Extensive genetic diversity and rapid population differentiation during blooms of *Alexandrium fundyense* (Dinophyceae) in an isolated salt pond on Cape Cod, MA, USA

Mindy L. Richlen¹, Deana L. Erdner², Linda A. R. McCauley¹, Katie Libera¹ & Donald M. Anderson¹

¹Woods Hole Oceanographic Institution, Woods Hole, Massachusetts, 02543
²Marine Science Institute, University of Texas at Austin, Port Aransas, Texas, 78373

Keywords
*Alexandrium*, Amoebophrya, dinoflagellate, Gulf of Maine, microsatellites, Nauset Marsh, paralytic shellfish poisoning.

Abstract
In Massachusetts, paralytic shellfish poisoning (PSP) is annually recurrent along the coastline, including within several small embayments on Cape Cod. One such system, the Nauset Marsh System (NMS), supports extensive marshes and a thriving shellfishing industry. Over the last decade, PSP in the NMS has grown significantly worse; however, the origins and dynamics of the toxic *Alexandrium fundyense* (Balech) populations that bloom within the NMS are not well known. This study examined a collection of 412 strains isolated from the NMS and the Gulf of Maine (GOM) in 2006–2007 to investigate the genetic characteristics of localized blooms and assess connectivity with coastal populations. Comparisons of genetic differentiation showed that *A. fundyense* blooms in the NMS exhibited extensive clonal diversity and were genetically distinct from populations in the GOM. In both project years, genetic differentiation was observed among temporal samples collected from the NMS, sometimes occurring on the order of approximately 7 days. The underlying reasons for temporal differentiation are unknown, but may be due, in part, to life-cycle characteristics unique to the populations in shallow embayments, or possibly driven by selection from parasitism and zooplankton grazing; these results highlight the need to investigate the role of selective forces in the genetic dynamics of bloom populations. The small geographic scale and limited connectivity of NMS salt ponds provide a novel system for investigating regulators of blooms, as well as the influence of selective forces on population structure, all of which are otherwise difficult or impossible to study in the adjacent open-coastal waters or within larger estuaries.

Introduction
The harmful algal bloom (HAB) syndrome known as Paralytic Shellfish Poisoning (PSP) is a significant and growing problem worldwide, with negative and sometimes devastating impacts on human consumers of contaminated seafood and local economies dependent on the shellfish industry. Paralytic Shellfish Poisoning outbreaks have also been linked to seabird and marine mammal mortalities, thus demonstrating the potential for serious ecosystem impacts (Anderson and White 1992; Shumway et al. 2003). Dinoflagellates within the genus *Alexandrium*, and in particular within the “tamarense” (*A. tamarense*, *A. fundyense*, and *A. catenella*) species complex (Fig 1), are responsible for many of these PSP outbreaks (Cembella 1998). In the US, PSP is prevalent along much of the west coast and more recently has also emerged as a major HAB problem in the northeastern Atlantic, particularly in...
the Gulf of Maine (GOM). Annual blooms of *A. fundyense* now impact most coastal areas of the northeastern United States and Canada, prompting the establishment of shellfish monitoring programs throughout the region.

In Massachusetts, PSP is annually recurrent along the coastline, including within a number of small embayments and salt ponds on Cape Cod. These embayments are characterized by restricted circulation and limited tidal flushing, receiving freshwater from groundwater flow and saltwater through one or more inlets to the sea. One such system on outer Cape Cod, the Nauset Marsh System (NMS), supports extensive marshes and thriving commercial and recreational shellfishing industries. This economically and socially important system also has a long history of PSP outbreaks associated with annual episodes of *Alexandrium* blooms, which are initiated and most intense in the landward extensions of the estuary.

Over the last several decades, the PSP problem in the NMS has worsened significantly, both in terms of intensity and duration. Anecdotal reports also suggest that PSP is occurring earlier and lasting longer than in the past. There are several explanations for these trends, including the increased retention and proliferation of toxic cells and dormant resting cysts in areas that are not well flushed in the system, as well as population stimulation by eutrophication pressures linked to the rapid development of the watershed through the years. Additionally, two major storms that occurred in 1991 may have altered flushing patterns and dispersed cysts within the NMS, thereby augmenting the magnitude of subsequent *A. fundyense* blooms.

Outbreaks in southern New England (i.e., south of Massachusetts Bay) are generally confined to salt ponds and embayments, suggestive of “point sources” of cells originating from locally germinated cysts (Anderson et al. 1983; Anderson and Stolzenbach 1985; Hattenrath et al. 2010). Although salt ponds in the NMS show strong characteristics of point-source *A. fundyense* habitats, it is clear that this estuary is in a transition area between the localized, estuarine populations of southern New England and the widespread, open-coastal populations of the GOM (e.g., Anderson et al. 2005a); thus, the system can thus potentially receive cells, and/or export them into the coastal flow. The implication is that the NMS may contain populations that originate in situ, as well as populations that have been introduced through Nauset Inlet from adjacent coastal waters during times of flood tide and downwelling-favorable winds. This exchange may also occur in the opposite direction as well – populations may be advected out of the system, into the coastal flow, thus mixing with southern GOM populations and affecting downstream shellfish resources.

This study investigated the genetic composition of *Alexandrium* blooms in the NMS to elucidate genetic linkages with coastal populations and to help determine the extent to which *Alexandrium* blooms within the salt ponds of the NMS are initiated solely within the system, and the degree to which cells from blooms are transported in and out of the NMS. In addition to examining the degree of genetic differentiation and connectivity between bloom populations in NMS and the GOM, we also hoped to gain insight into the potential effects of restricted geography on...
the genetic composition and diversity of populations within shallow, isolated embayments. Due to the extremely small geographic scale and limited connectivity with coastal populations in the GOM, this estuary provides a novel system for studying the genetic diversity and dynamics of *Alexandrium* blooms, as they appear to be largely restricted from genetic exchange with regional populations. These characteristics also make this study site an effective “natural laboratory” for the investigation of many important aspects of *Alexandrium* autoecology affecting bloom development and termination, such as life-cycle events as well as the influence of selective forces on population dynamics, all of which are otherwise difficult or impossible to study in the adjacent open-coastal waters, larger estuarine systems, or mesocosms.

To assess fine-scale spatial and temporal genetic diversity of *A. fundyense* populations, we employed rapidly evolving microsatellite markers. The ability of these markers to detect fine-scale variability among and sometimes within populations has provided researchers with an effective tool for exploring a diversity of questions regarding the population dynamics and molecular ecology of phytoplankton, at a resolution impossible to achieve with DNA sequencing. Microsatellites have been used in investigations of genetic differentiation across global and broad regional scales (Nagai et al. 2007, 2009; McCauley et al. 2009), to examine the genetic diversity and structure of bloom populations (Evans et al. 2005; Rynearson and Armbrust 2005; Rynearson et al. 2006) including the potential detection of cryptic species (Adams et al. 2009), infer the geographic origins of populations (Masseret et al. 2009), and to explore questions regarding life-cycle effects and phenotypic variability on population genetic structure (Iglesias-Rodriguez et al. 2006; Alpermann et al. 2009). Here, we used microsatellite markers to examine the genetic diversity and population structure of *Alexandrium* blooms that appear to be largely localized within the NMS. The study objectives were to: (1) characterize potential linkages among *Alexandrium* blooms within the NMS and with those in the wider GOM; (2) examine the effects of geographic restriction on the genetic diversity and population structure of cells within Salt Pond, where bloom residence times are high; and (3) examine population composition throughout a natural bloom.

