Invasion of Nostocales (cyanobacteria) to subtropical and temperate freshwater lakes – physiological, regional, and global driving forces

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Similar to the increased number of studies on invasive plants and animals in terrestrial and aquatic ecosystems, many reports were recently published on the invasion of Nostocales (cyanobacteria) to freshwater environments worldwide. Invasion and proliferation of Nostocales in new habitats have the potential to significantly alter the structure of the native community and to modify ecosystem functioning. But most importantly, they influence the water quality due to a variety of toxic compounds that some species produce. Therefore a special attention was given to the invasion and persistence of toxic cyanobacteria in many aquatic ecosystems. Here we summarize the currently published records on the invasion of two Nostocales genera, Cylindrospermopsis and Aphanizomenon, to lakes and water reservoirs in subtropical and temperate zones. These invading species possess traits thought to be common to many invasive organisms: high growth rate, high resource utilization efficiency and overall superior competitive abilities over native species when local conditions vary. Assuming that dispersion routes of cyanobacteria have not been changed much in recent decades, their recent establishment and proliferation in new habitats indicate changes in the environment under which they can exploit their physiological advantage over the native phytoplankton population. In many cases, global warming was identified as the major driving force for the invasion of Nostocales. Due to this uncontrollable trend, invasive Nostocales species are expected to maintain their presence in new habitats and further expand to new environments. In other cases, regional changes in nutrient loads and in biotic conditions were attributed to the invasion events.

Keywords: cyanobacteria, Nostocales, Cylindrospermopsis, Aphanizomenon, invasive species, climate change, eutrophication

INTRODUCTION

The entry, establishment, and spread of non-native species to a new ecosystem, is frequently described as biological invasion (Vitousek et al., 1997; Ricciardi and Cohen, 2007). In many cases, biological invasions are caused by human-induced environmental interference and have substantial economic impact, serious ecological consequences, and human health hazards (Pimentel et al., 2001). They also may threaten global biodiversity by altering the structure and function of ecosystems and disrupting key biological interactions (Levine et al., 2003; Traveset and Richardson, 2006). The invasion of freshwater and marine ecosystems by non-native species of various taxonomic groups, mostly crustaceans, mussels, fish, and aquatic macrophytes is well documented (Meinesz et al., 2001; Richardson, 2011). The invasion of free-living microorganisms to new aquatic habitats is rather cryptic and difficult to detect therefore invasions of these “invisible invaders” have been rarely reported (Litchman, 2010). Furthermore, information on prior and current composition of the microbial communities, which is required to determine if a given microorganism is an invader, is often missing. Algae and cyanobacteria are exceptions in that sense as they have visible characteristic of spectral signature and microscopic morphological features. This property partly contributed to the increased number of reports on cyanobacteria invasion to lakes and water reservoirs across Europe, North, and South America and the Middle East that were recently published (Dyble et al., 2002; Wiedner et al., 2007; Hadas et al., submitted).

Here we summarize recently published records on the invasion of cyanobacteria to subtropical and temperate lakes and water reservoirs and try to forecast different scenarios of ecological and management consequences. The invasion of two Nostocales genera, Cylindrospermopsis and Aphanizomenon is presented and analyzed based on the main concepts of invasion ecology. The following issues are critically addressed: recent variations in geographic distribution of invasive Nostocales; How does a Nostocales species become an invader? What are the special traits of the invading Nostocales? What are the level and the role of genetic variability in the invasion of Nostocales? Is invasion of Nostocales to new location further expected and under what conditions? What are the expected ecological effects of the invasion?
Biological invasion is considered as an extension of normal colonization processes such as succession (Elton, 1958) therefore it was proposed that the term “invasive species” will be restricted to describe non-native species that expanded their geographic range, became abundant and have environmental and economic impact (Colautti and Macisaac, 2004). Because the terminology associated with biological invasion is rather confusing and sometimes misleading (Colautti and Macisaac, 2004), here we define an invading or a non-native species as a species that spreads beyond its native range or a species that enters a habitat where it had not resided before. However, it should be kept in mind that many cyanobacteria species (as well as other microorganisms) can inhabit an aquatic ecosystem unnoticed, since it may remain at a rather low biomass concentration and it does not form a conspicuous fraction of the community. Furthermore, the lack of detailed water quality monitoring program or an insufficiently comprehensive surveillance do not pick up the presence of these species (Kling et al., 2011). Typically it is only when a non-native population is widespread and abundant, and causes ecological or economic harms, that it is named “invasive.” It is possible for a species to be invasive in one ecosystem, but non-invasive in another. This can be due to a variety of factors, such as the presence of less-than-ideal habitat conditions (Lockwood et al., 2007). We further suggest evaluating the invasion of free-living microorganisms within an appropriate timeframe, since an invader may become a common resident of a dynamically changing environment. Therefore native and non-native species may be viewed as part of a continuum with respect to their time of residency and the extent of evolutionary and ecological interactions they have within a given environment.

