A qPCR method for distinguishing biomass from non-axenic terrestrial cyanobacteria cultures in hetero- or mixotrophic cultivations

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Abstract
The cultivation of cyanobacteria with the addition of an organic carbon source (meaning as heterotrophic or mixotrophic cultivation) is a promising technique to increase their slow growth rate. However, most cyanobacteria cultures are infected by non-separable heterotrophic bacteria. While their contribution to the biomass is rather insignificant in a phototrophic cultivation, problems may arise in heterotrophic and mixotrophic mode. Heterotrophic bacteria can potentially utilize carbohydrates quickly, thus preventing any benefit for the cyanobacteria. In order to estimate the advantage of the supplementation of a carbon source, it is essential to quantify the proportion of cyanobacteria and heterotrophic bacteria in the resulting biomass. In this work, the use of quantitative polymerase chain reaction (qPCR) is proposed. To prepare the samples, a DNA extraction method for cyanobacteria was improved to provide reproducible and robust results for the group of terrestrial cyanobacteria. Two pairs of primers were used, which bind either to the 16S rRNA gene of all cyanobacteria or all bacteria including cyanobacteria. This allows a determination of the proportion of cyanobacteria in the biomass. The method was established with the two terrestrial cyanobacteria Trichocoleus sociatus SAG 26.92 and Nostoc muscorum SAG B-1453-12a. As proof of concept, a heterotrophic cultivation with T. sociatus with glucose was performed. After 2 days of cultivation, a reduction of the biomass partition of the cyanobacterium to 90% was detected. Afterwards, the proportion increased again.

Keywords Terrestrial cyanobacteria · qPCR · Heterotrophic cultivation · Mixotrophic cultivation · Phototrophic biofilms

Introduction
Cyanobacteria are an interesting source of natural bioactive compounds (Chlipala et al. 2011) that possess antibacterial (Gantar et al. 2011; Lamprinou et al. 2015), antimycotic (Swain et al. 2017) and anticancer (Kinghorn et al. 2016) activity. Terrestrial cyanobacteria, particularly, form an interesting subgroup of previously uninvestigated diversity (Garcia-Pichel et al. 2001). They produce energy via photosynthesis and are therefore normally cultivated in mineral salt medium (e.g. BG11 or Z8) (Allen 1968) under light. Phototrophic cultivation leads to slow growth rates (~ 0.1–0.5 day⁻¹) (Chojnacka and Noworyta 2004; Kuhne et al. 2014). In addition, the necessary light supply makes the upscaling of cultivation systems more difficult and requires special photobioreactors (Strieth et al. 2018). The production of biomass for the investigation of natural substances is thus severely restricted. One possibility to avoid this limitation is cultivation with the addition of carbohydrates (Yu et al. 2009; Trabelsi et al. 2013). If the cyanobacteria metabolize these in addition to the photosynthesis, they can produce extra biomass (mixotrophic cultivation) (Chojnacka and Noworyta 2004). Furthermore, heterotrophic cultivation without illumination would offer the possibility of cultivation in large scale in the widely used and well-characterized stirred tank bioreactors.

