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Avoiding quantification bias in metabarcoding: Application of a cell biovolume correction factor in diatom molecular biomonitoring

Valentin Vasselon1 | Agnès Bouchez1 | Frédéric Rimet1 | Stéphan Jacquet1 | Rosa Trobajo2 | Méline Corniquel1 | Kálmán Tapolczai1 | Isabelle Domaizon1

1CARRTEL, French National Institute for Agricultural Research (INRA), University of Savoie Mont Blanc, Thonon-les-Bains, France
2Aquatic Ecosystems, Institute for Food and Agricultural Research and Technology (IRTA), Catalunya, Spain

Correspondence
Valentin Vasselon
Email: valentin.vasselon@inra.fr

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Abstract
1. In recent years, remarkable progress has been made in developing environmental DNA metabarcoding. However, its ability to quantify species relative abundance remains uncertain, limiting its application for biomonitoring. In diatoms, although the rbcL gene appears to be a suitable barcode for diatoms, providing relevant qualitative data to describe taxonomic composition, improvement of species quantification is still required.

2. Here, we hypothesized that rbcL copy number is correlated with diatom cell biovolume (as previously described for the 18S gene) and that a correction factor (CF) based on cell biovolume should be applied to improve taxa quantification. We carried out a laboratory experiment using pure cultures of eight diatom species with contrasted cell biovolumes in order to (1) verify the relationship between rbcL copy numbers (estimated by qPCR) and diatom cell biovolumes and (2) define a potential CF. In order to evaluate CF efficiency, five mock communities were created by mixing different amounts of DNA from the eight species, and were sequenced using HTS and targeting the same rbcL barcode.

3. As expected, the correction of DNA reads proportions by the CF improved the congruence between morphological and molecular inventories. Final validation of the CF was obtained on environmental samples (metabarcoding data from 80 benthic biofilms) for which the application of CF allowed differences between molecular and morphological water quality indices to be reduced by 47%.

4. Overall, our results highlight the usefulness of applying a CF factor, which is effective in reducing over-estimation of high biovolume species, correcting quantitative biases in diatom metabarcoding studies and improving final water quality assessment.

KEYWORDS
benthic diatom, biovolume correction factor, freshwater ecosystems, gene copy number variation, quantitative metabarcoding
1 | INTRODUCTION

DNA metabarcoding allows species present in an environmental sample to be detected using a short DNA marker specific for a particular taxonomic group (Taberlet, Coissac, Hajibabaei, & Rieseberg, 2012). Combined with high-throughput sequencing (HTS), hundreds of samples can be analysed at the same time, offering an alternative to microscopy with higher resolution and accuracy, while being faster and cheaper (Stein, Martinez, Stiles, Miller, & Zakharov, 2014). This is particularly interesting for freshwater biomonitoring, in which thousands of river samples have to be analysed annually and management actions applied quickly (Keck, Vasselon, Tapolczi, Rimet, & Bouchez, 2017). The European Water Framework Directive (WFD, European Council, 2000) has implemented the use of benthic diatoms, among other biological indicators (fishes, macroinvertebrates, macrophytes and phytoplankton), for the assessment of aquatic ecosystem integrity. The different biotic diatom indices that have been developed are based on the relative abundances and the ecological values (sensitivity and tolerance to pollutants) of the species observed in rivers and lakes systems (e.g. Rimet, 2012). Different studies have already revealed the potential application of diatom metabarcoding in freshwater quality assessment (Apotheloz-Perret-Gentil et al., 2017; Kermarrec et al., 2014; Vasselon, Domaizon, Rimet, Kahlert, & Bouchez, 2017; Vasselon, Rimet, Tapolczi, & Bouchez, 2017; Visco et al., 2015). However, discrepancies between DNA metabarcoding and microscopy have been observed in species composition and relative abundance (Zimmermann, Glöckner, Jahn, Enke, & Gemeinholzer, 2015). This drawback is likely to affect the congruence between morphological and DNA metabarcoding quality index values and, in fine, the ecological assessment.