The invasion of cyanobacteria strains in Central Europe as high biomass of the typically tropical species *C. raciborskii* was first reported in the 1970s (Romo and Miracle, 1994; Padisák, 1997). Long term phytoplankton records in some European lakes demonstrate a clear increase of some cyanobacterial species that had been absent or overlooked before (Padisák, 1997). Although *C. raciborskii* significantly increased covering spatial and temporal distribution of persistent perennial blooms. Many of these bloom events are of toxic *Microcystis* species and strains that proliferate under current environmental alterations, including nutrient enrichment, global warming, and regional hydrologic changes (Paerl and Paul, 2011). However, not all blooms are associated with an invasion process as many species are of broad geographic distribution and rapidly respond to current environmental changes.
**racciborskii** was known in Europe since the 1930s, it was marginal to the southern European countries (e.g., Greece; see references in Mehnert et al., 2010). However, in the last two decades, this species has appeared in some European regions and is currently widespread at central latitudes (Kokocinski et al., 2009). Other Nostocales such as **Aph. aphanizomenoides** (currently designated as **Sphaeroperidum**) and **Aph. ovalisporum**, are also considered invasive species in Europe and have been reported at some locations as newcomers. **Aph. ovalisporum** is especially relevant because it is known to produce cylindrospermopsin, although several nontoxic strains were reported (Ballot et al., 2011). **Aph. ovalisporum** was described in different ecosystems of southern Europe (Bazzichelli and Abdelahad, 1994) and in the Iberian Peninsula where it produced massive blooms for the first time in 2005 (Quesada et al., 2006). Later, it has been found at least in seven water bodies in central Spain (Cires, 2012) and in several water bodies in the southern areas of the country (C. de Hoyos, Personal Communication). **Aph. ovalisporum** has been found in the Iberian Peninsula, both in deep stratified reservoirs and in very shallow (less than 1 m) ponds of urban gardens. The spread of **Aph. ovalisporum** throughout the peninsula is considered invasion since a solid reference for its presence and bloom prior to the current reports is missing.

While field studies on the adaptation of the European species and their invasive potential are very scarce, laboratory studies with isolated strains have demonstrated that some invasive species have strong competitive abilities under changing scenarios. Mehnert et al. (2010) compared several “native” strains with “invasive” species and found that under low temperature, native species showed a higher growth rate than the invasive ones. However, under higher temperatures, the trend was the opposite and invasive strains performed better. Using these data, the authors, in a modeling effort, demonstrated that under a scenario of climate change with an increase of 4°C in the water temperature, the invasive species **C. raciborskii** would outcompete the native species (**Aph. gracile**) and that the potentially invasive **Aph. ovalisporum** would be an important component in the community (Mehnert et al., 2010). Although akinete production and overwintering/seedling play an important role in the life cycle of Nostocales, it is not necessarily the only mechanism for seedling; for example **Aphanizomenon** can seed from vegetative cells and filaments overwintering under ice (Grossart HR personal communication). Temperature-dependent release of cylindrospermopsin was reported for **Aph. ovalisporum**, implicating management complications associated with global change scenarios (Cires et al., 2011).