Another problem during culturing cyanobacteria is adherent heterotrophic bacteria. Some of these bacteria cannot be completely removed from the cultures, even with complex isolation procedures (Ferris and Hirsch 1991; Palinska and Krumbein 1995; Temraleeva et al. 2016; Vasquez-Martinez...
et al. 2004). Many of these bacteria, such as Bacillus species, show very high growth rates under optimal nutrient conditions (μ > 1 h⁻¹). Therefore, if a carbohydrate is added to cyanobacteria cultivated under phototrophic conditions, it is possible that a heterotrophic bacterium will use most of it. This falsifies the measured growth rate for the cyanobacterium. The growth of cyanobacteria is usually measured via the cell dry weight or the optical density of the cell suspension. However, both methods do not distinguish between cyanobacteria and heterotrophic bacteria. This means that no reliable conclusions can be drawn about the growth of cyanobacteria, because the growth of heterotrophic bacteria also influences the measured parameters. Consequently, the composition of the biomass must be measured during cultivation to determine the level of contamination. The simplest variant is microscopic evaluation where the cells of the different species can be distinguished and counted to determine the composition of the biomass. However, this method is only suitable for the quantification of similarly sized cells such as microalgae and cyanobacteria (Arias et al. 2019). Yet cyanobacteria are about 10 μm in size and many heterotrophic bacteria are only about 2 μm. In addition, this is relatively labour- and time-intensive and requires high amounts of experience for the correct delineation between organisms. Moreover, bacteria within a biofilm of cyanobacteria cannot be quantified well. They are often located very close to and between the cells of cyanobacteria and therefore cannot be distinguished from them. Another technique for contamination determination is the use of automated fluorescence-based cell sorting (FACS), where each individual cell is sorted and subsequently counted automatically (Zhou et al. 2018). The differentiation when sorting is accomplished by the inherent auto-fluorescence of the cyanobacteria. This enables the small heterotrophic bacteria to be easily distinguished from the larger cyanobacteria. A FACS can only count single cells. Therefore, this method is limited to planktonically growing cyanobacteria and not suitable for filamentous- or biofilm-forming species.

An alternative to the aforementioned methods is the use of PCR. In the field of water monitoring, there have long been approaches to detect the occurrence of cyanobacteria, which are able to produce harmful cyanotoxins (Zhang et al. 2014). For this purpose, primers were established to detect genes necessary for toxin synthesis (Pacheco et al. 2016). A proven method for the detection of non-harmful strains has been established by the use of primers, targeting conserved regions of the 16S rRNA gene (Nübel et al. 1997). Although this method seems promising, there are inherent challenges that arise when extracting DNA from cyanobacteria. The most apparent difficulty is the production of mucilaginous polysaccharides by the cyanobacteria, which hinders cell lysis and interferes with most DNA extraction protocols (Porter 1988). One popular method, based on the use of xanthogenate, can lyse the cell walls effectively and forms insoluble aggregates from proteins and cell debris. These can then be removed immediately after extraction to achieve high initial sample purity (Tillett and Neilan 2000).

Other authors (Heck et al. 2016) presented a qPCR method to investigate the success of methods for the purification of Fischerella sp. CENA161. In this method, the 16S rRNA gene of cyanobacteria and bacteria were quantified using an external plasmid DNA standard. The ratio of the copy numbers was taken as a measure for the purity of the sample. It was shown that the additional steps of the treatment with sodium hypochlorite and detergent washing increased the DNA content of the cyanobacteria in the culture from 70 to 90%. However, there was no correlation to the proportion of cyanobacterial biomass.

This work presents a qPCR method for quantifying the biomass fraction of cyanobacteria in cultivations with a carbohydrate source. The DNA of two cyanobacteria (Trichocoleus sociatus and Nostoc muscorum) was purified with an advanced xanthogenate method and quantified with two 16S rRNA gene primer sets. The biomass in the samples was then quantified by correlation with a standard curve. The cyanobacterium T. sociatus was cultivated heterotrophic with glucose. The composition of the biomass after 2, 5, 7 and 14 days was determined.

**Material and methods**

**Strains and cultivation**

The cyanobacteria studied (Trichocoleus sociatus DSM 109967/SAG 26.92 and Nostoc muscorum SAG B-1453-12a; both obtained from the strain collection of Professor Büdel, Department of Plant Ecology and Systematics, TUK) were cultivated under phototrophic conditions at 100 μmol photons m⁻² s⁻¹, 27 °C and 120 rpm in an illuminated shaker (Multitron, Infors, Switzerland). Cultivation took place in 500-mL shaking flasks without baffles in BG11 medium (Allen 1968). The biomass was inoculated into flasks with fresh media at regular intervals of 4 weeks for strain maintenance. For the heterotrophic cultivation, 2 g L⁻¹ cell wet weight (cww) in a modified BG11 medium with 2.53 g L⁻¹ glucose was cultivated for 14 days at 27 °C without illumination.

