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Conservation genomics of anadromous Atlantic salmon across its North American range: outlier loci identify the same patterns of population structure as neutral loci

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Abstract

Anadromous Atlantic salmon (*Salmo salar*) is a species of major conservation and management concern in North America, where population abundance has been declining over the past 30 years. Effective conservation actions require the delineation of conservation units to appropriately reflect the spatial scale of intraspecific variation and local adaptation. Towards this goal, we used the most comprehensive genetic and genomic database for Atlantic salmon to date, covering the entire North American range of the species. The database included microsatellite data from 9,142 individuals from 149 sampling locations and data from a medium-density SNP array providing genotypes for >3,000 SNPs for 50 sampling locations. We used neutral and putatively selected loci to integrate adaptive information in the definition of conservation units. Bayesian clustering with the microsatellite dataset and with neutral SNPs identified regional groupings largely consistent with previously published regional assessments. The use of outlier SNPs did not result in major differences in the regional groupings, suggesting that neutral markers can reflect the geographic scale of local adaptation despite not being under selection. We also performed assignment tests to compare power obtained from microsatellites, neutral SNPs and outlier SNPs. Using SNP data substantially improved power compared to microsatellites, and an assignment success of 97% to the population of origin and of 100% to the region of origin was achieved when all SNP loci were used. Using outlier SNPs only resulted in minor improvements to assignment success to the population of origin but

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improved regional assignment. We discuss the implications of these new genetic resources for the conservation and management of Atlantic salmon in North America.

Introduction

The increased availability of genomic data in non-model organisms has the potential to revolutionize the use of genetic tools in conservation and management (Kohn *et al.* 2006; Primmer 2009; Avise 2009; Allendorf *et al.* 2010). At the most basic level, the greater number of loci available can increase the precision and accuracy of population parameters of interest formerly estimated with a small number of neutral markers (Allendorf *et al.* 2010). More fundamentally, however, genomic tools can also provide new types of information not available with traditional markers. For instance, they can provide information on the genomic architecture of inbreeding and/or outbreeding depression (Allendorf *et al.* 2010) and highlight regions of differential introgression in the genome of native taxa under threat from hybridization (Crispo *et al.* 2011; Lamaze *et al.* 2012; Hohenlohe *et al.* 2013). Furthermore, genomic data can be used to infer the presence of loci or genomic regions that show higher differentiation and are putatively under directional selection (Nosil *et al.* 2009; Stapley *et al.* 2010). Such outlier loci could therefore offer an opportunity to assess patterns of local adaptation among populations, thus informing conservation and management.

Establishing conservation units based on objective and practical criteria has always been a challenge (Moritz 1994; Waples 1995; Fraser & Bernatchez 2001). While neutral markers may provide appropriate tools to delimit reproductively isolated populations (Waples & Gaggiotti 2006), their utility in determining their adaptive potential or ecological distinctiveness has been questioned (Crandall *et al.* 2000; Fraser & Bernatchez 2001; McKay & Latta 2002). The inclusion of phenotypic or ecological data in the definition of conservation units has been advocated by some (Waples 1991; Crandall *et al.* 2000), but remained impractical for most taxa of conservation concern. The analysis of genomic data combined with genome scan approaches identifying putatively adaptive loci could provide a practical solution to this problem (Bradbury *et al.* 2013). Recently, Funk *et al.* (2012) proposed a framework to integrate adaptive genetic variation from genomic data into the

Accepted Article

definition of conservation units. In short, they propose a sequential assessment of population structure using in turn putatively neutral loci and putatively adaptive loci. In principle, the inclusion of information on putatively adapted loci can help resolve questions regarding the spatial scale of local adaptation, and help tailor conservation and management actions to specific conditions.

Anadromous Atlantic salmon (*Salmo salar*) is a species of biological as well as cultural and economic importance (Dodson *et al.* 1998; COSEWIC 2011). Population declines, however, have also made it a species of conservation concern (COSEWIC 2011; Chaput 2012). Historically, anthropogenic habitat destruction and overexploitation have severely reduced the range of the species in the United States (Spidle *et al.* 2001) and reduced abundance in southern Canada (COSEWIC 2011), with more recent declines (since the early 1990's) possibly resulting from increased mortality at sea (Friedland *et al.* 2005; Chaput 2012). In addition, the species is popular with the sport fishing industry, which targets adults returning from their marine migrations to spawn. Offshore fisheries also operate in Greenland, Labrador and on Saint-Pierre and Miquelon Islands targeting migrating adults in the marine phase of their life-cycle (Chaput 2012). In Greenland, for example, a subsistence fishery targets adults migrating from all regions of North America to the feeding grounds of West Greenland, making it an important mixed-stock fishery (Chaput *et al.* 2005; Gauthier-Ouellet *et al.* 2009). Improving genetic tools for mixed-stock fishery analysis and individual assignments to natal rivers or to their management unit of origin is therefore of interest. Furthermore, sampling as many source populations as possible is crucial to increase the precision of estimates of stock composition and of individual assignment. The use of a large number of SNPs, and especially 'outlier' SNPs, offers the promise to improve the power of mixed-stock fishery analysis and assignment tests (Russello *et al.* 2011).

An important paradigm in existing conservation and management strategies for Atlantic salmon is that populations are structured hierarchically (Dodson *et al.* 1998; COSEWIC 2011). At a small scale, the precise homing behaviour of the species (Stabell 1984; Hendry *et al.* 2004a; b), leads to genetic differentiation among rivers (Spidle *et al.* 2001; King *et al.* 2001; Dionne *et al.* 2008; Perrier *et al.* 2011; Bradbury *et al.* 2014), and in some cases within rivers (Garant *et al.* 2000; Vähä *et al.* 2007; Dionne *et al.* 2009). At a larger scale, dispersal occurs, but effective migration is spatially restricted, leading to genetically

differentiated regional groups (Dionne *et al.* 2008; Tonteri *et al.* 2009; Bourret *et al.* 2013a; Bradbury *et al.* 2014). These genetically differentiated regional groups form the basis of management and conservation units (COSEWIC 2011), because they represent units with restricted gene flow that are at least partially demographically independent (Waples & Gaggiotti 2006). Furthermore, there is increasing evidence that these genetic groups are differentially adapted to spatially varying environmental conditions (Taylor 1991; Garcia de Leaniz *et al.* 2007; Fraser *et al.* 2011), such as pathogen diversity (Dionne *et al.* 2007), water pH (Fraser *et al.* 2008), temperature, and geology (Bourret *et al.* 2013a). Anadromous Atlantic salmon would thus represent a good candidate species to evaluate the potential of adaptive markers to enhance the definition of conservation units.

