RESEARCH ARTICLE



Genetic heterogeneity of two bioeconomically important kelp species along the Norwegian coast

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Abstract

Knowledge of genetic diversity among wild populations is becoming increasingly important as more species are recognized for their bioeconomic value. Industrialization of natural resources, such as kelp in the marine shallow sublittoral zone through cultivation and wild-harvesting, may lead to extensive translocation and local population decimation. Without adequate resilience in the form of genetic diversity within and across populations and given the potential introduction of deleterious alleles from translocations, such anthropogenically pressured populations may not be able to sufficiently respond to future climate and other stressors. Here we provide an assessment of the genetic heterogeneity of two bioeconomically important kelp species, Laminaria hyperborea and Saccharina latissima, across the Norwegian coastal region from South (57°N) to North (78°N), by applying microsatellite genotyping. Isolation by distance was found for both kelp species when comparing genetic distance to geographic distance. L. hyperborea clustered into four distinct genetic groups corresponding to distinct geographical ecoregions, whereas S. latissima did not show equally strong geographical structuring but separated into three geographical clusters along the Norwegian coast. No genetic differentiation was found within the Norwegian Skagerrak region, corroborating previous findings. The identified genetic clustering of both kelp species supports the retention of established management regions along the Norwegian coast and argues for the continuation of a regional focused management plan for kelp resources. Further, the results demonstrate that care should be taken to prevent translocation of kelp between ecoregions in the ongoing industrialization of kelp cultivation, to maintain a healthy coastal ecosystem and sound natural population genetic diversity.

Keywords Genetic differentiation · Norway · Geographical heterogeneity · Laminariales · Laminaria hyperborea · Saccharina latissima

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Introduction

Anthropogenic pressure on coastal zones has contributed to dramatic habitat loss of submerged aquatic macrophytes on a global scale (Waycott et al. 2009; Krumhansl et al. 2016; Filbee-Dexter and Wernberg 2018). The loss of 'foundation species' (corals, kelp, seagrass, etc.) is especially problematic due to their key role in ecosystem functioning, threatening abundance and biodiversity of associated species (Kelp: Krumhansl et al. 2016; Filbee-Dexter and Wernberg 2018; Seagrass: Orth et al. 2006; Waycott et al. 2009; Coral: Pandolfi et al. 2003). Among these foundation species, kelp forests are highly productive marine coastal ecosystems creating three-dimensional forest-like habitats for multitudes of species, including juvenile fish important to commercial fisheries (Norderhaug et al. 2005; Christie et al. 2009). These kelp species have an emerging role in the bioeconomy, as they are being harvested and cultivated for their alginates and attractive nutritional content (Vásquez 2009; Kerrison et al. 2016). This industrialization of kelp in Europe has led to increased harvesting of wild populations and in cultivation of selected species along the coasts of Ireland, France, and Norway for production of a number of consumer goods (Draget et al. 2005; Broch et al. 2013; Kerrison et al. 2016).

Along the Norwegian coast, natural kelp forests cover more than 8000 km² (Gundersen et al. 2011), dominated by the species Laminaria hyperborea (Gunnerus) Foslie, and Saccharina latissima (Linnaeus) C.E. Lane, C. Mayes, Druehl, and G.W. Saunders. Since the 1970s, the Norwegian kelp forests have suffered large-scale population decline, likely due to increased sea urchin population size and failed recovery (Sivertsen 1997; Norderhaug and Christie 2009). However, this trend has partially reversed during the last 10 years, as sea urchin abundance and recruitment decreased as a consequence of increasing water temperatures, facilitating kelp forests' recolonization and regrowth in Mid-Norway (Norderhaug and Christie 2009; Fagerli et al. 2013; Rinde et al. 2014). The fast-growing S. latissima has particularly been shown to efficiently recolonize barren areas, in contrast to the slower-growing L. hyperborea (Leinaas and Christie 1996), potentially influencing the distribution of genetic diversity within and between these kelp species. Since the early 2000s, S. latissima kelp forests in Norway and globally have experienced degradation and potentially also a decline in genetic variation seemingly due to overgrowth of fine filamentous algae (turf algae), as reviewed by Filbee-Dexter and Wernberg (2018).

The changes in kelp forest distribution come at a time when science is just beginning to understand the population genetic dynamics of kelp forests (Nielsen et al. 2016; Wenberg et al. 2018; Luttikhuizen et al. 2018). Marine coastal ecosystems are generally assumed to be structured following isolation by distance (IBD), with increasing genetic differentiation between sites as a function of distance (Wright 1943; Guo 2012). However, this is not always found in nature due to potential long-range dispersal and the overall stochastic nature of coastal marine currents (Siegel et al. 2008; White et al. 2010). At a global scale, genetic patterns of kelp are structured by morphology (Valero et al. 2011), ocean currents (Billot et al. 1998; Tellier et al. 2009), distance (Alberto et al. 2010; Robuchon et al. 2014; Luttikhuizen et al. 2018) and occasional floating rafts (Fraser et al. 2010; Neiva et al. 2012). Moreover, in the northern hemisphere, diversity is expected to be highest at low latitudes as a result of glacial refugia in southern regions (Hewitt 2000; Maneiro et al. 2011; Neiva et al. 2012), whereas leading edge populations are expected to have less genetic diversity, as a consequence of founder effect (Hampe and Petit 2005).