**Methods**

**Study site**

The NMS is a coastal embayment of the Cape Cod National Seashore and is protected from the Atlantic by a rapidly changing barrier beach with a single connection through Nauset inlet (Fig 2). Its extensive marshes support thriving commercial and recreational shellfishing industries where quahogs, soft-shell clams, and razor clams are harvested along nearly all stretches of the shoreline and tidal flats. Terminal kettle ponds within the NMS include Salt Pond, Mill Pond, and Town Cove. Salt Pond is a circular, shallow embayment, approximately 82,200 m² in area, with average and maximum depths of approximately 3.4 and 7 m, respectively, at slack low tide. There are no stream or river inputs of freshwater, but groundwater does enter the pond via springs, which keeps the pond more brackish than the incoming tidal flow.

Freshwater flux to Salt Pond and the shallow inlet channel (0.5 km long, 30 m wide, 0.5 m deep at low tide) allows tidal exchange with the wider marsh system that connects to the ocean at Nauset inlet. Tidal mixing between Salt Pond and the NMS is efficient, equivalent to a maximum potential flushing rate of 0.4–0.7 day⁻¹, and is primarily due to an estuarine-like circulation in which flood-tide waters sink and vertically displace less dense pond water, which is then removed with the subsequent ebb tide (Anderson and Stolzenbach 1985).

**Sampling**

Phytoplankton samples were collected from near surface waters of Salt Pond and Goose Hummock in the NMS during the bloom seasons of 2006–2007, and from Massachusetts Bay (2006) and Nantucket Shoals (2007) in the
GOM (Fig 2). Single *Alexandrium* cells (contained in >20 and <64 μm Nitex-sieved field samples) were manually isolated via micropipette into the wells of a flat bottom 96-well plate containing 250 μL of f/2 -Si medium (Guillard 1975; Anderson et al. 1994). All cultures were maintained at 15 or 20°C at 250 μmol photons m⁻² s⁻¹ on a 14:10 light:dark (L:D) cycle. A total of 11 samples were collected during the course of the study: six in 2006 and five in 2007 (Table 1). In 2006, samples were collected from Salt Pond (herein termed 06SPA-06SPD), Goose Hummock (06GH), and Massachusetts Bay (06GOM). The 2007 sample set comprised four Salt Pond populations (07SPA-07SPD) and one from Nantucket Shoals (07GOM).

### Microsatellite analysis and ribotyping

For DNA extraction, mini-cultures were allowed to grow until each achieved a cell density of >100 cells per well. DNA was extracted from 200 μL of each culture using the Generation Capture Column Kit (Qiagen, Valencia, CA) following the manufacturers instructions, with an elution volume of 200 μL.

Of the microsatellite loci isolated by Nagai et al. (2004), 11 of 13 loci were tested with the small set of cultures from the GOM, after which the following four loci were selected for this study, based upon the number of alleles observed and amplification success (see Erdner et al. 2011): Atama 15, Atama 23, Atama 27, and Atama 39. PCR reactions contained ~5 ng of template DNA, 0.2 mmol/L of each dNTP, 0.5 μmol/L of each designed primer pair, with one primer labeled with 6FAM, NED, PET, or VIC, 1× PCR buffer (10 mmol/L Tris-HCl, pH 8.3, 500 mmol/L KCl, 15 mmol/L MgCl₂, 0.01% w/v gelatin), and 0.25 U of Ampli Taq Gold (Applied Biosystems Inc., Foster City, CA) to a total volume of 10 μL. The PCR cycling conditions were as follows: 10 min at 94°C, 38 cycles of 30 sec at 94°C, 30 sec at 60°C for the first 10 cycles and then at the primer-specific annealing temperature (Nagai et al. 2004) for the last 28 cycles, and 1 min at 72°C, and a final elongation for 5 min at 72°C. To determine if amplification was successful, PCR products were separated on a 1% TAE agarose gel adjacent to a 100 bp DNA ladder. For fragment analysis, PCR products were diluted 3–5× with nuclease-free water, and 1 μL of diluted product was mixed with 0.25 μL 500 LIZ Size Standard and 8.75 μL Hi-Di Formamide, and then analyzed using an ABI 3730xl DNA Analyzer (Applied Biosystems, Foster City, California). Allele sizes were determined using the program FPMiner (2005; Bioinformatics, Beaverton, OR).

Numerous studies examining the global biogeographical distribution of the *A. tamarense* species complex have observed only the NA ribotype in the Northwestern Atlantic (Anderson et al. 1994, 2005b; Scholin et al. 1995; Lilly et al. 2007; Brosnahan et al. 2010; Crespo et al. 2011). However, to ascertain species identification of the populations examined in this study, a subset of the 11 sample sets representative of each water body and study year (06GOM, 06SPC, 07SPD) used in the microsatellite analysis were also amplified using PCR primers specific to the *A. fundyense* NA ribotype as described in Dyhrman et al. (2006).

### Microsatellite data analysis

Genetic diversity at each locus was measured by the number of alleles and gene (haplotype) diversity (Nei 1973) using POPGENE (1.32) (Ellstrand and Roose 1987; Yeh

---

**Table 1.** Information regarding the sampling date, microsatellite loci, allelic characteristics, and clonal diversity of *Alexandrium fundyense* populations from Salt Pond, Cape Cod, MA, USA. Data are shown for each population sampled, and combined datasets for each project year. *N* = number of individuals genotyped, G:N = clonal diversity (Ellstrand and Roose 1987), and probabilities of identity (*P*ᵦᵦ and *P*ᵦᵦᵦ) across all loci.

