a single individual. **c**) Membership probability of each *S. latissima* individual to each of the six sampling localities (differentiated by colour). Individuals are represented by vertical bars. Each individual's sampling locality (i.e. population of origin) is specified along the x-axis

whereas on average 5.7 alleles per locus were obtained for nine EST-SSR loci (Liu et al. 2010).

To demonstrate the applicability of this newly developed set of *S. latissima* EST-SSR markers for effective management of this ecologically and economically important species, we investigated the patterns of within-population genetic diversity and connectivity between six localities spanning a latitudinal gradient from South Brittany, France to Spitsbergen, Norway using the reduced panel of 25 loci.

Our data indicate that the six studied populations were significantly genetically differentiated (significant F_{ST} ; Table 3), and formed genetically well-defined clusters even if there was some degree of connectivity between the populations that were closest geographically (Fig. 4). The clustering of the two Brittany populations (SB and NB) into two distinct but not completely isolated groups fits well with what was reported for two other kelps, *L. digitata* and *L. hyperborea* (Couceiro et al. 2013; Robuchon et al. 2014) in the same region. The Ushant tidal front, which constitutes a seasonal dispersal barrier, is probably one of the major factors that influence the observed pattern of kelp population structure in this region as discussed in Couceiro et al. (2013).

Further north along the French coast, the PDC sampled sporophytes exhibited substantially lower connectivity to the two Brittany localities than was detected within Brittany (i.e., between SB and NB) (Table 3, Fig. 4). The pattern of isolation between the Breton sites and PDC is likely explained by habitat discontinuity as the shore line on either side of the PDC sampling site is characterized by long stretches of sandy substrate. Lack of suitable (i.e., rocky) substrate has been proposed to be a major dispersal barrier for several other kelps (Fraser et al. 2010; Alberto et al. 2011; Couceiro et al. 2013; Robuchon et al. 2014). However, the status of PDC as a small, fragmented, and/or isolated population cannot be fully established from our data given that the expected decrease in the within-population gene diversity (H_S) and allelic richness (Ar) compared to the supposedly larger and better connected populations (i.e., SB and NB) was not significant (Fig. 2). Consequently, the genetic isolation of PDC from the Breton samples could simply be due to a combination of the distance between the two sites and the limited dispersal capabilities of *S. latissima* and not necessarily be due to habitat discontinuity. On the other hand, as the H_S and Ar estimates for the two Brittany sampling sites were particularly low, i.e., at least twice as low as those described for *L. digitata* (Billot et al. 2003; Couceiro et al. 2013; Robuchon et al. 2014) and *L. hyperborea* (Robuchon et al. 2014) sampled from nearby points along the Brittany coast, this could suggest that *S. latissima* populations are characterized by smaller population size in comparison to those of other kelps. Indeed, along the Brittany coast, *S. latissima* populations generally exhibited more patchy distribution when compared to the dense and continuous forests of the other two kelp species (Valero, personal observation). Genotyping of samples collected from additional localities should provide more information about *S. latissima* population structure along the coast of France, and in particular of Brittany.

Within Scandinavia, our data revealed moderate levels of connectivity between the *S. latissima* populations at the SWE and DEN sampling localities, and clear delimitation of the NOR population from those of the two former sampling sites. The observed admixture was less prevalent than between SB and

NB (Fig. 4). Overall, this intermediate degree of connectivity between the DEN and SWE populations suggests here also a role for currents in modulating *S. latissima* population structure in this section of the North Sea (the Skagerrak strait). There was almost no admixture between SWE/DEN and NOR (Fig. 4c). This near absence of connectivity between the NOR and the DEN/SWE sampling localities is in accordance with expectations, given that these sites are separated by more than 2000 km. Overall, these analyses validated the applicability of the 25 microsatellite loci in elucidating *S. latissima* population structure.

