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1 Influence of sampling strategies on the monitoring of cyanobacteria in

2 shallow lakes: lessons from a case study in France

3

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9 Abstract

10 Sampling cyanobacteria in freshwater ecosystems is a crucial aspect of monitoring programs 11 in both basic and applied research. Despite this, few papers have dealt with this aspect, and a 12 high proportion of cyanobacteria monitoring programs are still based on monthly or twicemonthly water sampling, usually performed at a single location. In this study, we conducted 13 14 high frequency spatial and temporal water sampling in a small eutrophic shallow lake that 15 experiences cyanobacterial blooms every year. We demonstrate that the spatial and temporal 16 aspects of the sampling strategy had a considerable impact on the findings of cyanobacteria 17 monitoring in this lake. In particular, two peaks of Aphanizomenon flos-aquae cell 18 abundances were usually not picked up by the various temporal sampling strategies tested. In 19 contrast, sampling once a month was sufficient to provide a good overall estimation of the 20 population dynamics of *Microcystis aeruginosa*. The spatial frequency of sampling was also 21 important, and the choice in the location of the sampling points around the lake was very 22 important if only two or three sampling points were used. When four or five sampling points 23 were used, this reduced the impact of the choice of the location of the sampling points, and 24 allowed to obtain fairly similar results than when six sampling points were used. These 25 findings demonstrate the importance of the sampling strategy in cyanobacteria monitoring, 26 and the fact that it is impossible to propose a single universal sampling strategy that is 27 appropriate for all freshwater ecosystems and also for all cyanobacteria.

28

Keywords: sampling strategy, cyanobacteria, spatiotemporal dynamic, *Microcystis aeruginosa*, *Aphanizomenon flos-aquae*

31

31 **1 Introduction**

32 Due to eutrophication and, to a lesser extent, to climatic changes (Markensten et al., 2010; 33 Paerl and Huisman, 2009) cyanobacterial blooms seem to be increasing in freshwater 34 ecosystems worldwide. These blooms severely disrupt the functioning of these ecosystems and potential water use. Furthermore, many cyanobacterial species are able to produce a 35 36 variety of toxic metabolites, which can be harmful to both human (Kuiper-Goodman et al., 37 1999) and animal (Codd et al., 2005) health. For these reasons, numerous attempts have been 38 made in the last 20 years to elucidate the factors that control cyanobacterial blooms and toxin 39 production, and thus to make it possible to evaluate better the health risks associated with 40 bloom events. From all these studies, it is clear that the spatial distribution of cyanobacteria in 41 freshwater ecosystems can display marked horizontal and vertical variations (Porat et al., 42 2001; Welker et al., 2003). Moreover, by means of a real time PCR analysis of a gene 43 involved in the biosynthesis of microcystins we have shown that considerable fluctuations can 44 also occur in the proportions of potentially microcystin-producing and non-producing cells 45 during the course of Microcystis aeruginosa blooms (Briand et al., 2009). Similar results have been found for various different *M. aeruginosa* populations located in the same geographic 46 47 area (Sabart et al., 2009), which makes it difficult to manage the health risks associated with 48 these events.

All these studies indicate that the sampling strategy used for monitoring cyanobacteria is a critical aspect, both in basic research on cyanobacteria, (e.g. investigation of the factors and processes involved in the development of the blooms), and in applied research, (e.g. implementing monitoring programs of these microorganisms in freshwater ecosystems used to provide drinking water or for recreational activities). In recent years, new tools have been tested with the intention of improving cyanobacterial sampling, for example, remote sensing reconnaissance to determine the horizontal distribution of cyanobacteria in freshwater
ecosystems (Hunter et al., 2009), or spectrofluorometric probes to reveal the vertical
distribution of these cyanobacteria in the water column (Leboulanger et al., 2002). Moreover,
these spectrofluorometric probes and other sensors have now been integrated into buoys, to
provide real-time monitoring of cyanobacteria in freshwater ecosystems (Le Vu et al., in
press).

