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2	The influence of anthropogenic nitrogen loading and meteorological conditions on
3	the dynamics and toxicity of Alexandrium fundyense blooms in a New York (USA)
4	estuary
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18 Abstract: The goal of this two-year study was to explore the role of nutrients and 19 climatic conditions in promoting reoccurring Alexandrium fundyense blooms in the Northport-Huntington Bay complex, NY, USA. A bloom in 2007 was short and small (3 20 weeks, 10<sup>3</sup> cells L<sup>-1</sup> maximal density) compared to 2008 when the A. fundyense bloom, 21 which persisted for six weeks, achieved cell densities  $>10^6$  cells L<sup>-1</sup> and water column 22 saxitoxin concentrations >2.4 x  $10^4$  pmol STX eq. L<sup>-1</sup>. During the 2008 bloom, both 23 24 deployed mussels (used as indicator species) and wild soft shell clams became highly 25 toxic (1,400 and 600 µg STX eq./100g shellfish tissue, respectively) resulting in the 26 closure of shellfish beds. The densities of benthic A. fundyense cysts at the onset of this 27 bloom were four orders of magnitude lower than levels needed to account for observed 28 cell densities, indicating in situ growth of vegetative cells was responsible for elevated bloom densities. Experimental enrichment of bloom water with nitrogenous compounds, 29 30 particularly ammonium, significantly increased A. fundyense densities and particulate saxitoxin concentrations relative to unamended control treatments. The  $\delta^{15}$ N signatures 31 32 (12 to 23‰) of particulate organic matter (POM) during blooms were similar to those of sewage (10 to 30%) and both toxin and A. fundyense densities were significantly 33 correlated with POM  $\delta^{15}$ N (p < 0.001). These findings suggest A. fundvense growth was 34 35 supported by a source of wastewater such as the sewage treatment plant which discharges 36 into Northport Harbor. Warmer than average atmospheric temperatures in the late winter 37 and spring of 2008 and a cooler May contributed to an extended period of water column temperatures optimal for A. fundvense growth  $(12 - 20^{\circ}C)$ , and thus may have also 38 39 contributed toward the larger and longer bloom in 2008. Together this evidence suggests

41 and toxic A. *fundyense* blooms in estuaries.

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43 Keywords: *Alexandrium*, anthropogenic nitrogen loading,  $\delta^{15}$ N, toxin, climate

45 **1. Introduction:** 

46 The intensity and impacts of harmful algal blooms (HABs) in coastal ecosystems have increased in recent decades (Anderson, 1994, Glibert et al., 2005; Anderson et al., 47 48 2008; Heisler et al., 2008). Blooms of the dinoflagellate *Alexandrium* are common to 49 many coastal regions around the globe and are particularly harmful because they produce 50 saxitoxins, the causative agent of paralytic shellfish poisoning (PSP) (Anderson, 1994; 51 Anderson, 1997; Glibert et al., 2005). Saxitoxins are a family of potent neurotoxins that block sodium channels and cause severe illness or death in humans who consume 52 53 saxitoxin-contaminated shellfish (Kvitek and Beitler, 1988; Anderson, 1994). The 54 frequency of Alexandrium blooms as well as the intensity of these events have been 55 increasing worldwide, and therefore so have PSP outbreaks (Anderson, 1994; Sellner et 56 al., 2003; Glibert et al., 2005). Although it is not certain whether these events can be 57 attributed to an increase in coastal monitoring or to increased anthropogenic nutrient 58 loading to coastal systems (Anderson, 1994; Anderson et al., 2002; Glibert et al., 2005; 59 Heisler et al., 2008), it is clear that these blooms have devastating economic impacts 60 (Anderson et al., 2000; Jin and Hoagland, 2008; Jin et al., 2008). Many consider PSP to 61 be the most widespread of all HAB poisoning syndromes (Hackett et al., 2004; Erdner et 62 al., 2008).

*Alexandrium fundyense* blooms are common along the northeast US coast.
Paralytic shellfish poisoning in the northeastern US was first documented in Maine in
1958 (Anderson, 1997). In 1972, a large *A. fundyense* bloom with cell densities
exceeding 10<sup>6</sup> cells L<sup>-1</sup> spread through the Gulf of Maine and affected coastal regions
from Maine to Massachusetts (Mulligan, 1975; Anderson, 1994; Anderson, 1997). Since

then these large-scale regional blooms and associated PSP-related shellfish bed closures
have been near-annual occurrences in this region (Anderson, 1994; Anderson, 1997;
Townsend et al., 2001) to the detriment of the shellfish industry. For example, during the
2005 *A. fundyense* bloom in New England, the seafood industry lost more than \$3 million
per week in revenue (Jin et al., 2008).

73 In contrast to these widespread coastal outbreaks, Alexandrium blooms also occur 74 in isolated embayments along the New England and Long Island coast. These are considered "point source" outbreaks in which localized cyst germination inoculates the 75 76 overlying waters, with deposition of new cysts at the end of blooms providing the means for the species to bloom again in subsequent years (e.g., Anderson and Morel, 1979; 77 78 Anderson et al., 1983; Anderson et al., 2008). There is thought to be no significant 79 connection between these small-scale blooms in estuaries and embayments and the large-80 scale regional blooms described above (Anderson, 1997; Anderson et al., 2008).

81 The presence of A. fundyense on Long Island was first documented during the 82 early 1980's (Anderson et al., 1982; Schrey et al., 1984). At that time, moderate densities of A. fundyense (>  $10^2$  cell L<sup>-1</sup>) were found on the north shore of Long Island in Northport 83 84 Bay and Mattituck Inlet (Schrey et al., 1984); these blooms, however, were not associated 85 with PSP events (e.g. toxic shellfish or human illness; Anderson et al., 1982; Schrey et 86 al., 1984). At the time, this was attributed to the low toxin content or quota of Long 87 Island isolates, which contain proportionally more of the low potency C toxins than other 88 more potent congeners (Anderson et al., 1994). The assumption was that very dense 89 blooms would be required for there to be dangerous levels of toxicity in shellfish. 90 Although there have been no studies of A. fundyense in NY waters since the 1980s, in

2006, the detection of elevated saxitoxin in shellfish by the New York State Department
of Environmental of Conservation (NYSDEC) prompted the closure of 2,000 acres of
shellfish beds in the Northport-Huntington Bay system of Long Island. Blooms recurred
in those waters in 2007 and 2008.