In this study, data from a geographically extensive microsatellite database is combined with thousands of SNP markers to apply a conservation genomics framework to anadromous populations of Atlantic salmon in North America. Our first goal was to provide a North American-wide assessment of population structure on the basis of both neutral and putatively adaptive genetic variation in order to inform the establishment of conservation units for management. Population structure has been examined previously for the Québec populations (Dionne *et al.* 2008), the Newfoundland populations (Bradbury *et al.* 2014), and the Nova Scotia populations (McConnell *et al.* 1997; Vandersteen Tymchuk *et al.* 2010). A population genomics assessment conducted on the entire North American range in a single analysis, however, is lacking (but see King *et al.* (2001) for a range-wide assessment of microsatellite DNA variation based on a small number of sampling locations and Verspoor *et al.* (2005) for a North-America wide study based on allozymes). By combining results from previous studies to new data, we now have the most extensive Atlantic salmon microsatellite database to date to infer population structure with high geographical resolution: the database is comprised of data from 149 sampling locations, with over 9,000 individuals genotyped at 15 microsatellite loci. In order to evaluate the usefulness of integrating adaptive variation in the definition of conservation units, we used SNP data generated with a medium-density SNP array from a subset of the populations (Bourret *et al.* 2013b). Population structure was assessed on both putatively adaptive and neutral markers identified with two genome scan approaches. Our second goal was to test the power of the database for assigning individuals to (1) regional groups defined using the microsatellite

Accepted Article

data, and (2) to their sampling location of origin. We did so with microsatellite markers and SNP markers separately, with the explicit aim of evaluating the relative performance of the two types of markers. For the SNP data, we also compared the performance of outliers vs. putative neutral markers in population assignment.

Methods

Microsatellite genotyping and analysis

A total of 9,142 anadromous individuals from 149 sampling locations from the entire North American range of the species were genotyped at 15 microsatellite markers (Fig S1 and Table S1 in online appendix). Life stages sampled include parr, smolts, and in most cases returning adults and the samples were collected between 2000 and 2010 (Table S1 in online appendix). The data represented a combination of previously analyzed datasets (see Bradbury *et al.* (2014) and Dionne *et al.* (2008) for regional analyses and further details of genotyping) and new data (Table S1). Note that some populations included have been supplemented by hatchery-reared fish and/or have been hybridized with farmed escapees (Carr *et al.* 2004; Bourret *et al.* 2011). While this will necessarily influence our results, our goal is to document present-day population structure regardless of its causes. Individuals were genotyped using an ABI 3130xl (or standardized from ABI 3100 following Gauthier-Ouellet *et al.* 2009) by three independent laboratories (Table S2). Allele scoring was standardized across laboratories in the Laval University laboratory using a panel of 10 standard individuals. Additional individuals (n=87 total; 46 individuals from Newfoundland and Labrador; 41 from Nova Scotia and New Brunswick) were genotyped to account for unrepresented allele values and assess the entire range of allele sizes. To account for discrepancies between ABI 3100 and ABI 3130xl, a panel of 64 individuals representing all allele sizes of Québec populations previously genotyped on the ABI 3100 were analyzed on the ABI 3130xl. Re-screening of samples or re-analysis of allele sizes were completed when discrepancies existed between labs or machines. Scoring patterns among laboratories were generally consistent and allowed standardization using simple rules (see Table S2 in online appendix).

We tested for departures from Hardy-Weinberg equilibrium (HWE) using the least-square method based on the AMOVA F_{IS} implemented in GENODIVE (Meirmans & van Tienderen

2004) with 999 permutations. We tested for linkage disequilibrium (LD) between each pair of loci in each population using Genepop (1000 iterations per batch for 100 batches; (Raymond & Rousset 1995)). We adjusted the p-values of the LD and HWE tests for multiple comparisons according to the false discovery rate (FDR) method in the function 'p.adjust' in R with an experiment-wide alpha of 0.05 (Team 2006). Observed (H_0) and expected heterozygosity (H_S ; (Nei 1987)) were calculated in GENODIVE (Meirmans & van Tienderen 2004). Allelic richness was calculated for each sampling location using the rarefaction method implemented in HP-rare (Kalinowski 2005). The Moisie River sample was removed from this analysis because it is missing data at three loci. The HP-rare analysis was repeated twice: once on all the sampling locations (except Moisie; minimal sample size of 16), and once with the KC6 and Georges Rivers samples removed because these two locations had small sample sizes and many missing values, respectively (minimal sample size of 36). Pairwise F_{ST} values were calculated in GENODIVE, significance was assessed using 999 permutations, and p-values were adjusted for multiple comparisons with the FDR method.

SNP-array genotyping, filters and basic statistics

A total of 1,080 anadromous individuals from 50 sampling locations across the North American range (Table S3) were genotyped at 5,568 loci using the SNP-array developed by the Centre for Integrative Genetics (CIGENE, Norway) following the manufacturer's instructions (Illumina, San Diego, CA, USA). The data from the Québec populations was previously published in (Bourret *et al.* 2013b), but data from the other regions is previously unpublished. Detailed methods for SNP discovery and quality control can be found in Bourret *et al.* (2013a,b). Ascertainment bias for the North American populations is minimal (Bourret *et al.* 2013a) and should not bias our results since most markers on the chip were discovered from populations not considered here (European). The quality of individual samples was assured by only using individuals genotyped at a greater than 0.95 call rate (CR: proportion of genotyped SNPs). Markers with minor allele frequencies less than 1% (MAF < 0.01) across all populations and markers missing in more than 5% of individuals were excluded from our analyses. Observed (H_0) and expected heterozygosity (H_S ; Nei (1987)) were calculated in GENODIVE (Meirmans & van Tienderen 2004). As with the microsatellite data, we tested for departures from HWE GENODIVE with 999 permutations,

and adjusted the p-values for FDR in R. Pairwise F_{ST} values were calculated in GENODIVE, and significance was assessed using 999 permutations.

Outlier markers detection

We used several alternative genome scan methods to detect loci with greater than expected levels of divergence among regional groups. First, we used BAYESCAN, a Bayesian approach that allows the estimation of the posterior probability of a given locus being under the effect of selection (Foll *et al.* 2008). It is based on the multinomial-Dirichlet model, which assumes that allele frequencies among demes are correlated through a common migrant gene pool, therefore allowing complex ecological scenarios to be modeled satisfactorily (Foll *et al.* 2008). Because previous studies suggested that local adaptation is likely important at the regional scale (Dionne *et al.* 2008; Bourret *et al.* 2013a), we ran BAYESCAN on the entire dataset by defining groups of individuals on the basis of the microsatellite-defined regional groups (not on populations; Table S1 and results) using all the defaults (tests runs with longer chain parameters gave identical results). Loci putatively under selection were defined as those with alpha-values significantly greater than 0 (i.e., with Q-values smaller than 0.05) while loci putatively under balancing selection had alpha-values significantly smaller than 0. All other loci were considered neutral. Second, we used hierarchical Fdist (Excoffier *et al.* 2009) implemented in Arlequin 3.5 (Excoffier & Lischer 2010). This method uses a hierarchical island model allowing for lower migration rates among groups compared to among populations within groups. This has been shown in some instances to reduce the prevalence of false positives compared to methods using the finite island model (Excoffier *et al.* 2009; Mita *et al.* 2013). We used the regional groups defined using the microsatellites (Table S1 and results) as the higher level of population structure. Third, as an alternative way to identify loci that maximize assignment to the sampling location of origin, we conducted a 'nested Fdist' analysis, where we used a non-hierarchical Fdist on each microsatellite-defined regional groups separately, and combined all the unique markers identified by each of the individual analyses. The goal with the nested analysis was to select markers that would maximize assignment power to the population, and are thus only discussed in this context. In both hierarchical and nested Fdist analyses, loci with significantly (at the 0.01 significance level) higher F_{CT} or F_{ST} values were classified as outliers potentially under directional selection among regional groups, loci with

Accepted Article

significantly lower F_{CT} or F_{ST} values were loci putatively under balancing selection, and all other loci were classified putatively neutral loci. The results of both the Bayescan and hierarchical Fdist analyses were plotted against the estimated genomic position of a subset of SNP loci for which map data was available (Brenna-Hansen *et al.* 2012) to assess whether specific genomic regions of linkage groups showed elevated amounts of divergence.