With respect to qualitative aspects, the incompleteness of the reference databases, the choice of the DNA marker and the efficiency of the PCR primers have been identified as important biases affecting species detection using DNA metabarcoding (Pawlowski, Lejzerowicz, Apotheloz-Perret-Gentil, Visco, & Esling, 2016). For benthic diatoms, the rbcL gene has been proved to be an appropriate taxonomic marker for biomonitoring (Kermarrec et al., 2013, 2014; Vasselon, Domaizon, et al., 2017; Vasselon, Rimet, et al., 2017) and a well-curated barcode reference library is already available in open-access to assign species names to rbcL sequences (R-Syst::diatom, Rimet et al., 2016). However, no clear relationship has yet been demonstrated between the relative species abundances obtained by DNA metabarcoding with the rbcL barcode and those obtained by morphological observations (Rimet et al., 2014). As quantification of diatom species is required by the WFD for quality index calculation, more investigation is needed to understand and correct biases affecting diatom quantification based on HTS data.

Species quantification based on HTS data can be estimated from the number of DNA sequences (i.e. reads) assigned to each species, from which relative abundances can be calculated. Previous studies have documented a variety of problems that may affect the proportions of DNA reads obtained with HTS (Amend, Seifert, & Bruns, 2010; Deagle, Thomas, Shaffer, Trites, & Jarman, 2013; Pawlowski et al., 2016; Tan et al., 2015; Thomas, Deagle, Eveson, Larsch, & Trites, 2016), including biological biases (e.g. gene copy number variation, tissue cell density, cell biovolume), technical biases (e.g. DNA extraction, PCR amplification) and biases linked to HTS itself (e.g. library construction, HTS technology used, bioinformatics treatments). Variation in gene copy number per cell constitutes a major bias known to affect the proportion of DNA read found for each species present in complex assemblages; this has been demonstrated for macroinvertebrates (Elbrecht, Peinert, & Leese, 2017), fish, amphibians (Evans et al., 2016), oligochaetes (Vivien, Lejzerowicz, & Pawlowski, 2016), foraminifera (Weber & Pawlowski, 2013) and microbial assemblages (Angly et al., 2014). However, to the best of our knowledge, no study has yet evaluated gene copy number variation bias on diatom metabarcoding quantification. While tissue cell density and species biomass are major biases likely to affect DNA metabarcoding quantification of multicellular organisms like macroinvertebrates (Elbrecht & Leese, 2015) or fish (Evans et al., 2016), diatoms are unicellular organisms for which gene copy number is mainly affected by the number of genomes and the number of gene copies per genome. This may be particularly true for non-nuclear markers like the chloroplast-encoded rbcL gene. Godhe et al. (2008) reported a clear correlation between the 18S gene copy number per cell with diatom cell length and biovolume, suggesting that the cell biovolume could be a proxy for the gene copy number. Keeping in mind that diatom biovolume varies from $10^4$ to $10^9$ μm$^3$ (Snoeijs, Busse, & Potapova, 2002), gene copy number may vary greatly between the smallest and the biggest diatom species, affecting metabarcoding quantification.

For all the reasons mentioned above, we hypothesized that a quantification correction factor (CF) based on diatom cell biovolume should be necessary to correct DNA read proportions to provide species quantification more comparable to microscopical counts. In order to confirm this hypothesis, we firstly conducted experiments on eight pure diatom cultures to examine whether variation in rbcL gene copy number per cell correlates with morphological characteristics (e.g. biovolume, cell length), from which a CF might be calculated. Secondly, the efficiency of the proposed CF was tested on (1) mock communities made by mixing known proportions of the eight diatom species cultures and (2) environmental diatom assemblages from rivers previously sequenced (Vasselon, Rimet, et al., 2017) and for which data are available online (Vasselon, Rimet, et al., 2017 dataset, https://doi.org/10.5281/zenodo.400160). Last, the capacity of the CF to improve the ecological assessment of rivers was tested by comparing water quality index values calculated from molecular data with corrected abundances to those calculated from classical morphological abundances.