95 Factors promoting toxic A. fundyense bloom events seem to vary with the 96 ecosystem within which blooms occur (Anderson et al., 2008). Decades of research in 97 the Gulf of Maine have led to the conclusion that the presence and dynamics of A. 98 *fundyense* benthic cyst beds and the physical transport of cells controls the dynamics of 99 the widespread regional blooms (Anderson, 1997; Anderson et al., 2005a,c; Stock et al., 100 2005; Anderson et al., 2008). The low levels of nutrients present during blooms in the 101 open waters of the Gulf of Maine (Townsend et al., 2001; Poulton et al., 2005; Townsend 102 et al., 2005b; Love et al., 2005) and the ability of A. fundyense dynamics to be 103 successfully modeled in the absence of a nutrient-dependent growth rate (Stock et al., 104 2005) suggests nutrients seem to have a smaller, secondary influence on these events 105 (Anderson, 1997; Anderson et al., 2005a,c). In contrast, anthropogenic nutrient loading 106 could have a larger impact on the development of A. fundyense blooms in coastal 107 embayments where nutrient concentrations and loads are substantially higher than the 108 Gulf of Maine (Anderson et al., 1983; Penna et al., 2002; Trainer et al., 2003; Poulton et 109 al., 2005; Anderson et al., 2008). Anthropogenic nutrient loading has been associated 110 with an increase in PSP incidences caused by *Alexandrium catenella* in multiple marine 111 ecosystems including shallow, poorly flushed coastal embayments of the northwest US 112 (Trainer et al., 2003). The degree to which A. fundyense populations in estuaries are 113 controlled by nutrient loading, cyst beds, or both factors is not well understood.

114 This study documented the dynamics of A. fundyense blooms in a coastal region 115 of New York in 2007 and 2008, including a bloom which persisted for six weeks, achieved densities of more than  $10^6$  cells L<sup>-1</sup>, and lead to the closure of more than 7,000 116 117 acres of shellfish beds. The spatial and temporal dynamics of the physical environment, 118 nutrients, toxins, A. fundyense cells, and A. fundyense cysts are presented in conjunction 119 with experiments examining the impacts of nutrient enrichment on the growth and 120 toxicity of A. fundyense populations. The role of nutrient loading and meteorological 121 conditions in the occurrence of A. fundyense blooms is subsequently assessed.

122 2. Materials & Methods:

#### 123 2.1. Field sampling and analyses

124 During 2007 and 2008 sampling was conducted at various locations across the 125 Northport-Huntington Bay complex, located on the north shore of Long Island, NY, USA 126 (Fig. 1, 40.9090°N, 73.4036°W). This system has previously hosted A. fundyense cells 127 (Anderson et al., 1982; Schrey et al., 1984; Anderson, 1997) and saxitoxin contaminated 128 shellfish (Karen Chytalo, NYSDEC, personal communication). Within this system, 129 Northport Harbor, located in the southeastern part of the Northport-Huntington Bay 130 complex, was sampled on a weekly basis from April through June at one site in 2007 (site 131 2) and at three locations in 2008 (Fig. 1; sites 2, 7, 8). Other sites, located in Huntington 132 Harbor (site 6) and Centerport Harbor (site 1) were sampled weekly, while 7 other sites 133 (sites 3, 4, and 5 in 2007; sites 3, 4, 10, 11, 16 and LIS in 2008) were sampled during the 134 pinnacle of blooms to document the spatial extent of these events (Fig. 1).

135At each station, a YSI© probe was used to record surface temperature, salinity136and dissolved oxygen. Subsurface water (~0.25m) was filtered for nutrient analysis using

137 precombusted (4 hr @  $450^{\circ}$ C) glass fiber filters (GF/F, 0.7 µm pore size) and frozen in 138 acid washed scintillation vials. Filtrate was analyzed colorimetrically for ammonium, nitrate, phosphate, and silicate (Jones, 1984; Parsons et al., 1984) using a 139 140 spectrophotometeric microplate reader. To determine the size distribution of 141 phytoplankton biomass, chlorophyll a was fractionated using GF/F (nominal pore size 0.7 142  $\mu$ m) and polycarbonate filters (2  $\mu$ m & 20  $\mu$ m) and measured using standard fluorometric 143 techniques described in Parsons et al. (1984). Whole water samples were preserved in 144 Lugol's iodine. Aliquots were settled in counting chambers and plankton were identified 145 and enumerated using an inverted light microscope (Hasle, 1978). Cells larger than 10 146 µm were identified to at least genus level and grouped as dinoflagellates, diatoms, and ciliates. To assess the  $\delta^{15}N$  signature of plankton communities dominated by A. 147 148 fundvense, replicate samples of particulate organic matter (POM) was filtered onto 149 precombusted (4h @ 450°C) GF/F filters, dried for 24 h at 60°C, pelleted, and analyzed 150 for  $\delta^{15}$ N via continuous flow isotope ratio mass spectrometry (IRMS) by David Harris at 151 the UC Davis Stable Isotope Facility.

152 A. fundyense cell densities were enumerated using a molecular technique 153 developed by Anderson et al. (2005b). In the field, 2 L of water was pre-sieved through a 154 200 µm mesh to eliminate large zooplankton from the sample and subsequently 155 concentrated onto a 20 µm sieve and backwashed into a centrifuge tube to a volume of 14 ml. Samples were preserved in ~2% formaldehyde and refrigerated at  $4^{\circ}$ C for at least 1 156 157 hour and no more than 24 hours. After refrigeration, samples were centrifuged at 3000 158 rpm for 11 minutes and the supernatant aspirated without disturbing the cell pellet. The 159 cell pellet was resuspended in 14 ml ice cold methanol and stored at  $-20^{\circ}$ C for up to six

160 months (Anderson et al., 2005b). An aliquot of preserved sample was filtered onto a 5 161 µm polycarbonate track-etched membrane (25mm in diameter). A pre-hybridization 162 buffer was incubated for 5 minutes with each sample and then filtered off of samples. A. 163 fundyense cells were labeled using oligonucleotide probe NA1 for the North American ribotype Alexandrium fundvense/catenella/tamarense with Nu-light <sup>™</sup> dye conjugated to 164 165 the 5' end (5'-/5Cy3/AGT GCA ACA CTC CCA CCA-3'). A hybridization buffer, 166 containing pre-hybridization buffer in addition to probe (a final probe concentration of 4.8 ng  $\mu$ l<sup>-1</sup>) was added to each sample and allowed to incubate for 1 hour at 50°C. 167 Following incubation, the hybridization buffer was filtered and samples were washed 168 169 with 0.2X SET for 5 minutes. Filters were then mounted onto a microscope slide and 170 glycerol was added to each filter to prevent fading of the probe. Cells were enumerated using a Nikon epifluorescence microscope with a Cy3<sup>™</sup> filter set (Anderson et al., 171 172 2005b). As a quality control, measured samples spiked with A. fundyense culture (clone 173 GTCA28 or ATNPD7) were hybridized with the oligonucleotide probe and quantified 174 during each analytical run. Oligonucleotide probe quantification of seawater spiked with 175 known densities of A. fundyense clone GTCA28 yielded mean recoveries of  $87 \pm 16\%$ . 176 Light microscope counts of Lugol's stained A. fundyense cells yielded large 177 overestimates of population densities compared to oligonucleotide quantification.

