Cyanobacterial Akinete Distribution, Viability, and Cyanotoxin Records in Sediment Archives From the Northern Baltic Sea

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Cyanobacteria of the order Nostocales, including Baltic Sea bloom-forming taxa Nodularia spumigena, Aphanizomenon flosaquae, and Dolichospermum spp., produce resting stages, known as akinetes, under unfavorable conditions. These akinetes can persist in the sediment and germinate if favorable conditions return, simultaneously representing past blooms and possibly contributing to future bloom formation. The present study characterized cyanobacterial akinete survival, germination, and potential cyanotoxin production in brackish water sediment archives from coastal and open Gulf of Finland in order to understand recent bloom expansion, akinete persistence, and cyanobacteria life cycles in the northern Baltic Sea. Results showed that cyanobacterial akinetes can persist in and germinate from Northern Baltic Sea sediment up to >40 and >400 years old, at coastal and open-sea locations, respectively. Akinete abundance and viability decreased with age and depth of vertical sediment layers. The detection of potential microcystin and nodularin production from akinetes was minimal and restricted to the surface sediment layers. Phylogenetic analysis of culturable cyanobacteria from the coastal sediment core indicated that most strains likely belonged to the benthic genus Anabaena. Potentially planktonic species of Dolichospermum could only be revived from the near-surface layers of the sediment, corresponding to an estimated age of 1–3 years. Results of germination experiments supported the notion that akinetes do not play an equally significant role in the life cycles of all bloom-forming cyanobacteria in the Baltic Sea. Overall, there was minimal congruence between akinete abundance, cyanotoxin concentration, and the presence of cyanotoxin biosynthetic genes in either sediment core. Further research is recommended to accurately detect and quantify akinetes and cyanotoxin genes from brackish water sediment samples in order to further describe species-specific benthic archives of cyanobacteria.

Keywords: cyanobacteria, resting stages, life cycles, microcystin, nodularin, resurrection, ndaF, mcyB
INTRODUCTION

Since the mid-twentieth century, anthropogenic eutrophication and climate variability have led to an increase of harmful cyanobacterial blooms in the Baltic Sea (Finni et al., 2001; Suikkanen et al., 2013; Funkey et al., 2014). Widespread cyanobacterial blooms now occur annually in the Baltic Sea and can negatively impact the local ecosystem, human health, and economy (Sivonen and Jones, 1999; Karjalainen et al., 2007; Burford et al., 2019). Despite the annual occurrence of cyanobacteria blooms, the life cycle of such bloom-forming cyanobacteria remains poorly understood (Hense and Beckmann, 2006; Suikkanen et al., 2010).

Cyanobacteria of the order Nostocales can produce resting stages, known as akinetes, under unfavorable conditions (Hori et al., 2003; Komárek, 2010). These akinetes persist in the sediment and germinate if favorable conditions return, simultaneously representing past blooms and contributing to the initiation of future blooms (Livingstone and Jaworski, 1980; Wood et al., 2009; Legrand et al., 2017b). Akinetes are characterized by their augmented size, thick cell walls, non-motility, and spheroid or oval shape (Kaplan-Levy et al., 2010; Sukenik et al., 2019). These characteristics lend toward the ability to withstand harsh environmental conditions. Therefore, akinetes often serve as an effective overwintering and recruitment strategy, similar to protist resting cysts, in temperate regions such as the Baltic Sea (Suikkanen et al., 2010). Many factors can influence akinete germination of different cyanobacteria species from benthic environments, including sediment resuspension, sufficient maturation time, light, temperature, oxygen, and nutrients (Huber, 1985; Adams and Duggan, 1999; Baker and Bellifemine, 2000; Hori et al., 2003; Kaplan-Levy et al., 2010; Myers et al., 2010).

Various techniques have been used to study sedimentary persistence, viability, and recruitment of cyanobacterial akinetes. Studies on cyanobacterial akinete sedimentation and records in the sediment have addressed akinete abundance, viability, and DNA from sediment samples in lakes (Livingstone and Jaworski, 1980; Wood et al., 2009; Legrand et al., 2016, 2017a,b, 2019) and brackish water environments (Suikkanen et al., 2010; Bormans et al., 2020). Via the incubation of sediment samples in culture media, germination experiments are commonly employed to assess the viability and recruitment conditions of akinetes (Livingstone and Jaworski, 1980; Wood et al., 2009; Suikkanen et al., 2010). Genetic methods have also been widely used to investigate cyanobacterial presence and potential cyanotoxin production in sediments by amplification of the 16S rRNA gene (e.g., Rinta-Kanto et al., 2009; Savichtcheva et al., 2011; Legrand et al., 2017a) and various cyanotoxin biosynthesis genes such as the microcystin synthetase encoding mcyA, mcyB, and mcyD (e.g., Rinta-Kanto et al., 2009; Savichtcheva et al., 2011; Monchamp et al., 2016; Legrand et al., 2017a, 2019; Pilon et al., 2019; Bormans et al., 2020) or ndaf, related to nodularin biosynthesis (e.g., Cegłowska et al., 2018). Recently, metabarcoding has been adopted to the selection of genetic tools used for the study of sediment cyanobacterial assemblages (e.g., Monchamp et al., 2016; Pilon et al., 2019; Yan et al., 2019). These studies have found akinetes to be viable up to more than a 1000 years after their sedimentation and identified the occurrence of microcystin (mcyB) and anatoxin (anaC) genes in 6,700 years old sediment layers (Legrand et al., 2019). Thus, sediments may represent archives of past blooms (Livingstone and Jaworski, 1980; Räsänen et al., 2006; Wood et al., 2009; Legrand et al., 2017b).

Plankton resting stages present in the sediment of aquatic environments create a reservoir of genetic diversity, possibly allowing for the adaptation to changing environmental conditions (Jones and Lennon, 2010; Kremp et al., 2016). Sediment archives of living resting stages and/or DNA are increasingly recognized as powerful tools for the study of climate change impacts on aquatic life (Ellegaard et al., 2020). In the Baltic Sea, the approach has been successfully used to trace the dynamics of dinoflagellates and their evolutionary responses to changing environmental conditions through the past century (Hinners et al., 2017; Kremp et al., 2018).

Baltic Sea cyanobacteria blooms are dominated by nostocalean, filamentous, and nitrogen-fixing cyanobacteria, represented by Nodularia spumigena, Aphanizomenon flosaquae, and Dolichospermum (ex. Anabaena) species. Baltic Sea strains of N. spumigena and certain Dolichospermum spp. produce the hepatotoxic nodularin and microcystins, respectively (Sivonen et al., 1989; Karlsson et al., 2005; Halinen et al., 2007). These cyanotoxins threaten the safety of drinking water and recreational water use due to their ability to damage liver cells in mammals, potentially causing death in acute doses (Sivonen and Jones, 1999). Understanding life cycle strategies, particularly akinete formation and recruitment, of these bloom-forming and potentially cyanotoxin-producing genera, provides insight into future bloom initiation and cyanotoxin outbreaks, which remains vital to enhancing bloom prediction and mitigation strategies (Suikkanen et al., 2010; Wasmund, 2017).