Phylogeographic studies have demonstrated the effects of glacial-interglacial cycles during the Quaternary (~2.6 Myr to present-times) on the patterns of contemporary genetic diversity, revealing several genetically rich refugia zones and a general trend of decrease in diversity with latitude (Hewitt 2004). In the northern Atlantic, several of these putative long-term climatic refugia have been identified for several marine species including seaweed. These refugia comprise Brittany/South Ireland (corresponding to the palaeo-Celtic Sea/Channel), northwestern Iberia, and for a smaller set of species, central-south Iberia and northwest Africa (Provan 2013; Neiva et al. 2016 for a recent review of North Atlantic Fucaceae). However, in the present study, we did not detect a latitudinal gradient in genetic diversity as none of the localities displayed significantly higher genetic diversity than others (Fig. 2) even if populations from Brittany, located in a putative refugia, were included in our data set. Erosion of genetic diversity may have occurred in this region due to low effective population size because Brittany was at the northern range limit of the species distribution during the colder periods of the Last Glacial Maximum. Reduced genetic diversity was observed for the kelp *Saccorhiza polyschides* in that region and was explained by models predicting marginal, although persistent, populations in the Brittany/Charentes regions (Assis et al. 2016). In addition, note that the *S. latissima* range extends as far south as Portugal (Bartsch et al. 2008), another predicted refugia of the northern Atlantic, and thus more genetically rich refugial populations might be found on the Iberian Peninsula. The three Scandinavian sampling localities all exhibited very low within-population gene diversity and allelic richness, similar to those of the French localities. This was a somewhat surprising result as the lowest genetic diversity might have been expected for the NOR population since it was sampled at the northern limit of the *S. latissima* distribution (Bartsch et al. 2008). This population is expected to show the same pattern of reduced diversity as has been reported for most North Atlantic Fucaceae (Neiva et al. 2016), which is the result of gene surfing at the leading edge of expansion during recolonization after the last glaciation period.

Relatively low within-population genetic diversity (measured in terms of A_r and H_S) was detected within each of the six *S. latissima* sampling localities as these values were twofold lower than the 4.24 alleles per locus and expected

heterozygosity of 0.550 averaged over 16 genomic SSRs analyzed in *S. latissima* populations originating from the North Sea-Baltic Sea transition zone (Møller Nielsen M, Paulino C, Neiva J, Krause-Jensen D, Bruhn A, Serrão EA: Genetic variability of *Saccharina latissima* along a salinity boundary in the North Sea-Baltic transition, submitted), and twofold lower than the values observed within the Brittany populations of *L. digitata* and *L. hyperborea* (Billot et al. 2003; Couceiro et al. 2013; Robuchon et al. 2014). As discussed above, the relatively low within-population genetic diversity observed along the French coast is most probably explained by the small effective population size since populations are small and patchily distributed in this region. In contrast, the low H_S estimates detected within the DEN and SWE sampling localities are counterintuitive as these sites are expected to form part of a large interconnected population(s) as they are located within the Skagerrak strait and the Norwegian side of the strait is known to harbor some of the largest kelp forests anywhere in Europe (Bekkby and Moy 2011). However, it has been observed that over the last 5 to 10 years, extensive areas of the *S. latissima* forests have all but disappeared from along the coast of Norway (including in the Skagerrak strait), as well as from the coastlines of Germany, Denmark, and Sweden (Pehlke and Bartsch 2008; Andersen et al. 2011; Bekkby and Moy 2011; Andersen 2013). Thus, it is conceivable that the fragmentation of populations, driven by anthropogenically induced kelp forest demise (Wahl et al. 2015), has occurred at such speed and to such an extent as to have resulted in a number of small, isolated patches that have retained only some fraction of their genetic diversity. However, we cannot exclude the hypothesis that the low within-population variability of the EST-derived microsatellites that was detected in our study is due to the fact that a proportion of the EST-SSR markers are located in coding regions and are thus generally less variable compared to randomly distributed genomic SSRs (see below).

The multilocus F_{IS} was not significant within any of the six localities, ranging from -0.066 in SB to 0.095 in NB (Supplementary Table 3). Therefore, it is not possible to conclude that inbreeding was occurring within any of the sampled localities. Moreover, we did not detect any identical multilocus genotypes indicating that vegetative propagation (clonality) was not predominant within the studied localities.

Comparison of the *S. latissima* genetic diversity at the European scale revealed very high levels of differentiation and a total absence of admixture between the three French and the three Scandinavian localities (Table 3, Fig. 4). The pairwise F_{ST} estimates are particularly high, resembling the between- rather than within-species estimates, and they are likely explained by the large geographic scale concerned in addition to the low dispersal capabilities of this seaweed species. However, there could also be other underlying factors, as discussed below.