Population structure

Initial tests using the widely used Bayesian clustering program STRUCTURE (Pritchard *et al.* 2000) determined that this approach was too computationally demanding even for the microsatellite dataset. Instead, we used BAPS (Corander *et al.* 2003; 2004; 2008), a model-based Bayesian clustering approach that, like STRUCTURE, infers genetic groupings that maximize Hardy-Weinberg and linkage equilibrium (Corander *et al.* 2003). Unlike STRUCTURE, however, BAPS infers the optimal number of clusters directly (instead of relying on *ad hoc* measures; (Corander *et al.* 2004)), and is computationally more efficient, therefore facilitating the analysis of large datasets. Since in this analysis we were interested in regional clustering of sampling locations, we used the 'clustering of groups of individuals' option in BAPS. This prior is justified by the homing behaviour of Atlantic salmon (Stabell 1984; Hendry *et al.* 2004b). For the microsatellite dataset, 20 replicate runs of BAPS were performed with a maximum number of K of 149 (the number of samples in the dataset). Runs with smaller values for maximum K (10, 20, 30, 40 and 50) were also performed to explore the effects of this prior on the final results, which were always identical.

After running BAPS on the entire dataset, we examined the possibility of further hierarchical population structure, by running both BAPS and STRUCTURE on the regional groups defined by BAPS with the full dataset. For these analyses on each regional groups, STRUCTURE was run with the admixture model with correlated allele frequencies, 50,000 burnin and 100,000 MCMC repetitions, K 1 to 10 with 10 iterations of each K . Alpha-values were examined to ensure that convergence had been attained after the burnin (Pritchard *et al.* 2000). For the BAPS analysis, we ran 20 independent runs with max $K = 50$ on each of the regional groupings separately.

For the SNP dataset, BAPS was run on all SNPs, on the neutral SNPs identified by Bayescan, on the outlier SNPs identified by Bayescan, and on the outlier SNPs identified by Fdist. In all cases, 20 independent runs were performed with a maximum number of K of 50 (again, the number of sampling locations).

Finally, in order to infer the relationship of regional groups among each other, we constructed an unrooted neighbor-joining tree of Cavalli-Sforza chord distances for the microsatellite data using the software package Phylip (Felsenstein 1993). The Moisie was excluded from this analysis because it is missing data at three loci. We used 1,000 bootstrap replicates to assess confidence in the nodes.

Individual assignment to regional groups and river of origin

We used two approaches to test and compare the relative power of the microsatellite and SNP datasets to assign individuals to their region of origin and their sampling location of origin. For the definition of regional groups, we used two hierarchical levels of population structure defined by the microsatellites (see Results). First, we used the leave-one-out approach implemented in ONCOR (Kalinowski *et al.* 2007). This approach sequentially removes each individual from the baseline and estimates its origin by determining the sampling location that has the highest probability of producing its genotype. Assignment success is then estimated as the proportion of individuals that were correctly assigned to their sampling location of origin. This approach does not allow for missing data. We therefore filled the missing data by drawing alleles randomly from the entire dataset in GENODIVE (Meirmans & van Tienderen 2004). This will reduce overall population differentiation and will in most cases provide conservative estimates of power of assignment. Second, we used the assignment test option in GENODIVE, which implements the approach of Paetkau *et al.* (1995; 2004). The assignment tests were performed on two different microsatellite datasets and on four different SNP datasets. For the microsatellites, analyses were first conducted on the entire database. However, to minimize biases due to sampling effects in the comparisons between SNPs and microsatellites, we combined the different sampling locations that represented several tributaries of the same river (see appendix for details). The second microsatellite dataset tested only included the sampling locations for which we had SNP data. This was done to further ensure comparability

between the microsatellite and SNP assignment tests in case the inclusion of many genetically similar populations in the full microsatellite dataset biased assignment success. For the SNPs, our goal was to compare the power of the entire dataset to other datasets which include only outlier loci, and which have been found elsewhere to be sufficient for powerful assignment (Russello *et al.* 2011). We therefore performed assignment tests on a dataset including all loci ($n = 3,192$), using only the outliers from the Bayescan analysis ($n = 106$), the outliers from the hierarchical Fdist analysis ($n = 61$), and from the nested hierarchical Fdist ($n = 293$). We further evaluated the effect of the number of SNPs included in the analysis on the power of assignment tests. We tested datasets with 50, 100, 250, 500 and 1000 randomly selected SNP markers and performed assignment tests in ONCOR for three replicate datasets for each number of random SNPs at both the population and regional level. In all cases, missing values were filled in GENODIVE according to the overall allele frequencies. Note that using the same samples to identify outliers and to assess their power of assignment can result in upwardly biased estimates of power (Anderson 2010). Our results, however, show that adding those outliers did not improve our assignment success compared to using all other markers (see Results), and the potential upward bias in power only reinforces this conclusion.

Results

Microsatellites

The microsatellite loci were highly polymorphic, with the number of alleles per locus ranging from 14 (D486) to 91 (SsaD144) (average 40; total 593). Missing values were rare (1.7% overall) and mainly the result of three loci (D486, D58, 2201) having a high failure rate for three populations (Moisie, George and Muddy Bay rivers; 22.2%, 16.9% and 10.1% missing values across all loci respectively). We found evidence for significant departures from HWE in 230 of 2,295 tests (at $\alpha = 0.05$), but none remained significant after adjustment for false discovery rate (FDR). Of the 230 significant tests before adjustments, 152 were heterozygote deficits. These heterozygote deficits were distributed more or less equally across populations and loci: the population with the most loci with heterozygote deficits (NL-SLR) had six and the loci with the most populations in heterozygote deficits (SsaD144) had 17. We also found evidence for significant LD in 1,773 of 15,863 tests (at $\alpha =$

0.05), but only 471 remained significant after adjustment for FDR. No pair of loci was consistently found to be in significant LD, but many populations showed elevated numbers of pairs of loci in significant LD. This was particularly true for the Nova Scotia populations (prefix NS in Table S1) that accounted for 226 of the 471 significant tests. Nova Scotia populations have been particularly affected by recent declines in abundance (COSEWIC 2011), which may partly explain this pattern. Pairwise population differentiation was moderate (average $F_{ST} = 0.043$) but values observed were typically greater than zero: only 45 pairwise F_{ST} estimates out of 11,628 tests were not statistically significant (at $\alpha = 0.05$ and after FDR adjustment). Non-significant tests mostly involved sampling locations with small sample sizes (GU-KC6 had 23 non-significant tests), or sampling locations found in the same watershed (e.g. the Miramichi watershed). Notably, comparisons involving sampling locations on Anticosti were either not significantly different (QC-CH was not different from both QC-SU and QC-JU), or marginally so ($p = 0.049$ for the QC-SU vs QC-JU comparison).