Toxin concentrations in plankton samples were determined by a competitive enzyme linked immunosorbent assay (ELISA). Several liters of seawater were pre-sieved through a 200 µm mesh and subsequently concentrated on a 20 µm sieve, backwashed into centrifuge tubes and pelleted. Cell pellets were acidified with 0.1 M HCl and subsequently analyzed for saxitoxin using ELISA kits from R-Biopharm© in 2007 and by

183 Abraxis<sup>©</sup> in 2008, with toxin concentrations reported in STX equivalents. Each of these 184 kits had varying degrees of cross-reactivities among saxitoxin congeners. Cross-185 reactivities for the ELISA kits from R-Biopharm<sup>©</sup> and Abraxis<sup>©</sup> were as follows: 100% 186 STX, 20% dcSTX, 70% GTX2,3 and 12% NEO, and 100% STX, 29% dcSTX, 23% GTX2,3, 23% GTX5B, 1.3% NEO, and <0.2% GTX1,4, respectively. Analysis of 187 188 replicated samples by both kits yielded statistically identical results. As a quality control 189 measure, for each analytical run, an Alexandrium fundyense culture (GTCA28) known to 190 produce saxitoxins was used as a positive control and Aureococcus anophagefferens 191 (CCMP 1984), which does not produce saxitoxins, was used as a negative control. Three 192 times the standard deviation of the negative control was used as the methodological 193 detection limit for each analytical run. Analysis of total saxitoxins in pelleted 194 Alexandrium fundyense cultures (clone ATNPD7) via high performance liquid 195 chromatography (HPLC) yielded statistically equivalent levels of total saxitoxin 196 concentrations on a per cell basis to those measured with the both ELISA kits.

197 During November 2007 and 2008 sediment samples were obtained from 17 198 locations across the Northport-Huntington Bay complex (Fig. 1). Surveys were timed to 199 occur following potential fall bloom events and thus quantified cysts represented 200 potential seed populations for the following year (Anderson et al., 2005c). Sediment 201 samples were obtained using a Ponar grab and several subcores from the top 3cm were 202 taken using a modified syringe. All samples were processed according to Anderson et al. 203 (2005c) and stained with primulin (Yamaguchi et al., 1995). Primulin stained cysts were 204 enumerated under an epifluorescent microscope using a 1 ml Sedgewick-Rafter slide. Cyst concentrations were reported in cysts  $cc^{-1}$  of sediment. 205

Meteorological data including wind intensity, wind direction, temperature, and precipitation were obtained from the National Weather Service's monitoring station in Islip, NY, USA which is ~20 km from Northport. For each of these parameters the monthly means for 2007 and 2008 were compared using t-tests. The degree to which all individual water column parameters were correlated to each other was evaluated by means of a Spearman rank order correlation matrix.

#### 212 2.2. Nutrient amendment experiments

213 To assess the impact of nitrogen (N) and phosphorus (P) loading on A. fundyense 214 growth and toxin production, a series of nutrient amendment experiments were 215 performed. Triplicate bottles (1.1 L in 2007 and 2.5 L in 2008) were filled with water 216 An unamended control was established along with four from Northport Harbor. treatments in 2007 including 20 µM nitrate, 20 µM ammonium, 10 µM urea (= 20 µM 217 218 N), and 2  $\mu$ M phosphate. Due to the response from reduced N in general and ammonium 219 in particular during 2007 experiments, experiments in 2008 included additional 220 treatments: 10 µM ammonium, 40 µM ammonium, 20 µM ammonium combined with 2 221  $\mu$ M phosphate, and 10  $\mu$ M glutamine (= 20  $\mu$ M N). All treatment concentrations were 222 chosen to match those which have previously elicited a growth response in *Alexandrium* 223 cells (Leong et al., 2004) and were similar to peak elevated levels found in Long Island 224 estuaries (Gobler et al., 2004). Bottles were incubated for  $\sim 48$  h at ambient light and 225 temperature after which A. fundyense cell enumeration, and toxin quantification were 226 performed via the aforementioned methods. Differences among treatments were elucidated by means of a One-Way ANOVA with multiple comparison tests (i.e. Student-227

Newman-Keuls) or with an appropriate non-parametric test when normality tests of logtransformed data failed.

### 230 2.3. Toxins in shellfish

231 During both 2007 and 2008, netted bags containing the blue mussel, Mytilus 232 *edulis*, from regions not exposed to PSP toxins were hung off piers located adjacent to 233 sampling sites in Northport Harbor and in Huntington Harbor. These mussel bags were 234 deployed in the early spring when temperatures were below those optimal for A. 235 fundvense growth (<  $10^{\circ}$ C). Mussel bags were collected weekly from each site and 236 mussels were shucked and extracts were prepared using standard techniques (Association 237 of Official Analytical Chemists (AOAC), 1990). Native soft shell clams (Mya arenaria) 238 from Northport Harbor were also harvested and extracts were prepared sporadically 239 during the months of April through May. Toxin levels in shellfish were quantified using 240 standard mouse bioassays (AOAC, 1990). Bioassays were performed by NYSDEC staff 241 at the Stony Brook University Health Sciences Center Division of Laboratory Animal 242 Resources by injecting shellfish extracts into mice (strain CD-1).

#### 243 **3. Results:**

#### 244 3.1. 2007 Northport Harbor <u>Alexandrium fundyense</u> bloom

During April of 2007 there was a bloom of non-*Alexandrium*, nanophytoplankton (2-20  $\mu$ m) in Northport Harbor which had chlorophyll levels exceeding 25  $\mu$ g L<sup>-1</sup> (Fig. 2) and was comprised primarily of diatoms (95±3% of cells enumerated). During May, as surface temperatures stabilized at ~ 15°C, the abundance of nanophytoplankton began to decline and a modest (>1,000 cells L<sup>-1</sup>) *A. fundyense* bloom developed (Fig. 2b). *A. fundyense* cells were detected in the water column from 8 May to 20 June with cell

densities peaking at 2,650 cells  $L^{-1}$  on 23 May (Fig. 2a, Table 1). Elevated toxin levels (> 251 2 pmol STX eq.  $L^{-1}$ ) in the water column were present through the bloom, with levels 252 peaking at 130 pmol STX eq.  $L^{-1}$  in unison with peak cell densities (Fig. 2a, Table 1). 253 254 The largest size fraction of chlorophyll (> 20  $\mu$ m) accounted for 23±0.8% of the total 255 chlorophyll during the bloom peak. Both ammonium and silicate concentrations 256 increased slightly during the bloom compared to before and after the A. fundyense bloom as did  $\delta^{15}N$  of the total plankton community which reached its annual maximum 257 258 (9.7±1.2‰) during the peak of the bloom (Fig. 2c, Fig. 3). During the week following peak cell densities, elevated levels of toxins were found in mussels deployed in Northport 259 260 Harbor (37 µg STX eq./100g shellfish tissue). The A. fundyense bloom ended in June as 261 temperatures exceeded 20°C (Fig. 2a,c).