Whereas several studies have investigated population genetic patterns of the smaller brown seaweeds along the Norwegian coastline (Hoarau et al. 2007; Olsen et al. 2010; Coyer et al. 2011), only a few have studied the population genetics of large kelp species in this geographic area (Guzinski et al. 2016; Nielsen et al. 2016; Luttikhuizen et al. 2018).

Kelp populations with sufficient genetic variation are considered more resilient to climatic stress compared to populations with low genetic variation (Wernberg et al. 2018). Identifying and mapping local as well as regional genetic variation is therefore of great importance to generate baseline information, which will enable efficient monitoring and sustainable use of wild kelp populations. This becomes increasingly important due to commercial interests in wild species, resulting in potential extensive translocations of organisms, and with that, the introduction of deleterious alleles hampering local adaptation. Along the Norwegian coast, extensive translocations of organisms are occurring for example as a biological measure to remove salmon lice from farmed Atlantic salmon (Salmo salar) by introducing fishes from the family of wrasses (Labridae) (Skiftesvik et al. 2014; Halvorsen et al. 2017a, b) and the lumpfish (Cyclopterus lumpus) (Powell et al. 2018) into affected areas. Translocations of organisms have proven to result in introgression of foreign genotypes into resident local wild populations (Glover et al. 2012; Jansson et al. 2017; Faust et al. 2018), which becomes even more problematic with increasing levels of genetic differentiation between the source and the resident population, possibly disrupting local adaptation if selection is not sufficiently strong to maintain locally beneficial alleles (Haldane 1930). Therefore, both the assessment of genetic diversity on a local and regional population level and the corresponding levels of genetic differentiation are needed before such translocations should occur for instance related to wide-spread industrial-scaled farming of kelp species.

This study provides documentation of genetic heterogeneity among populations and across ecoregions of the two most dominant and commercially important kelp species along the Norwegian coast, i.e. S. latissima and L. hyperborea. The study covers the entire Norwegian coast, from southern Norway to Svalbard, and encompasses six ecoregions based on climatic conditions and biogeographic patterns, with the aim to advise an ecosystem-based management of marine resources. Results are discussed in the context of current management plans and commercial exploitation of the species, and to help management preserve genetic diversity among Norwegian kelp populations, thereby securing ecological/genetic resilience against future climatic and anthropogenic pressures. The assessment of the level of genetic heterogeneity among Norwegian kelp together with the regional genetic diversity estimates provides a baseline for further studies on the genetic makeup of changing kelp populations. The results will assist the implementation of both a genetic database and a management tool for the safekeeping of healthy and sustainable kelp communities, both wild and farmed populations.

Materials and methods

Sample collection and preparation

As part of a national environmental monitoring program, a total of 106 *S. latissima* and 98. *L. hyperborea* were sampled across 16 locations in five of the six ecoregions along the Norwegian coast, including Svalbard (Table 1; Fig. 1). Emphasis was on the most densely populated regions, thus giving a good spatial representation of the Norwegian kelp forest. No samples were collected from the 'Norwegian Sea North' region as kelp forest is very sparsely present in this region due to over grazing by green sea urchins (Norderhaug and Christi 2009). As the samples for this study were collected alongside a monitoring program prioritizing a geographically wide sample collection over intensive local sample collection, sample sizes for some of the locations did not conform to recommendations for coverage of allele

frequencies within a population (Hale et al. 2012; Fung and Keenan 2014).

Tissue samples of individual sporophytes of *S. latissima* and *L. hyperborea* collected from 4 to 23 individuals per location were preserved and stored in silica gel at room temperature or stored in ethanol and freeze-dried prior to extraction. Samples used to initially test microsatellites were extracted with the DNeasy Plant Mini Kit (Qiagen, Hilden, Germany) with modifications from Snirc et al. (2010). Genomic DNA from all other samples was extracted from 2 to 10 mg of dried tissue with the cetyltrimethyl ammonium bromide (CTAB) protocol developed for plants (Murray and Thompson 1980), with modifications for brown algae (Hoarau et al. 2007; Coyer et al. 2009), and eluted in 100 µl AE buffer (Qiagen).

Microsatellite genotyping

Genotyping was done for eight and nine microsatellite markers, for *L. hyperborea* and *S. latissima* respectively (Table 2). Markers were selected from Robuchon et al. (2014) and Guzinski et al. (2016), in addition to four markers originally developed for other closely related species (CS34, CS12, CS13: Wang et al. 2011, SSR 261: Zhang et al. 2015).