61 However, despite the great potential interest of these tools, their cost will remain 62 prohibitive for their routine use in the foreseeable future, and most of the monitoring programs worldwide for the survey of cyanobacteria will continue to be based on more 63 64 conventional methods for some years to come. Taking discrete samples of various volumes of water taken from the shoreline of ecosystems is probably the method one most often used in 65 66 studies. Unfortunately, as a result of spatial and temporal differences in the distribution of 67 cyanobacteria, this approach can often provide a very poor estimation of cyanobacterial 68 abundance and, consequently, of the associated health risk. We therefore need to devise 69 simple sampling strategies for the low cost monitoring of cyanobacteria in shallow lakes. In 70 an attempt to do this, we performed intense spatiotemporal monitoring of cyanobacteria in a 71 shallow lake known to experience cyanobacterial blooms every year.

72

73 2 Materials & Methods

74 **2.1 Study site**

75 This study was performed in a shallow lake named Place (0.08 km², 2.5 m max depth,

76 45°43'N, 4°14'E) located in the plain of Forez (Central France), (Fig. 1). This lake is used for

extensive fish production and its trophic status is eutrophic to hypereutrophic (OCDE, 1982).

78 *Microcystis aeruginosa* blooms occur every summer.

79

80 **2.2 Data acquisition**

81 *2.2.1 Sampling strategy and cell counting*

82 In order to assess the variations in the horizontal distribution of cyanobacteria in this pond, we monitored six sampling points located around the lake at one meter from the shore (V1-V6; 83 84 Fig. 1). The water depth in each of these sampling points was around 1 meter. Samples were 85 taken every two days, between 09:00 and 10:00 a.m., from early June 2008 to early October 86 2008. The first 40-centimeters of the water column were sampled using a watersampler (Uwitech, Austria). This water sample was shacked and then divided into two 1-L bottles, one 87 88 liter being stored at room temperature with Lugol's iodine solution, and the other at 4°C. 89 In order to evaluate the vertical distribution of cyanobacteria in the water column, we performed a 22-hour survey (from 4:30 p.m. August 4, 2009 to 2:30 p.m. August 5, 2009), of 90 91 the variations of cyanobacterial biomass at five sampling points (A-E; Fig. 1) using a BBE 92 Algaetorch (Moldaenke, Germany). This torch is based on the same principle as the BBE 93 spectrofluorometric probe (Beutler et al., 2002), but provides only an estimation of the 94 concentrations of cyanobacteria and total chlorophyll in water. Every hour, the torch was 95 immersed to a depth of 20 centimeters at the five sampling points, and triplicate 96 measurements were performed in each point.

97 The cyanobacterial cell concentrations were estimated using a Nageotte cell and an
98 optical microscope, as described in Brient et al. (2008). For each rectangular area, we counted
99 at least 400 cells of each cyanobacterial species.

100

101 2.2.2 Meteorological data

The speed and direction of wind during our study were obtained from the Metéo France
meteorological station at St Etienne-Bouthéon (4°18'E – 45°32'N). The wind direction rose

104	for this station	is given	in Supplementa	al Figure 1,	and shows th	hat the two c	dominant wind

105 directions were NW and SE. The direction of winds blowing from $240^{\circ} - 60^{\circ}$ was classified

106 as NW, and that of winds blowing from $60^{\circ} - 240^{\circ}$ as SE.

107

108 **2.3 Data analysis**

109 The spatial distribution of cyanobacteria in the lake was represented using Surfer (v. 7.0,

110 Golden Software Inc.), and statistical analyses (Wilcoxon test, Spearman correlation) were

111 performed using the R package version 2.10 (R development core Team, 2010).