During the bloom in Northport Harbor, A. fundyense concentrations in Centerport 262 Harbor ranged from 8 to 50 cells  $L^{-1}$  with low pelagic particulate toxin concentrations 263 (1.42-3.73 pmol STX eq. L<sup>-1</sup>; Table 1). The remaining sites in Northport-Huntington 264 Harbor complex had < 12 cells L<sup>-1</sup> and toxin concentrations below 7.1 pmol STX eq. L<sup>-1</sup> 265 266 (Table 1). A. fundyense cells and toxins were not detected in the water column of the 267 Northport-Huntington Bay system from July through November. During an experiment 268 conducted on 15 May 2007, the addition of ammonium resulted in a 60% increase in A. 269 fundyense cell densities compared to unamended control treatments (Fig. 4). During a 270 second experiment (30 May), the addition of ammonium resulted in 25% higher 271 particulate toxin concentrations and 70% higher cell densities (Fig. 4).

272 3.2. Presence of cysts in the Northport-Huntington Bay area: 2007

During a sediment survey conducted on 14 November 2007, *A. fundyense* cysts were present at low levels in the Northport-Huntington Bay complex (0 - 50 cysts cc of sediment<sup>-1</sup>; Table 1). The highest concentrations of cysts were located in Northport Harbor with concentrations ranging from 18-50 cysts  $cc^{-1}$ (sites 2, 7 and 8; Table 1). Maximal cyst concentrations (50 cysts  $cc^{-1}$ ) were found at site 8 (Table 1) ~0.6km north of the site with maximal cell densities (site 2; Fig. 1). The remainder of the Northport-Huntington Bay system had relatively low cyst concentrations (0-13 cysts  $cc^{-1}$ ; Table 1).

# 280 3.3. 2008 Northport Harbor <u>Alexandrium fundyense</u> bloom

281 During April and May of 2008, an intense Alexandrium fundyense bloom 282 developed and persisted in Northport Harbor for six weeks, during which temperatures 283 ranged from 10-21°C (Fig. 5a,c). During the bloom, the  $> 20 \,\mu m$  size class accounted for 284  $45 \pm 1.2\%$  (up to 76% on 16 May) of total chlorophyll *a* (Fig. 5b). The first peak of the bloom occurred on 16 May when  $1.2 \times 10^6 A$ . fundvense cells L<sup>-1</sup> and 24,662 pmol STX 285 eq. L<sup>-1</sup> were present (Table 1). A secondary bloom peak occurred on 23 May (6 x  $10^5 A$ . 286 fundvense cells L<sup>-1</sup>) and a secondary toxin peak occurred three days later on 26 May 287 (7,300 pmol STX eq. L<sup>-1</sup>; Fig. 5a). Concentrations of nitrate, ammonium, and phosphate 288 289 were all significantly (p<0.01 for each, t-test) higher before and after the bloom 290 (phosphate  $1.5 \pm 0.3 \mu$ M, nitrate  $14.1 \pm 2.6 \mu$ M, ammonium  $7.0 \pm 2.0 \mu$ M) compared to 291 during the bloom peak (6 May through 29 May; phosphate 0.5 ±0.1µM, nitrate 5.0 292  $\pm 1.5 \mu$ M, ammonium 1.8  $\pm$  1.0 $\mu$ M; Fig. 5c). In contrast, silicate levels gradually rose 293 from 7µM to 32µM from April through June (Fig. 5c). Throughout the bloom period, the  $\delta^{15}$ N of particulate organic matter ranged from 12 to 23‰ (Fig. 3). Mussel toxin levels 294 exceeded the regulatory closure limit (80 µg STX eq./100g shellfish tissue) two weeks 295

after the first detection of *A. fundyense* cells and peaked on 27 May (1,400  $\mu$ g STX eq./100g shellfish tissue) 11 days after peak cell and water column toxin concentrations (Fig. 5a). Native soft shell clams from this area were also highly toxic (600  $\mu$ g STX eq./100g shellfish tissue). As such, 7,000 acres of shellfish beds in Northport and Huntington Bays were closed to shellfishing for most of May and June 2008. During the demise of the *A. fundyense* bloom, water column temperatures rose above 20°C and 2 – 20  $\mu$ m size fraction chlorophyll *a* levels increased nearly five-fold (Fig. 5b,c).

303 Although other sites in Northport Harbor had the highest levels of A. fundyense during the 2008 bloom (sites 7 and 8 cell densities and toxin concentrations =  $5.5 \times 10^5$ 304 cells  $L^{-1}$  and 4.5 x 10<sup>3</sup> pmol STX eq.  $L^{-1}$  and 8.8 x 10<sup>5</sup> cells  $L^{-1}$  and 1.9 x 10<sup>4</sup> pmol STX 305 eq.  $L^{-1}$ , respectively; Table 1), elevated cell densities and toxin concentrations were also 306 307 present throughout the Northport-Huntington Bay system (Table 1). Centerport Harbor (site 1), had peak cell densities of 7,170 cells  $L^{-1}$  and toxin concentrations of 183 pmol 308 STX eq. L<sup>-1</sup> (Table 1). A. fundyense cell densities in Huntington Harbor (site 6) peaked 309 at 24,900 cells  $L^{-1}$  with corresponding toxin concentrations of 312 pmol STX eq.  $L^{-1}$ 310 311 (Table 1). After the occurrence of peak cell densities in Huntington Harbor, high levels 312 of toxin were quantified in deployed mussels (161 µg STX eq./100g shellfish tissue). 313 Peak cell densities occurred across Northport-Huntington Bay between 16-26 May with  $>10^4$  cells L<sup>-1</sup> found throughout the system and over 8 x 10<sup>3</sup> cells L<sup>-1</sup> in Long Island 314 315 Sound (Table 1).