 Table 1
 Summary of samples per species

Ecoregion	ID	Sample position	#ind	Diversity		Heterozygosity	
				N _{A_Pop}	N _R	Не	Но
Laminaria hyperborea							
Skagerrak	1	58°52′22.2"N 9°36′35.6"E	10 (1.9%)	12	1.29	0.098 ± 0.060	0.103 ± 0.073
	2	58°30′47.5"N 8°56′39.5"E	23 (0.7%)	13	1.31	0.112 ± 0.062	0.093 ± 0.050
	3	58°16′23.5″N 8°32′13.9″E	9 (4.2%)	12	1.35	0.116 ± 0.067	0.102 ± 0.059
North Sea South	4	59°03′34.8"N 5°43′06.6"E	11 (1.7%)	19	2.10	0.404 ± 0.094	0.274 ± 0.093
North Sea North	5	60°09'43.2"N 5°02'16.7"E	16 (2.7%)	19	2.10	0.394 ± 0.088	0.259 ± 0.062
Norwegian Sea South	6	62°48′40.5"N 6°30′33.4"E	14 (3.3%)	22	2.15	0.356 ± 0.085	0.340 ± 0.084
-	7	65°41′24.4"N 12°07′51.5"E	7 (12.4%)	18	2.18	0.346 ± 0.104	0.099 ± 0.039
Barents Sea	8	70°41′18.2"N 29°22′31.5"E	8 (7.0%)	20	2.29	0.455 ± 0.071	0.140 ± 0.059
Saccharina latissima							
Skagerrak	1	58°59′36.9"N 9°48′30.7"E	18 (0.3%)	24	1.98	0.363 ± 0.079	0.323 ± 0.081
	2	58°15′16.3"N 8°31′22.1"E	22 (0.0%)	31	2.05	0.385 ± 0.066	0.288 ± 0.070
North Sea South	4	59°03′34.8"N 5°43′06.6"E	17 (2.0%)	28	2.05	0.382 ± 0.072	0.285 ± 0.070
North Sea North	5	60°16′11.1"N 5°13′18.8"E	9 (0.0%)	17	1.70	0.308 ± 0.071	0.296 ± 0.113
Norwegian Sea South	6	62°47′58.2"N 6°31′57.2"E	16 (2.4%)	30	2.29	0.468 ± 0.072	0.283 ± 0.072
	7	65°41′24.4"N 12°07′51.5"E	4 (5.6%)	25	2.60	0.420 ± 0.091	0.343 ± 0.064
Barents Sea	8	70°25'33.0"N 25°14'55.1"E	10 (0.0%)	24	1.89	0.292 ± 0.087	0.311 ± 0.103
	9	78°03′57.0"N 14°12′04.8"E	10 (1.1%)	30	2.23	0.357 ± 0.117	0.265 ± 0.102

Listed are: ecoregions, ID (sampling site numbering in a south-north direction), geographical sample position, #ind (number of individuals sampled at each location, and in parentheses the percentage of missing data from the microsatellite genotyping). Diversity indicators with number of genotyped alleles per population ($N_{A_{Pop}}$) including average rarefaction per population, rarefaction per sample (N_R), and expected (He) and observed (Ho) heterozygosity \pm SE



Fig. 1 Sampling maps covering Norwegian coastal territories with indications of ecoregions and sample positions (black dots) for the two studied kelp species *Laminaria hyperborea* (left) and *Saccharina latissima* (right). Pie carts indicate percentage proportions of a

defined number of genetic clusters to represent the genetic differentiation within and among ecoregions. For *L. hyperborea* and *S. latissima* four and three genetic clusters represent these, respectively. For precise sampling location positioning please see Table 1

 Table 2
 Overview of microsatellites used to genotype the collected samples of Laminaria hyperborea and Saccharina latissima (*Touchdown PCR), Multiplex (m) and Species Specific (SpSpecific)

Marker	Temp	М	Forward	Reverse	Developed by	SpSpecific
Laminaria	hyperbored	а				
CS34	50	А	CCAGTTCGCTCGTAGATTA	TTAAAGACGCCAGTAACCA	Wang et al. (2011)	No
Ld-148	55	В	TGGCGTGTTCCCTGATATG	TTTCTTGTCTAGGCCTCTCTGG	Billot et al. (1998)	No
Ld-158	55	В	CGACTAGAAGGGAGCGAGAA	CGTTTTTGCGCCTAACGTAT	Billot et al. (1998)	No
Ld-167	55	В	CGGACTCGATTTTAGCGATGGG	TCGGAAGCACGTGTTCTGTAT	Billot et al. (1998)	No
LOL-24	55	В	AAGTTTCGTCTCCGTTTCCTC	TTAACGTTCGTGCACCGTAG	Coelho et al. (2014)	No
LOL-15	55	С	AGAATCAGGACGGACACTGG	TCCGCGACTTATTCCCTCTA	Coelho et al. (2014)	No
LOL-23	55	С	AGAAAAAGCCTGCCGTGAC	CAGCCTGGAGCTTTCGAT	Coelho et al. (2014)	No
LOL-28	55	С	CACCAAGCTTGATTGGTTGA	TCGTGAGGTTATCGTGGTGA	Coelho et al. (2014)	No
Saccharina	ı latissima					
Sacl-68	65-55*	В	GGTGGGATTCTTTGGACGA	AAATGTGCTTGGGTCGGG	Guzinski et al. (2016)	Yes
Sacl-81	65-55*	С	ACTTTGGCTCGGTCTGCTT	CCTCCTCCCTTACCTACCTCC	Guzinski et al. (2016)	Yes
Sacl-19	65-55*	В	GCGTATTCCTAAACACCTCCC	CGATGACTGCCACCACAG	Guzinski et al. (2016)	Yes
Sacl-90	65-55*	С	ATTGTGTTGCTGGATGAGGAC	ACCTTCCGCTCTCTCGCT	Guzinski et al. (2016)	Yes
Sacl-65	65-55*	В	ATCTCCCAAACCACACAAG	CATCATCGTCAAGAACTCGG	Guzinski et al. (2016)	Yes
Sacl-95	65-55*	В	GGGAAGGAGGAAGAAGGTG	TAGCGGGAAAGAACGGGTAGT	Guzinski et al. (2016)	Yes
CS13	50	А	TTCTTGTTTCAGTGCCAGTTC	CGCAGCTTTACCCTCCTCTT	Wang et al. (2011)	No
CS12	50	А	GACGCTGCAAACCGACTTCT	GCTGGACCTTCACGTACCTTG	Wang et al. (2011)	No
SSR261	50	А	AGATGGAAGAAGACCTCG	ACATGAACCAAACCTACAGT	Zhang et al. (2015)	No