2 | MATERIALS AND METHODS

2.1 Evaluation of the quantification bias and development of a quantification correction factor (CF)

To evaluate whether the rbcL copy number per cell varies between diatom species, strains from eight freshwater diatom
species were selected from the Thonon Culture Collection (TCC; http://www6.inra.fr/carrtel-collection_eng/) (Table 1). The eight species were chosen for their contrasted morphological (size and cell biovolume), cytological (e.g. chloroplast number) and phylogenetic characteristics (Table 1). Cell dimensions (width, length, thickness) of the eight diatom species were measured under light microscopy (1,000 × magnification) using a minimum of 10 specimens per species. Then, appropriate geometrical models were applied to calculate their cell biovolume (Sun & Liu, 2003) (Table 1). The eight diatom cultures were cultivated in triplicate in 40 ml sterile DV medium (Rimet et al., 2014) using 50 ml Nunc™ EasYFlasks™ (Thermo Fisher Scientific, Waltham, Massachusetts). Flasks were placed on a rotating platter (4 rpm) in a controlled thermostatic room (21 ± 2°C, 14 hr light/10 hr dark cycle, light intensity of c. 100 μmol quanta m⁻² s⁻¹). Flasks were inoculated in order to reach a concentration of ≈100 cells/ml at the beginning of the experiment for each species, except for 

\[ \text{Pinnularia viridiformis} \] for which the survey lasted 73 days, due to its low growth rate. Cell concentrations, proportions of live/dead cells and \( \text{rbcL} \) gene copy concentrations per ml of media were measured for each culture at seven sampling times (referred to as T0 to T6) (Figure 1).

Diatom cell concentrations and proportions of live/dead cells were obtained by counting at least 400 specimens using inverted microscopy (×1,000 magnification) and the standard Utermöhl technique (European Committee for Standardization (CEN) 2006) (Figure 1). The proportion of live/dead cells was estimated by considering cells without visible intracellular contents as dead. Only living cells were taken into account to calculate the diatom cell concentration per ml of media. Flow cytometry using Sytox-Green was also used to confirm the microscopical data (not shown).

\( \text{rbcL} \) copy number per ml was estimated by qPCR. From each cultivation replicate, 10 ml of culture was centrifuged at 17,000 × g for 30 min (Figure 1). Total DNA was extracted from the resulting pellet using a protocol based on GenElute™-LPA DNA precipitation (Sigma-Aldrich, St Louis, Missouri) as previously described (Vasselon, Domaizon, et al., 2017). Then, qPCR assays were performed for each

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**TABLE 1** Characteristics of the eight diatom species selected in the Thonon Culture Collection (TCC) and used in this study

<table>
<thead>
<tr>
<th>Species</th>
<th>TCC code</th>
<th>Chloroplast (nb./cell)</th>
<th>Length (μm)</th>
<th>Width (μm)</th>
<th>Thickness (μm)</th>
<th>Biovolume (μm³)</th>
</tr>
</thead>
</table>
of the eight diatom species on DNA extracted at all seven sampling times and with each of the three replicates, using the QuantiTect SYBR Green PCR Kit (Life Technologies, Carlsbad, USA) and the Rotor-Gene Q (Qiagen, Hilden, Germany). A short 312 bp region of the rbcL gene (the same as was used for HTS sequencing) was targeted using primers used by (Vasselon, Rimet, et al., 2017) and described in Table S1. qPCR reactions were performed following the method used by Vasselon, Domaizon, et al. (2017), using a final volume of 25 μl using mix preparation and reaction conditions as described in Table S1. A fluorescence threshold of 0.01 was used to allow comparison of qPCR assays, denoising and determination of the cycles’ threshold (Ct). Data analysis was performed using the Rotor-Gene Q Series software (version 2.3.1) and the rbcL copy per ml of media was determined.

Finally, the number of rbcL gene copies per diatom cell was calculated for the eight diatom species by dividing the rbcL concentration (qPCR data) by the living cell concentration (microscopy data). A Kruskal–Wallis test was performed using r (R Development Core Team, 2013) to determine if the rbcL gene copy number per diatom cell varied significantly between the eight diatom species. Then, we tested the level of correlation between the number of rbcL gene copies per diatom cell and several morphological characteristics of the diatom cells (Table 1). Variables that did not approximate normal distributions were log transformed. Pearson correlation coefficients were calculated between the gene copy number per cell and the diatom cell morphological characteristics. This correlation was represented by a linear model.