## 316 *3.4. Nutrient amendment experiments: 2008*

317 The response of *A. fundyense* populations to nutrient amendments changed 318 through the course of the bloom. During experiments conducted at the beginning (30

319	April) and the demise of the A. fundyense bloom (2 June), there were no significant
320	changes in toxin concentrations in response to nutrient amendments (Fig. 6). However
321	during these same experiments, the addition of ammonium (10 $\mu M$ on 30 April; 20 $\mu M$
322	on 2 June) significantly increased A. fundyense densities compared to the control (p<0.01,
323	Student-Newman-Keuls; Fig. 6). On 6 May, the addition of ammonium (40 $\mu$ M) yielded
324	a significant (p<0.001, Student-Newman-Keuls) increase in both A. fundyense densities
325	and toxin concentrations by 4-fold and 8-fold, respectively, compared to controls. At the
326	same time, the addition of smaller concentrations of ammonium (10 and 20 $\mu$ M) yielded
327	smaller, but significant (p<0.01, Student-Newman-Keuls), increases in toxin (5-fold and
328	2-fold higher compared to controls, respectively) relative to the unamended control but
329	did not significantly alter cell densities. During the experiment conducted on 12 May the
330	enrichment of each nitrogenous compound produced significantly higher toxin
331	concentrations (3 - 10 fold increase compared to controls; p<0.001, Student-Newman-
332	Keuls; Fig. 6). During the same experiment, A. fundyense densities were also
333	significantly (p<0.05, Student-Newman-Keuls) enhanced by the additions of glutamine,
334	nitrate and ammonium (10 and 40 $\mu$ M); other N compounds (urea, ammonium (20 $\mu$ M),
335	and ammonium + phosphate) increased A. fundyense densities (60-80%), but not
336	significantly (Fig. 6). During late May (19 May, 26 May) the addition of N (all
337	nitrogenous compounds on 26 May, and only nitrate and urea on 19 May) yielded
338	modest, but non-significant increases $(10 - 60\%)$ in A. fundyense densities compared to
339	controls. During the 19 May experiment, the addition of ammonium (20 $\mu M$ ) and urea
340	resulted in modest (50% and 33%) increases in toxin, while toxin levels were

341 significantly (p<0.05, Student-Newman-Keuls) enhanced by the addition of nitrate and</li>
342 ammonium compared to the control during the 26 May experiment (Fig. 6).

343 Toxin concentrations normalized per cell were significantly increased by nutrient 344 enrichment in four of the six experiments conducted in 2008 (p < 0.05, Student Newman-345 Keuls; Fig. 7). The exceptions were the first (30 April) experiment during which cell-346 normalized toxin levels were unchanged and the final (2 June) experiment during which 347 the addition of N and P significantly decreased levels (p<0.05, Student Newman-Keuls; 348 Fig. 7). Experiments conducted on both 12 May and 26 May resulted in the most significant increases in toxin per cell for all N and P additions (2 - 4 times higher;349 350 p<0.05, Student-Newman-Keuls) with the exception of urea on 26 May (Fig. 7). In 351 contrast, only ammonium enrichment significantly increased cell-normalized toxin levels 352 during the 6 May and 19 May experiments (p < 0.05; Fig. 7).

### 353 *3.5. Presence of cysts in the Northport-Huntington Bay area: 2008*

354 The cyst survey conducted on 11 November 2008 indicated that A. fundyense 355 cysts were present at nearly every site in the Northport-Huntington Bay complex and 356 abundances were nearly an order of magnitude higher than those present in November 357 2007 (Table 1). Cyst concentrations were the highest in Northport Harbor with concentrations ranging from 220 to 745 cysts cc<sup>-1</sup>. As was the case in 2007, site 8 had 358 the highest cyst concentrations (745 cysts  $cc^{-1}$ , Table 1). Sites located just outside of 359 Northport Harbor also had elevated cyst concentrations compared to 2007 (20 - 115 cysts 360 361 cc<sup>-1</sup>, Table 1). The western part of the Northport-Huntington Bay complex generally had lower cyst concentrations (0-15 cysts cc<sup>-1</sup>, Fig.1, Table 1) compared to the eastern part of 362 363 the bay.

365 Atmospheric temperatures were significantly (p<0.001, t-test) warmer in February 366 2008 (1.3 $\pm$ 0.8°C) than February 2007 (-2.5 $\pm$ 0.7°C) as well as 1°C warmer than the long 367 term monthly mean (0.3°C) (Fig. 8). Furthermore, March 2008 (4.7±0.5°C) was 1.1°C warmer than March 2007 ( $3.6\pm1.0^{\circ}$ C) and slightly warmer than the long term monthly 368 369 mean (4.2°C). April 2008 (10.9±0.7°C) was significantly (p=0.05, t-test) warmer than 370 April 2007 (8.7 $\pm$ 0.9°C) as well as 1.5°C warmer than the long term monthly mean (9.4°C) 371 (Fig. 8). In May 2008, temperatures (14.1±0.6°C) were cooler than both May 2007 372 (16.0±0.8°C) and the long term monthly mean (15°C) (Fig. 8). During April of 2008, 373 winds blew persistently from the SE  $(160\pm18.7^{\circ})$ , whereas April 2007 winds came from 374 the SW  $(238\pm19.8^{\circ}; p=0.006, t-test)$ . There were no significant differences in 375 precipitation or wind intensity between 2007 and 2008 compared to long-term averages 376 for the months of January through June.

# 377 **4. Discussion:**

#### 378 4.1. 2007 & 2008 <u>Alexandrium fundyense</u> bloom toxicity and intensity

379 This study documented the dynamics of two contrasting blooms, one of which achieved cell densities greater than  $10^6$  cells L<sup>-1</sup> and resulted in the closure of 7000 acres 380 381 of shellfish beds in Northport, NY. The A. fundyense bloom in 2008 was dramatically 382 more intense and toxic than the bloom in 2007, with toxin and cell concentrations (means = 5,816 pmol STX eq.  $L^{-1}$ ; 353,184 cells  $L^{-1}$ ) in May 2008 being two and three orders of 383 384 magnitude higher (p<0.001, t-test) than those in May 2007; toxin levels were significantly correlated ( $r^2 = 0.942$ , p<0.001) with A. fundvense abundances during both 385 386 years. The sustained high densities of A. fundyense during the peak of the 2008 bloom

 $(>10^5 \text{ cells } \text{L}^{-1})$  were higher than those typically found in coastal embayments or open 387 388 waters of the Gulf of Maine where blooms are annual occurrences and cell densities are usually below  $10^4$  cells L<sup>-1</sup> (Townsend et al., 2001; Love et al., 2005; Poulton et al., 2005: 389 Townsend et al., 2005a, b). Similar concentrations (>10<sup>5</sup> cells  $L^{-1}$ ) of A. fundyense have 390 391 also been observed in the Nauset Marsh System on Cape Cod (D.M. Anderson, 392 unpublished data). While absolute toxin levels in Northport Harbor (up to 24,662 pmol STX eq.  $L^{-1}$ ) were also higher than those reported in Maine (400 pmol STX eq.  $L^{-1}$ ; 393 Poulton et al., 2005), the toxin contents or quotas in Northport Harbor (6.2 - 58.8 fmol)394 STX eq. cell<sup>-1</sup>; Fig. 9) were substantially lower than those of *Alexandrium* populations 395 from the Gulf of Maine  $(36 - 325 \text{ fmol cell}^{-1}; \text{Poulton et al., 2005})$ , a finding consistent 396 397 with the known north-south gradient in cell toxicity along the western Atlantic coast 398 (Maranda et al., 1985; Anderson et al., 1990a; Anderson et al., 1994; Bricelj and 399 Shumway, 1998), and with the dominance of low-potency saxitoxin congeners in 400 populations from Long Island and Connecticut waters (Anderson et al., 1994). Despite 401 the lower toxicity cells in NY, the large bloom in 2008 caused blue mussels (Mytilus 402 edulis), and native soft shell clams (Mya arenaria) in Northport Bay to become highly 403 toxic (1,400 and 600 µg STX eq./100g shellfish tissue, respectively) causing the closure 404 of >7,000 acres of shellfish beds for nearly two months (Karen Chytalo, NYSDEC, 405 Marine Division).