All bacteria (Bacillus subtilis DSM618, Bacillus sp. (own wild-type isolate from culture T. sociatus) and Escherichia coli K12 JM109 DSM3423) were cultivated in Mueller-Hinton-Bouillon (Carl Roth, Germany) for 17 h at 37 °C and 150 rpm. Saccharomyces cerevisiae DSM3799 was cultivated in YPG medium (yeast extract 10 g L⁻¹, 20 g L⁻¹ soy peptone, 20 g L⁻¹ glucose) at 30 °C and 24 h at 150 rpm.
DNA extraction

For DNA extraction from cyanobacteria the method developed by Tillet and Neilan (2000) was modified. This method is based on the use of an extraction buffer (1% potassium ethyl xanthogenate, 1% sodium dodecyl sulfate, 100 mM Tris, 20 mM EDTA, 800 mM ammonium acetate, pH 8.0). The buffer breaks down the cells and extracts the DNA directly; a separate cell disruption is not necessary. However, in order to extract the DNA from the pellet-like growing biomass, a short treatment in a ball mill was added in this work upstream. The ball mill disintegrates the pellet so that the buffer can penetrate the cell wall. To increase the purity of the DNA, also a phenol-chloroform extraction after the DNA extraction was added in this study. Approximately 50 mg cww of fresh cyanobacteria was suspended in 400 μL TE buffer (10 mM Tris, 1 mM EDTA, pH 8.0) and a stainless steel ball (3.75 mm diameter, Heinrich Amelung GmbH, Germany) was added. The sample was then crushed for 1 min at a frequency of 30 s⁻¹ in the ball mill (MM 301, Retsch, Germany). The cells were separated by centrifugation at 12,000×g for 10 min. The supernatant was collected. The pellet was resuspended with 600 μL freshly prepared extraction buffer and incubated for 60 min at 70 °C and 900 rpm in a thermo shaker (TS-100, Biosan, Latvia). Afterwards, it was cooled on ice for 30 min. The remaining cell debris were separated by centrifugation at 12,000×g for 10 min. Both supernatants were pooled and the DNA was extracted with 1 mL phenol-chloroform-isooamyl alcohol mixture pH 8.0 (product number 0883, VWR, PA, USA). For this, the two phases were mixed for 1 min in the ball mill with a frequency of 30 s⁻¹. The two phases were then separated in the centrifuge for 10 min at 12,000×g. Eight hundred microliters of the aqueous top phase was re-extracted with 800 μL chloroform using the same procedure. Subsequently, the DNA in 700 μL of the aqueous top-phase was precipitated with 700 μL of ice-cold 2-propanol for 18 h at −20 °C. The DNA was separated by centrifugation at 12,000×g for 10 min at 4 °C. The pellet was washed with ice-cold 70% ethanol, then air-dried and dissolved in 100 μL TE buffer.

The DNA of the bacteria and yeasts for the analysis of the primer specificity were extracted with the GenElute Bacterial Genomic DNA Kit (Sigma-Aldrich, USA) according to the manual. The concentration and purity of DNA was determined using a NanoDrop 2000 spectrophotometer (Thermo Fisher Scientific, USA).