SNP-array genotyping, filters and basic statistics

A total of 3,192 loci were retained for the analyses. We found evidence for significant departures from HWE in 2,467 of 159,600 tests (at $\alpha = 0.05$), but none remained significant after adjustment for false discovery rate (FDR). Departures appeared roughly equally distributed among populations and loci. Expected heterozygosity averaged 0.179 across all populations, but varied by almost two-fold within population: from 0.128 (NS-MED) to 0.215 (NL-RKR) (Table S3). Average heterozygosity also varied among regions, and was highest in the Avalon Peninsula populations (0.215) and lowest on the East Coast of Nova Scotia (0.138). All pairwise population differentiation estimates were statistically significant ($p < 0.01$), with an average pairwise F_{ST} of 0.108, and ranging from 0.004 (QC-MAT vs. QC-SA) to 0.222 (QC-AF vs. GU-CRO).

Population structure: microsatellites

BAPS identified $K = 29$ as the most likely number of clusters (Figure 1). The clusters were consistent with geography and are largely consistent with previously published regional groups (Dionne *et al.* 2008; Bradbury *et al.* 2014). Of the 29 clusters, 17 were singletons, i.e.

populations that clustered in their own cluster (identified with asterisks in Figure 1). Note that the only population from the United States (Penobscot River) clustered on its own, which is consistent with previous studies suggesting U.S. populations are distinct from Canadian populations (Spidle *et al.* 2001). Of the other 12 clusters, 5 consisted of pairs of populations (one pair in Labrador, two pairs in the Inner Bay of Fundy, one pair grouping two populations in Prince Edward Island, and one pair on the Avalon Peninsula). The seven remaining clusters identify major regional groups: St. Lawrence North Shore, Gaspésie / Anticosti, Gulf of St. Lawrence, Nova Scotia East Coast, Newfoundland, Labrador, and Ungava. A detailed description of the regional groupings is provided in the online appendix.

The neighbor-joining tree of chord distances was largely consistent with the results of the BAPS analysis (Fig. 2). Few of the groups, however, were monophyletic, and none of the internal branches defining regional groups were well supported by bootstrap values. The location of the singleton populations identified by BAPS in the tree was consistent with geography (except for GU-KC6; but this populations had the smallest sample size with $n = 9$), and we therefore include them in the nearest regional clusters for all further analyses on regional groups.

We also identified further levels of hierarchical population structure within some of the regional groupings defined by the BAPS analysis (details in online appendix). A STRUCTURE analysis provided evidence of two sub-regional groups within the St. Lawrence North Shore regional grouping (upper vs. lower; Fig 1 and online appendix), and both STRUCTURE and BAPS showed that the three Anticosti populations were distinct from the Gaspésie populations (Fig 1 and appendix). There was also some evidence for a separation between the north and south portions of the Nova Scotia East Coast regional group, but BAPS and STRUCTURE disagreed regarding the location of the boundary (see online appendix for details). In all other regions, there was no clear evidence for further hierarchical structuring, thus suggesting that the previous analysis was sufficient in capturing population structure, at least at the regional scale (see online appendix for details).

Outlier markers detection

The Bayescan analysis conducted on the regional groups identified 106 loci putatively under divergent selection, 1,413 loci under balancing selection, and 1,673 putatively neutral

loci. The hierarchical Fdist analysis identified 61 loci putatively under directional selection, 32 loci putatively under balancing selection and 3,099 putatively neutral loci. Of the 61 loci putatively under directional selection, 36 were also identified as such by Bayescan (Fig. 3). The outlier loci were distributed fairly evenly across the genome, and were mapped to several linkage groups (Fig. 3). The nested analysis in Fdist identified a total of 293 loci putatively under directional selection within at least one of the regional groups.

Population structure: SNPs

The BAPS analyses conducted on different subsets of the SNP datasets did not substantially differ from the analysis conducted on the microsatellite dataset (Fig. 4). When all SNPs were used, BAPS uncovered 15 genetic clusters, three of which were formed of a single population. These clusters were very similar to the one based on microsatellites. Consistent with the hierarchical analysis performed on microsatellites, the SNP analysis identified two clusters in the St. Lawrence North Shore and concluded that the three Anticosti populations cluster apart from the Gaspésie sampling locations. Unlike the microsatellite analysis, however, the SNP analysis divided the Newfoundland sampling locations in two clusters, with the Northern Peninsula populations appearing distinct. The analysis on neutral loci (Bayescan) identified 20 clusters, but the increase was only due to more numerous singletons (8). The clusters formed of more than one populations were otherwise identical to those with all SNPs. Surprisingly, the analysis based on the Bayescan outlier loci gave identical results to that based on neutral SNPs. Using the hierarchical Fdist outliers, 20 clusters were identified, 7 of which were singletons. The clusters observed were similar to those from the other analyses, with the following differences: the Avalon Peninsula cluster disappeared; two new clusters of two locations were identified in the Labrador cluster and the Gulf of St. Lawrence cluster.

Individual assignment to regional groups and river of origin

The analyses conducted using ONCOR (ON) and GENODIVE (GD) gave comparable results (Fig. 5). Assignment success was high for all sets of markers at both hierarchical levels of regional groupings explored, and we therefore only discuss the results on the level of the

region. Microsatellites achieved a reasonable level of success in assigning individual to their region of origin both when all sampling locations were included (ON: 77%; GD: 64%); or when only the sampling locations with comparable SNP data were included (ON: 76%; GD: 73%). Note that the reduction in the number of sampling locations only improved the results of the GENODIVE analysis. Assignment success to the regional level was higher than 80% for all the SNP datasets examined, and was maximal when all SNPs were included in the analysis (ON: 100%; GD: 97%). Assignment success to the population of origin was low for microsatellite loci when all populations were included (ON: 51%; GD: 52%), but increased to a level comparable to that achieved with the outlier SNP when we used only the populations for which SNP data was available (Fig. 5). Filling missing data with overall allele frequencies did not seem to have a large influence on the ONCOR assignment success, even for populations with large amounts of missing data (e.g., Moisie River: ON: 32%; GD: 32%; George River: ON: 66.7%; GD: 77%). Assignment success to the population level also increased for the outlier loci from the nested genome scan (ON: 85%; GD: 84%), and was best when all loci were included (ON: 94%; GD: 97%).

Evaluation of the number of SNP loci included in the analysis on power of assignment suggested that the SNP dataset was sufficiently large to reach maximal power (Fig. 6). For randomly selected loci, assignment success at the regional level reached a plateau at around 250 loci (~95% assignment success), while at the population level, a plateau in assignment success seemed to be attained around 500 loci. Interestingly, using outlier loci only seemed to result in marginal increases in power compared to randomly selected loci. Indeed, the use of outlier loci only resulted in important increases in assignment success when the number of randomly selected loci was less than about 100 (Fig. 6).