| Population ID | Sampling date | Number of alleles | Size range of alleles | G:N | Pᵦᵦ | Pᵦᵦᵦ |
|---------------|---------------|-------------------|-----------------------|-----|------|-------|
| 06SPA         | 4/26/06       | 29                | At15:11, At23:7, At27:15, At39:7 | 240–266 176–204 160–168 136–150 | 0.97 | 3.26E-05 | 2.72E-02 |
| 06SPB         | 5/03/06       | 36                | At15:11, At23:7, At27:15, At39:7 | 240–250 176–204 160–168 132–142 | 0.85 | 9.27E-05 | 3.35E-02 |
| 06SPC         | 5/10/06       | 30                | At15:11, At23:7, At27:15, At39:7 | 238–266 176–202 160–168 138–142 | 0.95 | 1.14E-04 | 3.48E-02 |
| 06SPD         | 5/17/06       | 17                | At15:11, At23:7, At27:15, At39:7 | 240–250 178–202 162–168 138–144 | 0.82 | 1.30E-04 | 5.05E-02 |
| 06GH          | 5/23/06       | 33                | At15:11, At23:7, At27:15, At39:7 | 238–252 176–200 160–168 136–150 | 0.94 | 1.43E-04 | 3.34E-02 |
| 06GOM         | 5/17/06       | 61                | At15:11, At23:7, At27:15, At39:7 | 224–267 174–182 160–164 138–144 | 0.64 | 3.16E-03 | 1.05E-01 |
| 07SPA         | 5/09/07       | 31                | At15:11, At23:7, At27:15, At39:7 | 238–265 176–202 162–170 138–144 | 0.97 | 8.76E-05 | 3.40E-02 |
| 07SPB         | 5/16/07       | 30                | At15:11, At23:7, At27:15, At39:7 | 240–250 178–208 154–166 136–146 | 0.83 | 5.10E-04 | 5.17E-02 |
| 07SPC         | 5/23/07       | 32                | At15:11, At23:7, At27:15, At39:7 | 238–250 176–202 160–172 132–148 | 0.81 | 3.14E-04 | 3.71E-02 |
| 07SPD         | 6/06/07       | 37                | At15:11, At23:7, At27:15, At39:7 | 238–251 176–202 160–164 136–144 | 0.92 | 1.82E-04 | 3.85E-02 |
| 07GOM         | 5/30, 5/31/07 | 77                | At15:11, At23:7, At27:15, At39:7 | 238–266 176–182 160–166 138–146 | 0.47 | 8.01E-03 | 1.17E-01 |
| GOM combined  |               | 138               | At15:11, At23:7, At27:15, At39:7 | 224–267 174–182 160–166 136–146 | 0.46 | 5.11E-03 | 1.09E-01 |
| NMS combined  |               | 274               | At15:11, At23:7, At27:15, At39:7 | 238–266 176–208 154–172 132–150 | 0.76 | 8.15E-05 | 2.70E-02 |
and Boyle 1997). Clonal diversity was calculated as the ratio of the number of unique four-locus genotypes (G) to the total number analyzed (N) (Ellstrand and Roose 1987). Additional tests to calculate the probability of identity (P(ID) and P(ID) unbiased) for each locus and over all loci were performed using GIMLET v1.3.3 (Valière 2002) to assess the resolving power of the four microsatellite loci used in this study. For these tests, haplotype genotype data were first converted to diploid format using the data conversion utility in GENEPOP (4.0) (Raymond and Rousset 1995; Rousset 2008). P(ID) unbiased (herein referred to as P(ID)) provides a measure of the probability of observing identical multilocus genotypes between two individuals randomly sampled from a population, with a sample size correction. P(ID) unbiased provides a more conservative measure and estimates match probabilities among sibs. According to guidelines provided by Waits et al. (2001), P(ID) values between 0.01 and 0.0001 indicate sufficient resolving power by the microsatellite markers.

Pairwise genic comparisons using Fisher’s exact test (G test) to assess genetic differentiation among samples, and tests for linkage disequilibrium were performed using GENEPOP (4.0) (Raymond and Rousset 1995; Rousset 2008) with Markov chain parameters of 2000 for dememorization number, 200 batches, and 2000 iterations per batch. Estimates of gene flow (FST) were measured using Arlequin v3.5.1.2 (Excoffier et al. 2005). Bonferroni-corrected permutation tests were used to determine the significance of statistical values of pairwise comparisons.

Genetic structuring and admixture were further investigated using Bayesian cluster analysis as implemented in STRUCTURE 2.3.2 (Pritchard et al. 2000). The number of potential clusters (K) was first assessed using 30 runs performed at each K value from 1 to 10, each with a burn-in period of 10,000 steps and 10,000 Markov chain Monte Carlo repetitions. The log likelihoods of these data were used to calculate the statistic ΔK, to infer the uppermost level of population structure (Evanno et al. 2005). Simulations were performed using the admixture model with and without the locprior option, assuming correlated allele frequencies among populations. Clustering among samples was also inferred using principal components analysis (PCA) of FST values, performed using PCA-GEN (1.2) (Goudet 1999). Significance of each axis was determined using 10,000 randomizations of genotypes.

Finally, partitioning of genetic variance within and among samples was examined using analysis of molecular variance (AMOVA), with distance measures based on the number of different alleles, as implemented in Arlequin v3.5.1.2. Tests were first performed without groups, using: (1) the complete 2006–2007 GOM/NMS sample set; (2) 2006 NMS samples; and (3) 2007 NMS samples. Two additional tests were then performed in which the samples were organized into population groupings. The first test compared two groups: the GOM samples and the NMS samples. The second test included NMS samples only, and assessed interannual variation between 2006 NMS samples and 2007 NMS samples.

Table 2. Extent of genetic differentiation among Alexandrium fundyense populations from the Nauset Marsh System and the Gulf of Maine (GOM) sampled in 2006–2007. Differentiation estimators FST are shown in the lower half of the matrix and Fisher’s combined test (F values) are shown in the upper half of the matrix. Bold numbers indicate significant values (P < 0.05) after sequential Bonferroni correction. H.S., highly significant.

| Sampling date | 06SPA | 06SPB | 06SPC | 06SPD | 06GHA | 06GOM | 07SPA | 07SPB | 07SPC | 07SPD | 07GOM |
|---------------|------|------|------|------|------|------|------|------|------|------|------|
| 06SPA         |      |      |      |      |      |      |      |      |      |      |      |
| 06SPB         | 0.1573 |      |      |      |      |      |      |      |      |      |      |
| 06SPC         | 0.1207 | 0.0967 |      |      |      |      |      |      |      |      |      |
| 06SPD         | 0.0040 | 0.0013 | 0.2209 |      |      |      |      |      |      |      |      |
| 06GHA         | 0.9749 | 0.0013 | 0.0091 | 0.0391 |      |      |      |      |      |      |      |
| 06GOM         | 0.0000 | 0.2209 | 0.0091 | 0.3091 | 0.6618 |      |      |      |      |      |      |
| 07SPA         | 0.8228 | 0.3623 | 0.0000 | 0.0000 | 0.5064 | 0.0000 |      |      |      |      |      |
| 07SPB         | 0.1185 | 0.3553 | 0.0000 | 0.0000 | 0.5064 | 0.0000 | 0.0902 |      |      |      |      |
| 07SPC         | 0.0735 | 0.0000 | 0.0000 | 0.0000 | 0.5064 | 0.0000 | 0.0000 | 0.0000 |      |      |      |
| 07SPD         | 0.6618 | 0.0000 | 0.0000 | 0.0000 | 0.5064 | 0.0000 | 0.0000 | 0.0000 | 0.0000 |      |      |
| 07GOM         | 0.1410 | 0.2984 | 0.0762 | 0.6618 | 0.6618 | 0.6618 | 0.6618 | 0.6618 | 0.6618 | 0.6618 |      |

Results

Molecular identification of A. fundyense populations in the GOM and NMS

To ascertain that the genetic separation we observed among samples reflected population dynamics and not species dynamics, we assessed of a subset of samples used in the microsatellite analyses with primers specific to the NA ribotype of A. fundyense. PCR amplification of GOM and NMS sample subsets using these primers specific to
the NA ribotype produced a single band of approximately 174 bp (Fig S1), confirming the species identification of these samples.