**MIDDLE EAST**

An exceptional bloom of **Aph. ovalisporum** first appeared in Lake Kinneret (Sea of Galilee), Israel in July 1994 and dominated the summer phytoplankton population for a few weeks (Pollingher et al., 1998). This toxic species (a producer of cylindrospermopsin) remained a regular member of the summer phytoplankton community since then and variably contributes between 10 and 40% to the summer phytoplankton biomass (Zohary, 2004). The reason for **Aph. ovalisporum**’s first appearance and its later establishment in Lake Kinneret was a subject for debate (Berman, 1997, 2001; Berman and Shteinman, 1998; Goplen et al., 1999) but it is clear now that this species took advantage of a new set of conditions that were imposed by global and regional changes (Hadas et al., submitted). The origin of the **Aph. ovalisporum** strain that established in Lake Kinneret remained unclear. A long and detailed record of the phytoplankton population in Lake Kinneret (since 1964) clearly shows its absence, although its presence at a level which is below the detection limit cannot be excluded. Interestingly, the **Aph. ovalisporum** strain isolated from Lake Kinneret was genotypically related to strains of **Anabaena bergii** isolated from constructed man-made lakes in Queensland, Australia (Shaw et al., 1999). Nevertheless, **Anabaena bergii** was not reported to bloom in the lake. The invasion process was repeated once again in Lake Kinneret as **C. raciborskii** was first reported in 2000 as a minor component of the summer phytoplankton community but dominated the community in summer 2005 with biomass as high as 68 g (ww) m⁻² (Hadas et al., submitted). Again, it is rather difficult to trace the origin of Lake Kinneret **C. raciborskii** but the fact that it is a non-cylindrospermopsin producing strain suggests that it may be related to European strains. These isolates do not produce cylindrospermopsin as opposed to Brazilian strains known to produce cylindrospermopsin and to Brazilian strains that have been reported to produce paralytic shellfish poisoning toxins (Neilan et al., 2003).

Other reports on Nostocales species in the Middle East include the Caspian Sea and Egypt. **C. raciborskii** was reported as a dominant phytoplankton species in the open and freshwater basin in the west part of the Anzali Lagoon (Caspian Sea, Iran). It was also dominant for short periods after flooding events in rivers flowing into the lagoon (Ramezanpoor, 2004). Toxins of **C. raciborskii** and Raphidiopsis mediterranea were reported in an Egyptian shallow (4 m depth) freshwater El-Dowyrat fish pond for the first time in 2002. Human activities at the end of 2001, including removal of trees and elimination of macrophytes around the fish pond have led to an increase in temperature at the surface waters of the pond (25–30°C). Both species showed seasonal variations with highest densities recorded in August each year (Mohamed, 2007). The loss of macrophytes was reported as one of the variables promoting the proliferation of **C. raciborskii** also in New Zealand lakes (Ryan et al., 2003). Thus local maintenance activities may be involved in an increase in temperature of the water column of temperate lakes and expand the distribution of **C. raciborskii** (Briand et al., 2004).

**NORTH AND SOUTH AMERICA**

The record of **C. raciborskii**-like Nostocales in North America goes back to 1955 when **Anabaenopsis seriata** Prescott, was reported from Kansas. Other reports of morphologically similar species originated from Minneapolis and Lake Erie (Kling, 2009). It was only in the 1990s when **C. raciborskii** was identified in many Florida lakes and rivers (Chapman and Schelske, 1997). Since then, **C. raciborskii** has been found in many water bodies of the central and eastern United States (Kling, 2009) including in many lakes and reservoirs in Indiana (Jones and Sauter, 2005). These sites tended to be shallow and turbid with high epilimnetic phosphorus and high chlorophyll a values (Jones and Sauter, 2005). In 2004, **C. raciborskii** was identified in water samples from the Assiniboine River, Manitoba Canada. Variable morphological features
were reported in accordance with earlier reports from Europe, and predictions based on the adaptability of that species to eutrophic conditions, elevated temperatures and turbid waters suggested its expansion in Lake Winnipeg and replacing the original phytoplankton assemblage (Leavitt et al., 2006; Patoine et al., 2006). Indeed summer blooms development and phytoplankton succession was recently reported. Lake Winnipeg shows reduced taxonomic diversity and an increased predominance of nitrogen-fixing cyanobacteria (Kling et al., 2011).

In South America C. raciborskii is well known in tropical waters of Brazil (Huszar et al., 2000), although the information about its distribution in southern latitudes is disperse and incomplete (Komárek, 2002). Vidal and Kruk (2008) analyzed spatial distribution and relative frequency of C. raciborskii in southern Uruguay, probably its southernmost distribution. C. raciborskii was found in several polymictic shallow eutrophic lakes as a dominant summer population and variable morphology. It was suggested higher occurrence of C. raciborskii in that climatic zone was associated with shallow, well-mixed eutrophic systems. Furthermore, morphological variations in C. raciborskii were not associated with latitudinal, continental, or even country distribution and independent of climatic origin.