This study investigated cyanobacterial akinete distribution, viability, and potential for cyanotoxin production in >40 to >400-year-old sediment archives from the northern Baltic Sea in order to assess cyanobacteria bloom expansion, the extent of cyanotoxins in past blooms, and species-specific life cycle strategies. We hypothesized that akinete abundance and presence of cyanotoxins would correspond with degradation of akinetes over time and the expansion of cyanobacterial blooms in the northern Baltic Sea since the mid-twentieth century. Cyanotoxin gene examination and phylogenetic analysis of culturable cyanobacteria strains was used to further investigate species-specific life cycle strategies by associating germinated strains with known Baltic Sea strains. Furthermore, akinete survival and germination potential was studied, to better understand their function as long-term seed banks and species-specific life cycle strategies of cyanobacteria in the northern Baltic Sea. This project aimed to contribute to a greater understanding of cyanobacteria and akinete resilience, diversity, and viability in the fluctuating northern Baltic Sea environment.
MATERIALS AND METHODS

Sediment Core Collection and Processing
Sediment cores were obtained from the Gulf of Finland at a coastal site (35 m depth) near Tårnminne Zoological Station in July 2018 and at an open-sea site (approximately 100 m depth), LL7, in January 2019 (Figure 1) using a GEMAX gravity corer. Two replicate cores of approximately 40 cm length were taken at the coastal site, with one core reserved for sediment dating. A single core of 30 cm length (LL7-2019) was collected at the open-sea site. A replicate core was not taken from the open-sea site due to the availability of previous sediment dating data specific to the LL7 site (LL7-2015; Kremp et al., 2018). All cores were stored in their original tubes under cold and dark conditions until processing. Sediment layers were selected for analyses according to high dry to wet weight ratios, relative age, and for consistency across analyses.

Cores were extruded from sampling tubes using a piston and sliced into 1- to 2-cm layers. To minimize contamination, the outer 5 mm of each slice was removed using a small inner corer. Individual slices for dating were stored in plastic containers at 4°C and dark conditions. Individual slices for biological analysis were stored in plastic bags submerged in water and dark conditions in order to ensure the preservation of akinetes (Kremp et al., 2018). Wet and dry weight of sediment slices was determined by placing aliquots (5 mL) of wet sediment in crucibles, weighing them and drying them in an UNE 400 Memmert Universal oven at 70°C to remove air at 4°C and dark conditions in order to ensure the preservation of akinetes. Wet and dry weight of sediment slices was determined by placing aliquots (5 mL) of wet sediment in crucibles, weighing them and drying them in an UNE 400 Memmert Universal oven at 70°C for approximately 24 h, after which the mass of dried sediment samples was recorded.

Sediment Dating
Wet sediment slices of one coastal core reserved for gamma-spectrometric dating were weighed and freeze-dried for 3 days in a Christ Beta 2–8 LD plus freeze dryer at 1.0 mbar and −60°C. Dry material was then reweighed. Dry sediment was analyzed using gamma-spectrometry at the Finnish Radiation and Nuclear Safety Authority (STUK) Laboratory in Helsinki, Finland. The relative age of each 2-cm sediment layer was determined using Cs-137 radionuclide analysis and certified National Physical Laboratory reference samples. Markers for Cs-137 dating were peaks corresponding to calendar years 1963 and 1986. These peaks were produced by nuclear weapon testing, accidents, and the Chernobyl disaster.

Dating of sediment core LL7-2015 from the open-sea site was conducted for previous research using Cs-137 and Pb-210 concentrations, constant initial concentration and constant rate of supply models (Kremp et al., 2018). Cores LL7-2015 and LL7-2019 were correlated based on clear physical sediment features such as variations in water content and wet-to-dry-weight ratio.

Determination of Vertical Akinete Abundance
Akinetes present in the 20–63 μm fraction of wet sediment samples were quantified using light microscopy. Every 2- to 4-cm depth of sediment was counted from both cores to determine vertical akinete abundance. The 20–63 μm sediment fraction was selected because it likely contained the majority of cyanobacterial akinetes, which are on average one order of magnitude bigger than the average vegetative cell size (Adams and Duggan, 1999; Kaplan-Levy et al., 2010; de Tezanos Pinto et al., 2016). The 20–63 μm fraction also eliminated fine silt particles present in the 10–63 μm fraction. These particles were excluded to maximize light microscopy visibility and counting accuracy.

To obtain the 20–63 μm fraction, aliquots (2 mL) of sediment were diluted with 6 psu sterile-filtered seawater and sonicated on a constant duty cycle for 30 s at 20–40 kHz (Brandelin Sonoplus Sonicator HD 2200). The sonicated sediment slurry was sifted through 63 and 20 μm stainless steel mesh sieves with 6 psu sterile-filtered seawater. The remaining material on the 20 μm sieve was collected and stored in 15-mL polypropylene centrifuge tubes with 6 psu sterile-filtered seawater under dark conditions at 4°C. Akinetes from each triplicate sample were quantified using an inverted light microscope (Leica DMI 3000 B) with a 1-mL Gridded Sedgewick Rafter chamber (Wildlife Supply Company) at 200X magnification. Identification of akinetes followed Komárek (2013, 2016). A minimum of 400 cells (or all cells in the sample, if <400) were counted in each 1-mL sample. Recorded akinete counts and wet to dry weight sediment ratios were used to calculate the average abundance of akinetes present in dry sediment samples.

DNA Extraction From Sediment
Genomic DNA (gDNA) was extracted from selected wet sediment samples from the coastal and open-sea core corresponding with sediment layers containing akinete abundance data. Up to 250 mg of wet sediment was transferred to PowerBead tubes (Qiagen) with a sterile, disposable Pasteur pipette. The mass of all samples was recorded. Tubes were bead-beat using a FastPrep-24 5G (MP Biomedicals) sample disruption instrument for two cycles of 6 m s⁻¹ for 45 seconds with a 5 s break between cycles. DNA extraction was conducted using the DNaseasy PowerLyzer Power Soil Kit (Qiagen) according to the manufacturer’s instructions, eluted with 100 μL molecular biology grade water, and stored at −20°C. Concentration and purity of extracted DNA was verified (NanoDrop 2000, Thermo Fisher Scientific).

Cyanotoxin Extraction From Sediment and Detection by Non-competitive Immunoassay
Particulate cyanobacterial hepatotoxins microcystin and nodularin were measured from the 20–63 μm sediment fraction. Sediment core layers selected for hepatotoxin content analysis corresponded with those selected for determination of vertical akinete abundance. This included nine open-sea sediment core samples and 11 coastal sediment core samples.

From each selected sediment sample, 6 mL of wet sediment was sieved to obtain the 20–63 μm size fraction which was suspended in approximately 10 mL of sterile-filtered 6 psu seawater. This material was centrifuged (3,500 rpm for
10 min) and the supernatant was discarded. The resulting pellets were washed three times with 5 mL Milli-Q water to remove accumulated salts. Each pellet was suspended in 100 µL Milli-Q water, frozen at −20°C, and freeze-dried for approximately 12 h at 1.0 mbar and −60°C. Freeze-dried material was suspended in 1.2 mL 75% (v/v) methanol and sonicated in an ultrasonic bath for 15 min. The methanol extract was then transferred to a new vial for evaporation. The methanol extraction was repeated twice, and extracts were combined into one vial per sample. Methanol extracts were evaporated to dryness at room temperature under nitrogen flow for approximately 12 h. Samples were then reconstituted in 400 µL Milli-Q water and centrifuged to remove any remaining solids (3,500 rpg for 10 min). The supernatants were transferred into glass vials and stored at −20°C until used for detection of hepatotoxins by non-competitive immunoassay. Sample preparation procedure was based on methods previously described (Kankaanpää et al., 2001, 2009).