Additional methodology and results for cross amplification tests are available in the supplementary material (Table S1).

Laminaria hyperborea Final amplification volume was 5 μ l, containing 2.5 μ l 2× Multiplex Master Mix (Qiagen) with HotStarTaq DNA Polymerase, 0.08 μ l forward primer (5 μ M) with M13 tail, 0.33 μ l fluorescent-labelled M13 tail (5 μ M, FAM, PET, VIC, or NED), 0.33 μ l reverse primer (5 μ M), 0.76 μ l Milli-Q water and 1 μ l 10× diluted template DNA. PCR conditions included an initial denaturation step at 95 °C for 15 min and two rounds of cycles: 30 cycles of denaturation at 94 °C for 30 s, annealing at 50 or 55 °C for 45 s (Table 2) and extension at 72 °C for 45 s, followed by seven cycles of denaturation at 95 °C for 45 s. The cycles were followed by an extension at 72 °C for 20 min and a 10 °C hold.

Saccharina latissima Final amplification volume was 10 μ l, containing 5 μ l 2× Multiplex Master Mix (Qiagen) with HotStarTaq DNA Polymerase, 0.4 μ l fluorescentlabelled forward primer (5 μ M, FAM, YaYe, 565, or VIC), 0.4 μ l reverse primer (5 μ M), 3.2 μ l Milli-Q water and 1 μ l 10× diluted template DNA. PCR conditions included an initial denaturation step at 95 °C for 15 min, 10 cycles of touchdown with denaturation at 94 °C for 30 s, annealing at 65 °C (-1 °C for each cycle) for 30 s and extension at 72 °C for 30 s, followed by 30 cycles of denaturation at 94 °C for 30 s, annealing at 50 or 55 °C for 30 s (Table 2), and extension at 72 °C for 30 s. The cycles were followed by an extension at 72 °C for 10 min and a 10 °C hold.

All PCR amplifications were run on a Mastercycler nexus (Eppendorf, Germany) thermal cycler with PCR conditions as specified above. DNA fragments from both species were separated by capillary electrophoresis using an ABI-3130 sequencer (Applied Biosystems, USA). PCR products were pooled according to Table 2 and 1 μ l was mixed with 10.5 μ l of HiDi formamide (Life Technologies, USA) and 0.5 μ l of GeneScan 500 LIZ (for *L. hyperborea*) and GeneScan 600 LIZ (for *S. latissima*) size standard (ABI). Peaks were scored manually using GENEMAPPER 4.0 (ABI).