qPCR

Two primer pairs were used for the qPCR. Primer pair (PP) Cya (Cya359F: GGG GAA T(C/T)T TCC GCA ATG GG and Cya781R: GAC TAC TGG GGT ATC TAA TCC CAT T (Nübel et al. 1997)) binds specifically to the 16S rRNA gene sequence of cyanobacteria. Primer pair Bak (COR-F: CCAATGAATCGGAA TCGCTAG and COR-R: ACTCCCATGGGTGTCACGACGG (Corless et al. 2000)) binds specifically to the 16S rRNA gene sequence of all bacteria including cyanobacteria. The primers were ordered Lab-Ready (100 μM in IDTE buffer, pH 8.0) and standard desalted (Integrated DNA Technologies, USA). The Cya forward primer was supplied as an equimolar mixture of C and T, respectively, at position 8. The PCR was performed in 96-well MTPs (catalogue number HSP9601, Biorad, USA), which were manually sealed with adhesive foils (catalogue number MSB1001, Biorad, USA). The entire qPCR preparation was performed manually under a fume hood at room temperature. Approximately 5 ng DNA, 0.1 μM primer and 10 μL Mastermix (SsoAdvanced Universal SYBR Green Supermix, Biorad, USA) were used and filled up with DNA-free water for a final volume of 20 μL. The qPCR was performed in a CFX Connect System (Bio-Rad, USA) with the following parameters: Initial 3 min 98 °C; 40 cycles 15 s 98 °C, 10 s 53 °C, 50 s 60 °C. The cycle threshold values (Cq) were determined by the CFX Maestro software (Version 4.1.2433.1219; Biorad, USA) using the single threshold method. The qPCR products were tested by default with a melting curve (65–95 °C, 0.5 °C steps). The melting point of PP Cya product was 86.5 °C, that of PP Bak 87.5 °C. In method development, the product length was regularly checked using a 1.5% agarose gel in TAE buffer using a low-range ladder (GeneRuler Low Range DNA; Thermo Fisher Scientific, USA). The amplicon length was determined for the example of T. sociatus (NCBI accession number EF654080.1) in silico. Amplicon Cya starts at bp 303 and ends at bp 728, is 425 bp long and has a GC content of 53.4%. Amplicon Bak starts at bp 1252 and ends at bp 1343, is 91 bp long and has a GC content of 60.4%. The amplicons were checked for secondary structures via the mfold web server (Markham and Zuker 2005; Keith 2008). Using the relevant PCR parameters (53 °C, concentrations of 50 mM Na⁺ and 0 mM Mg⁺²), structures with a free energy of −4.77 kcal mol⁻¹ for PP Cya and −3.24 kcal mol⁻¹ for PP Bak were predicted (see Supplemental S3 and S4). The Cya primers used were examined for their specificity using NCBI-BlastN. The entire nucleotide collection was tested, excluding cyanobacteria (ID 1117). Additionally, uncultured/environmental sample sequences were excluded and otherwise the standard parameters for short sequences were used. The forward primer was not specific, e.g. it annealed to B. subtilis DSM618. For the reverse primer, only hits were found
against chloroplasts that are related to cyanobacteria and mycoplasmas that are also related at the 16sRNA gene level.

**Primer efficiency**

To determine the efficiency of the primers, a dilution series was prepared and the Cq values for both primer pairs were determined by the qPCR method as a technical triplicate. For this purpose, the DNA of *T. sociatus* (initial concentration in qPCR 20 ng μL⁻¹) was diluted with water in 1:10 steps. The efficiency (E) of the primers was calculated in the linear range using the following equation (Svec et al. 2015):

\[
E = \left(10^{-\frac{1}{\Delta Cq}} - 1\right) \times 100
\]

where \(X\) is the slope of the linear regression of the qPCR threshold (Cq) value over the logarithmic template concentration. The slope was determined using a linear fit in Origin 2019 (version 9.6.0.172; OriginLab Corporation, USA).

**Calibration curve**

For calibration, mixtures of 70 to 100% dry weight percent (wt%) of *T. sociatus* respectively 60 to 100% for *N. muscorum* together with *B. subtilis* DSM 618 were used (e.g. 60% wt% cyanobacterium means that 1 g cell dry weight (cdw) contains 600 mg of the respective cyanobacterium and 400 mg *B. subtilis*). *Bacillus subtilis* was cultivated overnight in 100 mL LB medium (Carl-Roth, Germany) in a 500-mL shaking flask with baffles at 160 rpm and 37 °C for 12 h. The DNA of the mixtures was extracted and purified with the protocol for cyanobacteria. Then, a qPCR was performed with the DNA preparations using both primer pairs. Biological triplicates were measured in technical triplicate.

The Cq for the amplification of the Cya primer was corrected by the Cq of the Bak Primer by subtraction (Eq. 2) to compensate for possible differences in the absolute DNA concentration in the PCR well according to Livak and Schmittgen (2001). This could arise from measuring the DNA concentration in the sample and from pipetting error.

\[
\Delta Cq = Cq_{Cya} - Cq_{Bak}
\]

The resulting \(\Delta Cq\) was then used to generate the calibration function. Equation 3 was used to fit the data points:

\[
\Delta Cq = a \cdot e^{\frac{\text{partition cdw cyanobacteria}}{\text{cdw cyanobacteria}}} + c
\]

with \(a\), \(b\) and \(c\) as empirical parameters and

\[
\text{partition cdw cyanobacteria} = \frac{\text{cdw cyanobacteria}}{\text{cdw cyanobacteria} + \text{cdw B. subtilis}} \times 100
\]

The empirical parameters were determined using the ExpDec1 function with Origin 2019.