**Microsatellite data**

The microsatellite loci employed in these studies were developed by Nagai et al. (2004) to examine *A. tamarense* strains from Japanese and Korean coastal waters (Nagai et al. 2007), and have also been employed to examine the genetic structure and dynamics of populations in the GOM (Erdner et al. 2011). A total of 412 isolates were genotyped for this study, 205 in 2006 and 207 in 2007 (Table 1). All loci examined were moderately or highly polymorphic, Atama15 and Atama23 displayed maximum numbers of 22 and 15 alleles, respectively, whereas maximum allele numbers displayed by Atama16 and Atama23 were lower, 8 and 9, respectively. A total of 27 alleles were restricted to samples from either the NMS or GOM; most of these alleles (24/27) were unique to samples collected from the NMS. Gene diversity (Hₑ) at each allele was slightly lower than Nagai et al. (2004, 2007), but comparable to Erdner et al. (2011); with the exception of Atama39, allele numbers were higher in this study (Table S1). Linkage disequilibrium was not detected at any of the loci.

**Genetic diversity and marker resolution**

Clonal diversity of bloom populations in the NMS, calculated as the proportion of unique four-locus genotypes (G:N), was remarkably high in both project years (Table 1), ranging from 81% to 97% genetically unique cells (mean = 88 ± 6%), and was highest in the early bloom samples (97% unique genotypes, 2006 and 2007). Clonal diversity was also comparable between the combined NMS yearly datasets from 2006 versus 2007 (83% and 86%, respectively; data not shown) and slightly lower in the total combined dataset (76%). However, clonal diversity was much lower in the GOM populations. Samples collected from Massachusetts Bay in 2006 and from Nantucket Shoals in 2007 comprised 64% and 47% unique genotypes, respectively, and clonal diversity of the combined GOM dataset was 46%.

Probability of identity (P(ID)) values of the NMS samples were within the guidelines suggested by Waits et al. (2001), indicating that the four microsatellite markers had sufficient resolving power for identifying individuals. However, values of the more conservative P(ID)Sub values were slightly above the upper limit (Waits et al. 2001), ranging from 0.0272 to 0.0517 (combined dataset, 0.0272), thus limiting our ability to distinguish true clones. In contrast with the NMS samples, P(ID) and P(ID)Sub values of samples from the GOM (combined dataset) were 0.0051 and 0.109, respectively, and were comparable between years (Table 1). These values were higher than those in the NMS, and exceeded the upper limit of 0.01, indicating that more loci would provide greater resolving power (also see Erdner et al. 2011).

**Genetic differentiation among populations**

Pairwise comparisons using the Fisher’s combined test of genotypic variation (Raymond and Rousset 1995) and Wright’s Fometric estimator (Wright 1951) detected significant spatial and temporal genetic differentiation among *A. fundyense* blooms in different years, as well as between populations in the NMS and the GOM. In both project years, all populations sampled from the NMS were genetically distinct from those in the GOM; differentiation was statistically significant using both differentiation estimators, with F values indicating moderate to high differentiation among the populations sampled (Table 2).

Within the NMS, we observed population differentiation among several samples collected from Salt Pond over the course of the bloom season in 2006 and again in 2007. In the 2006 dataset, F values indicated significant genetic differentiation between the late bloom sample (06SPD), and those collected earlier in the bloom (Table 2). Results of the Fisher’s combined test of genetic differentiation were slightly different, in that significant differentiation was only detected between 06SPC and 06SPD, and between 06SPD and 06GHA. These differences arise in part to the reduced statistical power associated with the Bonferroni-corrected P-value for the multiple pairwise comparisons (55) in the sample set (Bonferroni’s adjustment, P = 0.00091). For example, when samples from 2006 were analyzed separately from the 2007 samples (10 pairwise comparisons), 06SPD was significantly different from all other early bloom populations (P = 0.005, data not shown). The genetic composition of the bloom in 2006 changed rapidly, as differences were detected between samples collected approximately 7 days apart. With the exception of 06SPD in the Fisher’s test, the sample collected from Goose Hummock (06GHA), located in an area of the NMS known as Town Cove, was not significantly different from the samples collected in nearby Salt Pond.

Again in 2007, we observed temporal genetic differentiation among samples collected from Salt Pond (Table 2). Pairwise comparisons using Fisher’s combined test indicated that the late bloom samples 07SPC and 07SPD were genetically distinct from 07SPB (P < 0.05), but not from 07SPA, which was collected during bloom initiation. However, comparisons using Wright’s F estimator revealed a slightly different pattern of genetic structure in which 07SPB was differentiated from 07SPC, but not from 07SPA. The F
value between 07SPB/07SPC was 0.074, indicating moderate genetic differentiation. In both analyses, the sample collected during bloom initiation, 07SPA, was not differentiated from samples collected subsequently during bloom development, or from the late bloom sample. As in 2006, changes in the genetic composition of _A. fundyense_ populations collected from Salt Pond in 2007 occurred rapidly; again, on the order of 6–7 days.

In addition to temporal differentiation observed during the bloom, comparisons of the 2006 Salt Pond samples to those collected in 2007 revealed significant differences between sampling years. Both Wright’s _F_ statistic and the Fisher’s combined test revealed significant differences between 07SPB and 06SPB-06SPD (Table 2). Significant differences were also detected between the late bloom sample 07SPD and 06SPB-06SPC. Notably, early bloom samples from both 2006 and 2007 were largely undifferentiated from mid- and late-bloom samples, both within and between years.

Results of the AMOVA analysis indicated that the highest proportion of genetic variation (88–100%) was within samples, regardless of the population groupings examined (Table 3). In the overall dataset (GOM and NMS), among-group variation was highest when GOM and NMS samples were grouped (9.4%), reflecting the genetic separation among samples from these water bodies. Among-sample genetic variation was lower, but still significantly different when the NMS samples were examined separately from GOM samples (2.3%), but not when these samples were grouped according to study year (2006 vs. 2007). An AMOVA was also performed on NMS samples from each study year. In 2006, differences in genetic variation were significant among samples, but not in 2007 (Table 3).

Relationships among samples were further examined using STRUCTURE analysis as well as PCA of pairwise _F_ values. STRUCTURE analysis plots of the _ΔK_ statistic as a function of _K_ showed maximum values at _K_ = 2 for the entire GOM and NMS datasets. In these simulations, samples were divided into two clusters (Fig 3A), comprised of GOM and NMS samples, respectively. GOM samples were assigned to the “blue” cluster, with a proportional membership of > 0.88. NMS samples were assigned to the “grey” cluster, and the proportional membership of these samples ranged from 0.66 (07SPB) to 0.96 (06SPD). In general, admixture was higher in the 2007 NMS samples than in the 2006 NMS samples. Examination of the NMS dataset separately showed maximum values of the _ΔK_ statistic as a function of _K_ at _K_ = 3 (Fig 3B). In this analysis, all samples were admixed, although allele proportions are notably different among samples collected in 2006. The 06SPB and 06SPC samples had the highest proportional membership to the “green” cluster (> 0.65), whereas 06SPD had the highest proportional membership to the “orange” cluster (> 0.75). In the 2007 samples, differences in allele proportions were again observed. 07SPB had the highest proportional membership to the “blue” cluster (0.65) and similar to 06SPB and 06SPC, 07SPC had the highest proportional membership to the “green” cluster (0.42).