Data analyses

MicroChecker v2.2.3 (Van Oosterhout et al. 2004) was used to analyze the genotyped microsatellites for null alleles and scoring errors. The number of alleles genotyped at each locus and for each sampling location was calculated with HP-RARE (Kalinowski 2005), using the rarefaction with eight genes for *L. hyperborea* and six genes for *S. latissima*. The rarefaction was thereby run based on the smallest representation of genes in the samples being four and three, respectively, due to missing data in some of the loci. Observed and expected heterozygosity, and departure from Hardy–Weinberg equilibrium (HWE) were calculated using ARLEQUIN v3.5 (Excoffier et al. 2005). Linkage disequilibrium (LD) was tested in GENEPOP v4.0 (Raymond and Rousset 1995; Rousset 2008). The power of the set of microsatellites to detect genetic differentiation (both χ^2 and Fisher's exact tests) among all samples, for both species independently, was estimated in POWSIM v4.1 (Ryman and Palm 2006) running 1000 simulations using empirical sample sizes and allele frequencies, and loci numbers. Global and pairwise F_{ST} (Θ_{ST} ; Weir and Cockerham 1984), with statistical significance tested by 10,000 permutations, was calculated using MSA v4.05 (Dieringer and Schlötterer 2003). This program was also used to calculate genetic distances between population pairs by computing Cavalli-Sforza and Edwards (1967) genetic chord distances (D_{CE}), and bootstrapping 2000 times (Hedges 1992). These genetic distances were used to construct a Neighbor–Joining (NJ) tree (Takezaki and Nei 1996) with the PHYLIP software package (Felsenstein 2005) and visualized in SPLITSTREE v4.14.4 (Huson and Bryant 2006). All tests of statistical significance were adjusted for multiple tests by the false discovery rate (FDR) correction (Benjamini and Yekutieli 2001). Genetic relationship among individuals and sampling locations was assessed by applying a discriminant analysis of principal components (DAPC) using the adegenet v.2.0.1 (Jombart 2008) package in R v3.3.2 (R Development Core Team 2010). IBD in a northward direction along the coastline, using sampling location 1 of both kelp species as the starting point, was calculated in two ways: (i) comparing either genetic distance (D_{CE}) or $F_{ST}/(1-F_{ST})$ using the ape v2.3-1 package against geographic distance and (ii) testing for statistical significance in a Mantel test run in R (R Development Core Team 2010). Genetic clustering of sampled individuals was assessed using STRUCTURE v3.4.2 (Pritchard et al. 2000; Falush et al. 2007; Hubisz et al. 2009) performing 100 000 burn ins and 300,000 iterations with 20 replicates per K for K 1–10 assuming an admixture model and correlated allele frequencies (Falush et al. 2003). The best representation of each dataset was evaluated using both Ln P(K) (Falush et al. 2003) and Delta K (Evanno et al. 2005) calculated using STRUCTURE HARVESTER (Earl and vonHoldt 2012). To explore the potential presence of subpopulation structure, additional STRUCTURE runs were conducted for each K-cluster in a hierarchical manner. To maximize the accuracy of the twenty independent runs, the program CLUMPP v1.1.2 (Jakobsson and Rosenberg 2007), using the greedy function, was used and finally the results were visualized using DISTRUCT v1.1 (Rosenberg 2004). Final evaluation of K was compared with significant bootstrapping [i.e. > 70% (Hillis and Bull 1993)] in the NJ tree, population clustering in the DAPC, and positioning of the sampling locations in the IBD plots, as STRUCTU RE should not stand alone (Anderson and Dunham 2008; Thaulow et al. 2013).

Results

Four microsatellite markers (CS34, CS12, CS13, SSR261) were successfully cross-amplified from other closely related species (Table 2). For all markers combined, a total of 34 alleles were genotyped in L. hyperborea ranging from 22 (sample 6) to 12 (samples 1 and 3) among sampling locations, and 59 for S. latissima ranging from 31 (sample 2) to 17 (sample 5) (Table 1). The rarefaction allele count showed an increasing number (decimal numbers) from the southern samples and northward for L. hyperborea. For S. latissima, rarefaction allele count was variable but with an indication of more alleles with increasing latitude (Table 1). Expected and observed heterozygosity for populations of L. hyperborea ranged from 0.10 to 0.46 and 0.10 to 0.34, respectively, and for S. latissima from 0.29 to 0.47 and 0.27 to 0.34, respectively (Table 1). Locus specific calculations within each population for L. hyperborea, and S. latissima are presented in Tables S2 and S3, respectively. None of the microsatellite markers, for either species, contained null alleles or LD between the same pair of loci, in any of the sampled locations. Departure from HWE was not pronounced for any locus in L. hyperborea, whereas locus Sac190 in S. latissima showed significant departure in five of the eight sampling locations. Poor tissue preservation or suboptimal extraction may have been responsible for the 6% and 2% failed microsatellite genotyping in the L. hyperborea and S. latissima data sets, respectively (Table 1). The power of the two data sets to detect true population differentiation at an F_{ST} value of 0.030 was supported by a 99% probability by Fisher's exact test and the χ^2 test of 100% for both species (Fig. 2). Since the smallest significant F_{ST} value



Fig. 2 Power analysis of the genotyped microsatellites, showing Fisher's exact test and χ^2 test, for both *Laminaria hyperborea* (LH) and *Saccharina latissima* (SL)

for each species was above 0.030 (Table 3), these are well within the supported detection limit.

STRUCTURE clustering of the locations of L. hyperborea showed a clear separation into four K-clusters (Fig. S1) in accordance with ecoregions (Figs. 1, 3). All three Skagerrak sampling locations showed over 95% genetic identity to the first cluster (Fig. 1). The two North Sea sampling locations assigned mainly to the second cluster, which showed 89% and 66% membership, respectively. The two Norwegian Sea sampling locations showed 95% and 75% identity, respectively, to the third cluster. Cluster four was the most dominant cluster in the Barents Sea sample with 89% representation. The clustering into four groups corresponding to sampling locations and ecoregions is well corroborated by the bootstrap values in the NJ tree analysis and by the DAPC, which showed a clear visual separation of the Barents Sea from the remaining L. hyperborea samples along the second eigenvalue axis (Fig. 3).