406 4.2. The relative importance of nitrogen, cysts, and meteorological conditions promoting
407 New York <u>Alexandrium fundyense</u> blooms:

The dynamics of *A. fundyense* blooms in Northport Harbor and the differences between the magnitude of the 2007 and 2008 blooms might be controlled by multiple

factors including cyst beds, meteorological conditions, and nutrient loading. Benthic cyst 410 411 concentrations in November 2008 were an order of magnitude greater than those present 412 in November 2007 (p < 0.001, t-test) and the spatial extent of cysts also expanded in 2008 413 likely due, in part, to the larger bloom that year compared to 2007. In the Gulf of Maine, 414 cyst seed bed distribution and cyst densities in combination with physical circulation 415 patterns are used to model blooms since cysts provide the inocula for future events 416 (Anderson, 1997; Anderson et al., 2003, 2005a,c; Stock et al., 2005; McGillicuddy et al., 417 2005). The cyst densities found in Northport Harbor during 2007 were more than an 418 order of magnitude lower than those found in the Gulf of Maine and the Bay of Fundy 419 (Anderson et al., 2005c), suggesting that cysts may be less important to bloom dynamics 420 This hypothesis is affirmed by comparing the density of cysts in in this system. 421 November 2007 to the abundance of cells in May 2008. The highest cyst densities in 2007 (50 cvsts cc<sup>-1</sup>) would yield a vegetative population of only 125 cells  $L^{-1}$  if all cvsts 422 423 in the top cm of sediment emerged successfully and simultaneously into the 4 m water 424 column. Since this cell abundance is four orders of magnitude smaller than vegetative cell densities observed in 2008 (10<sup>6</sup> cells L<sup>-1</sup>), in situ growth of vegetative cells likely 425 426 played an important role in the development of the 2008 bloom (Anderson 1998).

427 Meteorological conditions likely affected bloom dynamics in Northport Harbor. 428 Vegetative *A. fundyense* cells are known to grow maximally from 12 to 20 °C (Yentsch et 429 al., 1975; Anderson et al., 1983) and during 2007 and 2008, *A. fundyense* blooms 430 developed when Northport Harbor temperatures were between 10 and 20°C, with 431 temperatures close to 15°C yielding the highest cell densities. The spring of 2008 was 432 warmer than 2007 as during 2007, temperatures persisted between 15°C and 20°C for 433 only three weeks whereas in 2008, temperatures stabilized near 15°C for almost six weeks 434 (mid-April – June), giving the 2008 population more time to bloom. In contrast to early 435 spring, May 2008 temperatures were cooler than both May 2007 and the long term May 436 mean, which likely aided in keeping water temperatures in the optimal range for A. 437 fundvense growth allowing the large A. fundvense bloom to develop. In addition to 438 influencing pelagic cell dynamics, warmer temperatures during early spring 2008 likely 439 stimulated the germination of A. fundvense cysts (Anderson and Morel, 1979; Anderson, 440 1998) earlier than in 2007. Wind patterns may have also influenced the 2008 A. 441 fundvense bloom. During April of 2008, winds blew from the SE, whereas April 2007 442 winds came from the SW. While the SW winds in 2007 might have kept water within 443 Northport Harbor, winds in April 2008 may have spread cells throughout the Northport-Huntington Bay complex and thus may have contributed to the more widespread bloom 444 445 in that year. Atmospheric conditions such as wind direction have often been found to 446 control the spread and persistence of *Alexandrium* blooms (Anderson and Morel, 1979; 447 Garcon et al., 1986; Anderson, 1997; Townsend et al., 2005a,b).

448 N played a central role in supporting A. fundyense blooms in Northport Harbor. 449 During the 2008 bloom, there were significant (p < 0.01, t-test) declines in phosphate, 450 nitrate and ammonium concentrations during the A. fundyense bloom (6 May through 29 451 May) compared to before and after the bloom, suggesting that there was a larger nutrient 452 demand due to the higher biomass and more prolonged bloom in 2008. Furthermore, 453 nitrate concentrations were significantly (p<0.01, t-test) lower in 2008 ( $5.12 \pm 1.58 \mu$ M) 454 compared to 2007 (12.4  $\pm$  1.86  $\mu$ M) and ammonium concentrations were also lower in 455 2008 ( $0.58 \pm 0.17 \mu$ M; 6 May to 26 May) compared to 2007 ( $1.34 \pm 0.51 \mu$ M; 8 May to 5

June). These observations suggest N was more likely to be limiting to the *A. fundyense* bloom in 2008 compared to 2007. High biomass *A. taylori* blooms in the Mediterranean which are influenced by anthropogenic N loading have caused a drawdown of nutrients similar to that observed in Northport in 2008 (Penna et al., 2002).

460 Nutrient amendment experiments performed during 2007 and 2008 demonstrated 461 that N loading can affect A. fundyense densities and toxicity, and affirms that N was important in supporting the large 2008 bloom. Overall, the addition of N (glutamine, 462 nitrate, ammonium and/or urea) resulted in increased A. fundvense densities and/or toxin 463 464 concentrations compared to control treatments during every 2008 experiment. These 465 increases were frequently significant in 2008 (83% of experiments), when ambient 466 inorganic N concentrations were lower, suggesting this bloom was N stressed. On 467 average, the additions of ammonium and glutamine, specifically, resulted in the highest 468 A. fundyense densities and toxin concentrations when compared to the addition of other N 469 species when pooling together all experiments conducted in both 2007 and 2008. 470 However, the addition of ammonium most frequently yielded statistically significant 471 increases in A. fundyense densities and toxin concentrations compared to control treatments (66% and 50% of experiments in 2008), suggesting that ammonium may 472 473 promote the formation of toxic A. fundyense blooms. The strong response to glutamine 474 also suggests that dissolved organic N and amino acids such as glutamine may play an 475 important role in supporting *Alexandrium* blooms as they are known to do for other 476 HABs (Mulholland et al., 2002; Gobler et al., 2004).