THE INVASION PROCESS

Successful biological invasions involve complex interactions between the invading species and the physical and biological characteristics of the recipient environment. Often, invasive cyanobacteria species owe their success in colonizing new ecosystems to unique physiological characteristics (Figure 2). The invasion may be initiated with dispersion to new zones and is facilitated by human activities by migrating animals or birds or by winds. Obviously, aeolian transport is an efficient means to transfer free-living microorganisms, prokaryotes, and protists (Smith et al., 2011). But reaching a new environment is not sufficient and the invader needs a variety of traits that support its establishment and proliferation. Two distinctive traits of Nostocales are: (1) The ability to form dormant cells (akinetes) that may survive long and extreme dispersion routes, on one hand and serve as overwintering form that survive unfavorable conditions and assure perennial germination and proliferation, on the other hand. (2) The ability to fix atmospheric nitrogen in the absence of combined inorganic sources. This last trait opens invasion opportunities for Nostocales in oligotrophic aquatic ecosystems, besides ecosystems temporarily limited in combined nitrogen and extends the spectrum of ecosystems to which they can invade.

Invasive cyanobacteria have growth rates that are relatively high compared with those of native species (Isvánovics et al., 2000). They may have high resource use efficiency or possess traits allowing them to gain access to resources unavailable to other species, such as the capacity to fix molecular nitrogen. C. raciborskii has high competitiveness under fluctuating nitrogen availability due to efficient uptake capacity for ammonium (Figueroedo et al., 2007) and it also can fix atmospheric nitrogen, which allows it to escape N limitation (Briand et al., 2004; Moisander et al., 2011). This species may also utilize other limiting resources, such as phosphorus, more efficiently than other cyanobacteria due to high affinity and P storage capacity (Isvánovics et al., 2000; Wu et al., 2011) and, thus, be a superior nutrient competitor. Padisák (1997) suggested that C. raciborskii, tends to invade lakes with very high nutrient concentrations and low phytoplankton diversity, in addition to its ability to form akinetes allowing easy dispersal and environmental resistance. An important feature of Cylindrospermopsis is its wide thermal tolerance (Briand et al., 2004), which is essential to maintain the populations during cold winters. Detailed information on the effect of temperature on Nostocales life cycle processes such as akinete formation, their survival and germination, and the recruitment of vegetative filaments is currently scarce, but Cirés (2012) indicated that akinete production of Aph. ovalisporum is rather restricted to a relative narrow temperature range. Padisák (1997) reported that akinetes of C. raciborskii germinate at temperatures <24°C. Further elucidation of life cycle processes responding to temperature is required in order to determine the invasiveness capabilities and limitations of various Nostocales species.

In many cases, floating Nostocales populations create a thick scum that blocks light penetration through the water column and inhibits the growth of other species, thus under high cyanobacterial growth, the populations remain established for longer periods minimizing competition by other algal groups. A scum formation is a dynamic process that depends on a delicate balance between flotation, cellular ballast and water turbulence (Calandrino and Paerl, 2011). Filaments floating at the water surface gain advantage over sub-surface phytoplankton populations. They directly intercept CO₂ diffusing into the water from the atmosphere, thus minimizing inorganic carbon limitation of photosynthetic growth. Buoyant species contain UV-absorbing compounds such as mycosporine-like amino acids (MAAs) and scytonemin that ensure survival under extremely high irradiance conditions (Paerl
and Paul, 2011). However, in some occasions scum formation represents a trap for the organisms that lose their dynamic buoyancy behavior and they are exposed to full sunshine (including high UV radiation in summer), as well as to nutrient deprivation. Frequently they die and remain on the water body shores with extremely low metabolic activities (i.e., C and N uptake, Oliver and Ganf, 2000 and references therein).

Another trait commonly found among Nostocales but also in other cyanobacteria (i.e., *Microcystis*) is the ability to synthesize a suite of secondary metabolites some of which were identified toxic to humans and animals (Stewart et al., 2011) and others may affect community structure functioning as allelo-chemicals that inhibit other phytoplankton species or deter and reduce grazing (Fastner et al., 2007; Paerl et al., 2011). Allelopathy was suggested as a beneficial trait of *C. raciborskii* that contributes to its stable dominance and geographic expansion (Figueroa et al., 2007) but also for other cyanobacteria species (Kaplan et al., in review). A unique feature of *Aph. ovalisporum* to acquire phosphate under oligotrophic conditions was recently reported by Bar-Yosef et al. (2010). Excreted chemicals including the toxin cylindrospermopsin cause cells of some other phytoplankton species to make extracellular alkaline phosphatase whose inorganic phosphate product can be used by *Aph. ovalisporum* due to its high affinity to phosphate, thus contributing to its proliferation and domination. *Aph. ovalisporum* has also been found to outcompete other cyanobacteria at high temperatures due to higher growth rates than other Nostocales under these circumstances (Mehnert et al., 2010; Cirrés et al., 2011).