Non-competitive immunoassays were used to detect the presence of microcystins and/or nodularin in the sediment extracts. Each sample was initially analyzed with an assay that detects both microcystins and nodularin (Akter et al., 2016), with slight modifications. Using 25 µL well−1 of sample, the assay was performed in 75 µL well−1 reaction volume. The detection limit (expressed in microcystin-LR equivalents) of the immunoassay was 0.04 µg L−1, based on the average + 3 SD of the blank measurements. Similarly, each sample was also analyzed to detect any possible nodularin-R according to the method described by Akter et al. (2017) in reaction volume of 75 µL well−1 using 25 µL well−1 of sample. The detection limit of the nodularin-R immunoassay was 0.011 µg L−1. Finally, all hepatotoxin concentrations were converted to pg of hepatotoxin per g of dry weight. For more detailed description of the immunoassays, see Supplementary Data 1.

Cyanobacterial Akinete Germination and Establishment of Clonal Cultures

Live cyanobacterial akinetes present in sediment layers were enumerated and quantified using the serial dilution culture-most probable number (SDC-MPN) method (Throndsen, 1978). Three and eight sediment layers were selected for analysis from varying depths within the coastal and open-sea cores, respectively. Aliquots (3 mL) of wet sediment were initially diluted 1:10 in 6 psu sterile-filtered seawater and sonicated on a constant cycle at 20–40 kHz for 30 s to separate akinetes from sediment particles. Sonication of the sediment slurry was completed on ice to approximately maintain the 4°C sediment storage conditions. The original 1:10 subsample was then serially diluted by 1:10 seven times, with 2 mL of the previous dilution and 18 mL sterile, nitrogen-free, 6 psu Z8xS media (Kotai, 1972). Only 1:103 through 1:107 dilutions were further incubated and each of these dilutions contained five replicates.

Apart from four open-sea core sediment layers, this procedure was completed in duplicate for all samples to undergo two temperature treatments, 4 and 16°C. These temperature conditions were used to simulate winter/early spring and summer conditions in the northern Baltic Sea. The open-sea core sediment layers 8–9, 10–11, 12–13, and 14–15 cm were only incubated in 16°C. Incubating samples were stored in 50-mL sterile tissue culture flasks under the selected temperature treatment and constant light conditions, approximately 100 µmol photons m−2 s−1.

FIGURE 1 | Map of the study area showing the location of the sediment core collection sites.
Sediment slurry incubations were checked for cyanobacteria germination after approximately 4 and 8 weeks using an inverted light microscope with a 1-mL Sedgewick Rafter chamber at 200X magnification. The presence or absence of any cyanobacteria along with the specification of Nostocales taxa was recorded. Most probable number calculations were completed using the presence numbers recorded from the 1:10^3 to 10^5 dilution replicates and Thronsden (1978) Most Probable Number table. As the MPN calculations only required three dilutions, the 1:10^6 and 1:10^7 were excluded.

Identification of single filaments with morphological characteristics of Dolichospermum and Nodularia spp. followed Komárek (2013). Selected single filaments were isolated and cultured from selected SDC-MPN samples of the coastal sediment core containing these taxa. Approximately 1 mL of the germinated SDC-MPN sample was transferred to a 3-mL sedimentation chamber. The sample was examined for Dolichospermum and Nodularia spp. filaments using an inverted light microscope at 200X magnification. Single filaments of interest were selected using a sterile glass micropipette, washed three times with Milli-Q water, and transferred to a 24-well culture plate (Eppendorf) filled with 1:3 dilution of sterile, 6 psu Z8xS media and sterile-filtered 6 psu seawater. Well plates were then incubated at 16°C under constant light conditions, approximately 100 µmol photons m^{-2} s^{-1}.

**DNA Extraction, 16S rRNA Gene Amplification and Phylogenetic Analysis of Clonal Cultures**

Ten successful clonal cultures were selected for further analysis. Cells were collected approximately 48 h after the cultures were inoculated with fresh 6 psu Z8xS culture medium. A 10 mL subsample of each selected culture was concentrated (<100 mg), pelleted and then ground. gDNA was extracted using a Thermo Scientific GeneJET Plant Genomic DNA Purification Mini Kit according to the manufacturer's instructions and stored at −20°C. DNA concentration and quality was determined spectrophotometrically (Nanodrop 2000).

PCR amplification of cyanobacterial 16S rRNA gene was conducted in 20 µL reactions containing 2 µL template (gDNA or molecular biology grade water for negative control), 1X Phusion HF Buffer, 0.4 U Phusion DNA Polymerase, 0.2 mM dNTP mix (all reagents Thermo Scientific), 0.4 µM 16S27F and 23S30R primers (Table 1; all primers manufactured by Integrated DNA Technologies). Thermal cycling (C1000 Thermal Cycler, Bio-Rad) was carried out as follows: 98°C, 30 s followed by 30 cycles of 98°C, 5 s; 66°C, 30 s; 72°C, 30 s followed by final 72°C for 5 min. Amplification success was verified using gel electrophoresis (Invitrogen E-Gel, General Purpose Agarose Gel, 1.2% w/v agarose; Invitrogen E-Gel Power Snap Gel Electrophoresis Device). PCR products were purified using the QIAquick PCR Purification Kit (Qiagen) and the quantity and quality of purified PCR products were checked using the Qubit fluorometer and a Qubit dsDNA HS kit (Thermo Scientific), both according to the respective manufacturers’ instructions.

Sanger sequencing of the 16S rRNA gene was performed by Macrogen Inc. using the primers listed in Table 1. The sequences were trimmed with a phred score cutoff of 30, assembled in Unipro UGENE v36 (Okonechnikov et al., 2012) and are available under accessions MW491275-MW491284. Similar Anaabaena/Dolichospermum sequences (0.0 e-value, > 90% identity) were identified using the NCBI BLAST (Altschul et al., 1990) and downloaded from the GenBank nr database. An alignment was prepared in UGENE, redundant (100% identical) sequences were removed and phylogenetic analyses were conducted in MEGAX (Kumar et al., 2018; Stecher et al., 2020). Two Microcystis aeruginosa strains were selected as an outgroup. Maximum Parsimony, Maximum Likelihood, and Neighbor Joining methods were used with the Tamura-Nei model (Saitou and Nei, 1987; Tamura and Nei, 1993; Nei and Kumar, 2000).

**Detection of 16S rRNA Gene and Cyanotoxin Genes in Sediment and Clonal Cultures**

Cyanobacterial 16S rRNA gene, hepatotoxin biosynthesis genes mcyE/ndaF, and nodularin biosynthesis gene ndaF were amplified using DNA extracted from selected sediment layers and clonal cultures. gDNA, extracted as described in section “DNA Extraction, 16S rRNA Gene Amplification and Phylogenetic Analysis of Clonal Cultures,” from Anaabaena cylindrica PCC 73105, and Microcystis aeruginosa PCC 7005 were used as negative controls; Nodularia sp. PCC 7804 was used as a positive control for nodularin; Microcystis aeruginosa PCC 7820, and Dolichospermum flos-aquae NIVA-CYA 267/4 were used as positive controls for microcystin. The strains were obtained from the Pasteur Culture Collection of Cyanobacteria and the Norwegian Culture Collection of Algae. Culture conditions and cyanotoxin data have been described earlier by Hautala et al. (2013).