2.2 Validation in the quantification CF to mock and environmental HTS data

2.2.1 Mock communities

The calculated CF was applied to metabarcoding data obtained from controlled diatom mock communities. Five mock communities (M1 to M5) were created by mixing DNA extracted from each of the eight diatom species sampled during their exponential growth phase, and for which the correspondence between cell abundances (microscopy) and qPCR counts was known. For each of the five mock communities, the volume of DNA used for seven species was kept unchanged (1 μl) and only the volume of DNA of P. viridiformis varied as followed: M1 = 0.2 μl, M2 = 0.4 μl, M3 = 0.8 μl, M4 = 1.6 μl, M5 = 3.2 μl. This resulted in contrasted rbcL proportions of the eight species among the five mock communities. Then, HTS sequencing of the rbcL 312 bp fragment was performed on three replicates of the five mock communities. The 15 corresponding libraries were prepared following the method described by Vasselon, Domaizon, et al. (2017) with the same primers and PCR reaction conditions as those used for rbcL qPCR (Table S1), changing only the cycle number to 30. Each library was diluted to 100 pm and all 15 were pooled together for one HTS run performed on the PGM Ion Torrent machine by the “Plateforme Génome Transcriptome” (PGTB, Bordeaux, France).

The sequencing platform provided a unique fastq file for each of the 15 libraries containing demultiplexed DNA reads without the sequencing adapters. Quality filtering of DNA reads was performed using the mothur software (Schloss et al., 2009) and bioinformatics process described previously (Vasselon, Domaizon, et al., 2017; Vasselon, Rimet, et al., 2017). Finally, a taxonomy was assigned to each DNA read with the “classify.seqs” command (Mothur) using default parameters with a confidence threshold of 85% and the R-Syst::diatom library (Rimet et al., 2016, version updated in January 2015 and available upon request) as a rbcL reference library. A molecular taxonomic list with the associated read numbers assigned to each of the eight diatom species was obtained for each of the five mock communities and used for subsequent analysis.

The quantification CF defined for the rbcL gene was then applied to the molecular taxonomic lists for the five mock communities by dividing the read number for each species by its corresponding CF. Both the uncorrected and corrected HTS relative abundances of species from the five mock communities were then compared to the relative abundances obtained using microscopy.

2.2.2 Environmental diatom assemblages

To evaluate the efficiency of the CF to improve metabarcoding quantification from environmental samples, we used rbcL HTS data obtained from (Vasselon, Rimet, et al., 2017), corresponding to 80 benthic diatom samples collected from rivers in tropical island of Mayotte, Indian Ocean (Vasselon, Rimet, et al., 2017 dataset, https://doi.org/10.5281/zenodo.400160). A CF was calculated for each species (or genus when the species level was not reached) detected in molecular inventories of the rivers of Mayotte island using a generalized average of the morphological information (e.g. biovolume, length) available in the R-Syst::diatom library. A Specific Pollution-sensitivity Index (SPI) used for ecological assessment was calculated for each sample based on the corrected diatom molecular inventories using the omindia 5 software (Leclonte, Coste, & Prygiel, 1993; library 5.3 2015) and compared to the morphological SPI values for all river samples (Vasselon, Rimet, et al., 2017). Pearson correlation was used to evaluate the strength of correlations between original or corrected molecular SPI values and the morphological SPI values. Wilcoxon Signed Rank tests were conducted to determine whether the difference between the molecular and the morphological SPI (ΔSPI) varied significantly when using the original or the corrected molecular data for the molecular SPI calculation.

3 RESULTS

3.1 Variation in rbcL gene copy number between diatom species

Cell and rbcL gene concentrations were measured, by inverted microscopy and qPCR, respectively, for the eight diatom species at different cultivation stages corresponding to seven sampling points (T0 to T6).
Information has been summarized in Tables S2 and S3. As the eight diatom species reached the beginning of the stationary phase at the sampling time T2 (i.e. between 13 and 31 days of cultivation), only the [cell] and the [gene copy] values obtained for the T0, T1 and T2 sampling times were used for further analysis. The calculated mean values of the rbcL gene copy number per cell for each diatom species varied between 0.5 and 130 copies per cell (Figure 2). The Kruskal–Wallis test revealed that the rbcL copy number per cell was significantly different ($p < .001$) between the eight diatom species.