**Results and discussion**

**DNA extraction**

The method according to Tillett and Neilan (2000) was extended with a chloroform phenol extraction to increase the purity of the obtained DNA. Although we successfully extracted DNA with this method, the reproducibility and robustness were insufficient. The studied cyanobacteria form pellets of up to 0.5 cm diameter, and therefore, the cells inside the pellet cannot be penetrated well for DNA extraction with XS buffer. A short pre-treatment in a ball mill was therefore applied. This disrupts the pellets and allows the XS buffer to effectively extract the DNA from the separated cells. The treatment in the ball mill is kept as brief as possible in order not to break up cells. But because this cannot be definitely excluded, the supernatant was kept and then combined with the extract for purification. For phototrophic culture of *T. sociatus*, a DNA yield of 2.29 ± 0.02 μgDNA mg⁻¹ cdw or 151 ± 1.6 ngDNA mg⁻¹ cdw and for *N. muscorum* a yield of 3.49 ± 0.74 μgDNA mg⁻¹ cdw or 230 ± 49 ngDNA mg⁻¹ cdw can be achieved. For mixotrophic and heterotrophic cultures of *T. sociatus*, median yields of 149 ngDNA mg⁻¹ cdw were achieved. With the initial method, higher overall yields of 12.7 ± 6.0 μgDNA mg⁻¹ cdw were achieved (Tillett and Neilan 2000). However, other cyanobacteria were examined. The yields are strain and order specific. For the strains which, like the *N. commute* examined here, belong to the order of Nostocales, Tillet and Neilan (2000) found smaller yields of 9.25 ± 3.27 μgDNA mg⁻¹ cdw. In this work, a purification with phenol-chloroform extraction was also implemented. This probably reduces the yield. In any case, the DNA yields achieved were sufficient for the qPCR analyses performed. The purity of the purified DNA was determined by the quotient of adsorption at 260 and 280 nm in the NanoDrop. The value of 1.9964 ± 0.1395 can be considered sufficiently pure.

**Primer and calibration curve**

The aim was to develop a method to determine the exact composition of the biomass of a co-culture of cyanobacteria and heterotrophic bacteria. However, two characteristics of
the non-axenic cultures prevented a standard absolute quantification. Firstly, there are no pure cultures of the quantified cyanobacteria, so that no direct standard curves can be generated. Second, the exact composition of the heterotrophic co-cultures is unknown, so no specific primers can be selected for them. The composition could certainly be determined relatively precisely for the individual cultures by means of complex microbiological and genetic methods, but this was not the approach of this work. The aim was to develop a method that could be quickly transferred to any other cyanobacterial culture. Therefore, two primer pairs were selected based on the literature. Primer pair Cya (Nübel et al. 1997) binds specifically to the 16sRNA sequence of all cyanobacteria. Primer pair Bak (Corless et al. 2000) binds to the 16S rRNA gene sequence of all bacteria including cyanobacteria. Better would be an additional pair of primers that bind all bacteria except cyanobacteria, but we do not know of any such primers. Both can be annealed at the same temperature (53 °C) and thus analyzed in the same PCR run. The specificity of the two primer pairs was tested against two cyanobacteria (T. sociatus and N. muscorum), two Gram-positive bacteria (B. subtilis and B. spec.), one Gram-negative bacterium (E. coli K12) and one yeast strain (S. cerevisiae, as negative control) (see Fig. 1).