In an attempt to follow the invasion route of *C. raciborskii* to new ecosystems Neilan et al. (2003) studied genetic variations between strains isolated from freshwater rivers and reservoirs in Australia, Brazil, Germany, Hungary, Portugal, and the USA by characterizing their 16S rRNA gene nucleotide sequences and by analyzing cyanobacterium-specific short tandem repeat sequence (HIP1). Three distinct groups of *C. raciborskii* strains were clustered: (1) a group consisting of strains from the USA and Brazil; (2) a group comprised of European strains (Germany, Hungary, and Portugal); (3) a group of strains from Australia. A subsequent study that examined other isolates of *C. raciborskii* revealed the same continental distribution based on the 16S–23S internally transcribed spacer (ITS1) sequences (Gugger et al., 2005). This study suggests that the current expansion of *C. raciborskii* in Europe and in Central – and North America did not result from recent invasion and colonization by African or Australian strains (Padisák, 1997) but rather represent local strains that maintained “cryptic” populations over time and only recently proliferated due to climate change and variations in other environmental conditions. In a recent study, Piccini et al. (2011) proposed that phenotypic and genetic variability of *C. raciborskii* populations is linked to the existence of different ecotypes whose success is subject to the local environmental conditions.

Nevertheless, analysis of the 16S rRNA gene nucleotide sequences revealed 99.1% similarity between the strains of *C. raciborskii* collected from a range of global locations and continental clustering was not supported by statistically significant bootstrap values (Neilan et al., 2003). Therefore it cannot be excluded that *C. raciborskii* was transferred from the Australasian region to Europe unintentionally by humans or by migrating birds, thus providing the source of this cyanobacterium which is now reported in many previously unaffected temperate and subtropical water bodies. It is further speculated that some physiological characteristics unique to *C. raciborskii* support its proliferation in newly invaded ecosystems as they are currently exposed to higher temperature and environmental perturbations.

**GLOBAL VERSUS REGIONAL DRIVING FORCES**

In order to exploit its wide spectrum of physiological traits and to successfully outcompete native species, the conditions in a new habitat have to fit the invader’s needs. Increased water temperatures, mostly during the summer, in subtropical, and temperate zones, is considered one of the major factors supported the invasion of *C. raciborskii* and other Nostocales (Figure 2). (Padisák, 1997; Briand et al., 2004; Rücker et al., 2009). Both *C. raciborskii* and *Aph. ovalisporum* are known to proliferate at temperatures higher than 20°C (Hadas et al., 1999, 2002; Mehnert et al., 2010), thus moderate elevation in water temperatures may open new ecological habitats for them. The persistence of Nostocales in their new locations is further maintained by their ability to form dormant cells, akinetes, which stay dormant but viable in the sediment for extended periods of time (Hense and Beckmann, 2006; Kaplan-Levy et al., 2010). This seed bank may reside in the sediment and wait for a warm summer to germinate. Consequently, bloom events may follow local climatic patterns. It is important to note that freshly settled akinetes are more likely to germinate during the next summer relative to those settled in earlier blooms as the latter are already covered deeper in the sediments and probably have lower germination potency.

Nostocales species are recently reported to reside in, and dominate eutrophic as well as oligotrophic aquatic systems (Figure 3). Eutrophic conditions provide ample nutrients that can be easily and competitively utilized by opportunistic invaders (Pichler et al., 2009). Such eutrophic systems maintain high internal nutrient recycling processes that fortify primary production and biomass accumulation. Although *N* fixation is commonly recorded during Nostocales blooms in eutrophic aquatic systems (mainly indicated by the presence of heterocysts, specialized cells formed under N limiting conditions), this process is responsible for up to 2% of N acquisition by the phytoplankton community (Ferber et al., 2004; Ariosa et al., 2006). The ability of many Nostocales to efficiently use low light, allows their growth in turbid waters as well as at high biomass concentrations.