The effects of nutrients on the 2008 *A. fundyense* bloom was also evident fromcell normalized toxin concentrations found in the field and during experiments. Variation

479 in toxin content per cell of natural bloom populations and isolates from the Gulf of Maine 480 has been previously attributed to nutrient limitation, with N limited cells generally 481 displaying lower levels of toxin (Anderson et al., 1990a,b; Poulton et al., 2005). During the 2008 A. fundyense bloom, cell toxicity was high (34.5 - 58.8 fmol STX eq. cell<sup>-1</sup>) at 482 483 the beginning and end of the bloom (April and June) but was significantly lower during the peak of the bloom (15.2±5.1 fmol STX eq. cell<sup>-1</sup>; 6 May - 29 May; p<0.001, Student-484 Newman-Keuls; Fig 9). Since values of  $51.9 \pm 29.5$  fmol STX eq. cell<sup>-1</sup> were measured in 485 486 nutrient replete cultures of A. fundyense strains (n=3) isolated from Northport, this field 487 pattern supports the hypothesis that A. fundyense populations were nutrient replete at the 488 end and beginning of the bloom, but nutrient stressed during May. Nutrient amendment 489 experiments displayed similar variations in toxin concentrations normalized per cell, with 490 significant increases in toxin per cell during experimental N loading in general and 491 ammonium loading in particular. The ability of ammonium to consistently increase 492 cellular toxin content has also been observed in A. tamarense cultures (Leong et al., 493 2004), supporting the hypothesis that ammonium promotes toxic A. fundyense blooms. 494 The decreases in toxin per cell during the bloom peak could indicate N-stress causing a 495 cellular partitioning of resources (Leong et al., 2004), with more N put toward growth 496 and less toward toxin production during the peak of the bloom since saxitoxin is a N-rich 497 molecule, containing 7 N atoms (with the decarbamoyl derivatives having 6 N atoms; 498 Samsur et al., 2006).

N played an important role in the development and toxicity of *A. fundyense*blooms in Northport, and the Scudder Beach Sewage Treatment Plant, which discharges
0.4 million gallons of effluent daily into Northport Harbor, may have been an important

502 N source which supported these blooms (discharge pipe at 40.8965°N, 73.3567°W, Fig.1; 503 Paul Harding, NYSDEC, personal communication). During periods when chlorophyll a 504 levels and presumably nutrient demands were low, DIN concentrations in Northport 505 Harbor frequently exceeded  $25\mu$ M, suggesting there is a strong source of N in this region. The active uptake of sewage-derived N was evident in the isotopic signatures of 506 particulate organic nitrogen (PON) from Northport Harbor as  $\delta^{15}$ N values ranged from 12 507 508 to 23‰ during large A. fundyense blooms. This range overlaps with wastewater derived 509 N (10 to 30 ‰; Kendall, 1998; Bianchi, 2007), and is significantly higher than levels 510 measured in particulate organic matter (POM) of the adjacent waters of Long Island 511 Sound (7 to 9 %). Furthermore, toxin and A. fundyense densities were significantly correlated to  $\delta^{15}$ N of POM (r<sup>2</sup>=0.63 and 0.68, respectively; p<0.001) indicating POM was 512 the most enriched in <sup>15</sup>N during bloom events. These findings, combined with the ability 513 514 of N enrichment to significantly increase the abundance and toxicity of A. fundyense 515 supports the hypothesis that N from the Scudder Beach wastewater treatment plant or 516 some other sources of highly enriched wastewater supported the proliferation of these 517 blooms. Similarly, anthropogenic nutrient loading has been associated with an increase 518 in PSP incidences caused by A. catenella in multiple marine ecosystems including 519 shallow, poorly flushed coastal embayments of the northwest US (Trainer et al., 2003).

Nutrient loading has been cited as a factor responsible for promoting multiple HABs around the world (Anderson et al., 2002; Penna et al., 2002; Trainer et al., 2003; Poulton et al., 2005; Glibert et al., 2006; Anderson et al., 2008; Heisler et al., 2008). However, the degree to which *A. fundyense* blooms are related to anthropogenic nutrient loading to coastal systems has been unclear (Anderson, 1994; Anderson et al., 2002, 525 2008; Glibert et al., 2005). This study demonstrated that N enrichment was capable of 526 significantly increasing A. fundyense cell densities, particulate toxin levels, and the levels 527 of toxin per cell. Moreover, the isotopic N signature of POM during blooms was 528 consistent with those found in wastewater. This data set combined with the proximity of 529 a sewage treatment plant to the occurrence of this bloom indicates that estuarine A. 530 fundyense blooms can be promoted by anthropogenic N loading. It is possible that 531 anthropogenic nutrient loading plays a similar role in the development of A. fundyense 532 blooms in coastal embayments around the world, although this phenomenon has not been 533 well studied.

534

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768	during May of 2007. Bars are means while error bars represent SD of triplicate					
769	measurements.					
770	<b>Figure 5.</b> Dynamics of: A) Pelagic toxin (pmol STX eq. $L^{-1} \times 10^{3}$ ), Alexandrium					
771	<i>fundyense</i> densities (cells $L^{-1} \times 10^6$ ) and toxin concentrations (µg STX eq. 100g <sup>-1</sup> )					
772	$x 10^3$ ) in deployed blue mussels ( <i>Mytilus edulis</i> ) as determined by mouse					
773	bioassay, off-scale values are indicated by the black arrow, B) size fractioned					
774	chlorophyll a ( $\mu$ g L <sup>-1</sup> ), and C) inorganic nutrient concentrations ( $\mu$ M) and					
775	temperature (°C) in Northport Harbor (site 2) during spring 2008. Points are					
776	means while error bars represent SD.					
777	<b>Figure 6.</b> Alexandrium fundyense densities (cells L <sup>-1</sup> ) and toxin concentrations (pmol					
778	STX eq. L <sup>-1</sup> ) following experimental nutrient amendments during April - June					
779	2008. Bars are means while error bars represent SD of triplicate & duplicate					
780	(saxitoxin concentrations) measurements. C= control, P= phosphate, N= nitrate,					
781	U= urea, A= ammonium (10, 20 and 40 indicate different concentrations added in					
782	$\mu$ M), G= glutamine, and A+P= ammonium + phosphate.					
783	<b>Figure 7.</b> Toxin per cell (fmol STX eq. cell <sup>-1</sup> ) following experimental nutrient					
784	amendments during April - June 2008. Bars are means while error bars represent					
785	SD of duplicate measurements. C= control, P= phosphate, N= nitrate, U= urea,					
786	A= ammonium (10, 20 and 40 indicate different concentrations added in $\mu$ M), G=					
787	glutamine, and A+P= ammonium + phosphate.					
788	<b>Figure 8.</b> Atmospheric temperatures (°C) observed during the winter and spring of 2007					
789	and 2008 compared to long term monthly means from Islip, NY, USA. Bars are					
790	monthly means while error bars represent SE.					
791	<b>Figure 9.</b> Alexandrium fundyense densities (cells $L^{-1} \times 10^{\circ}$ ) and toxin concentrations per					
792	cell (fmol STX eq. cell <sup>-1</sup> ) for Northport Harbor (site 2) in 2008. Points are means					
793	while error bars represent SD (error bars for toxin concentrations per cell					
794	represent propagated SD). The area highlighted in grey represents the range of					

total toxin concentrations per cell (fmol STX eq. cell <sup>-1</sup>) measured in nutrient
replete cultures of *Alexandrium fundyense* isolated from Northport Bay.