K = 3 was the most likely clustering pattern for the S. latissima samples after visual inspection despite Delta K indicating K = 2 (Fig. S2). STRUCTURE showed a genetic clustering of samples in relative accordance with geographic positioning (Fig. 3). However, the S. latissima samples did not cluster at an equally structured scale as the L. hyperborea samples (Fig. 3). The two Skagerrak samples shared the same cluster with the North Sea South sample, whereas the North Sea North sample clustered together with the most southern of the Norwegian Sea samples. The two northernmost samples from the Barents Sea were assigned to the third cluster (Fig. 1). The most northern of the Norwegian Sea samples (7) showed a mixed assignment with equal representation from the two southern clusters ($\approx 43\%$) and only 14.4% from the northern cluster (Fig. 1). Also, the NJ tree and DAPC (Fig. 3) grouped the Skagerrak sampling locations closely together at one end of the latitudinal gradient and the two Barents Sea sampling locations at the other end. The North Sea and Norwegian Sea samples, however, were not separated according to ecoregions, but rather as a mix between the northern and southern samples (Fig. 3).

The genetic relationship among sampling locations as a function of geographic distance (i.e. isolation by distance, IBD) was identified to be statistically significant (Mantel test) and with good data representation, calculated based on both genetic distance (p-value < 0.0001, $R^2 = 0.72$, Fig. 4a) and F_{ST} (p-value = 0.0050, $R^2 = 0.59$, Fig. 4b) for the *L. hyperborea* samples in a northward direction. Significant IBD was also identified among the *S. latissima* sampling locations when calculated based on genetic distance (p-value < 0.0050, $R^2 = 0.63$, Fig. 4c), however, not when using F_{ST} (p-value = 0.0590, $R^2 = 0.04$, Fig. 4d).

Lamina	aria hyperbore	а						
	1	2	3	4	5	6	7	8
1		NS	NS	***	***	***	***	***
2	0.000		NS	***	***	***	***	***
3	0.000	0.000		*	***	***	***	***
4	0.261	0.320	0.229		*	***	***	***
5	0.267	0.327	0.242	0.076		***	***	***
6	0.534	0.587	0.500	0.339	0.239		*	***
7	0.594	0.656	0.563	0.308	0.216	0.096		***
8	0.527	0.613	0.505	0.328	0.362	0.441	0.366	
Saccha	rina latissima							
	1	2	4	5	6	7	8	9
1		*	***	***	***	***	***	***
2	0.032		*	***	***	*	***	***
4	0.072	0.052		***	***	*	***	***
5	0.271	0.232	0.152		***	*	***	***
6	0.281	0.254	0.169	0.091		NS	***	***
7	0.178	0.166	0.084	0.126	0.013		***	**
8	0.419	0.394	0.372	0.381	0.183	0.262		***
9	0.331	0.320	0.315	0.308	0.167	0.209	0.154	

Table 3 Pairwise F_{ST} values (below diagonal) among sampled locations (numbers from Table 1) of *Laminaria hyperborea* (upper table) and *Saccharina latissima* (lower table) locations, with tested statistical significance (above diagonal)

NS not significant

*5%, **1%, ***0.1%

Discussion

The present study provides the first screening of genetic diversity, geographical heterogeneity and genetic differentiation among the two most dominant and commercially important kelp species along the Norwegian coast, i.e. *S. latissima* and *L. hyperborea*. Both species demonstrated genetic heterogeneity along the Norwegian coast and clustered into three (*S. latissima*) and four (*L. hyperborea*) different genetic groups in accordance with defined ecoregions and with geographic distance from South to North, i.e. showing IBD.

Geographical heterogeneity and genetic diversity

The Norwegian coastal ecosystem is divided into six ecoregions (Fig. 1) for management purposes based on climatic conditions, ocean currents and biogeographic patterns of biologically important species and other biological quality elements (Gundersen et al. 2017). The ecoregions are defined to fulfil the requirements of the Norwegian Water Management Regulation (Water Regulation 2016) and the European Water Framework Directive (Jncc.defra.gov.uk 2010), which aim to ensure comprehensive ecosystem-based management of marine resources. The ecoregions are also used to determine restrictions related to aquaculture and kelp farming.

Both species of kelp showed strong signatures of IBD when using chords distance D_{CE} compared to the more traditional regression of $F_{ST}/(1-F_{ST})$ (Fig. 4), in accordance with a recent study by Séré et al. (2017). IBD based on F_{ST} / $(1-F_{ST})$ has been found for L. hyperborea along the coast of France (Robuchon et al. 2014) and for S. latissima in the Irish Sea (Mooney et al. 2018). In contrast, larger studies of S. latissima across Europe have not found IBD based on $F_{ST}/(1-F_{ST})$ (Guzinski et al. 2016), which is also true for smaller scale studies along the coast of Maine, USA (Breton et al. 2018). Different genetic distance estimates for the calculation of IBD should therefore in each case be evaluated. The relatively strong differentiation among ecoregions in Norway, designated to IBD (Fig. 4) and genetic clustering (Fig. 1), indicates limited range dispersal of zoospores or colonization success by both species. Sea urchin populations along the Norwegian coast show a weaker pattern of IBD compared to kelp (Norderhaug et al. 2016), which could be explained by the higher duration and dispersal potential of the sea urchin pelagic larval stage compared to the kelp spores (see Fredriksen et al. 1995; Sogn Andersen 2013). Despite differences in the strength of population genetic Fig. 3 Wild populations of both Laminaria hyperborea and Saccharina latissima significantly separated into distinct ecoregions along the Norwegian coast. STRUCTU RE, Neighbor-Joining tree (with significance from 2000 bootstraps), and DAPC analyses of L. hyperborea (left) and S. latissima (right) individuals from nine sampling locations along the Norwegian coast line, including Svalbard. Numbering of sampling locations correlate to Table 1. Colours in the three different analyses correspond to ecoregions as specified for the L. hyperborea STRUCTU RE results. However, in the S. latissima STRUCTURE results, the clustering of the North Sea South and the Norwegian Sea South sampling locations does not conform as consistent as L. hyperborea, since only three clusters were identified