The 16S rRNA gene PCR reactions were completed as described above in section “DNA Extraction, 16S rRNA Gene Amplification and Phylogenetic Analysis of Clonal Cultures.” The mcyE/ndaF PCR reactions were conducted in 20 µL volumes containing 8 µL template (DNA or molecular biology grade water for negative control), 1X Phire Hot Start II DNA Polymerase, 0.2 mM dNTP mix (all Thermo Fisher Scientific) and 0.4 µM of HEFP and HEPR primers (Table 1). Thermal cycling was carried out as follows: 98°C, 30 s followed by 30 cycles of 98°C, 5 s; 66°C, 5 s; 72°C, 20 s followed by a final 72°C, 1 min.

The ndaF PCR reactions were conducted as described above for mcyE/ndaF except with 0.4 µM of ndaF8452 and ndaF8640 primers (Table 1) and an annealing temperature of 58°C. Amplification success was examined using gel electrophoresis as described above in section “DNA Extraction, 16S rRNA Gene Amplification and Phylogenetic Analysis of Clonal Cultures.”

In order to delineate between potential microcystin and nodularin producers, the presence of the microcystin biosynthesis gene mcyB in sediment DNA samples was examined using PCR with primers mcyBHF03A and mcyBHR04A (Table 1).
TABLE 1 | Primer sequences, target sites, and respective references.

| Primers  | Target genes                      | Sequence (5′–3′)                          | References                       |
|----------|-----------------------------------|-----------------------------------------|----------------------------------|
| 16S27F   | 16S rRNA                          | AGAGTTTGATCCTGGCTCAG                    | Taton et al., 2003; Rajaniemi-Wacklin et al., 2005; Hrouzek et al., 2005 |
| 23S30R   |                                   | CTCGCGCTCGTGGCGCTAGGT                   |                                  |
| 16S979F  |                                   | CGATGCACACGGCGAAGAAC                    |                                  |
| 16S544R  |                                   | ATCCGCGGATACCCGCTGCG                   |                                  |
| 16S1029R |                                   | GGGCTGCGGGCGGCTGCGG                    |                                  |
| HEPF     | Microcystin/nodularin synthetase   | TITGGCGGTAATTTTATTGTGGCATTGCT           | Jungblut and Neilan, 2006        |
| HEPR     | genes mcyE/ndaF                    | AAATCTTGGACACTAGTGAATCGGCTTT            | Koskenniemi et al., 2007         |
| ndaF8452 | Nodularin synthetase gene ndaF    | GTGATGTGAATTTGCTGGCG                    | Koskenniemi et al., 2007         |
| ndaF8640 |                                   | GGAAATTTCTATGTCTGACTC                   | Koskenniemi et al., 2007         |
| mcyBHR03A| Microcystin synthetase gene mcyB  | GTTTTAATACGAAAGCTGTTTTAAGTGGC          | Hautala et al., 2013; mcyBHR04A modified from mcyBHR04 |
| mcyBHR04A|                                   | CTGGTGACCTCTGACTGAAAGACTGACT            |                                  |

in the same way as described above for mcyE/ndaF and ndaF, with the following modifications: 2 µL template was used, the thermal cycling protocol was modified to consist of 35 cycles, and annealing was performed at 64°C. The specificity of the mcyBHR04A reverse primer, modified from a previously published primer mcyBHR04 (Hautala et al., 2013) by changing two nucleotides was assessed using gDNA from the culture collection strains listed above and found to yield the same results as reported in the original publication.

Statistical Analysis

Statistical analysis was conducted using R 4.0.2 software (R Core Team, 2020) and RStudio (version 1.3.1093; RStudio Team, 2020). Spearman’s rank correlation was used to investigate the relationships between sediment depth, median sediment layer age, average akinete abundance, and hepatotoxin concentration. The significance level was set at $p < 0.05$.

RESULTS

Sediment Chronology

Sediment chronology was determined in order to estimate sediment and akinete age. The Cs-137 concentrations indicated that the dates of the vertical layers of the coastal sediment core ranged from late 2018 to 1977 ± 4, which corresponds with ages up to 42 ± 4 years old (Figure 2).

The open-sea site was dated previously (Kremp et al., 2018) using Pb-210 and Cs-137. Reliable age estimation extended to 16 cm depth due to the small amount of unsupported Pb-210 in the 16–18 cm region (Kremp et al., 2018). The sediment age at 16 cm depth was 105 ± 8 years, corresponding to calendar years 1914 ± 8. This is equivalent to the depth of 12 cm of the core LL7-2019 (Table 2, Supplementary Table 1, and Supplementary Figure 1), which has a slightly slower sedimentation rate (mean sedimentation rate 0.12 cm year$^{-1}$) compared to the sedimentation rate of the core LL7-2015 (0.17 cm year$^{-1}$; Kremp et al., 2018). Beyond the dated section, ages were estimated assuming a constant linear sedimentation rate, using a sedimentation rate of 0.05 cm year$^{-1}$ based on the sedimentation rate of the deepest successfully dated section. Thus, it must be kept in mind that sediment ages below the depth of 12 cm (1914 AD) are only rough estimations.

Vertical Akinete Abundance and Cyanobacterial 16S rRNA Genes

Only intact and distinct cyanobacterial akinetes were counted to determine vertical akinete abundance. All cyanobacterial akinetes were counted, regardless of suspected species. Akinetes resembling Nodularia spp. were only noted in the surface layers (2–4 and 6–8 cm) of the coastal core; the rest of the akinetes found were of the Anabaena/Dolichospermum type (Komárek, 2013, 2016). The 2–4 cm depth was considered the surface layer of the coastal core due to the high water content (15.2 wet to dry weight ratio) of the 0–2 cm layer, which was approximately three times the average of all the layers.