The PCA separated the samples into groups (Fig 4): (1) 06GOM, 07GOM, (2) 06SPB, 06SPC, 07SPC, (3) 06SPD, and (4) the five remaining NMS samples. The first three axes of the PCA explained 63%, 15%, and 9% of the total variation in _F_ statistic, respectively. Variation was only significant along the first axis (_P_ < 0.05), reflecting the separation of samples into two groups (GOM and NMS) in the STRUCTURE analysis. Subtle population structure was also apparent among the NMS samples. The grouping of 06SPB, 06SPC, 07SPC appeared to reflect the differentiation between these three samples and the remaining NMS samples, although differences detected in the pairwise comparisons were only statistically significant between this group and 07SPB and 07SPD (Fisher’s exact test, _P_ < 0.05). This group also shared the highest proportional membership to the “green” cluster in the STRUCTURE analysis (Fig 3B). Comparisons of genetic differentiation and STRUCTURE analysis also supported the separation of 06SPD from the other NMS samples observed in the PCA.

### Table 3. Analysis of molecular variance within and among samples from the Gulf of Maine (GOM) and Nauset Marsh System (NMS).

| Analysis                        | df  | SSD      | Variance | Percent of variation |
|---------------------------------|-----|----------|----------|----------------------|
| Without groups (all samples)    |     |          |          |                      |
| Among samples                   | 10  | 46.063   | 0.09346  | 7.45*                |
| Within samples                  | 402 | 466.588  | 1.16067  | 92.55                |
| GOM versus NMS (two groups)     |     |          |          |                      |
| Among groups                    | 1   | 35.643   | 0.12328  | 9.37*                |
| Among samples                   | 9   | 20.420   | 0.03201  | 2.43*                |
| within groups                   |     |          |          |                      |
| Within samples                  | 402 | 466.588  | 1.16067  | 88.20*               |
| Without groups (NMS samples only)|    |          |          |                      |
| Among samples                   | 8   | 17.832   | 0.03039  | 2.28*                |
| Within samples                  | 266 | 346.829  | 1.30387  | 97.71                |
| 06NMS (five samples, no groups) |     |          |          |                      |
| Among samples                   | 4   | 10.727   | 0.0454   | 3.18*                |
| Within samples                  | 140 | 193.280  | 1.38057  | 96.82                |
| 07NMS (four samples, no groups) |     |          |          |                      |
| Among samples                   | 3   | 2.951    | -0.00706 | -0.59                |
| Within samples                  | 126 | 152.802  | 1.21272  | 100.59               |
| 06NMS versus 07NMS (two groups)|     |          |          |                      |
| Among groups                    | 1   | 3.993    | 0.01449  | 1.08                 |
| Among samples                   | 7   | 13.839   | 0.02224  | 1.66*                |
| within groups                   |     |          |          |                      |
| Within samples                  | 266 | 346.829  | 1.30387  | 97.26*               |

* _P_ < 0.05.
Discussion

The four microsatellite loci employed in this study exhibited a high degree of diversity, and proved to be effective markers for detecting genetic differentiation in *A. fundyense* blooms in the NMS and GOM. These loci are a subset of 11 developed for *A. tamarense* in Japan (Nagai et al. 2004), which belong to the same *Alexandrium* clade as *A. fundyense* in the northeastern Atlantic (=North American/Group 1 ribotype). In this study, all four loci were polymorphic, and gene diversity was moderate or high at all loci, comparable to Nagai et al. (2004). Interestingly, differences in their resolving power were observed between the two water bodies, and were reflected in the lower $P_{IID}$ values calculated for the NMS samples and higher levels of clonal diversity (Table 1).

Bloom populations in the NMS exhibited remarkably high levels of clonal diversity in both project years, ranging from 81% to 97% genetically unique cells (mean = 88 ± 6%). The discovery of such high levels of diversity in a geographically restricted area was unexpected, as the study area largely comprised a shallow, isolated salt pond <1000 m in diameter, and a nearby cove. In contrast with the NMS, diversity was lower in samples collected from the GOM: 64% in the 2006 sample collected from Massachusetts Bay and 47% in the 2007 sample collected from Nantucket Shoals. These levels are also lower than those documented previously by Erdner et al. (2011), which employed five loci to investigate GOM bloom populations in 2005. Clonal diversity of samples collected in 2005 ranged from 83% to 92%, thus providing further indication that four loci used in this study have less resolving power for the GOM populations.

One of the major objectives of this investigation was to determine the degree of connectivity between *A. fundyense* populations in the NMS and the GOM. Due to the restrictive nature of the Nauset inlet, which connects the terminal endpoints of the NMS to the open ocean through a complex maze of channels in the central marsh, *A. fundyense* in these endpoints appear to be contained within the estuary. Paralytic shellfish poisoning outbreaks in the NMS exhibit a pattern of localization and are generally confined to the terminal salt ponds and embayments, suggestive of “point sources” of cells originating from locally germinated cysts (Anderson et al. 1983; Anderson and Stolzenbach 1985). Paralytic shellfish poisoning on
Cape Cod also occurs much earlier than in the GOM; within the NMS, Salt Pond is one of two key locations in which toxicity first appears, the second being Mill Pond, a terminal endpoint in the southernmost area of the system (Anderson et al. 1982; Crespo et al. 2011; Fig 2). These two locations feature the lowest flushing rates of the entire system, as well as the highest residence time of water, estimated to be between 5 and 10 days, with the longer values for deeper water (D. Ralston, pers. comm.).

The vertical migration behavior of *A. fundyense* has been shown to be an important mechanism limiting advective cell loss from Salt Pond, and may be equally as important as the hydrodynamic characteristics of this location in promoting the selective retention of *Alexandrium*. Studies of diel migration patterns by Anderson and Stolzenbach (1985) showed that *A. fundyense* exhibits surface-avoiding vertical migration behavior whereby the cells typically stop swimming upwards when they reach a light level of \( \sim 150 \, \mu E \, m^{-2} \, sec^{-1} \), effectively separating them from the low salinity surface water lens that is flushed out of the salt pond with each ebb tide. This retention of cells would facilitate the local deposition of cysts that re-inoculate blooms in subsequent years (Anderson et al. 1983). Differences in the timing of blooms between these water bodies as well as the apparent retention of *A. fundyense* in the salt ponds of the NMS would prevent sexual reproduction between cells from the NMS and GOM, thus serving to reduce gene flow and reinforce the genetic differentiation between these populations.

We hypothesized that the localization of *A. fundyense* cells in Salt Pond from the combination of these behavioral and hydrographic factors restricts genetic exchange with GOM populations that may be present in the inlet channel and other well-mixed areas of the embayment.