797	List of Tables
798 799 800 801 802 803 804	<ul> <li>Table 1. Peak Alexandrium fundyense densities (cells L<sup>-1</sup>) and pelagic toxin concentrations (pmol STX eq. L<sup>-1</sup>) in Northport- Huntington Bay, NY for 2007 (May 15<sup>th</sup>-30<sup>th</sup>) and 2008 (May 16<sup>th</sup>-26<sup>th</sup>), and mean cyst concentrations (cysts cc<sup>-1</sup>) in Northport-Huntington Bay, NY sediments during November of 2007 and 2008. Values in parentheses are standard deviations.</li> </ul>
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814 **Figure 1.** Site locations in Northport- Huntington Bay complex; located on the north

shore of Long Island, NY, USA. Cyst sampling locations include sites 1-17 whereas

- 816 pelagic samples were obtained from sites 1-8, 10, 11, 16 and LIS, SD= Sewage discharge
- 817 pipe from Scudder Beach Sewage Treatment Plant.
- 818



**Figure 2.** Dynamics of: A) Pelagic toxin (pmol STX eq.  $L^{-1}$ ) and *Alexandrium fundyense* densities (cells  $L^{-1} \times 10^2$ ), off-scale values are indicated by the black arrow, B) size fractioned chlorophyll *a* (µg  $L^{-1}$ ), and C) inorganic nutrient concentrations (µM) and temperature (°C) in Northport Harbor during spring 2007. Points are means while error bars represent SD.



**Figure 3.**  $\delta^{15}N$  (‰) values of particulate organic nitrogen from Northport Harbor during spring 2007 and 2008. The ranges of levels measured in particulate organic matter in Long Island Sound are depicted by the grey bar. Nitrogen from wastewater typically ranges from 10-30‰ (Kendall 1998, Bianchi 2007). Points are means while error bars represent SD.



**Figure 4.** Alexandrium fundyense densities (cells  $L^{-1} \times 10^2$ ) and toxin concentrations

839 (pmol STX eq. L<sup>-1</sup>) at the end of nutrient amendment experiments conducted during May

840 of 2007. Bars are means while error bars represent SD of triplicate measurements.







Alexandrium fundyense (cells L<sup>-1</sup>)



**Figure 6.** Alexandrium fundyense densities (cells  $L^{-1}$ ) and toxin concentrations (pmol) 849

STX eq. L<sup>-1</sup>) following experimental nutrient amendments during April - June 2008. 850

Bars are means while error bars represent SD of triplicate & duplicate (toxin 851

concentrations) measurements. C= control, P= phosphate, N= nitrate, U= urea, A= 852

- 853 ammonium (10, 20 and 40 indicate different concentrations added in  $\mu$ M), G= glutamine,
- 854 and A+P= ammonium + phosphate.



Toxin per cell (fmol STX eq. cell<sup>-1</sup>)

855 856 Figure 7. Toxin per cell (fmol STX eq. cell<sup>-1</sup>) following experimental nutrient

amendments during April - June 2008. Bars are means while error bars represent SD of 857 duplicate measurements. C= control, P= phosphate, N= nitrate, U= urea, A= ammonium 858

859 (10, 20 and 40 indicate different concentrations added in  $\mu$ M), G= glutamine, and A+P=

860 ammonium + phosphate.



864

Figure 8. Atmospheric temperatures (°C) observed during the winter and spring of 2007 and 2008 compared to long term monthly means in Islip, NY, USA. Bars are monthly means while error bars represent SE.



Figure 9. Alexandrium fundyense densities (cells  $L^{-1} \times 10^6$ ) and toxin concentrations per cell (fmol STX eq. cell<sup>-1</sup>) for Northport Harbor (site 2) in 2008. Points are means while error bars represent SD (error bars for toxin concentrations per cell represent propagated SD). The area highlighted in grey represents the range of total toxin concentrations per cell (fmol STX eq. cell<sup>-1</sup>) measured in nutrient replete cultures of *Alexandrium fundyense* isolated from Northport Bay. 

Table 1. Maximal *Alexandrium fundyense* densities (cells L<sup>-1</sup>) and pelagic toxin
concentrations (pmol STX eq. L<sup>-1</sup>) in Northport- Huntington Bay, NY for 2007 (May
15<sup>th</sup>-30<sup>th</sup>) and 2008 (May 16<sup>th</sup>-26<sup>th</sup>), and mean cyst concentrations (cysts cc<sup>-1</sup>) in
Northport-Huntington Bay, NY sediments during November of 2007 and 2008. Values
are means with standard deviations in parentheses.

	Northport-Huntington Bay					
Site	A. fundyense (cells L <sup>-1</sup> )		water column toxin (pmol STX eq. $L^{-1}$ )		A. fundyense cysts $(cc^{-1})$	
	2007	2008	2007	2008	2007	2008
1	50 (9)	7,166 (983)	3.73 (0.68)	183 (60.8)	3 (3)	25 (7)
2	2650 (81)	1,199,567 (435,248)	130 (3.90)	24,662 (564)	18 (12)	345 (35)
3	9 (4)	4,429 (578)	3.04 (0.14)	98.6 (0.57)	13 (10)	20 (14)
4	0 (0)	13,580 (2,623)	2.62 (0.06)	399 (31.8)	0	10 (14)
5	11 (8)	-	3.01 (0.31)	-	0	0
6	12 (0)	24,850 (1,072)	7.14 (0.66)	312 (22.7)	5 (7)	0
7	-	554,167 (41,908)	-	4,483 (11.3)	26 (4)	220 (28)
8	-	887,600 (352,422)	-	19,521 (3152)	50 (21)	745 (176)
9	-	-	-	-	20 (21)	285 (35)
10	-	31,675 (16,581)	-	379 (36.1)	3 (3)	115 (35)
11	-	14,733 (0)	-	449 (63.9)	8 (3)	75 (7)
12	-	-	-	-	1(1)	35 (21)
13	-	-	-	-	3 (3)	25 (7)
14	-	-	-	-	1(1)	35 (21)
15	-	-	-	-	0	30 (42)
16	-	28,178 (10,019)	-	335 (36.0)	0	15 (7)
17	-	-	-	-	0	10 (0)
LIS	-	8,244 (82)	-	422 (26.9)	-	-