Vertical akinete abundance in the coastal sediment core ranged from approximately 22,000 to 70 akinetes g$^{-1}$ dry weight...
TABLE 2 | Most probable number (MPN) estimates of viable akinete abundance (akinetes g\(^{-1}\) dry weight) in three coastal and eight open-sea core sediment layers based on germination presence or absence after 4- and 8-week incubation periods.

| Site         | Depth (cm) | Approximate dates | Taxa                          | MPN 4\(^{\circ}\)C | MPN 16\(^{\circ}\)C |
|--------------|------------|-------------------|-------------------------------|---------------------|--------------------|
|              |            |                   |                               | 4 weeks 8 weeks     | 4 weeks 8 weeks    |
| Coastal      | 2–4        | 2015–2017          | Anabaena/Dolichospermum spp.  | 313 57,000+         | 57,000+ 57,000+    |
|              |            |                   | Nodularia spp.                | 0 6,013             | 0 178              |
|              | 20–22      | 1998–2000          | Anabaena/Dolichospermum spp.  | 2,270 14,917        | 38,000+ 8,755      |
|              |            |                   | Nodularia spp.                | 0 389               | 178 178            |
|              | 38–40      | 1975–1977          | Anabaena/Dolichospermum spp.  | 271 149             | 420 4,742          |
|              |            |                   | Nodularia spp.                | 0 112               | 0 189              |
| Open sea     | 2–3        | 2008–2011          | Anabaena/Dolichospermum spp.  | 48 8,408            | 12,811 12,972      |
|              |            |                   | Nodularia spp.                | 0 1,177             | 1,041 769          |
|              | 8–9        | 1965–1971          | Anabaena/Dolichospermum spp.  | – –                 | 107 1,239          |
|              |            |                   | Nodularia spp.                | – –                 | 0 0               |
|              | 10–11      | 1932–1952          | Anabaena/Dolichospermum spp.  | – –                 | 154 2,173          |
|              |            |                   | Nodularia spp.                | – –                 | 0 0               |
|              | 12–13      | 1897–1916          | Anabaena/Dolichospermum spp.  | – –                 | 261 261            |
|              |            |                   | Nodularia spp.                | – –                 | 0 0               |
|              | 14–15      | 1857–1876*         | Anabaena/Dolichospermum spp.  | – –                 | 354 354            |
|              |            |                   | Nodularia spp.                | – –                 | 0 0               |
|              | 16–17      | 1816–1836*         | Anabaena/Dolichospermum spp.  | 0 0                 | 6.5 358            |
|              |            |                   | Nodularia spp.                | 0 0                 | 47 0              |
|              | 20–21      | 1737–1756*         | Anabaena/Dolichospermum spp.  | 0 0                 | 0 0               |
|              |            |                   | Nodularia spp.                | 0 0                 | 0 0               |
|              | 28–29      | 1577–1596*         | Anabaena/Dolichospermum spp.  | 0 0                 | 0 0               |
|              |            |                   | Nodularia spp.                | 0 0                 | 0 0               |

The open-sea core layers 8–9, 10–11, 12–13, and 14–15 cm were only incubated at 16\(^{\circ}\)C. Numbers with an indefinite MPN value (e.g., 57,000+) indicate all dilution series replicates showed growth for that taxa. The open-sea 2–3 cm layer 16\(^{\circ}\)C 4-week incubation MPN estimate for Aphanizomenon sp. was 5,765 akinetes g\(^{-1}\) dry weight. The dates below 13 cm (*) are estimations based on a constant sedimentation rate.

in all examined sediment layers (Figure 3A). The highest akinete abundance (22,000 akinetes g\(^{-1}\) dry weight) was determined from the 18–20 cm layer, with an age corresponding to roughly 16–18 years. Aside from this maximum akinete abundance, the vertical abundance decreased from the surface layer abundance (5,200 akinetes g\(^{-1}\) dry weight) with depth and age with minimal variation. The oldest intact akinetes measured from the sediment core were 42 ± 4 years old. Spearman’s correlation analysis indicated a significant negative correlation between akinete abundance and both sediment depth and age, which yielded the same value ($\rho = -0.82$, $p = 0.003$).

Similarly, vertical akinete abundance in the open-sea core ranged from zero to approximately 21,000 akinetes g\(^{-1}\) dry weight in all examined layers of sediment (Figure 3C). The greatest akinete abundance occurred at the surface layer (0–1 cm, 0–6 years old) and decreased with depth and age. Within the accounted for 20–63 μm sediment fraction, no akinetes were present in any replicate at 12–13, 20–21, or 28–29 cm depth. The oldest intact akinetes measured directly from sediment in the open-sea core were from the depth of 18–19 cm, i.e., up to 240 years old. Spearman’s correlation analysis indicated a significant negative correlation between akinete abundance and both sediment depth and age ($\rho = -0.92$, $p < 0.001$).

Overall, akinete abundance was maximal in the open-sea sediment core at the surface and decreased with depth and age; whereas, in the coastal sediment core, the maximal akinete abundance occurred at the mid-depth (18–20 cm) of the core, though the akinete abundance otherwise decreased with depth and age.

DNA extraction from sediment yielded >4 ng/μL with an average $A_{260}/A_{280}$ of 1.74. All sediment layers of both cores showed positive results for the presence of cyanobacterial 16S rRNA genes (Supplementary Table 2).

**Cyanotoxin Concentration and Biosynthesis Genes in Sediment**

Cyanobacterial cyclic peptide hepatotoxins (total microcystins and nodularin) were present in the 20–63 μm sediment fraction from sediment layers of both cores, even the deepest sediment layers (Figures 3B,D). For the coastal sediment core, six out of 11 analyzed samples contained concentrations above the detection limit, ranging from 2.9 to 968 pg g\(^{-1}\) dry weight. The sediment sample with the highest hepatotoxin concentration was the 30–32 cm-layer, which corresponds to calendar years 1985 to 1987. For the open-sea sediment core, eight out of nine analyzed samples contained concentrations above the detection limit, ranging from 2.7 to 1,137 pg g\(^{-1}\) dry weight. The sediment sample with the highest hepatotoxin concentration was the 8–9 cm-layer, which corresponds to calendar years 1965 to 1971.
All the samples were also analyzed by the nodularin-specific assay to detect any presence of nodularin-R in the samples. Nodularin was only detected in one of the 20 sediment samples examined: the 30–32-cm coastal sediment core layer was weakly positive for the presence of nodularin (0.12 µg L⁻¹ by nodularin-specific immunoassay, equivalent to 8.7 pg g⁻¹ dry weight), just above the detection limit according to the assay. Hepatotoxin concentrations did not correlate significantly with sediment depth, age, or akinete abundances at either the coastal or the open sea site (p > 0.05 for all).

Microcystin and nodularin biosynthesis genes mcyB and ndaF were only detectable in the 2–4 cm surface layer of the coastal sediment core (Figure 3A and Supplementary Table 2). Amplification of mcyE/ndaF was also observed in this layer. The mcyB gene was also detected at the open sea sediment core 0–1 and 2–3 cm layers (Figure 3C and Supplementary Table 2).

**Akinete Germination and Identification of Germinated Filaments**

After an 8-week incubation period at 16°C, viable akinetes had germinated from all tested sediment layers in both sediment cores (Table 2 and Figure 4). A greater number of viable akinetes germinated from the coastal sediment core than open-sea core, despite a higher akinete abundance in the upper layers of the open-sea core (Figures 3A,C). *Nodularia* spp., *Anabaena/Dolichospermum* spp., and *Aphanizomenon* sp. filaments germinated from at least one sediment layer. Germinated *Aphanizomenon* sp. filaments were only present...
in the 2–3 cm layer of the open-sea core in 16°C after the 4-week incubation period (MPN estimate 5,765 akinetes g⁻¹ dry weight). Generally, fewer Anabaena/Dolichospermum spp. akinetes germinated under 4°C than 16°C conditions, excluding the four open-sea core sediment layers only incubated at 16°C.