This hypothesis was supported by data collected during both years of the study. In both 2006 and 2007, we observed moderate to high genetic differentiation between GOM and NMS populations (Table 2); AMOVA tests also showed the highest levels of among-group variation between populations from these bodies (Table 3). Two distinct clusters, comprising populations from each water body, were also apparent in the STRUCTURE analysis (Fig 3A). These data clearly indicate considerable genetic isolation between *A. fundyense* in the NMS and populations in the wider GOM, providing evidence that bloom dynamics and outbreaks of toxicity are indeed localized, driven by hydrographic and ecological factors specific to the embayment.

Within the NMS, we observed temporal genetic differentiation, both within and among sampling years. In both study years, successive sampling was timed to capture the genetic composition of populations at bloom initiation, development and progression, and during bloom decline; and in both 2006 and 2007, we observed temporal population differentiation. In 2006, the late-bloom sample 06SPD was significantly different from those collected in Salt Pond during bloom initiation and progression (Table 3; also evident in STRUCTURE analysis, Fig 3). Rapid differentiation was again observed in 2007; however, the population structure was more subtle. Although the Fisher’s combined test showed significant genetic differentiation between each of the two late-bloom samples (07SPC, 07SPD) and 07SPB, these two late-bloom samples were not significantly different from each other, nor from the sample collected at bloom initiation (Table 2). Interannual comparisons of genetic differentiation and STRUCTURE analysis suggest that NMS bloom populations are comprised of at least three largely admixed subpopulations.

Considering the rapidity with which the genetic composition of populations in Salt Pond changed over the course of the bloom season, one explanation is mixing of genetically distinct cells from elsewhere in the NMS during the bloom. However, evidence regarding differences in the timing and population dynamics of the blooms among salt ponds in the NMS (Crespo et al. 2011), along with hydrodynamic data showing a long residence time of water within these salt ponds, do not support this hypothesis and instead suggest that blooms in the terminal endpoints of the system (e.g., Salt Pond, Mill Pond) are initiated locally and are largely retained within the endpoints of the system. Given the lack of information regarding the genetic composition of blooms from the remaining terminal endpoints in the NMS (e.g., Mill Pond), population mixing cannot be entirely ruled out, and is the subject of a system-wide investigation currently underway.

Other factors contributing to the genetic structure we observed include the effects of life-cycle transitions, such as differences in cyst maturation intervals and germination characteristics, which are important aspects of *Alexandrium* autoecology in Cape Cod salt ponds (Anderson et al. 1983; Anderson 1998). *Alexandrium* blooms appear earlier in these salt ponds than in the GOM and there are even differences among ponds in the NMS, in which blooms (and toxicity) are first observed in the southern extreme (Mill Pond), followed by the northern extreme (Salt Pond), potentially due to differences in cyst abundance as well as physiological differences in cyst maturation intervals and timing of germination (Crespo et al. 2011). Studies of life-cycle dynamics in Salt Pond also revealed that the first planozygotes appear when populations as a whole were increasing rapidly, suggesting that differentiation and fusion of gametes occurred quite early in the stages of bloom development, well before the size of the bloom population peaks (Anderson et al. 1983). This process was remarkably consistent among populations from embayments on Cape Cod; in three different
salt ponds, zygotes appeared approximately six divisions after the first vegetative cells were seen, suggesting endogenous or “clock”-related sexuality.

The initiation of sexuality in the early bloom population could curtail vegetative growth, leading to the formation of cysts and potentially to population decline. Differences in the timing of germination along with the early induction of sexuality may impede sexual recombination among physiologically (and genetically) distinct subpopulations that may be present in the terminal endpoints of the system, and those advected from the GOM, thus contributing to the clonal diversity and genetic divergence we observed. Anderson et al. (1983) posited that the encystment/excystment cycle temporally restricts the occurrence of the vegetative population and may not be optimized for rapid or sustained vegetative growth and bloom formation in shallow embayments. In this scenario, population succession could occur in which early germinating populations undergo vegetative growth, form gametes within a short period of time, and are then replaced by later germinating populations. This is supported by the 2006 data, which documented a genetically distinct late-bloom population. In 2007, however, population differentiation occurred earlier, and the late-bloom sample was not significantly different from the sample collected at bloom initiation. Thus, the relative importance of germination timing and early encystment relative to other factors influencing the population structure of blooms in Salt Pond still remains unclear.

A final consideration is that of grazers and parasites, which strongly influence the dynamics of Alexandrium in the NMS, and may also affect population structure. Watras et al. (1985) previously estimated grazing rates utilizing radioactive tracers (14C bicarbonate) and A. fundyense growth rates in Salt Pond, providing evidence that rates of grazing often exceed growth rates, thus regulating the magnitude of the bloom and contributing to rapid bloom decline. These studies showed that historically, mortality due to zooplankton grazing has the potential to remove 100% of Alexandrium populations within a single day, and has been a key regulatory mechanism of population dynamics of A. fundyense in Salt Pond and other Cape Cod embayments (Turner and Anderson 1983; Watras et al. 1985). In addition to grazing, parasite-induced mortality by dinoflagellates in the genus Amoebophrya is a powerful “bottom-up” control of bloom-forming dinoflagellates in coastal environments, and has also been shown to drive annual cycles of dinoflagellate parasitoid successions observed in a coastal estuary in France (Chambouvet et al. 2008). A separate investigation conducted during the same period, as this study showed that Amoebophrya sp. parasitism is prevalent in Salt Pond, with estimates of host infection rates as high as 90% (Sengco and Coats, unpubl. ms.). Thus, there is historical and current evidence that both grazing pressure and parasitism can strongly influence Alexandrium dynamics in Salt Pond, and therefore may contribute to the population structure. Microbial food web models incorporating parasitism by Amoebophrya spp. can result in the near extirpation of dinoflagellate blooms (Montagnes et al. 2008), thus influencing the genetic dynamics of the successive populations rebounding from infection. Such strong selective pressure may consequently help maintain genotypic diversity in the population.

The results described herein clearly highlight the need for further investigation of the roles of population mixing and selective forces in the genetic dynamics of bloom populations in the NMS. To begin addressing these factors, an expanded investigation is underway of the genetic composition of A. fundyense populations from the other terminal endpoints of the NMS. Continued studies of the population and genetic dynamics of A. fundyense in the NMS, coupled with future investigations of host–parasite interactions and grazing pressure, will provide a better understanding of the evolutionary ecology of blooms and potential role of top-down controls in driving genetic dynamics. Due to their small geographic scale and limited connectivity, NMS salt ponds provide a novel system for investigating regulators of bloom development and termination, such as life-cycle events, as well as the influence of selective forces on population dynamics, all of which are otherwise difficult or impossible to study in the adjacent open-coastal waters or larger estuaries.