112

113 **3 Results**

3.1 Change over time in the population dynamics of the two dominant cyanobacterial species

116 Two cyanobacterial species, *Microcystis aeruginosa* and *Aphanizomenon flos-aquae*,

117 dominated the phytoplankton community during the summer of 2008. The population

118 dynamics of these two species displayed very contrasting patterns (Fig. 2). The population

119 dynamics of *Microcystis aeruginosa* was characterized by a steady increase in the cell

abundance from June to August, apart from a brief dip in the middle of July. The maximum

121 population was reached on August 21 (264,000 cells/mL), and subsequently the cell

122 concentration remained stable until the end of September, and then decreased in October. In

123 contrast, the population dynamics of *Aphanizomenon flos-aquae* were much more chaotic,

124 with the cell abundance reaching two very high and short-lived peaks in July

125 (400,000 cells/mL on July 17, and 560,000 cells/mL on July 23).

126

127 **3.2 Influence of sampling frequency on the estimation of the population dynamics**

128 Our assessment of the changing population dynamics of the two cyanobacteria were obtained 129 using a very frequent high temporal sampling regime (every two days), which would not be 130 practicable in the context of normal monitoring programs. In order to evaluate the impact of 131 the sampling frequency, we simulated weekly, twice-monthly and monthly sampling 132 frequencies to our data set. The results of these simulations are shown in Fig. 3 and 4. From 133 this figure, we can see that changes in *M. aeruginosa* cell abundance over time would have 134 been fairly accurately estimated at all these sampling frequencies. Moreover, for all sampling 135 frequencies, the quality of the estimation of the *M. aeruginosa* population dynamics was not 136 influenced by choice of the first sampling date (Fig. 3). In contrast, the population dynamics 137 of A. flos-aquae would have been badly or even very badly estimated by using weekly, twice-138 monthly and monthly sampling frequencies (Fig. 4). We would only have detected both 139 A. flos-aquae peaks in one of the three trials testing the weekly sampling strategy, and we 140 would never have detected these peaks with twice-monthly and monthly sampling 141 frequencies.

142

143 3.3 Evolution of the horizontal distribution of cyanobacteria in the lake during the 144 bloom

As shown in the video (Supplemental Fig. 2), the horizontal distribution of both cyanobacteria displayed marked variations during the course of the study. Moreover, when the spatial distributions of the two species at the same sampling dates were compared, it could be seen that similar or contrasting patterns in the horizontal distribution of *M. aeruginosa* and *A. flosaquae* cells would have been found, depending on the dates chosen (some examples are provided in Fig. 5).

151 In order to obtain a better picture of this spatial variability in the cell concentrations of 152 the two species, we estimated the coefficients of variation in the mean abundance for each sampling date and for each species from the results obtained at the six sampling points (Fig. 6). These coefficients were usually higher for *A. flos-aquae* than for *M. aeruginosa* (Wilcoxon test, p=- $3.25.10^{-05}$), suggesting that the horizontal distribution of *A. flos-aquae* was more variable. Finally, there was no correlation (Spearman coefficient) between the coefficient of variation and the mean cell abundance for *Aphanizomenon*, and only a weak correlation was found for *Microcystis* (Spearman coefficient, p=0.003 r =-0.4; Supplemental Fig. 3).

160 In order to find out whether wind speed/direction could account for the variations in 161 the horizontal distribution of cyanobacterial cell abundance in the lake, we recorded in a first 162 time, for each species and for each sampling date, the sampling point (out of the six) at which 163 the highest cell abundance was detected. We then constructed a table in which we related 164 these findings to the wind direction and speed in the five hours before the sampling, knowing 165 that only data with wind speed values ≥ 2.0 m/s were taken into consideration. For M. 166 aeruginosa, the detection of the highest cell abundances in the southernmost sampling points 167 V2 and V3 were associated with winds blowing from the NW (Table 1), whereas those at the 168 V1 and V4 sampling points were more surprisingly associated with winds from the SE. High 169 cell abundances in the northern most sampling points V5 and V6 were equally associated with 170 winds from NW and SE. For A. flos aquae, the results were more complicated, and no 171 obvious link could be seen between the direction of the wind and the distribution of the 172 cyanobacteria (Table 1). The same analyses were performed by taking into account the wind 173 data one and two days before sampling (instead 5-10 hours before sampling), but no obvious 174 relationship was detected (data not shown).