The Cya primer achieved a Cq value significantly below the no template control (NTC) for the two tested cyanobacteria. The remaining bacteria and the yeast had a signal at the level of the NTC. It can therefore be confirmed that the Cya primers only amplifies cyanobacterial DNA. Nübel (Nübel et al. 1997) already successfully tested the specificity of the primers with a larger number of cyanobacteria (23) and bacteria (11). The primer specificity was also successfully tested in silico by Mühling et al. (2008). The Bak primer generated a signal in all bacteria including cyanobacteria below the NTC but not for S. cerevisiae. The Bak primer therefore amplifies the DNA of all tested bacteria. This includes the dominant heterotrophic contamination (B. sp.) isolated from T. sociatus. The melting curves and subsequent gel electrophoresis confirmed the purity of the PCR products. The melting point for the amplicon of primer pair Cya is 86.50 °C, for the amplicon of primer pair Bak 87.50 °C (see S. 2). The in silico amplicon length for T. sociatus for Cya is 425 bp, for Bak 91 bp, as was estimated from the gel (see S.1). The efficiency of the primers was determined with a 1:10 dilution series of DNA extracted from T. sociatus for Cya with 96% and for Bak with 107% (see Fig. 2a). The intra-assay variance between the technical triplicates was 0.28% for Cya and 0.32% for Bak. At the lowest still detectable DNA concentration (0.0002 ng μL⁻¹), the standard deviation was 0.24% for Cya and 1.21% for Bak. The presence of a signal at the NTC was caused by primer dimers or similar fragments. The melting curve of the NTC usually showed an undefined peak around 70 °C (data not shown). To reduce this problem, different primer pairs or an optimized qPCR setup could be tested (Brownie et al. 1997; Elnifro et al. 2000). For the Bak primers, it is also possible that small amounts of DNA from the bacterial production strain of the polymerase are present in the master mix. These would be amplified by the primers (Corless et al. 2000).

To quantify unknown mixtures of bacteria, a standard function of known mixtures was created. Biomass from phototrophic-cultivated cyanobacteria was used, with the assumption that the proportion of heterotrophic bacteria in the biomass was negligible (not the absolute amount of cells). These were mixed with biomass of B. subtilis. Bacillus subtilis was chosen because a Bacillus species was identified as the dominant heterotrophic bacterium in the culture of T. sociatus (own unpublished results, identified by 16S rRNA gene sequencing after cultivation on LB-Agar). The DNA of the samples was extracted and purified and the Cq values for both primer pairs were determined by qPCR. It was not possible to normalize CqCya with CqBak, because the latter does not remain constant over the biomass concentration. This was also seen in the measured data (not shown) and can also be explained theoretically. The basic assumption is that the CqBak depends on the number of 16S rRNA gene sequences (both from cyanobacteria and B. subtilis) in the sample. The same absolute amount of DNA was applied. Thus, CqBak depends on the amount of 16S rRNA sequences per megabase pair (Mbp). The genome size of cyanobacteria is around 5.66 ± 2.45 Mbp (Louca et al. 2018). The gene copy number of the measured 16S rRNA gene of cyanobacteria is 2.3 ± 1.2 (Větrovský and Baldrian 2013). Simplified, this results in

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**Fig. 1 Cq values of the primers (Cya and Bak) for different microorganisms and a no template control (NTC).** The horizontal lines mark the value of the NTC and provide a better overview. N. muscorum and T. sociatus as mean and standard deviation of biological quadruplicate in technical triplicate, B. subtilis and NTC as mean of technical triplicate, the others are single measurements.
about 0.3 copies per Mbp. The genome of *B. subtilis* is about 4.165 Mbp in size and contains 5 copies, or about 1.2 copies per Mbp (Farrelly et al. 1995). Thus, with an increasing number of *B. subtilis* in the mixture, the amount of gene copies per Mbp in the DNA sample also increases and thus the Ct value decreases. The Ct value depends only on the cyanobacterial DNA and thus decreases with an increasing amount of cyanobacteria in the mixture (not shown). However, it is not possible to use only the Ct value of the cyanobacterium or the Ct value alone to determine the biomass fraction of the cyanobacterium. The absolute Ct value of the individual biological samples is subject to certain variability. This is caused by the amount of DNA used, which fluctuates due to possible errors in the DNA measurement as well as pipetting errors. In addition, small amounts of inhibitors or a fluctuating DNA quality/integrity can play a role in the Ct value. Therefore, we use the ΔCt value according to Livak and Schmittgen (2001). The ΔCt varies with the biomass fraction (see Fig. 2b). At a certain biomass fraction, ΔCt should always assume a certain value. We assume that the variation of the samples has an equal influence on the measurements of CtCya and CtBak. Thus, the difference between the two is therefore always the same. This compensates the variability of the samples. Thus, we can use ΔCt over the biomass fraction to generate a standard function. The data are described using a simple empirical function.