Conclusions

This study examined A. fundyense populations within a shallow embayment on Cape Cod, MA, to investigate the genetic characteristics of localized blooms and assess their connectivity with regional GOM populations. Comparisons of genetic differentiation showed that A. fundyense blooms in the NMS were genetically distinct from populations in the GOM, indicating that the localization of A. fundyense populations by behavioral and hydrographic factors effectively restricts genetic exchange with GOM populations. A. fundyense blooms in the NMS were characterized by extensive genetic diversity and rapid temporal genetic differentiation. On the basis of preliminary data from other investigations, we hypothesize that the genetic dynamics observed in Salt Pond may be influenced by one or more of the following factors: (1) life-cycle stages, including early onset of sexuality, and differences in cyst maturation intervals and timing of germination; (2) strong selection pressure exerted by Amoebophrya sp. parasitism and potentially from zooplankton grazing; and, (3) the mixing of distinct genotypes from a separated population elsewhere in the NMS.
Acknowledgments

The authors gratefully acknowledge the hard work of Bruce Keafer, Kerry Norton, and Dave Kulis in obtaining the samples used in this study. We thank Hannah Blossom for helping to collect the microsatellite data and Cecile Jauzien for providing micrographs. We also thank three anonymous reviewers for their very helpful suggestions and comments. This study was funded through the Woods Hole Center for Oceans and Human Health, National Science Foundation OCE-0430724 and National Institutes of Health 1 P50 ES012742-01, and National Science Foundation OCE-0911031. Funding was also provided by NOAA Grant NA06NOS4780245. This is contribution #706 from the US ECOHAB program.

Conflict of Interest

None declared.

References

Adams, N., V. Trainer, G. Rocap, R. Herwig, and L. Hauser. 2009. Genetic population structure of *Pseudo-nitzschia pungens* (Bacillariophyceae) from the Pacific Northwest and the North Sea. J. Phycol. 45:1037–1045.

Alpermann, T., B. Beszteri, U. John, U. Tillmann, and A. Cembella. 2009. Implications of life-history transitions on the population genetic structure of the toxigenic marine dinoflagellate *Alexandrium tamarense*. Mol. Ecol. 18:2122–2133.

Anderson, D. 1998. Physiology and bloom dynamics of toxic *Alexandrium* species, with emphasis on life cycle transitions. Pp. 29–48 in D. Anderson, A. Cembella and G. M. Hallegraeff, eds. The physiological ecology of harmful algal blooms. Springer, Heidelberg, Germany.

Anderson, D. M., and K. D. Stolzenbach. 1985. Selective retention of two dinoflagellates in a well-mixed estuarine embayment: the importance of diel vertical migration and surface avoidance. Mar. Ecol. Prog. Ser. 25:39–50.

Anderson, D. M., and A. W. White. 1992. Marine biotoxins at the top of the food chain. Oceanaus 35:55–61.

Anderson, D., D. Kulis, J. Orphanos, and A. Ceurvels. 1982. Distribution of the toxic dinoflagellate *Gonyaulax tamarensis* in the southern New England region. Estuar. Coast. Shelf Sci. 14:447–458.

Anderson, D. M., S. W. Chisholm, and C. J. Watras. 1983. The importance of life cycle events in the population dynamics of *Gonyaulax tamarensis*. Mar. Biol. 76:179–183.

Anderson, D. M., D. M. Kulis, G. J. Doucette, J. C. Gallager, and E. Balech. 1994. Biogeography of toxic dinoflagellates in the genus *Alexandrium* from the northeast United States and Canada as determined by morphology, bioluminescence, toxin composition, and mating compatibility. Mar. Biol. 120:467–478.

Anderson, D. M., B. A. Keafer, D. J. McGillicuddy, M. J. Mickelson, K. E. Keay, and P. Scott Libby, et al. 2005a. Initial observations of the 2005 *Alexandrium fundyense* bloom in southern New England: General patterns and mechanisms. Deep Sea Res. Part II 52:2856–2876.

Anderson, D. M., D. M. Kulis, B. A. Keafer, K. E. Gribble, R. Marin, and C. A. Scholin. 2005b. Identification and enumeration of *Alexandrium* spp. from the Gulf of Maine using molecular probes. Deep Sea Res. Part II 52:2467–2490.

Brosnanah, M. L., D. M. Kulis, A. R. Solow, D. L. Erdner, L. Percy, J. Lewis, and D. M. Anderson. 2010. Outbreeding lethality between toxic Group I and nontoxic Group III *Alexandrium tamarense* spp. isolates: predominance of heterotypic encystment and implications for mating interactions and biogeography. Deep Sea Res. Part II 57:175–189.

Cembella, A. D. 1998. Ecophysiology and metabolism of paralytic shellfish toxins in marine microalgae. Pp. 381–403 in D. M. Anderson, D. A. Cembella and G. M. Hallegraeff, eds. Physiological ecology of harmful algal blooms. Springer, Berlin/Heidelberg/New York.

Chambouvet, A., P. Morin, D. Marie, and L. Guillou. 2008. Control of toxic marine dinoflagellate blooms by serial parasitic killers. Science 322:1254–1257.

Crespo, B. G., B. A. Keafer, D. K. Ralston, H. Lind, D. Farber, and D. M. Anderson. 2011. Dynamics of *Alexandrium fundyense* blooms and shellfish toxicity in the Nauset Marsh System of Cape Cod (Massachusetts, USA). Harmful Algae 12:26–38.

Dyhrman, S. T., D. Erdner, J. L. Du, M. Galac, and D. M. Anderson. 2006. Molecular quantification of toxic *Alexandrium fundyense* in the Gulf of Maine using real-time PCR. Harmful Algae 5:242–250.

Ellstrand, N. C., and M. L. Roose. 1987. Patterns of genotypic diversity in clonal plant species. Am. J. Bot. 74:123–131.

Erdner, D. L., M. Richlen, L. A. R. McCauley, and D. M. Anderson. 2011. Diversity and dynamics of a widespread bloom of the toxic dinoflagellate *Alexandrium fundyense*. PLoS ONE 6:e22965.

Evanno, G., S. Regnaut, and J. Goudet. 2005. Detecting the number of clusters of individuals using the software structure: a simulation study. Mol. Ecol. 14:2611–2620.

Evans, K. M., S. F. Kühn, and P. K. Hayes. 2005. High levels of genetic diversity and low levels of genetic differentiation in North Sea *Pseudo-nitzschia pungens* (Bacillariophyceae) populations. J. Phycol. 41:506–514.

Excoffier, L., G. Laval, and S. Schneider. 2005. Arlequin ver. 3.0: an integrated software package for population genetic data analysis. Evol. Bioinform. Online 1:47–50.

Goudet, J. 1999. PCA-GEN for Windows, V. 1.2. University of Lausanne, Lausanne, Switzerland.

Guillard, R. R. L. 1975. Culture of phytoplankton for feeding marine invertebrates. Pp. 29–60 in W. Smith and M. Hance, eds. The physiological ecology of harmful algal blooms. Springer, Heidelberg, Germany.

Hamer, P. 1987. Genetic differentiation among *Alexandrium tamarense* populations of the northeast United States and Canada. PhD dissertation, Woods Hole Oceanographic Institution, Woods Hole, Massachusetts.