175

3.4 Influence of the number of sampling points on the estimated cyanobacterial cell
abundances in the lake

178 The cyanobacterial cell abundances in the shallow lake were estimated by calculating the 179 average value for the six sampling points (see Fig. 1). In order to determine the number of 180 sampling points required to obtain a good estimation of cyanobacterial cell abundances in the 181 lake, we compared the estimations of cell abundance based on using samples from just one, 182 two, three, four or five sampling points with that based on all six. To do this, we calculated 183 the correlation coefficients (Spearman) between the estimations based on the six sampling 184 points and those based on one to five sampling points for each species (Fig. 7). We considered 185 all possible combinations of points, and the results are classified in the figure on the basis of 186 increasing order of r values within each combination of groups. For both species, we found 187 that the estimations of cell abundances based on only one or two sampling points were 188 generally rather badly correlated with those obtained using all six sampling points. On the 189 other hand, it appeared that good correlations (around or > 0.9) were obtained when at least 190 three sampling points were used, but also that the variations due to the choice of the sampling 191 points was still considerable when only three sampling points were used.

In order to find out which combinations of sampling points provided the best results when only two or three sampling points were used, we classified all the possible combinations of points. To do this, we added the rank of each combination of sampling points obtained for the two species (*M. aeruginosa* and *A. flos-aquae*). From Figure 8, we can see that the best estimations obtained using only two or three sampling points were provided by combinations in which the sampling points used were on the shore opposite to the prevailing wind direction over the lake.

199

200 **3.5** Diel variations in the subsurface cyanobacterial biomass in the lake

201 Finally, we carried out a 24-hour estimation of the variations in the total cyanobacterial

202 biomass in the subsurface water (20 cm depth) of the lake, at five sampling points using the

203 BBE torch (A-E, see Fig. 1). As shown in Fig. 9, there was a steady fall in the cyanobacterial 204 biomass at all sampling points during the afternoon and evening, and conversely an increase 205 late at night and in the morning. Moreover, the differences in biomass between the five 206 sampling points were smaller during the night than during the day, as was the standard error 207 (three measurements per sampling point). A multidimensional scaling analysis performed on 208 the same values confirmed these observations, with all the night sampling times being 209 grouped together, whereas the sampling times during the day were much more scattered (Data 210 not shown).

211

212 **4 Discussion**

213 As far as we are aware, this is the first attempt to investigate the influence of sampling 214 strategies on the evaluation of spatial and temporal variations in cyanobacterial abundances in 215 shallow lakes, which constitute unstable and complex ecosystems. These lakes are used by 216 humans for numerous activities, including recreational activities and the supply of drinking 217 water, which makes the monitoring of cyanobacteria in such ecosystems of particular 218 importance, especially as part of the evaluation of the health risks linked to cyanobacterial 219 blooms and their toxins. Sampling strategy is also very important in the context of basic 220 studies, because the quality of sampling has a major impact on the quality of the final results. 221 In this study, we found that the sampling frequency required to obtain a good 222 estimation of the temporal evolution of the cyanobacterial abundance depends on the 223 blooming species, M. aeruginosa or A. flos-aquae. Twice-monthly or monthly sampling 224 provided good results for *M. aeruginosa*, whereas this was not often enough to monitor the 225 chaotic population dynamics of A. flos-aquae. These findings are in contradiction with the 226 recommendations of Codd et al. (1999), who proposed weekly or a twice-monthly sampling

227 for species that do not form scum (A. *flos-aquae* for example), and more frequent sampling 228 for scum-forming species (such as *M. aeruginosa*), because they can display more rapid 229 changes in concentration. On the other hand, in agreement with these authors, our findings 230 also demonstrate that a reactive approach to cyanobacterial sampling is called for, and that 231 appropriate monitoring programs must be devised for each ecosystem based on what is known 232 about how these systems function. It is clear that sampling only once or twice a month can 233 lead to a very considerable under-estimation of cyanobacterial concentrations, and thus of the 234 health risks associated with the bloom. As a result, a weekly sampling frequency seems to be 235 required for cyanobacteria in small freshwater ecosystems.