In both the coastal and open-sea sediment cores, more Anabaena/Dolichospermum spp. akinetes germinated than akinetes belonging to the genus Nodularia. Nonetheless, the 2–4 cm coastal sediment core layer had the highest MPN viable akinete estimates for both Anabaena/Dolichospermum and Nodularia genera. Germinated Anabaena/Dolichospermum spp. filaments were present in samples from each examined sediment layer from both sediment cores. In the three deepest layers of the open-sea core, Anabaena/Dolichospermum spp. did not germinate in 4°C conditions. Germinated Anabaena/Dolichospermum spp. filaments were present in 16°C incubation samples even in the deepest layers of the open-sea core, dating back to the late eighteenth century. Of the coastal core sediment samples, Nodularia spp. filaments had generally germinated after 8 weeks of incubation. In the open-sea core, germinated Nodularia spp. filaments were only present in the 2–3 and 16–17 cm layers, the latter dating back to the late nineteenth century; whereas, in the coastal core, they were present in each examined layer after 8 weeks.

Germinated cyanobacteria filaments from the coastal sediment core were successfully grown in clonal culture. Nine suspected Anabaena or Dolichospermum sp. strains (Figures 4A–D,F) and one Nodularia-like strain (Figure 4E) were selected for Sanger sequencing and phylogenetic analysis. The Nodularia-like 20–22 cm strain showed a positive result for the mcyE/ndaF genes and a weak positive result for the ndaF gene. No amplification of either mcyE/ndaF or ndaF alone was observed in the other putative Anabaena/Dolichospermum strains.

The Maximum Parsimony, Maximum Likelihood, and Neighbor Joining phylogenetic trees showed consistent topology, and therefore only the Maximum likelihood tree is shown (Figure 5). The cyanobacteria strains isolated from the coastal sediment core separated into distinct groups. Two well-supported clades (bootstrap values 98 and 95%, respectively) consisted of Nodularia spp. (Figure 5, clade 4) or benthic Anabaena spp. (Figure 5, clade 3). A larger clade consisting of Anabaena spp. such as Anabaena sp. strain BIR272, BIR169 and Anabaena cf. cylindrica XP6B (Halinen et al., 2008) with often similar morphology but with both benthic and planktonic lifestyles, and a planktonic Dolichospermum sp. strain was further divided into smaller subclades that included the remaining strains isolated in this study (Figure 5, clades 1 and 2).

Clade 1 contained suspected Anabaena or Dolichospermum sp. strains from the 20–22 and 38–40 cm-layers. The support for this cluster was moderate (65%), and the strains were found to be most closely related to one another, and then to non-toxic but planktonic Anabaena sp. (BIR272, BIR169). The second clade (Figure 5, clade 2) included putative Anabaena or Dolichospermum sp. strains from the sediment surface (2–4 cm-layer strains 1 and 2) as well as planktonic Anabaena and Dolichospermum strains. The strains isolated in this study were most closely related to planktonic Anabaena sp. BIR50, with good bootstrap support (97%). Anabaena sp. BIR50 has a 16S rRNA sequence identical to other Gulf of Finland strains (BIR19, BIR28, BIR30, BIR50, BIR232, BIR300, Halinen et al., 2008) which were initially identified as closely related to the sequences from this study but omitted from the analysis as redundant. D. circinale strain 1tu33s12, also included in clade 2, has been shown to harbor mcy genes (Rajaniemi et al., 2005). However, none of the strains from this study were very closely related to the planktonic, toxin-producing taxa isolated from the Baltic Sea and various lakes in Finland (Lyra et al., 2001; Rajaniemi et al., 2005).

The third clade comprised suspected Anabaena or Dolichospermum sp. strains from all three examined sediment core layers, including 2–4 cm-layer strain 3 (Figure 4C), 20–22 cm-layer strain 1 (Figure 4D), and 38–40 cm-layer strain 3, and Anabaena oscillarioides strains (BECID22, BECID32) isolated from epiphytic and epilithic habitats (Rajaniemi et al., 2005; Halinen et al., 2008; Shishido et al., 2015). Overall, this branch was well supported (95%).

In the fourth clade, the Nodularia-like 20–22 cm-layer strain 2 (Figure 4E) formed a well-supported (93%) cluster with N. sphaerocarpa strains (HKVV and Up16f) over other N. spumigena strains (309, AV63, HEM). N. sphaerocarpa strains are non-toxic and lack gas vesicles, though N. sphaerocarpa Up16f was isolated from the planktonic environment (Lehtimäki et al., 2000; Lyra et al., 2005).

DISCUSSION
Vertical Akinete Abundance and Germination
This is the first study to investigate benthic archives of planktonic cyanobacterial akinetes in the Baltic Sea. Thus far, other studies hindcasting the presence of cyanobacteria in the Baltic Sea have focused only on cyanobacterial genetic and chemical markers (Bianchi et al., 2000; Funkey et al., 2014; Cegłowska et al., 2018). In this study, vertical cyanobacterial akinete abundance and germination were used to determine the persistence, viability, and species diversity of cyanobacterial akinetes in coastal and open sea regions of the northern Baltic Sea.

In the coastal sea core, the presence of cyanobacterial akinetes throughout the vertical sediment layers indicates that cyanobacteria have been continually present in the archipelago zone of the southern coast of Finland since the mid-1970s. Akinete abundance of the upper layer (2–4 cm) was roughly four times less than that of the open-sea core, which may be a result of differences in bloom magnitudes between both sites but possibly also reflect the timing of sampling. The settled late summer and early autumnal bloom was not accounted for in 2018, a significant bloom year, as coastal samples were collected well before the annual bloom started to settle. The highest akinete abundance occurred at 18–20 cm, corresponding with calendar years 2000 ± 2 to 2003 ± 2. This peak akinete abundance may represent the above average cyanobacterial blooms in the Gulf of Finland during summers of 1999 and 2002 (Kownacka et al., 2020).
In the open-sea sediment core, the presence of cyanobacterial akinetes in the depths of the core corroborates that cyanobacteria were present in open waters of the Gulf of Finland in as early as the late eighteenth century (Finni et al., 2001; Zillén and Conley, 2010; Hällfors et al., 2013). Greater akinete abundance in the upper 11 cm, dating back to the calendar year 1952, is consistent with the expansion of cyanobacterial blooms in the open Baltic Sea since the mid-twentieth century (Finni et al., 2001; Zillén and Conley, 2010; Suikkanen et al., 2013). In an investigation of Baltic Sea sediment cores, hypoxic conditions in the Baltic Sea beginning in the mid-twentieth century were linked to elevated cyanobacteria abundance (Funkey et al., 2014). The high akinete abundance (ca. 21,000 akinetes g\(^{-1}\) dry weight) of the surface layer, approximately double that of the subsequent 2–3 cm-layer, corresponds with the highest intensity bloom of the current decade in summer 2018 (Kownacka et al., 2020). At sediment depth below 11 cm a sudden drop in akinete abundance as well as MPN of viable akinetes indicates a change in deposition patterns, less intense blooms and/or reduced preservation. Generally, caution should be exercised when looking at the individual dates of the open-sea sediment core, because the dates are obtained by correlation with an earlier core from the same site. However, the dating and correlation confirm that sediment succession at the site is unmixed and the precision of the dates is adequate for the present research questions.