Hart, R. A., and D. M. Anderson. 2010. Outbreeding lethality between toxic Group I and nontoxic Group III *Alexandrium tamarense* spp. isolates: predominance of heterotypic encystment and implications for mating interactions and biogeography. Deep Sea Res. Part II 57:175–189.

Hofmann, P., V. Trainer, G. Rocap, R. Herwig, and L. Hauser. 2009. Genetic population structure of *Pseudo-nitzschia pungens* (Bacillariophyceae) from the Pacific Northwest and the North Sea. J. Phycol. 45:1037–1045.

Jauzien, C. 2011. Molecular quantification of toxic *Alexandrium fundyense* in the Gulf of Maine using real-time PCR. Harmful Algae 5:242–250.

Lavoie, S., and J. Goudet. 1999. Detecting the number of clusters of individuals using the software structure: a simulation study. Mol. Ecol. 14:2611–2620.

Libby, D. S., and D. M. Anderson. 2004. Molecular characterization of toxic and non-toxic *Alexandrium fundyense* isolates: predominance of heterotypic encystment and implications for mating interactions and biogeography. Deep Sea Res. Part II 51:7–55.
Chanley, eds. Culture of marine invertebrate animals. Plenum, New York.
Hattenrath, T. K., D. M. Anderson, and C. J. Gobler. 2010. The influence of anthropogenic nitrogen loading and meteoro
tological conditions on the dynamics and toxicity of Alexandrium fundyense blooms in a New York (USA) Estuary. Harmful Algae 9:402–412.
Iglesias-Rodriguez, M. D., O. M. Schofield, J. Batley, L. K. Medlin, and P. K. Hayes. 2006. Intraspecific genetic diversity in the marine coccolithophore Emiliania huxleyi (Prymnesiophyceae): the use of microsatellite analysis in marine phytoplankton population studies. J. Phycol. 42:526–536.
Lilly, E. L., K. M. Halanych, and D. M. Anderson. 2007. Species boundaries and global biogeography of the Alexandrium tamarense complex (Dinophyceae). J. Phycol. 43:1329–1338.
Masseret, E., D. Grzebyk, S. Nagai, B. Genovesi, B. Lasserre, and M. Laabir, et al. 2009. Unexpected genetic diversity among and within populations of the toxic dinoflagellate Alexandrium catenella as revealed by nuclear microsatellite markers. Appl. Environ. Microbiol. 75:2037–2045.
McCauley, L., D. Erdner, S. Nagai, M. Richlen, and D. Anderson. 2009. Biogeographic analysis of the globally distributed harmful algal bloom species Alexandrium minutum (Dinophyceae) based on rRNA gene sequences and microsatellite markers. J. Phycol. 45:454–463.
Montagnes, D., A. Chambouvet, L. Guillou, and A. Fenton. 2008. Responsibility of microzooplankton and parasite pressure for the demise of toxic dinoflagellate blooms. Aquat. Microb. Ecol. 53:211–225.
Nagai, S., C. Lian, M. Hamaguchi, Y. Matsuyama, S. Itakura, and T. Hogetsu. 2004. Development of microsatellite markers in the toxic dinoflagellate Alexandrium tamarense (Dinophyceae). Mol. Ecol. Notes 4:83–85.
Nagai, S., C. Lian, S. Yamaguchi, M. Hamaguchi, Y. Matsuyama, and S. Itakura, et al. 2007. Microsatellite markers reveal population genetic structure of the toxic dinoflagellate Alexandrium tamarense (Dinophyceae) in Japanese coastal waters. J. Phycol. 43:43–54.
Nagai, S., G. Nishitani, S. Sakamoto, T. Sugaya, C. K. Lee, and C. H. Kim, et al. 2009. Genetic structuring and transfer of marine dinoflagellate Cochlodinium polykrikoides in Japanese and Korean coastal waters revealed by microsatellites. Mol. Ecol. 18:2337–2352.
Nei, M. 1973. Analysis of gene diversity in subdivided populations. Proc. Natl. Acad. Sci. USA 70:3321–3323.
Pritchard, J. K., M. Stephens, and P. Donnelly. 2000. Inference of population structure using multilocus genotype data. Genetics 155:945–959.
Raymond, M., and F. Rousset. 1995. GENEPOP (version 1.2): population genetics software for exact tests and ecumenicism. J. Hered. 86:248–249.
Rousset, F. 2008. Genepop’007: a complete reimplementation of the Genepop software for Windows and Linux. Mol. Ecol. Resour. 8:103–106.

Rynearson, T., and E. Armbrust. 2005. Maintenance of clonal diversity during a spring bloom of the centric diatom Ditylum brightwelli. Mol. Ecol. 14:1631–1640.
Rynearson, T., J. Newton, and E. Armbrust. 2006. Spring bloom development, genetic variation, and population succession in the planktonic diatom Ditylum brightwelli. Limnol. Oceanogr. 51:1249–1261.
Scholin, C. A., G. M. Hallegraef, and D. M. Anderson. 1995. Molecular evolution of the Alexandrium tamarensis "species complex" (Dinophyceae): dispersal in the North American and West Pacific regions. Phycologia 34:472–485.
Robust Euro-wildlife loss and harmful algal blooms: sporadic victims or under-reported events? Harmful Algae 2:1–7.
Turner, J. T., and D. M. Anderson. 1983. Zooplankton grazing during dinoflagellate blooms in a Cape Cod embayment, with observations of predation upon tintinnids by copepods. Mar. Ecol. 4:359–374.
Valière, N. 2002. gimlet: a computer program for analysing genetic individual identification data. Mol. Ecol. Notes 2:377–379.
Waits, L. P., G. Luikart, and P. Taberlet. 2001. Estimating the probability of identity among genotypes in natural populations: cautions and guidelines. Mol. Ecol. 10:249–256.
Watras, C. J., V. C. Garcon, R. J. Olson, S. W. Chisholm, and D. M. Anderson. 1985. The effect of zooplankton grazing on estuarine blooms of the toxic dinoflagellate Gonyaulax tamarensis. J. Plankton Res. 7:891–908.
Wright, S. 1951. Genetical structure of populations. Annals of Eugenics 15:247–249.
Yeh, F. C., and T. J. B. Boyle. 1997. Population genetic analysis of co-dominant and dominant markers and quantitative traits. Belg. J. Bot. 129:157.

Supporting Information

Additional Supporting Information may be found in the online version of this article:

Table S1. Characteristics of the microsatellite loci used to assess genetic diversity of Alexandrium fundyense populations from the GOM and the NMS on Cape Cod, MA, USA. Results of this are compared with those observed by Nagai et al. (2004, 2007) in Japanese toxic A. tamarense. N_i individuals, N_a maximum number of alleles, R-size ranges of the alleles, H_e gene diversity, ND, no data.

Figure S1. PCR products from (A) 06GOM, (B) 06SPC, and (C) 07SPD. Amplification performed using PCR primers specific to the NA ribotype of Alexandrium fundyense.

Please note: Wiley-Blackwell are not responsible for the content or functionality of any supporting materials supplied by the authors. Any queries (other than missing material) should be directed to the corresponding author for the article.