236 Our data on the variability of the spatial distribution of cyanobacteria in the lake 237 indicate that at least three sampling points were needed to obtain a good estimation of the 238 abundance, based on a comparison with estimations based on six sampling points. It appeared 239 also that if only three sampling points are used, the choice of the location of these sampling 240 points is very important for the quality of the estimation. The most reliable results were 241 obtained using sampling points located on the opposite side of the lake shore to the main axis 242 of the wind direction, and that adding more sampling points reduces the impact of the choice of the location of the sampling points. Such horizontal variability in the distribution of 243 244 cyanobacteria has been previously documented for many ecosystems, and also for many 245 cyanobacterial species. For example, in a recent study, Briand et al. (2009) showed that the 246 spatial distribution of *M. aeruginosa* in a large freshwater reservoir on a given date could vary from 7.10^3 cells/mL to 2.10^8 cells/mL, depending on the location of the sampling points in the 247 248 reservoir. Many factors and processes can influence the horizontal distribution of 249 cyanobacteria in a freshwater ecosystem. Among them, wind and surface currents seem to 250 have the greatest impact. For example, the distribution of *Microcystis* spp. in lake Taihu (see 251 the review paper of Qin et al., 2010) and in Lake Ontario (Hotto et al., 2007) is clearly

influenced by both winds and currents. Similarly, Moreno-Ostos et al. (2009) have shown that 252 253 in a Spanish reservoir currents have a marked effect on the distribution of cyanobacteria, and 254 more globally on the phytoplankton community. In this study, we found that the horizontal 255 distribution of *M. aeruginosa* in the lake was influenced more by wind direction than that of 256 A. flos-aquae. This could be explained by the fact that M. aeruginosa colonies are located at 257 the surface of the lake at the end of the night, and thus are more subjected to the influence of 258 the wind than A. flos-aquae filaments, which are distributed over the entire water column. We 259 found also that two sampling points in the lake (V5 and V6) were less influenced by wind 260 direction that the others. This could be explained by the fact that these two sampling points 261 are protected from the influence of winds blowing from the NW by an embankment located in 262 the North part of the lake. Finally, we also demonstrated that in such a small lake, the impact 263 of wind occurred at the scale of a few hours, in contrast to the previous findings of Welker et 264 al. (2003) showing that the distribution of cyanobacteria was influenced by winds that had 265 been blowing one or two days earlier.

266 In addition to this variability in their horizontal distribution; the vertical distribution of 267 cyanobacteria was also variable. Indeed, during the 24 h for which we used the BBE Torch to 268 monitor the concentrations of cyanobacteria, we found that they were lower in the subsurface 269 layer early at night than during the day. The greatest variations in biomass were recorded 270 during the daytime, both at the scale of one sampling point when the three measurements 271 were compared, and at the scale of the five sampling points monitored during this study. 272 These findings also suggest that several sampling points are necessary to obtain an accurate 273 assessment of the cyanobacterial biomass and that integrated sampling of the first meter of the 274 water column reduces the variability in the estimation of the biomass due to the position of 275 cyanobacteria in the water column. This finding is consistent with data reported by Ahn et al. 276 (2008) showing that an integrated method was the most appropriate sampling method for

Oscillatoria and Microcystis blooms. The causes of these variations in the position of
cyanobacteria in the water column have been studied for different species. Several papers
(Porat et al., 2001; Rabouille and Salençon, 2005; Rabouille et al., 2005; Visser et al., 2005;
Walsby, 1994) have shown that migrations of cyanobacteria in the water column are probably
due to the dynamics of the carbon-reserve metabolism, and are strongly influenced by light,
temperature, and water mixing.