structure, the genetic patterns found for kelp are consistent with ocean current larval dispersal in a northward fashion as seen for sea urchins. This indicates that the dispersal possibility also exists for kelp, but that other ecological barriers probably limit the dispersal and mixing rate, especially for *L. hyperborea*.

Overall, genetic diversity of *S. latissima* along the coast of Norway was similar if not slightly higher than reported for populations in Maine, USA (Breton et al. 2018) and lower than genetic diversity for populations within Europe, including one sample from Greenland (Nielsen et al. 2016, Paulino et al. 2016). Genetic diversity of *L. hyperborea* was lower for most Norwegian populations in comparison to populations along the French coast (Robuchon et al. 2014). In the southern ecoregion of Norway, *L. hyperborea* displayed even lower genetic diversity and strong differentiation from the remaining sampling locations in northern Norway. This points to the Skagerrak ecoregion being isolated (Höglund 2009) for *L. hyperborea*, a pattern that was also observed in a disconnected region on the French coast (Robuchon et al. 2014). Such low genetic diversity may be the consequence of fragmented and low density sites compared to what has been found for *S. latissima* (Norderhaug et al. 2011), potentially resulting in limited opportunities for genetic mixing with close-by populations. However, if some mixing does occur among unsampled (hence unaccounted for) populations, it could lead to the Wahlund effect (Wahlund 1928). This could explain the here observed heterozygosity deficiency in



Fig. 4 Genetic differentiation of *Laminaria hyperborea* and *Saccharina latissima* as a function of geographical distance from the sampling location closest to Sweden and in a northward fashion, cal-

culated based on genetic distance (a, c) or $F_{ST}(b, d)$. Sample numbering is explained in Table 1

the northernmost populations and further the partial deviation from HWE expectations. For *L. hyperborea* exchange of gametes may be further limited by slower growth in sheltered areas compared to exposed areas (Sjøtun et al. 1993) leading to a delay in spore production (Kain and Jones 1975) and thereby reducing overall fitness.

Some differences in the regional patterns of genetic structure and connectivity between the two kelp species exist and can most likely be explained by differences in dispersal abilities. Spore dispersal of *L. hyperborea* has been found to be distance-limited (Fredriksen et al. 1995, see also; Nielsen et al. 2016) while spores of *S. latissima* stay longer in the water masses (Sogn Andersen 2013) and therefore also travel farther (Kain and Jones 1975). This is reflected in the more opportunistic life strategy of

the short-lived *S. latissima* (Moy and Christie 2012) and can be observed in the sea urchin removal experiment of Leinaas and Christie (1996). In this experiment, when sea urchins were removed from a small isolated island far from any known kelp beds, *S. latissima* appeared as dense beds within the first year while *L. hyperborea* took at least four years to settle. Similarly, *S. latissima* is the first to recolonize more recent sea urchin depleted areas (own unpublished observations). The dispersal ability of *S. latissima* may also explain differences in connectivity between the two species in the Skagerrak and North Sea region, given that a long coastline of sand (Jæren) divides these regions of rocky shores available for kelps, as also pointed out by Luttikhuizen et al. (2018) discussing dispersal barriers.

Ecological trends

We found that the genetic diversity of S. latissima in the Skagerrak oceanic region was comparable to other regions in Norway (Table 1). This trend is surprising given the fact that the region has experienced large declines in S. latissima biomass, in the order of 51-80%, during the last fifteen years (Bekkby and Moy 2011; Moy and Christie 2012). Most of the decline has occurred in sheltered areas, at shallow depths, due to anthropogenic stressors (e.g. increased land run-off and nutrient loads from rivers) and elevated water temperatures. Warmer temperatures, in conjunction with increased shading by epibionts and decreased water transparency, have been identified as the main drivers for this substantial kelp forest loss (Sogn Andersen et al. 2011; Moy and Christie 2012; Sogn Andersen 2013). This largescale disappearance was observed in 2002, and the severe reduction in biomass (demographic bottleneck) may have resulted in reduced allelic richness. A bottleneck analysis showed, however, no indication thereof (data not shown). The consequences of a demographic bottleneck are expected to be reduced, if connectivity among habitat patches was high (Jangjoo et al. 2016).