Under at least one of the incubation conditions for each sediment layer, the MPN viable akinete estimates exceeded akinete counts in both the open-sea and coastal sediment cores. This inconsistency is perhaps unsurprising, given that measuring akinete abundance in sediments has been cited as challenging (Pham and Utsumi, 2018; Bormans et al., 2020). This discrepancy could be the result of counting akinetes only from the 20–63 \(\mu\)m fraction and the MPN method bias toward culturable strains. Akinete size has not been extensively studied and tends to be quite variable, ranging from 4 to 40 \(\mu\)m diameter and up to ten times the size of a vegetative cell (Kaplan-Levy et al., 2010; de Tezanos Pinto et al., 2016; Sukenik et al., 2019). As such, some akinetes were likely lost in the less than 20 \(\mu\)m fraction, but still present in the SDC-MPN sediment slurries. Implementing an akinete-specific stain, such as SYTOX-green used by Legrand et al. (2016) or the CARD-FISH method by Ramm et al. (2012), might have helped in akinete enumeration.

Despite the inconsistency between akinete counts and MPN estimates, both values decreased with sediment age and depth in both sediment cores. This suggests that akinetes present throughout each core remained viable, but the number of akinetes and viability decreased considerably with depth. Previous studies have successfully identified akinetes from sediment samples up to 6,700 years old and germinated akinetes up to 1,830 years old from relatively shallow (<30 m) lake environments (e.g., Livingstone and Jaworski, 1980; Wood et al., 2009; Legrand et al., 2017a, 2019). Therefore, it remains plausible that akinetes from the open Gulf of Finland could be intact and viable up to >400 years old, given germination took place in even the deepest layers after 8 weeks. Only Legrand et al. (2019) has quantified akinetes and tested their ability to germinate over a 1000-year time period. Of the akinetes identified as 318 ± 40 years old in the respective study, only 6–8% germinated and were therefore considered viable, in comparison with 70% for surface sediment (Legrand et al., 2019). This decreasing rate of germination with sediment depth and age has been further corroborated by other studies and corresponds with the decrease seen in the MPN viable akinete estimates (Tsujimura and Okubo, 2003; Legrand et al., 2017b, 2019).
FIGURE 5 | Maximum likelihood phylogenetic tree of the cyanobacterial 16S rRNA gene including the Anabaena sp. strains and one Nodularia sp. isolated in this study (shown in bold). The GenBank accession numbers are included in parenthesis with each strain. The 1196 nt alignment consisted of 28 non-redundant sequences. Bootstrap values (500 replicates) ≥ 70 are shown next to the nodes. Baltic Sea strains, other than the strains from this study, are indicated with a circle with no fill, known cyanotoxin producers (microcystins or nodularin) are indicated with filled circles. A positive result from the hepatotoxin biosynthesis gene (mcyE/ndaF) and ndaF PCRs (this study) is indicated with a triangle.

Various environmental factors affect cyanobacterial akinete germination, including sediment resuspension, temperature, salinity, nutrient availability, and light (Kaplan-Levy et al., 2010; Sukenik et al., 2019). The open-sea location has been reported as continually hypoxic and stratified for many decades (Raateoja and Setälä, 2016). The coastal site, on the other hand, receives a higher degree of land-based and riverine input and undergoes seasonal hypoxia and vertical mixing (Lyra et al., 2005; Gammal et al., 2017). Sediment resuspension and mixing caused by wind and wave action, such as upwelling and vertical mixing reaching up to 100 m depth, likely occurs at both sampling sites (Stipa, 1999; Suikkanen et al., 2010). Therefore, it remains feasible that akinetes from surface layers of both sediment cores could contribute to bloom initiation.

The akinete counts and MPN viable akinete estimate data indicated that akinetes play a more significant role in the life cycle of Anabaena/Dolichospermum spp., when compared with Nodularia and Aphanizomenon spp., as postulated in other studies (Livingstone and Jaworski, 1980; Suikkanen et al., 2010; Wasmund, 2017). However, the high MPN viable akinete estimates for Anabaena/Dolichospermum spp. throughout both cores are possibly due to the fact that they include both benthic and planktonic akinetes and that the MPN method favors culturable strains, such as benthic Anabaena spp.

Nodularia spp. akinetes were only observed in the upper layers (2–4 and 6–8 cm) of the coastal sediment core, which was in line with the positive hepatotoxin and ndaF PCR results for these layers. The negative results for the remainder of the sediment core layers, for both the coastal and open-sea sites, indicate a true absence of N. spumigena akinetes, as all strains are known to carry the ndaF gene (Laamanen et al., 2001; Moffitt et al., 2001). Based on the nodularin-specific assay all samples except for one coastal sediment core sample were negative for presence of any detectable nodularin, indicating that the hepatotoxins detected in the generic assay were predominantly microcystins. On the other hand, the very weak positive result...
of the open-sea sediment core. When compared with the biovolume germinated filaments, present only in the upper layer (2–3 cm) (Suikkanen et al., 2010).

benthic N. spumigena previous research that found planktonic from the deep, open-sea core. This result, however, contradicts because N. spumigena akinetes did not persist past the surface layer of the shallow, coastal sediment core and were absent from the deep, open-sea core. This result, however, contradicts previous research that found planktonic N. spumigena to germinate from sediment taken from open-sea locations and benthic Nodularia spp. to germinate from coastal locations (Suikkanen et al., 2010).

Aphanizomenon sp. was the least abundant genus among germinated filaments, present only in the upper layer (2–3 cm) of the open-sea sediment core. When compared with the biovolume of N. spumigena and Dolichospermum spp., Aphanizomenon sp. had the greatest average biovolume at two HELCOM monitoring stations, including the open-sea site station LL7, in the open Gulf of Finland during the summers between 1979 and 2016 (Olofsson et al., 2020). Therefore, due to high density, it remains plausible that some filaments were exported and then persisted in the bottom sediment of the open-sea site. In comparison, the absence of germinated Aphanizomenon sp. filaments from the coastal sediment core is surprising given the waters are less saline—and therefore favor Aphanizomenon sp. growth—and that blooms have been detected also in the studied coastal area previously (Laamanen et al., 2001). It is possible that the MPN values underestimated the true abundance of Aphanizomenon sp. akinetes because their germination was not favorable under the culture conditions. Though, it seems equally probable that Baltic Sea Aphanizomenon sp. do not overwinter in the form of akinetes, because previous research has found that they overwinter as vegetative filaments in the water column and are present in the water column year-round (Laamanen et al., 2002; Palińska and Surosz, 2008; Suikkanen et al., 2010; Wasmund, 2017).

Qualitative Cyanotoxin Gene PCR, Cyanotoxin Concentration, and Phylogeny of Germinated Culturable Cyanobacteria

Hepatotoxin biosynthetic gene detection from sediment samples and germinated cyanobacterial cultures was conducted to determine the potential hepatotoxin production of vegetative strains and akinetes. The aim of this qualitative hepatotoxin gene examination in conjunction with phylogenetic analysis of culturable cyanobacteria strains was to further investigate species-specific life cycle strategies by associating germinated strains with known planktonic, benthic, or epiphytic Baltic Sea strains. The presence of cyanobacterial 16S rRNA genes throughout both sediment cores further confirms the sedimentation of cyanobacteria in both the coastal and open-sea regions of the northern Baltic Sea for the past >40 and >200 years, for coastal and open-sea sites, respectively. It is important to note, however, that the presence of the 16S rRNA gene does not necessarily indicate the sedimentation of cyanobacterial akinetes due to the presence of picocyanobacteria in the Baltic Sea.