283 From all these findings, guidelines should be proposed for the monitoring of 284 cyanobacteria in shallow lakes Codd et al. (1999) propose that the choice of sampling 285 frequency and the choice of the number and location of the sampling sites should depend on 286 the purpose of monitoring. For example, sampling near public bathing sites was 287 recommended in freshwater ecosystems used for recreational activities. However, this 288 strategy might generate data relevant only to the immediate vicinity of the bathing area, which 289 do not reflect the global distribution of cyanobacteria in the lake. This is especially true when 290 this distribution is very varied, and could make it difficult to prevent or manage blooms. On 291 the basis of our findings, we proposed a different sampling strategy, which does not depend 292 on the purpose of the monitoring. In order to minimize the cost of the cyanobacteria survey, 293 twice-monthly sampling could be the norm for monitoring, but only if it is complemented by 294 regular visual surveys. Changes in the appearance of the water (e.g. its color) between two 295 successive dates would lead to an immediate increase in the sampling frequency. If it is not 296 possible to carry out this visual survey, only a weekly sampling strategy can ensure that a 297 sporadic cyanobacterial bloom is not missed. With regard to the number of sampling points, 298 we found that at least three sampling points were necessary to obtain an accurate assessment 299 of the cyanobacterial biomass (based on comparison with six sampling points). However, 300 even when three sampling points were used, we found that the choice of the location of the 301 sampling points was also very important (Fig. 8), even though the lake was fairly rectangular

302 in shape and its perimeter small (around 1.3 Km). These findings suggest that for large lakes 303 and also for lakes with a more complex shape, a large number of sampling points would be 304 necessary to obtain a good estimation of the cyanobacterial abundance. Clearly such sampling 305 is time consuming and expensive. One way to reduce these costs would be to collect a large 306 number of samples and then pool equal volumes of these samples in the same flask, before 307 carrying out a single analysis. In this study, as in most of the monitoring programs performed 308 in small lakes, all samples were taken from the shoreline of the lake. This kind of sampling is 309 suitable for small lakes, but it has been shown that for large lakes (Rogalus and Watzin, 2008) 310 shoreline sampling may miss early warning signs of bloom development, and also lead to the 311 overestimation of the concentration of microcystins, when compared to data obtained from 312 offshore samples. For bigger lakes, therefore, the sampling strategy must include offshore 313 samples.

314 Different programs worldwide are testing alternatives to water sampling for the 315 monitoring of cyanobacteria in freshwater ecosystems. Two main approaches have been 316 investigated. The first one is based on the use of remote sensing, which has long been in use 317 in marine ecosystems (see for example Bracher et al., 2009). In freshwater ecosystems, the 318 paper of Hunter et al. (2008) has shown the potential of high resolution images for the 319 assessment of the spatial distribution of *M. aeruginosa* in a shallow eutrophic lake. However, 320 the cost of these images and the impact of meteorological conditions are limiting factors for 321 envisaging the use of this tool in routine cyanobacteria monitoring programs. One alternative, 322 lower-cost solution could be based, in the future, on the use of drones to take aerial 323 photographs of freshwater ecosystems, but these tools are still in development. Moreover, 324 they will be only useful for cyanobacterial species that live in the surface water of lakes. 325 The second way of monitoring of cyanobacteria without sampling the water being 326 investigated is the use of buoys equipped with a variety of sensors, including, for example, a

submersible spectrofluorometer to quantify the biomass of the cyanobacteria. This kind of
tool permits the real-time monitoring of phytoplankton, including cyanobacteria, as shown for
example in the paper of Le Vu et al. (in press). The two obstacles to their use in routine
cyanobacteria monitoring programs are the high price of these systems, and the fact that they
only provide estimations for one sampling point. Despite this, the possible use of such buoys,
combined with the spatial monitoring of cyanobacteria by water sampling looks very
promising for surveying cyanobacteria in freshwater ecosystems.