### 3.2 Development of quantification CFs

The $rbcL$ copy number per cell was highly correlated with cell biovolume ($r = .97$, $p < .001$), length ($r = .82$, $p < .001$), width ($r = .94$, $p < .001$) and thickness ($r = .96$, $p < .001$). The correlation between the $rbcL$ copy number per cell and the cell biovolume followed a linear model (Figure 3). Assuming that this linear relation based on eight diatom species is applicable to all diatom species, the equation of this model allows calculation of an estimate of the relative $rbcL$ copy number per cell as soon as the biovolume of the cell is known, and thus to define a CF specific to each species. Such quantification CFs were calculated for each of the eight diatom species of the mock communities (Table 2) and varied from 0.6 for *Achnanthidium minutissimum* to 78.5 for *P. viridiformis*. For each of the diatom taxa found in the environmental samples, CFs were also calculated using the biovolume information available for each taxa (from Rsyst::diatom library) (Table S4) and varied over a wider range, from 0.03 for *Fistulifera saprophila* to 649.8 for *Rhopalodia gibba*.

### 3.3 Application of CFs to mock and environmental HTS data

953,082 DNA reads were produced from the 15 libraries corresponding to the five DNA mock communities (3 replicates per mock). Following the bioinformatics quality filtering steps, 385,367 DNA reads were retained. A molecular taxonomic list was then created by removing DNA reads which remained unclassified (0.43% of the reads) or assigned to different taxa than the eight diatom species present in the mock communities (0.004% of the reads) (Table S5). The proportions of *P. viridiformis* reads in the five mock communities varied from 9% in M1 to 57% in M5 (Figure 4a) while observed cell proportions were lower; ≈0.03% in M1 and 0.55% in M5 (Figure 4b). The application of the CF on DNA reads counts of the eight species changed their relative abundances in the five mock communities (Figure 4a). The rank of the eight species was also affected; for example in M5 the application of the CF changed the proportion of *P. viridiformis* from 57% to 4% and the proportion of *A. minutissimum* from 4% to 42%. The correspondence between morphological and molecular relative abundances was highly improved by applying the CF on the HTS data (Figure 4a,b).
From the 80 environmental samples previously sequenced (Vasselon, Rimet, et al., 2017), a molecular taxonomic list based on assigned DNA reads was produced including 23 families (75.1% of total reads assigned), 39 genera (72% of total reads assigned) and 66 diatom species (40.7% of total reads assigned). From this list, 84 diatom taxa, including taxa assigned at the genus and the species level, were used to calculate the SPI freshwater quality index. CFs calculated from cell biovolumes for those 84 taxa were then applied to correct the quantification of the environmental molecular inventories (Table S4). The proportions and ranks of the dominant taxa were affected by the application of the CFs (Figure 5). For example the application of CFs reduced the relative abundances of *Eunotia* and *Ulnaria* from 31.9% to 3.3% and 11.7% to 2.3%, respectively, making them more congruent with cell proportions observed with microscopy (3.1% for *Eunotia* and 0.4% for *Ulnaria*). The correlation between the morphological and the molecular SPI values for all river samples previously described ($r = .72$, $p < .001$) was slightly improved using SPI values based on inventories with corrected abundances ($r = .77$, $p < .001$). The application of the CF to correct the HTS quantification reduced significantly ($p < .001$) the differences between the molecular and morphological SPI values by 47% ($\Delta$SPI reduced to 1.9 on average compared to 3.6 before correction, corresponding to 37.3% and 21.2% of error respectively) (Figure 6).