The detection of microcystin and nodularin biosynthetic genes was limited to the surface sediment layers of both sediment cores, and therefore, the highest abundance of akinetes with hepatotoxic potential likely existed in these layers. Otherwise, however, the majority of sedimentary DNA was likely from cyanobacteria strains that did not contain microcystin or nodularin biosynthesis genes and therefore did not produce hepatotoxic blooms. Similar studies found cyanobacterial microcystin (mcyA, mcyB, mcyD, mcyE), nodularin (ndaF), and anatoxin (anaC, anaF) coding genes in up to 1000-years-old lake and river sediment samples using classic PCR, nested PCR, and qPCR methods (Legrand et al., 2016, 2017a, 2019; Monchamp et al., 2016; Magonono et al., 2018; Pilon et al., 2019). Consequently, it could also be useful to adapt a nested or qPCR method for lowering the detection limit of cyanotoxin genes.

Total microcystin/nodularin measurements from the 20–63 µm sediment fraction did not consistently correspond with akinete abundance or hepatotoxin gene detection. This measurement was likely affected by the timing of the analysis, which took place over a year after sediment core collection, as hepatotoxins have short half-lives in brackish sediment and are subject to microbial degradation (Kankaanpää et al., 2009). Thus, the microcystin/nodularin measurements presented here possibly reflect intracellular akinete concentrations, which is largely unknown on a per cell basis.

The immunoassay method itself is unlikely to have suffered from interference with organic materials since the 20–63 µm sediment fraction was used, rather than the raw sediment sample. Sample processing included several washing steps, which removed much of the water-soluble organic matter. Furthermore, the non-competitive immunoassay applied here is based on two-site specific recognition of microcystins/nodularin and the assay is less prone to the matrix interference compared to the competitive assay.

Culturable cyanobacteria strains germinated from the coastal core grouped together primarily with other Baltic Sea strains, including planktonic Anabaena/Dolichospermum spp., benthic and epiphytic Anabaena spp., N. spumigena, and N. spiorea. Of all 10 cultured strains, the 2–4 cm-layer strains 1 and 2 formed a group with exclusively planktonic Anabaena and Dolichospermum species, suggesting that a greater number of benthic or epiphytic cyanobacteria deposit akinetes at the coastal location. It is also possible that revived planktonic akinetes were not represented in the selected strains or in fact were less resistant to microbial degradation, as they did not persist in
the sediment beyond the surface layer in this study. Further research needs to be conducted to confirm the presence of gas vacuole gene clusters in planktonic Anabaena spp. strains to corroborate their planktonic habitat and taxonomic classification to Dolichospermum (Wacklin et al., 2009; Komárek, 2010).

The mixing of benthic and planktonic strains in the second clade indicates that germinated strains could correspond with either habitat. This finding also agrees with the grouping of cyanobacteria in accordance with their morphology over their ecological niche (Lyra et al., 2005; Rajaniemi et al., 2005; Halinen et al., 2008). The third cluster of Anabaena/Dolichospermum spp. included strains from all three sediment layers. The monophyly, despite a 40-year age difference, could indicate a high degree of seasonal akinete recruitment from the sediment at this location among benthic strains.

Of all the cultured strains, only the 20–22 cm-layer Nodularia-like strain showed the potential for nodularin production. Therefore, the 20–22 cm-layer Nodularia-like strain may in fact be hepatotoxic N. spumigena, despite being more closely related to N. sphaerocarpa strains than N. spumigena included in the tree. This finding is consistent with clustering among Baltic N. spumigena and N. sphaerocarpa strains, hypothesized to be the result of similar akinete shape and size (Lyra et al., 2005; Rajaniemi et al., 2005). The detection of potential cyanotoxin production in the clonal culture but not the 20–22 cm sediment layer further supports that other PCR methods, such as nested PCR or qPCR, could allow for lowering the detection limit of cyanotoxin genes in the sediment.

Ultimately, the phylogenetic analysis indicated that the majority of culturable cyanobacteria strains from the coastal sediment core likely belonged to the benthic genus Anabaena. Two strains from the upper sediment layer (2–4 cm) may be Dolichospermum spp. because they grouped with known planktonic Anabaena and Dolichospermum taxa. Only the Nodularia-like strain showed positive results for hepatotoxin biosynthetic genes and grouped most closely with a Baltic Sea strain of N. spumigena. For many strains, phylogenetic grouping possibly resulted from similar morphology and bloom season, rather than ecological niche.

**SUMMARY AND CONCLUSION**

In this study, cyanobacterial akinetes persisted in and germinated from northern Baltic Sea sediment up to >40 and >400 years old. Akinete abundance and viability decreased with age and depth of vertical sediment layers. Increases in akinete abundance largely corresponded with the historical expansion of anthropogenic eutrophication-fueled blooms of cyanobacteria in the northern Baltic Sea, beginning in the mid-twentieth century. The detection of potential microcystin or nodularin production from akinetes was minimal and restricted to surface sediment layers. Phylogenetic analysis of culturable cyanobacteria from the coastal sediment core indicated that most strains likely belonged to benthic species of Anabaena. Of the culturable cyanobacteria strains, suspected planktonic species of Dolichospermum only germinated from near-surface sediment layers, with an estimated age of 1–3 years. Findings also supported the notion that, in comparison with Nodularia and Aphanizomenon spp. akinetes, Anabaena/Dolichospermum spp. akinetes play a more significant role in their life cycle and bloom initiation strategies. Overall, there was minimal congruence between akinete abundance, cyanotoxin concentration, and the presence of cyanotoxin biosynthetic genes in either sediment core. Further research is recommended to accurately quantify all akinetes and lower the detection limit of cyanotoxin genes from brackish water sediment samples in order to further describe species-specific benthic archives of cyanobacteria.

This is the first study to research benthic archives of cyanobacterial akinetes in Baltic Sea sediment cores. Measuring cyanobacterial akinete abundance, germination experiments, and genetic methods can be effectively used to determine akinete persistence, viability, and potential cyanotoxin production in brackish water sediment samples. This study highlights the prolonged survival of cyanobacterial akinetes in northern Baltic Sea sediment. Though the present study demonstrates the higher likelihood of reviving benthic cyanobacterial species, further research should be done to confirm that Baltic Sea sediment cores can be used as a proxy to hindcast blooms of planktonic cyanobacteria. Contrasting viable akinete estimates found in this study for species of bloom-forming genera Anabaena/Dolichospermum, Nodularia, and Aphanizomenon corroborate that akinetes do not play an equally significant role in the life cycles of all nostocalean cyanobacteria in the northern Baltic Sea.

**DATA AVAILABILITY STATEMENT**

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: https://www.ncbi.nlm.nih.gov/genbank/, MW491275-MW491284.

**AUTHOR CONTRIBUTIONS**

SSu and AK conceived the study. SSu, AK, and HS designed the research. AK and SW organized the fieldwork and performed the sampling. SW, HS, and SA performed molecular and experimental work. SW, V-PV, SA, and SSa analyzed the data and SSu and HS helped with data interpretation. SW wrote the manuscript. All authors discussed the results and commented on the manuscript.

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SUPPLEMENTARY MATERIAL
The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2021.681881/full#supplementary-material
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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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