5 Conclusion

335 The sampling of cyanobacteria in freshwater ecosystems is a hot topic, in particular in the 336 context of programs for surveying these toxic microorganisms in ecosystems used for the 337 production of drinking water or for recreational activities. Paradoxically, fewer studies deal 338 with the impact of sampling strategies on the estimation of cyanobacterial cell abundances in 339 freshwater ecosystems. In this study, we demonstrate that the choice of sampling strategy can 340 lead to very different estimations of the cell abundances of two blooming species in a shallow 341 lake and also that, depending on the cyanobacterial species involved, different sampling 342 strategies are required to obtain a good estimation of their population dynamics. All these 343 findings suggested that monthly or twice-monthly sampling strategies at just one sampling 344 point do not allow to provide an accurate estimation of cyanobacterial abundances, and thus 345 of the health risks associated with the presence of toxic species in aquatic ecosystems. 346 Moreover, although promising new technologies are being developed for monitoring 347 freshwater cyanobacteria, their cost and some other drawbacks mean that at present they 348 cannot replace water sampling, which will remain the basis of most of these monitoring 349 programs for the foreseeable future.

- 350
- 351

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355

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Table 1: Relationship between wind direction and high cell abundance recorded for
 Microcystis aeruginosa and *Aphanizomenon flos-aquae* at the different sampling points. We
 446

Fig. 1: Geographical location of the study site in France (left), and of the sampling points inthe lake (right)

449

Fig. 2: Changes over time of the concentrations of *Microcystis aeruginosa* (top) and *Aphanizomenon flos-aquae* (bottom). These concentrations were estimated by calculating the average cell count for the six samples at each date. The error bars indicate the standard deviation.

454

Fig. 3: Simulation of the change over time of *Microcystis aeruginosa* cell concentrations
found using a weekly (top), twice-monthly (middle) or monthly sampling frequency (bottom),
with lags for the first sampling day of zero days (-), 2 days (--) and 4 days (....) comparing to
our first sampling day. The gray curve corresponds to the reference data.

459

460 Fig. 4: Simulation of the change over time of the biomass of *Aphanizomenon flos-aquae*461 found using a weekly (top), twice-monthly (middle), or monthly sampling frequency
462 (bottom), and with lags for the first sampling day of zero days (-), 2 days (--) and 4 days (....)
463 comparing to our first sampling day. The gray curve corresponds to the reference data.

464

465 Fig. 5: Spatial distribution of two cyanobacteria, *Microcystis aeruginosa* and *Aphanizomenon*466 *flos-aquae*, in the lake at four sampling dates (July, 9, 17 & 23; August, 8)

467

468 Fig. 6: Change over time in the coefficients of variation of the mean cell abundances of
469 *Microcystis aeruginosa* (black triangle) and *Aphanizomenon flos-aquae* (white square)
470 estimated at all six sampling points.

471

472 Fig. 7: Spearman correlation values between *Microcystis aeruginosa* (top) and
473 *Aphanizomenon flos aquae* (bottom) cell abundances estimated from the mean values for all
474 six sampling point values, and those estimated from only one, two, three, four or five of these
475 six sampling points.

476

Fig. 8: Location of the sampling points providing the best (left) and worst (right) estimations of cyanobacterial cell abundances, compared to estimations based on six sampling points. We give the combinations for two (top) and three (bottom) sampling points. The polar plot shows the direction of the maximum daily wind speed during the study. The different line types permit to distinguish the two best or the two worst combinations of sampling points, using two or three sampling points.

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Fig. 9: Cyanobacterial biomass in the subsurface water of the lake over a 24-hour period at five sampling points (\blacklozenge point A, \blacksquare point B, \blacktriangle point C, \times point D, and \diamond point E). The error bars indicate the standard deviation.

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Supplemental Figure 1. Distribution of the wind directions at the St-Etienne-Bouthéon meteorological station during this study (June, 13 to October, 10, 2008). The curve and the bars indicate respectively the mean speed and the occurrence per hour of the wind in each direction. 493

494 Supplemental Fig. 2. Evolution of the spatio-temporal distribution of *Microcystis aeruginosa*495 (left) and *Aphanizomenon flos aquae* (right) in the lake during our study (the scale is the same
496 than in Fig. 5).

497

- 498 Supplemental Fig. 3. Relationship between the cell concentration and the coefficient of
- 499 variation for *Microcystis aeruginosa* (top) and *Aphanizomenon flos-aquae* (bottom)

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