The functions described the data well with a corrected R-squared value of 0.98 for *T. sociatus* and 0.99 for *N. muscorum*. The calibration functions for the two strains under investigation differ. This means that a separate calibration must be created for each additional cyanobacterium strain. For both strains, a biomass fraction of 100–60/70 wt% of the respective cyanobacterium was quantified using the functions. The standard series was not tested further than 60% biomass of the cyanobacterium. This seems acceptable since cultures of less than 60% are probably not interesting for a bioprocess.

Especially for *T. sociatus*, it can be seen that even a small proportion of *B. subtilis* biomass has a great influence on the signal. We reason that the cells of *T. sociatus* and *N. muscorum* are relatively large and therefore presumably contain little DNA amounts in relation to their biomass.

Both cyanobacteria are coccoidal and have a diameter of 5–6 µm, which corresponds to a volume of about 65 µm³. A rod-shaped bacterium like *B. subtilis* with a length of about 2 µm and a diameter of 1.2 µm only has a volume of about 2.2 µm³. Both cyanobacteria and *B. subtilis* probably have a similar amount of DNA per cell but different gene copy number of the 16S rRNA gene sequence (see above). A defined amount of biomass of *B. subtilis* thus contains a much higher amount of DNA with a higher concentration of target gene than the same amount of biomass from a cyanobacterium.

Small amounts of biomass therefore already have a large impact on the ΔCt value.

**Heterotrophic cultivations**

The evolution of biomass composition over time was investigated. For this purpose, *T. sociatus* was cultivated under heterotrophic conditions with glucose for 14 days. After 2, 5, 7 and 14 days, the biomass composition was determined by qPCR and the cdw was determined (Fig. 3). After 2 days, the proportion of *T. sociatus* in the biomass is reduced to 90%. The cdw increases from 0.75 to 1.64 g L⁻¹ by 0.9 g L⁻¹ during the
first 2 days of cultivation. Using the qPCR, it can be determined that about 0.73 g L\(^{-1}\), or 80%, of the newly produced cdw consists of \textit{T. sociatus}. From this, it can be assumed that about 4/5 of the glucose used is utilized by the cyanobacteria and 1/5 by the heterotrophic bacteria. The results are in good agreement with other studies that show that cyanobacteria grow well on glucose (Markou and Georgakakis 2011; Francisco et al. 2014).

After 2 days, glucose is almost completely consumed in the cultivation media (data not shown). Subsequently, the cyanobacteria displace the heterotrophic bacteria (below 2% of cdw on day 5). \textit{Trichocoleus sociatus} has shown antibacterial activity in previous studies (Strieth et al. 2017). With this activity, it seems to be possible to control the growth of the heterotrophic bacteria below the detection limit. It is unclear why the cyanobacterium does not completely eliminate the heterotrophic bacteria. Perhaps this is not its intention at all. Many studies have shown that cyanobacteria live in symbiosis with heterotrophic bacteria. These supply the cyanobacteria with important secondary metabolites and in return receive carbohydrates or nitrogen compounds (Thompson et al. 2006; Chojnacka et al. 2019).

**Conclusion**

For the first time, a qPCR method was presented which is capable of measuring the quantitative composition of cultivations of terrestrial cyanobacteria with a carbohydrate source. By using biomass mixtures as a calibration series, the proportion of cyanobacterial biomass in cultivation can be determined. Due to the universal primers, the method can be easily transferred to new cyanobacterial cultures. However, a new standard curve must be recorded for each strain. This new method is particularly useful in the development of bioprocesses in order to assess the benefits of carbohydrate supplementation. In future cultivations, the development of the biomass composition over the process period can be examined in more detail. In combination with a sugar analysis, it is expected that further knowledge about the consumption of sugar by the different species in cultivation can be gained.

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