4 | DISCUSSION

Species quantification based on DNA metabarcoding is challenging for most of taxonomic groups as technical and biological biases affect DNA reads proportions. In order to limit those biases, several attempts were done to apply a CF on metabarcoding data, as shown for fishes (Thomas et al., 2016), bacteria and archea (Angly et al., 2014) or oligochaetes (Vivien et al., 2016). For those studies, application of the CF, whether for correcting single (Angly et al., 2014) or multiple sources of quantification biases (Thomas et al., 2016), improved taxa quantification from metabarcoding data compare to morphological one. The result is generally a change in the ranks of the dominant taxa which affect directly the community structure and can lead to different ecological interpretations. For example the application of a CF on metabarcoding data obtained from aquatic oligochaetes samples improved the freshwater quality assessment based on molecular index calculation (Vivien et al., 2016). However, the development of CF can be challenging depending on the organism studied, as it requires finding a clear relationship between DNA reads and specimen proportions. This may be impossible due to accumulation of quantification biases (e.g. cell density, cell biomass, gene copy number). Nevertheless, the use of CF can be advantageous for organisms with a high variation in the DNA reads proportions between taxa (e.g. several log) and where a limited number of biases are involved like diatoms.

4.1 | Correlation between rbcL gene copy number and diatom cell biovolume: Impacts on HTS quantification

The copy number of the rbcL gene present in one diatom cell is affected by three parameters: (1) the number of chloroplasts per cell, (2) the number of genomes per chloroplast and (3) the number of...
For example, while unicellular algae generally exhibit a lower number of copies per cell (Bendich, 1987; Rauwolf, Golczyk, Greiner, & Herrmann, 2010) can contain up to thousands of copies of chloroplast genome per cell. Regarding the chloroplast genome number per cell, higher plants have no maximum of 130 copies estimated for Olisthodiscus luteus (Raphidophyceae), Thalassiosira pseudonana (centric diatom) contain, respectively, around 650, 80, 137 and 55 genome copies per cell (von Dassow, Petersen, Chepurnov, & Virginia Armbrust, 2008; Erland et al., 1981; Gruber, 2008; Koop, Herz, Golds, & Nickelsen, 2007). (3) Finally, there is only one copy of the rbcL gene per chloroplast genome. This was also confirmed with the Mayotte river samples, for which the quantification CF resulted in a better congruence between DNA reads and cells proportions, reducing the over-representation of high biovolume diatom species with high cell biovolume are over-represented (e.g. P. viridiformis) and diatom species with low cell biovolume are underestimated (e.g. A. minutissimum). The relative abundance of P. viridiformis in the mock communities was negligible compared to other species, and doubling its proportion did not change its rank: the species remained the least abundant taxon within the morphological inventory. However, due to its high cell biovolume ($10^4 \mu m^3$) and relatively high rbcL copy number per cell, a marked over-representation of this species within the molecular inventory was observed. A CF was thus defined to correct these quantitative biases and was verified on mock communities and environmental samples.

![Figure 6](image.png)

**Figure 6** Distribution of the differences between the molecular and the morphological SPI ($\Delta$SPI) for all Mayotte samples using original molecular SPI values (left) and new molecular SPI values based on molecular inventories corrected with the biovolume CF (right).  

<table>
<thead>
<tr>
<th>rbcL copies</th>
<th>10.0</th>
<th>7.5</th>
<th>5.0</th>
<th>2.5</th>
<th>0.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>Without CF</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>With CF</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