Discussion

We used a conservation genomics framework to inform the definition of conservation units for anadromous Atlantic salmon in North America. We first analyzed the most extensive microsatellite dataset generated to date for the species to provide a North American-wide assessment of population structure based on neutral markers. This analysis confirmed with confidence previous regional assessments of population structure while also providing

interesting nuances. Contrary to expectations, however, the use of SNPs (and especially outlier SNPs) did not substantially change the overall population structure identified. This result contrasts with several previous studies, and we discuss technical and biological factors that may explain these discrepancies. We argue that such patterns may be common and that putatively adaptive markers identified from genome scans are unlikely to provide a practical alternative to the use of ecological or phenotypic data when defining conservation units. The SNP dataset, on the other hand, did provide greatly increased assignment success compared to microsatellites, achieving assignment success of up to 100% at the regional level and more than 90% at the population level. We discuss the utility of these tools in management, as well as strategies to achieve optimal assignment power with minimal costs.

Population structure: microsatellites vs. neutral SNPs

The re-analysis of North American-wide population structure based on microsatellites did not substantially differ from previous assessments that only considered parts of the range (McConnell *et al.* 1997; Spidle *et al.* 2001; Dionne *et al.* 2008; Bourret *et al.* 2013a; Bradbury *et al.* 2014). Contrary to the analysis of Dionne *et al.* (2008) for the Québec populations, our initial analysis on the entire dataset clustered all St. Lawrence North Shore sampling locations together, and grouped the Anticosti locations with the other Gaspésie locations. The differences, however, were resolved when the hierarchical population structure analysis was conducted and unambiguously recovered these groupings, as well as when SNP loci were used for clustering (see below). Multiple levels of hierarchical population structure is common in anadromous Atlantic salmon (Vähä *et al.* 2007; Perrier *et al.* 2011). Note also that the three populations from Anticosti Island are less genetically differentiated than other comparably distant populations, adding confidence in the reality of this genetic grouping. Our analysis was also consistent with that of Bradbury *et al.* (2014) and Palstra *et al.* (2007) in showing that the Labrador and Newfoundland populations were distinct, and that within Newfoundland, the Avalon Peninsula sampling locations were also distinct. Several hypotheses were suggested by Palstra *et al.* (2007) to explain the distinctiveness of the Avalon Peninsula samples, including smaller population sizes, differences in run timing, and increased prevalence of mature male parr (Myers *et al.* 1986). Both studies, however, found evidence for further genetic regional groupings within Newfoundland (Palstra *et al.* 2007; Bradbury *et al.* 2014), something our microsatellite analysis, including the

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hierarchical population structure analysis, failed to uncover. Another notable result is the increased prevalence of singletons in Newfoundland from the BAPS analysis, which appears consistent with previous conclusions that drift was one of the most important forces structuring populations in Newfoundland (Bradbury *et al.* 2014). Our analysis was also consistent with previous studies in suggesting the Inner Bay of Fundy populations were evolutionarily distinct, most likely due to a different re-colonization history following the last glaciation (McConnell *et al.* 1997; Verspoor *et al.* 2002; Vandersteen Tymchuk *et al.* 2010). While the populations did not cluster together in the BAPS analysis, they were clearly related in the NJ tree (Fig. 2). The BAPS analysis, in fact, suggested that two clusters were present, a result consistent with previous assessments that found differentiation between Chignecto Bay (NS-PWF and NS-BSR) and Minas Basin (NS-GRV and NS-STW) populations (Verspoor *et al.* 2002; Vandersteen Tymchuk *et al.* 2010). Other sampling locations clustered as singletons, which was perhaps explained by high drift due to small population sizes. Effective numbers of breeders of 12 anadromous Atlantic salmon populations from Québec have been found to vary between 48 and 876 (mean = 252) (Perrier *et al.* 2013). If typical, such small effective sizes could result in important genetic drift. The Prince Edward Island (PEI) rivers mainly clustered with the other Gulf of St. Lawrence populations, which is expected given that many of these rivers have been stocked heavily with Miramichi River fish (Cairns *et al.* 2010). The fact that two rivers clustered apart in the BAPS analysis, however, suggest that some locations may retain ancestral allele frequencies and that PEI may have constituted a different genetic group prior to stocking. Finally, our analysis included only a single population from the US, but its clustering in a group by itself was consistent with previous results suggesting US populations (including the Penobscot River) were genetically differentiated from Canadian populations (Spidle *et al.* 2003).

In short, the analysis of the microsatellite dataset supported the definition of twelve genetic regional groups for anadromous Atlantic salmon in North America: (1) upper North Shore of St. Lawrence River, (2) lower North Shore of St. Lawrence River, (3) Gaspésie Peninsula, (4) Anticosti Island, (5) Gulf of St. Lawrence (i.e. New Brunswick, Nova Scotia, Prince Edward Island), (6) Nova Scotia East Coast, (7) Newfoundland (excluding Avalon Peninsula), (8) Labrador, (9) Bay of Ungava, (10) Inner Bay of Fundy, (11) Avalon Peninsula

and Southeast Newfoundland, (12) United States (Penobscot River) (Fig. 1). Detailed justifications for these conclusions can be found in the online appendix. These genetic regional groups differ in some notable ways from the designatable units (DUs) currently recognized for Atlantic salmon in Canada by COSEWIC (2011). There are currently four recognized DUs in Newfoundland, but the microsatellite analysis suggested only two conservation units for the island (including the Avalon Peninsula, which was not previously recognized as distinct from the other south east Newfoundland populations). The Gaspé Peninsula and the Gulf populations were previously grouped in a single DU, but our analyses suggested that they are in fact distinct. The eastern Cape Breton Island populations were recognized as a DU, but our analysis suggested that they are part of the Gulf of St. Lawrence conservation unit. Three DUs were recognized on the St. Lawrence North Shore, which is partly consistent with the recognition of a separate group formed of a pair of populations at the boundary between our upper North Shore and Labrador groups. We concluded that these populations should be grouped with the Labrador populations, but we recognize that this grouping is somewhat ambiguous. Note that some of these DUs were defined on the basis of ecological or demographic information and may therefore represent useful conservation units despite the lack of concordance with the genetic data.

The BAPS analysis on the SNP datasets containing all loci and only neutral loci only differed in a minor way from the analysis on microsatellites. The SNP datasets uncovered all the geographical clusters identified with the microsatellites, including the sub-regional groupings in the St. Lawrence North Shore and Gaspésie/Anticosti that required hierarchical analyses to be evident with the microsatellite dataset. Main differences were the identification of a new grouping along the North Shore close to the Labrador grouping, as identified by Dionne *et al.* (2008) (therein called 'Higher North Shore'), and a grouping on the Northern Peninsula sampling locations in Newfoundland. This latter grouping was consistent with geography, and to a certain extent with the analysis of Bradbury *et al.* (2014), where the Northern Peninsula rivers appeared distinct in a NJ tree, but less so in a STRUCTURE analysis.