Regional inappropriate management of watersheds and high pollution impact induce eutrophication, and are certainly involved in the Nostocales geographic expansion. But these invading species can survive and propagate also under oligotrophic conditions (Figure 3). This is achieved due to their various efficient phosphate acquisition capabilities (Jukinovics et al., 2000; Spröber et al., 2003; Posselt et al., 2009; Bar-Yosef et al., 2010) and nitrogen fixation which seems to be more relevant under oligotrophic than under eutrophic conditions. Thus, in oligotrophic systems, nitrogen cannot be considered as a limiting factor. Phosphate availability controls the development of the *N* fixing population and its growth. Consequently, management efforts to control
eutrophication by reducing N loads, apparently effective in reduc-
ing blooms of *Microcystis* and other Chroococcales species, may
actually support the growth of Nostocales which easily outcom-
pete native species in such N limited ecosystems (Schindler et al.,
2008).

**CONSEQUENCES OF NOSTOCALES INVASIONS**

The immediate consequence of Nostocales invasion, besides the
ecological impairment, is the expansion of toxic blooms as vari-
uous strains produce harmful substances. Many *Aph. ovalisporum*
and *C. raciborskii* strains produce the cytotoxic compound cylin-
drospermopsin whereas several Brazilian *C. raciborskii* strains
have been reported to produce paralytic shellfish poisoning tox-
ins: neosaxitoxin, saxitoxin, and gonyautoxins (Lagos et al., 1999).
Interestingly, many strains isolated from temperate lakes in Europe
and Northern America did not produce any known toxin, although
they were found toxic in various bioassay experiments (Saker
et al., 2004). Recent studies indicated that a given population
may be composed of toxic and non-toxic strains (Ballot et al.,
2011). The bloom of toxic Nostocales presents an immediate
threat to human and animals that use the water for drinking,
bathing and recreational activities (Carmichael, 2001). Local and
政府 authorities are requested to provide solutions to
ing pressure due to different grazer population (e.g.,Winder and
Sauter, 2005) or nutrient depletion that may follow an early
variations expected after this earlier germination, as different graz-
tions to conditions which advance population establishment and
growth. However, these models do not consider the ecological
variations expected after this earlier germination, as different graz-
ing pressure due to different grazer population (e.g.,Winder and
Sander, 2004) or nutrient depletion that may follow an early
bloom. These ecological variations are hard to predict and should
be further investigated to improve models prediction.

While the possibilities to control and reduce the current trend
of global climate change are rather limited, the management of
eutrophication processes is feasible. Since a synergistic effect
of nutrients and climate was frequently indicated in many sites
invaded by Nostocales, it is important that nutrient concentrations
in many temperate and subtropical lakes be reduced substantially
from present values if cyanobacterial dominance is to be controlled
(Kosten et al., 2012). Based on long term experimental manipula-
tion, Schindler et al. (2008) concluded that N-fixing cyanobacteria

**PROSPECTIVE AND PREDICTIONS**

The current trend of increasing bloom events of cyanobacteria,
including Nostocales, will probably be further enhanced as global
warming continues. Any further temperature increase would pro-
mote the growth and development of Nostocales species in general,
and that of the invasive species in particular, and would enable
further expansion in the temperate zone (Mehnert et al., 2010).
This trend was further confirmed by a coupled biological–physical
model that predicted that high temperatures favor cyanobacteria
over other phytoplankton taxa through increased growth rates that
boost the development of blooms (Juehnk et al., 2011). A similar
conclusion was reached by Wiedner et al. (2007) who evaluated a
case of an earlier rise in water temperature associated with climate
change. They suggested that such a scenario will prompt further
spread of *C. raciborskii* to the temperate zone as earlier warming
permits earlier germination, thereby shifting the pelagic popu-
lations to conditions which advance population establishment and
growth. However, these models do not consider the ecological
variations expected after this earlier germination, as different graz-
ing pressure due to different grazer population (e.g.,Winder and
Schindler, 2004) or nutrient depletion that may follow an early
bloom. These ecological variations are hard to predict and should
be further investigated to improve models prediction.
cannot be limited by a shortage of dissolved N and instead are competitively favored. Thus reducing N inputs could actually intensify the dominance of N-fixing cyanobacteria thus enhancing the expansion of invasive Nostocales. The alternative approach to control and reduce blooms of invasive Nostocales species and their further expansion is to control and reduce external and internal sources of phosphorus, a complex task by itself.

ACKNOWLEDGMENTS

This work was partly supported by the German Ministry of Research and Technology (BMBF) and Israel Ministry of Science and Technology (MOST) under contracts FKZ 02WT0985 and WR803. Assaf Sukenik and Ora Hadad acknowledge the continuous support of the Israel Water Authority provided to the Kinneret Limnological Laboratory, IOLR.

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Conflict of Interest Statement: 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.