| ΔSPI (SPI null−SPI morph) | 3.6 | 1.9 |

4.2 | Current potential and limits of the quantification CF

The use of the same rbcL primers for the qPCR assays and the HTS enabled us to generate a specific CF well suited to correct rbcL metabarcoding quantifications. Its application to the HTS data of the mock communities allowed us to obtain comparable species proportions in morphological and molecular based approaches of mock communities. This was also confirmed with the Mayotte river samples, for which the quantification CF resulted in a better congruence between DNA reads and cells proportions, reducing the over-representation of high biovolume Eunotia and Ulnaria species. Furthermore, SPI calculation based on corrected metabarcoding data gives SPI values more comparable to SPI values obtained from morphological data, suggesting that it may be possible to replace morphological by molecular monitoring for the ecological assessment of Mayotte rivers. In the same way, (Vivien et al., 2016) have shown that application of a CF to correct DNA reads proportions allows a more accurate estimation of oligochaete proportions, improving quality index calculation and quality assessment of watercourse sediments. Our results confirm that water quality index based on diatom metabarcoding and DNA read proportions are directly affected by gene copy number variation, and show the potential value of integrating CFs into molecular SPI calculation. However, as the biovolume–copy number relationship was based on only eight diatom species and the efficiency of the resulting CFs validated on only one HTS dataset, further experiments including more species and larger datasets will be required to develop and fully validate CFs for use in molecular biomonitoring.
The CF developed in this study assumes that gene copy number is constant in each taxon. However, gene copy number may vary with the physiological status of the cell and stage of the life cycle, since in most diatoms cell volume decreases during the vegetative phase. The physiological status varies with cell cycle progression; additionally several factors may affect the physiological status of diatoms like changes in environmental conditions (e.g. nutrient availability, pollutants, temperature, etc.) (Pandey et al., 2017). Altered physiological status of a given population is generally characterized by a higher proportion of damaged cells. The compromised/damaged cells are characterized by alteration of membrane integrity, degradation of the photosynthetic pigments or fragmentation of genomic DNA (Zetsche & Meysman, 2012; Znachor, Rychtecký, Nedoma, & Visocká, 2015). Variations in DNA integrity and chloroplast physiology between cells of a given population can impact directly the rbcL gene copy number per cell and thus DNA metabarcoding quantification. (Eberhard, Drapier, & Wollman, 2002) showed that chloroplast genome copy number is reduced when the green alga *Chlamydomonas reinhardtii* is cultivated under phototrophic conditions compared to cultivation in mixotrophic conditions. Limitation by mineral nutrients may also have an impact; for instance iron limitation can reduce the number of the chloroplast per cell (from 4 to 2) and their size in the marine diatom *Thalassiosira oceanica* (Hustedt) Hasle et Heimdal (Lommer et al., 2012). Variation in the cell physiological state was not taken into account in developing CFs for diatom metabarcoding. However, during our experiments we discriminated live and dead cells; we observed that their respective proportions did not affect significantly the correlation between the gene copy number per cell and the cell biovolume (Figure S1). Further experiments should be performed to evaluate the impact on the final CFs of rbcL gene copy number variation linked to physiological status.

The biovolume of each diatom species is required to apply the CF and hence correct the quantification in metabarcoding datasets. Several reference databases provide biovolume information for a lot diatom species (e.g. Rimet et al., 2016), but they do not generally account for biovolume variability, which is a complicating factor in diatoms because of the peculiarities of the life cycle. Diatom cell size within a population is not constant due to the method of vegetative reproduction, which leads to a progressive cell size reduction in the population (Crawford, 1981), followed by restoration of cell size via a sexual event. For this reason, different cell sizes can be observed in the same diatom population, either in pure cultures (e.g. in the marine diatom *Thalassiosira weissflogii* Grunow; Armbrust & Chisholm, 1992) or in environmental populations (e.g. the freshwater species *Sellaphora pupula* (Kützing) Mereschk: (Mann, Chepurnov, & Droop, 1999)). However, although the range of cell sizes within a given diatom population may vary by a factor of 2 to 5 in the environment (Hense & Beckmann, 2015), natural populations usually have a rather narrow range of sizes and larger cells form a negligible fraction of the population (Mann, 2011). Furthermore, the distribution of cell size within environmental populations is often close to being normal (Mann et al., 1999; Spaulding, Jewson, Bixby, Nelson, & McKnight, 2012). The balance between small and big individuals in the same population will therefore limit errors associated with the use of a mean biovolume. Hence, we propose to use the mean of biovolume to calculate CFs; without considering other potential HTS quantification biases, its application to DNA reads of environmental material should allow a good correction of their proportions.

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**AUTHORS’ CONTRIBUTIONS**

V.V., A.B., F.R., S.J., M.C., K.T. and I.D. contributed to the study designed. V.V., M.C. and S.J. conducted the laboratory work. V.V. analysed the data and wrote the manuscript. All the authors contributed to the discussions and to manuscript editing.

**DATA ACCESSIBILITY**

All PGM raw sequence data are available for the 15 libraries, corresponding to the five DNA mock communities with three replicates, on the Zenodo repository website (https://doi.org/10.5281/zenodo.807178).

**ORCID**

Valentin Vasselon http://orcid.org/0000-0001-5038-7918

**REFERENCES**


SUPPORTING INFORMATION

Additional Supporting Information may be found online in the supporting information tab for this article.