Defining conservation units with adaptive genetic variation

Analysis of population structure on the outlier loci from the Bayescan (n=106) and Fdist (n=61) analyses led to roughly the same results as that conducted on the putatively neutral loci (n=1673). This contrasted with previous results obtained from other species where putatively adaptive markers uncovered population structure at a finer scale (Hohenlohe *et al.* 2010; André *et al.* 2011; Bradbury *et al.* 2013; Hemmer-Hansen *et al.* 2013; Hess *et al.* 2013). The failure of putatively adaptive markers to uncover further levels of population structure in North American anadromous Atlantic salmon may be due to both technical and biological factors. First, studies have shown that results of genome scans should be interpreted with caution and that false positives are likely to be common (Narum & Hess 2011; Mita *et al.* 2013), especially for species that recently recolonized their range and are thus violating assumptions of equilibrium (Lotterhos & Whitlock 2014). If false positives were a random subset of the entire pool of loci, they would not be expected to lead to differences in population structure compared to neutral markers. The fact that we found limited overlap between the identities of outlier loci generated with Bayescan and hierarchical Fdist provided some support for this explanation. Second, outlier loci could be associated with several environmental variables that vary along different geographical axes. Their joint consideration as outlier loci (instead of considering alleles that are associated with specific environmental variables) would then blur potential population structure along a single axis of variation. This could be especially important for North American Atlantic salmon given the complexity of the coastline. Third, there is evidence of potentially polygenic adaptive traits in Atlantic salmon which are underlain by many genes of small effects (Bourret *et al.* 2014), thus making them difficult to identify with a traditional genome scan approach. The even distribution of outlier loci identified in the genome scan analyses presented here lends support to this explanation. This contrasted with cases from marine fishes where genomic islands of divergence were identified (Hemmer-Hansen *et al.* 2013). Fourth, neutral markers could have been sufficient to represent the scale of local adaptation even if they are not themselves under selection. Previous work on Atlantic salmon concluded that regional groupings identified with neutral markers were associated with environmental differences (Dionne *et al.* 2007; Bourret *et al.* 2013a; Bradbury *et al.* 2014). This could lead to selection against migrants from different regional groups (Nosil *et al.* 2005). It has also been proposed that straying is more likely to occur among rivers with similar characteristics, for example among rivers with similar chemical cues used for

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olfaction during homing (Dittman & Quinn 1996; Perrier *et al.* 2011; Bourret *et al.* 2013a). It is possible that selection effectively limits gene flow among regional groups, therefore resulting in similar patterns of allelic variation at both neutral and adaptive markers (Thibert-Plante & Hendry 2010). Finally, many of the studies that found drastically different population structure with outlier vs. neutral markers focused on species with high gene flow and large population sizes (i.e. low genetic drift) such as marine fishes (André *et al.* 2011; Bradbury *et al.* 2013; Hemmer-Hansen *et al.* 2013). In such cases, alleles conferring an adaptive advantage to certain environmental conditions would be expected to (1) increase in frequency rapidly, and (2) spread rapidly to all parts of the range experiencing these conditions. In the case of Atlantic salmon, however, low gene flow and small population sizes (i.e., high drift) could result in much more idiosyncratic patterns of adaptation at the molecular level, perhaps making genome scan approaches less likely to identify the loci underlying adaptation. Since many species of conservation concern are likely to be species with small population size and restricted gene flow (e.g., through habitat fragmentation), this could be an important factor to consider for a broad range of species.

In summary, the use of SNPs for delineating conservation units did not change substantially the conservation units defined on the basis of the microsatellite data. This is likely to be a common feature of such data sets and genome scans may not offer the hoped-for solution to the problem of incorporating adaptive information in the definition of conservation units. This also means that the framework proposed by Funk *et al.* (2012) may only be applicable in a minority of cases, unless better methods for identifying adaptive variation from genomic data are developed. In the meantime, the use of genomic data in conjunction with other types of data may offer more potential. Future work could, for instance, identify outlier loci from correlations with environmental data (e.g. Bourret *et al.* 2013) instead of from genome scans to alleviate some of the problems discussed above. Alternatively, markers under selection and involved in adaptation may often correspond more to markers that covary but involve only subtle differences in allele frequencies and are thus not detectable by genome scans (Le Corre & Kremer 2012; Bourret *et al.* 2014).

Individual assignments: SNPs vs microsatellites

An important management and conservation concern with Atlantic salmon is the operation of mixed-stock fisheries that target migratory individuals on their feeding grounds far from their natal rivers (Gauthier-Ouellet *et al.* 2009; Chaput 2012). Determining the origin of harvested salmon is therefore an important management priority. There are two main approaches utilizing genetic tools that can be used by managers to address this question: genetic mixture analysis and assignment tests (Manel *et al.* 2005). Bradbury *et al.* (in press) have already evaluated the power of the microsatellite database in a mixed-stock fishery context and found an average mixture analysis accuracy of 97.2% at the regional level. In some cases (e.g., management actions required at the scale of a single river, forensics), however, a mixture approach evaluating the proportional representation of regional groups may be insufficient. We therefore focused on the problem of assigning individuals to their region and river of origin. We were aided in this by intensive sampling of all possible regions of origin of Atlantic salmon in North America.

The use of a large number of SNP loci greatly increased assignment success in our study compared to microsatellites. Because microsatellites have far more alleles per locus than SNPs, previous studies based on a small number of SNPs rarely outperformed microsatellites in the context of population or parental assignment (Narum *et al.* 2008). Our use of >3,000 SNP loci definitively showed that numerous SNPs could outperform microsatellites, probably a result of both an increase in the number of independent loci and in the total number of alleles (Bernatchez & Duchesne 2000). Very few studies have used large SNP datasets in assignment tests. In one example, Larson *et al.* (2014) were able to assign individual Chinook salmon to their river of origin with 91% to 100% success using 10,944 RAD-generated SNPs. The study, however, only included five populations from a restricted portion of the Chinook salmon's range. In the present study, we demonstrate comparable success rate in assignment with a far greater number of populations distributed throughout the geographical range of Atlantic salmon in North America.

Despite high assignment success with the entire SNP dataset, assignment success based on outlier SNPs was only marginally better than that based on a similar number of randomly selected SNPs. At first glance, this result contrasted with several existing studies that have demonstrated increased assignment success with outlier SNPs compared to neutral markers (O'Malley *et al.* 2007; Ackerman *et al.* 2011; Freamo *et al.* 2011; Russello *et*

al. 2011). Most of these studies, however, used a small number of loci (< 100). For instance, Freamo *et al.* (2011) found that 14 outlier SNPs were more accurate than 67 neutral SNPs at assigning Bay of Fundy Atlantic salmon to their group of origin (outer vs. inner bay of Fundy). At a small number of loci, we also find that using outlier SNPs could perhaps improve assignment success: assignment to the population was ~20% higher when using the 61 Fdist outlier SNPs than when using a random subset of 50 SNPs. Few studies using a comparable number of SNPs for assignment exist to compare our results. Glover *et al.* (2010) used 300 SNPs in Atlantic salmon and found that the use of outlier SNPs did not result in great increases in assignment success when the overall number of SNPs used was > 100. Although it has been shown that some methods used to select markers for population assignment may result in upwardly biased estimates (Anderson 2010), the method we used here seems unlikely to play a major role in determining assignment success. Indeed, the fact that using outliers obtained from different genome scan methods led to similarly marginal increase in assignment success over random loci suggested that it had little bearing on the end result. Other methods exist for selecting markers that provide increased assignment power. For instance, software BELS (Bromaghin 2008) appeared to select loci that increased assignment success in Atlantic salmon, but performed poorly with more than 100 markers (Glover *et al.* 2010).