Although SSRs are generally considered neutral genetic markers, EST-derived microsatellites are more likely to be involved in regulatory processes and to influence the differential expression of genes than microsatellites from other parts of the genome (Li et al. 2004). Thus, the substantial genetic differentiation within *S. latissima* that was detected at the European level may not reflect only neutral genetic diversity but also adaptive genetic diversity (Holderegger et al. 2006). In other words, if some of the newly developed loci are important for the regulation of genes vital for adaptation to different environmental regimes, then these loci would be under diversifying selection and hence become more differentiated between the disparate sampling localities than neutral SSR loci. Several studies have provided evidence of ecotypic differentiation between *S. latissima* populations from temperate (North Sea, Helgoland) or Arctic (Spitsbergen, Svalbard) regimes (Müller et al. 2008; Olischläger et al. 2014) with strong indications that differential expression of genes plays a major role in the adaption of the distinct *S. latissima* ecotypes to the local conditions (Heinrich et al. 2012). Taking together (1) the fact that our samples originated from across a latitudinal gradient and from populations that have been exposed to different environmental conditions, (2) the evidence for local adaptation within *S. latissima* populations, and (3) the potential role of EST-derived SSRs in regulating gene expression, it is conceivable that the high level of inter-population genetic differentiation detected in this study could be influenced by these factors rather than simply reflect the degree of inter-population connectivity. This scenario could also explain in part the low overall (i.e., over all loci) within-population heterozygosity, as it is conceivable that selection has acted to conserve the number of repeats for at least some of the loci within the localities. Indeed, a recent study that used 13 *S. latissima*-specific genomic SSRs to genotype 225 *S. latissima* sporophytes collected from one Swedish, one Norwegian, and six Danish sites uncovered within-population gene diversity to range from 0.451 to 0.613 (Møller Nielsen M, Paulino C, Neiva J, Krause-Jensen D, Bruhn A, Serrão EA: Genetic variability of *Saccharina latissima* along a salinity boundary in the North Sea-Baltic transition, submitted), thus two to three times higher than what we have detected with the EST-SSRs. Therefore, the fact that the SSRs presented herein are EST-derived should be taken into account when interpreting our findings. A comparative study of both sets of markers on a larger sample size would allow outlier loci to be detected. In addition, the genomic location of our EST-derived SSRs could be investigated in order to find out which genes they are associated with and thus evaluate whether they potentially have a role in the adaptivity of the disparate *S. latissima* populations to the local conditions.

The six *S. latissima* sampling localities did not follow the IBD pattern, with no significant positive correlation detected between the genetic and geographic distance measurements

(Fig. 3). This is a surprising result in light of a recent outcome of a meta-analysis, which found IBD for all of the macroalgae species analyzed (Durrant et al. 2014). The macroalgae species included in the study by Durrant et al. (2014) originated from multiple taxonomic groups, exhibited various life histories, and inhabited varied environments. In the present study, the IBD signal may be obscured due to the DEN/SWE samples being more genetically differentiated from the Brittany localities than the levels of genetic differentiation between NOR and the Brittany localities (Table 3), even though the geographic distance between DEN/SWE and Brittany is much shorter than the geographic distance between the NOR locality and Brittany (Fig. 1). Moreover, for all the Scandinavian versus French comparisons, the furthest locality (SB) was the least differentiated genetically whereas the closest locality (PDC) was the most diverse (Table 3). We have already noted that habitat discontinuity probably isolates the PDC population from other localities, resulting in increased pairwise F_{ST} values that are greater than expected given the geographic distance. However, the reasons underlying the patterns involving the two Breton and three Scandinavian populations are open to speculation. Historical events could also have shaped the current population substructure, in particular vicariance due to the environmental and climatic changes that occurred during the Last Glacial Maximum. As the spatial arrangement of the habitat patches will influence the IBD patterns (van Strien et al. 2015), another reason for the lack of detection of the IBD signal could be that the relative distances between the sampled populations are poorly replicated (for instance small distance between the two Brittany populations) and thus there is an uneven spread of the data points. Moreover, we cannot exclude the hypothesis that recent human impact may have modified *S. latissima* population structure, in particular in the Scandinavian region (see references above), or that the effect of selection on EST-derived microsatellites prevents the detection of IBD signal (see above).

Based on the results we present here, we are confident that this newly developed set of *S. latissima*-specific SSRs will prove to be a very useful genetic tool for this commercially important kelp species. These markers will find applicability in elucidating the population structure and dispersal patterns of this species, information that will be crucial for making informed and effective conservation and population management decisions (i.e., avoiding risk of translocation by taking care to avoid the transfer of strains between different geographic regions). Moreover, this set of markers could be used to identify strains/genotypes of interest (for instance strains displaying commercially important traits), to measure the environmental impact of cultivation and harvesting of wild *S. latissima* populations on biodiversity (Klinger 2015), and to construct genetic linkage maps for this species.

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