Due to the ability of S. latissima zoospores to survive for several days in ocean currents (Kain and Jones 1975), high interbreeding within the Skagerrak oceanic region is possible. Further, Moy and Christie (2012) indicated that S. latissima is a species with more opportunistic traits and dispersal abilities than other kelps, leading to shorter-term disappearance and reappearance and thus higher connectivity within the region. Connectivity along the southern coastline of Norway could potentially be explained by longrange dispersal of zoospores, as indicated by genetic clustering of the southern sampling location in the North Sea region with the Skagerrak samples and single individuals from other northern populations. Indeed, no genetic differentiation was observed between a Norwegian and a Swedish population collected close to the Norwegian boarder, within the Skagerrak basin (Nielsen et al. 2016). Other causes than geographical distance exist explaining the genetic differentiation among S. latissima populations (Mooney et al. 2018). Evidence for connectivity therefore seems to reside with water currents within and across regions, counteracted by dispersal barriers in the form of unfavourable bottom substrate, freshwater efflux and open water (Breton et al. 2018; Luttikhuizen et al. 2018; Mooney et al. 2018).

Once the kelp species disappear, filamentous algae and sediment become dominant and may inhibit recolonization of kelps as shown by seaweed species in several regions (Gorman and Connell 2009; Sogn Andersen et al. 2011; Sogn Andersen 2013). Efforts to minimize nutrient and sediment fluxes seem to be of great importance for the preservation of kelp in the region. Genetic resilience was indeed proven to play a significant role in a marine heat wave extirpation of a kelp species (Wernberg et al. 2018). In the present recovery process of the Norwegian kelp forests, after sea urchin depletion, it is important to gain knowledge on the baseline population structures, genetic diversity, and other stressors before a large-scale reforestation takes place.

Genetic diversity within and differentiation among sampling locations of L. hyperborea was higher in the northern regions compared to the Skagerrak locations. This could indicate more abundant and relatively isolated (sub)populations with minimal, yet sufficient, genetic exchange to cluster together, compared to what was found in the south of Norway. The extensive areas grazed by sea urchins (Sivertsen 1997; Norderhaug and Christie 2009) have created longer distances between kelp sub-populations. The possibility of genetic input from un-sampled 'ghost' populations within the regions could be feasible yet hard to document, since kelp populations have been decimated for more than 45 years. However, range expansion of the crustaceans Cancer pagurus and Carcinus maenas crabs (Fagerli et al. 2013) and warming ocean temperatures (Fagerli et al. 2014) have in recent years led to collapse and northward retraction of sea urchins. In the north, these are also experiencing population decimation due to increases in king crab, Paralithodes camtschaticus, leading to kelp recovery on both the Russian and Norwegian coasts (Gudimov et al. 2003; Christie and Gundersen 2014). Despite their limited current scope, future populations of L. hyperborea and S. latissima might experience even higher genetic diversity within the region due to recolonization from multiple founder populations, as suggested for the brown seaweed, Fucus distichus, recolonizing an area after an oil spill (Coyer et al. 2011). Founder populations exist within the region harbouring in exposed habitats (Rinde et al. 2014) not utilized by sea urchins.

Recommendations for management and Outlook

Marine management must strive to preserve genetic heterogeneity among wild populations and a rich local and regional genetic diversity within species to ensure healthy kelp forests that can withstand natural and anthropogenic pressures. Genetic diversity and the integrity of differentiated populations are needed to preserve robust ecosystems and maintain natural resilience properties despite human harvesting and cultivation efforts. The results from this study are intended to serve as a baseline for follow-up studies to monitor wild kelp forests along the Norwegian coast and to assist management agencies in large-scale kelp reforestation in areas previously dominated by sea urchins. A detailed understanding of kelp genetic heterogeneity across ecoregions is important due to the high interest in large-scale cultivation of kelp, particularly of S. latissima, all along the Norwegian coast. The results presented here will serve to assist formulation of knowledge-based guidelines for sustainable wild-harvesting and large-scale cultivation of kelp. Guidelines should include recommendations to exclusively cultivate kelp strains of local origin to preserve local genetic structure and diversity. Thus, the present study supports the continuation of the precautionary principle strategy recommended for kelp cultivation; that only local ecotypes of kelp should be cultivated and that kelp strains should not be transported between fjords and across ecoregions for cultivation (Fredriksen and Sjøtun 2015).

Despite the rather low numbers of samples from some of the locations, the power analysis showed sufficient strength and significance to support the degree of genetic differentiation and heterogeneity (Fig. 2). Overall genetic variance among the Norwegian samples presented here are in accordance with a recent study that sampled individuals from multiple regions across Europe and North America (Luttikhuizen et al. 2018; but see Neiva et al. 2018). However, to fully investigate the transition zones between genetic clusters, additional samples should be collected and analysed with higher genome coverage to identify areas of special concern for the establishment of kelp and seaweed cultivation, avoiding unintended introgression from cultured conspecifics into wild populations, as was observed previously in the salmon farming industry (Glover et al. 2012; Faust et al. 2018). Additionally, a more intensive sampling program including higher sample density is needed to obtain a full understanding of the genetic diversity of kelp along the Norwegian coast and to draft appropriate management strategies for future large-scale seaweed and kelp cultivation.

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