In summary, managers wishing to decrease costs by selecting a small subset of SNPs can carefully select highly differentiated markers to increase assignment success relative to a comparatively sized random subset of SNPs, but they are unlikely to attain the assignment success possible with a large number of SNPs. The use of the present medium-density SNP chip for assignment of a large number of individuals will, however, entail significant financial costs. If the goal is to assign individuals to their river of origin, our analysis suggest that a large number of SNPs is required (>1,000), and we suggest that using low-cost genotyping by sequencing technology (Davey *et al.* 2011) is probably the better strategy to lower costs. If the goal is to assign individuals to their region of origin, our analysis showed that using ~100 carefully selected SNPs allows assignment success of >90%. Investing in the development of a lower-density SNP array, such as the Fluidigm array used extensively in Pacific salmon (Ackerman *et al.* 2011; Larson *et al.* 2013), could therefore be more advantageous in this case.

Conclusions

The combination of a microsatellite database with high geographic resolution with data from thousands of SNPs allowed us to perform the most complete assessment of population structure of anadromous Atlantic salmon in North America to date. While the results were largely consistent with previous regional assessments, some important differences emerged from the range-wide analysis. Notably, our analysis suggested that some currently recognized conservation units did not coincide with genetic boundaries, something managers will want to take into consideration in future conservation efforts. Contrary to expectations, the definition of conservation units on the basis of microsatellites and SNPs, even those putatively under selection, did not differ all that much. This suggested that microsatellites are still a useful tool for defining conservation units, and that neutral markers may satisfyingly reflect the scale of local adaptation in anadromous salmon even if they are not under selection (e.g., via selection against migrants). Nevertheless, there are still several advantages to using a large number of SNPs. First, SNPs provide a more repeatable platform when genotyping is done across laboratories (standardization across laboratories for the present article required large efforts; Table S2 in online appendix). Second, they provided increased confidence in the conservation units defined on the basis of microsatellites, and were able to recover regional groupings that required hierarchical analyses to identify on the basis of microsatellites alone. Third, the equivalency of the outlier SNPs and the microsatellites for defining conservation units may stem from methodological issues related to the identification of putatively selected SNPs on the basis of genome scans alone. Future work may benefit from using alternative ways of identifying loci under selection, for example with methods that directly correlate allelic frequencies with specific environmental variables (Bourret *et al.* 2013a). There was no doubt, however, that using thousands of SNP loci for assignment of individuals to their population of origin provided greater assignment success compared to analyses based on microsatellites. The geographically extensive microsatellite and SNP databases presented here will therefore provide a diverse and powerful set of tools for managers attempting to achieve various research and management goals.

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Figure captions

Figure 1.

North American-wide population structure of anadromous Atlantic salmon showing the 149 sampling locations analyzed using the microsatellite dataset. Locations correspond to the mouths of sampled rivers; in cases where locations are inland, they correspond to a tributary where it intersects the main-stem. The sampling locations are colored according to the results of the BAPS clustering analysis. BAPS clusters containing only one sampling location are denoted by asterisks (*). Two of the BAPS groupings were further divided based on the hierarchical population structure analysis and are shown with different symbols (triangles and circles). Major regional groups discussed in the text are labeled: (1) upper North Shore of St. Lawrence River, (2) lower North Shore of St. Lawrence River, (3) Gaspésie Peninsula, (4) Anticosti Island, (5) Gulf of St. Lawrence (including Prince Edward Island (P.E.I.), see text for details), (6) Nova Scotia East Coast, (7) Newfoundland (excluding Avalon Peninsula), (8) Labrador, (9) Bay of Ungava, (10) Inner Bay of Fundy (comprised of two BAPS clusters; see text for details), (11) Avalon Peninsula, (12) United States (only one

sample location included in the analysis). Further details on the regional group designations can be found in the online appendix.

Figure 2. Un-rooted neighbor-joining (NJ) tree of Cavalli-Sforza Chord distances obtained using software Phylip (note that the Moisie River population is not included because it is missing data for all individuals sampled at three loci, something Phylip does not allow). Groups of populations belonging to the same regional cluster as identified in BAPS are highlighted with the same colors as in Fig. 1. Populations identified as singletons in BAPS are still denoted with the asterisk. Bootstrap values (1,000 replicates) above 50% are shown. The stars denote the only geographically inconsistent relationships (GU-KC6 and NL-BVB; see text for details). Black dotted lines group sampling locations that form regional groups on the basis of geographical proximity and genetic similarity but that did not group together in the BAPS analysis. Colored dotted lines group sampling locations that were found to form distinct clusters in the hierarchical analysis of population structure: Anticosti and the two sub-regional groups within the St. Lawrence North Shore regional group.

Figure 3. Genome scan results for the 2,607 SNP loci genotyped in this study for which genetic map position is available (out of a total of 3,192). (a) Among-region F_{ST} per locus based on the Bayescan analysis. (b) Among region F_{CT} per locus based on the hierarchical Fdist analysis. In both panels, black circles are putatively neutral loci (or loci putatively under balancing selection), red circles are loci identified as outliers by each analysis alone, and blue circles are loci identified as outliers in both analyses. Alternating white and shaded background shows boundaries between linkage groups.

Figure 4. North American-wide population structure of Atlantic salmon in 50 sampling locations inferred using BAPS from four different SNP datasets including: (a) all SNP loci retained after filters, (b) the putatively neutral markers from the Bayescan analysis, (c) the putatively adaptive loci from the Bayescan analysis, and (d) the putatively adaptive loci from the hierarchical Fdist analysis. The colors and shapes of the symbols denote the different BAPS-determined genetic clusters (we used the same color-coding as in figure 1 and used triangles for new clusters to highlight differences with the microsatellite-based analysis). Asterisks denote clusters that contain only a single sampling location (i.e., singletons). The number of clusters inferred from the BAPS analysis is indicated in the upper-right corner of each panel as well as the number of singleton populations.

Figure 5. Dot plots showing the results of the assignment tests performed in GENODIVE (top) and ONCOR (bottom) for different sets of markers. The x-axis represents the proportion of individuals that has the highest likelihood of being assigned to the sampling location where they were sampled. Assignment tests were performed at the population level (filled black symbols), and the regional level (filled grey symbols) and at a finer scale of regional differentiation (hollow symbols; see methods for details). Assignment tests were performed using the entire microsatellite data set (“all pops”), a subset of the microsatellite dataset including only populations for which we also had SNP data (“SNP pops”), a dataset including all the available SNP markers (“all loci”; $n = 3,192$), the Bayescan outliers ($n = 106$), the hierarchical Fdist outliers ($n = 61$), and the nested Fdist outliers ($n = 293$).

Figure 6. Effect of the number of randomly selected SNP loci on assignment success at the population (small black circles) and regional levels (grey dots). Assignment success for the three sets of outlier loci previously used are also shown for comparison, with the full symbols representing assignment to the population, and the hollow symbols representing assignment to the regional: Fdist outliers (blue, $n = 61$); Bayescan outliers (green, $n = 106$), nested Fdist outliers (red, $n = 293$). Michaelis-Menten curves were fitted to the data from randomly selected markers to help visualization.

Author contributions

J.-S.M. performed most of the analyses and wrote the manuscript. V.B. coordinated the sample collection and microsatellite genotyping for the database, including the standardization across laboratories, and participated in data analysis and writing. P.O. also helped coordinate sample collection and provided samples and help with the writing. M.K. coordinated the SNP genotyping. M.D., I.B., and G.C. provided samples and help with the writing. L.B. designed and supervised the study and was involved in the analysis and writing phases.

Data accessibility

The raw microsatellite and SNP genotypes (GENEPOP format) have been deposited in Dryad: doi:10.5061/dryad.sb601. The different subsets of outlier and neutral SNPs (also in GENEPOP format), the NJ tree file and the PHYLIP infile, the Bayescan parameter file, an

example STRUCTURE param file from the hierarchical structure analysis, the genetic map data for the loci, and the code to generate the random loci in figure 6 are also accessible on Dryad.

Supporting information

Table S1. Sampling locations included in the microsatellite database, along with information on the year and life stages sampled, and summary statistics for the microsatellite data.

Table S2. Standardization rules for comparison of microsatellite genotype data collected by the three laboratories and within Laval University.

Table S3. Pairwise F_{ST} values among all populations included in the microsatellite database.

Table S4. Sampling locations included in the SNP database, along with summary statistics for the SNP data.

Table S5. Pairwise F_{ST} values among all populations included in the SNP database.

Figure S1. Maps showing the approximate locations of the mouths of the rivers where samples of Atlantic salmon were collected for the microsatellite database.

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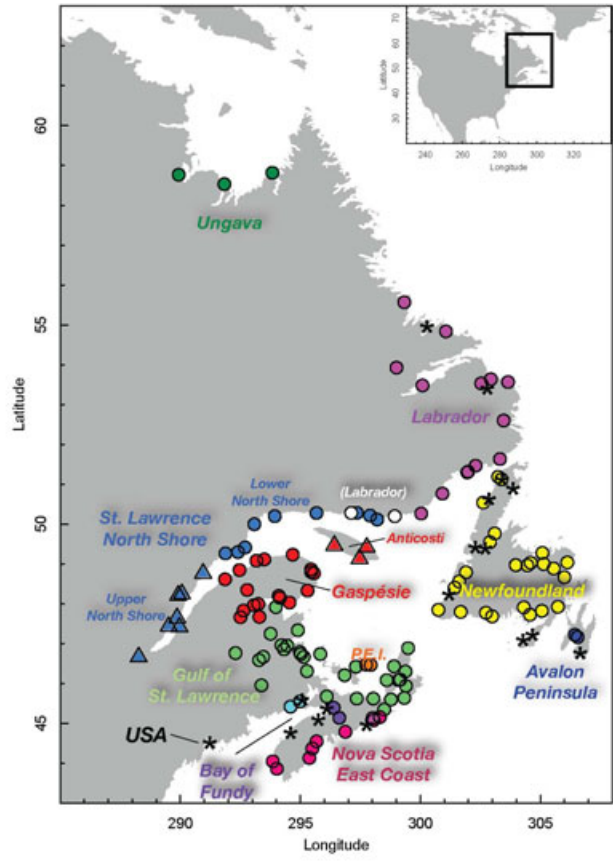


Figure 1

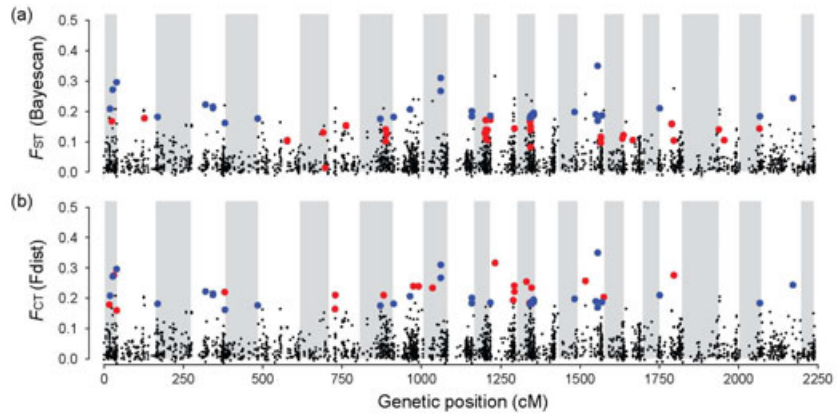


Figure 3

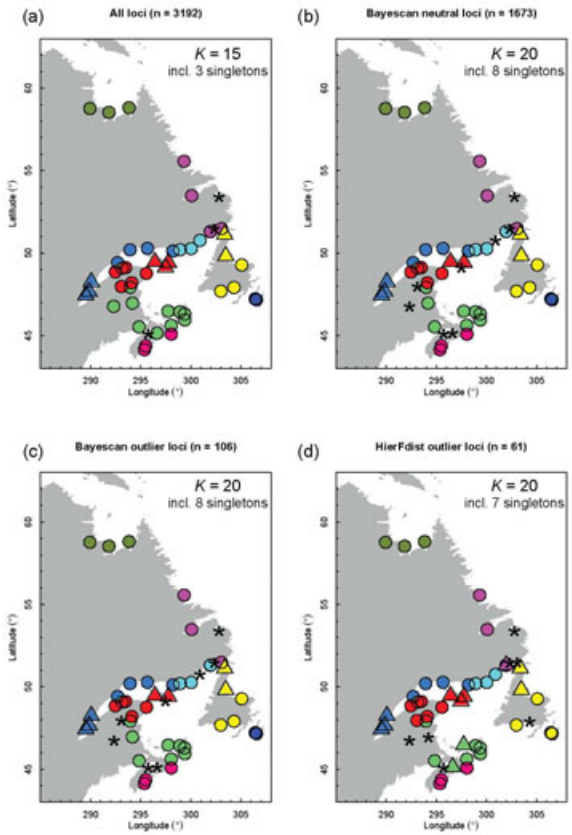


Figure 4

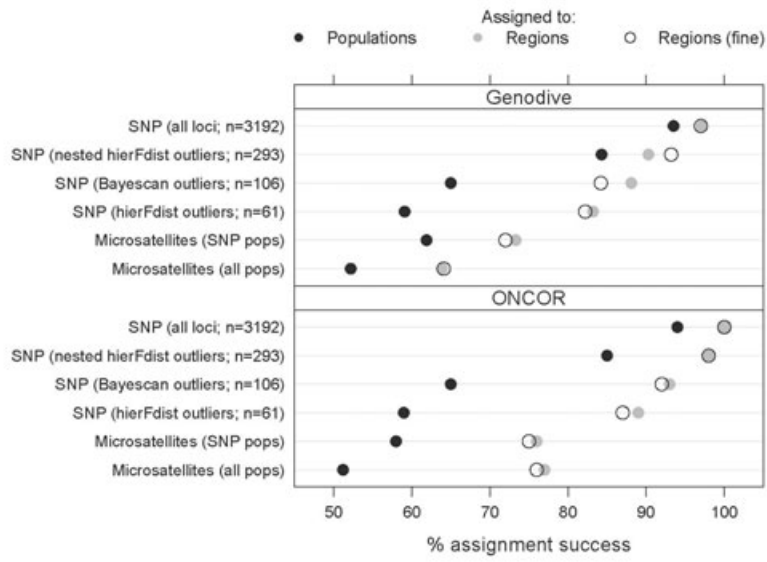


Figure 5

