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Introduction

Legionella is a group of rod-shaped human pathogens found in various engineered water systems like hot water tanks,^{1,2} cooling towers^{3,4} and evaporative cooling systems.⁵ By law, *Legionella* prevalence must be monitored in these systems.⁶ There are around 60 known different species (spp.) of *Legionella*, while *Legionella pneumophila* (*L. pneumophila*) is responsible for over 90% of legionellosis diseases.⁷ *L. pneumophila* can be divided into 16 serogroups (Sg) based

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Standard addition method for rapid, cultivation-independent quantification of *Legionella pneumophila* cells by qPCR in biotrickling filters[†]

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Cultivation-independent molecular biological methods are essential to rapidly quantify pathogens like Legionella pneumophila (L. pneumophila) which is important to control aerosol-generating engineered water systems. A standard addition method was established to quantify L. pneumophila in the very complex matrix of process water and air of exhaust air purification systems in animal husbandry. Therefore, cryopreserved standards of viable L. pneumophila were spiked in air and water samples to calibrate the total bioanalytical process which includes cell lysis, DNA extraction, and gPCR. A standard addition algorithm was employed for gPCR to determine the initial concentration of L. pneumophila. In mineral water, the recovery rate of this approach (73%-134% within the concentration range of 100-5000 Legionella per mL) was in good agreement with numbers obtained from conventional genomic unit (GU) calibration with DNA standards. In air samples of biotrickling filters, in contrast, the conventional DNA standard approach resulted in a significant overestimation of up to 729%, whereas our standard addition gave a more realistic recovery of 131%. With this proof-of-principle study, we were able to show that the molecular biology-based standard addition approach is a suitable method to determine realistic concentrations of L. pneumophila in air and process water samples of biotrickling filter systems. Moreover, this guantification strategy is generally a promising method to guantify pathogens in challenging samples containing a complex microbiota and the classical GU approach used for gPCR leads to unreliable results

> on different lipopolysaccharide (LPS) structures on their membrane surface.⁸ According to the legislation, cultivation is the only accepted method for quantifying Legionella spp.⁶ However, the long time between sampling and obtaining results of 9-15 days,9 the potential for overgrowth of other microorganisms, and the presence of viable but non-culturable cells (VBNC)10 have increased the demand for culture-independent methods.¹¹ These methods use either antigens on the cell membrane of bacteria or genomic DNA to quantify rapidly bacteria cells. The serotype-specific LPS of L. pneumophila can be used as an antigen for antibody-based methods like ELISA,¹² lateral flow-immunoassays,¹³ or chemiluminescence sandwich microarray immunoassays.¹⁴ Immunomagnetic separation coupled with flow cytometry (IMS-FCM) is a technique that is able to quantify intact bacterial cells of either L. pneumophila Sg 1 or Sg 1–15, depending on the used panel of antibodies.¹⁵ The advantage of antibody-based methods is that no additional sample pretreatment is needed. Nevertheless, no information for Legionella spp. is gained with these methods.

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[†]Electronic supplementary information (ESI) available: IMS-FCM and FCM data for the biofilter samples. Fig. S1 shows TLC and ILC dot plots of a clean gas sample. Fig. S2 shows TCC and ILC dot plots of a process water sample. See DOI: https://doi.org/10.1039/d3an02207b

Molecular biological methods, such as quantitative polymerase chain reaction (qPCR) assays, are widely used and have a high sensitivity and the ability to simultaneously quantify both Legionella spp. and L. pneumophila.¹⁵ Several companies provide qPCR assay kits, including Biotecon, Bio-Rad, Promega, Minerva Biolabs, NZYTech, Creative Biogene and Thermo Scientific. The assay kits are all designed and validated according to the DIN ISO 12869:2019¹⁶ guideline for the quantification of Legionella spp. with qPCR. According to this guideline, the quantification is based on Genomic Units (GU) standards which is stated to work in all types of water, except when the accompanying flora interferes with the determination.¹⁶ Environmental samples with a very high microbial load are a challenge in many ways. First, false positive results can be obtained when non-specific amplification of non-target DNA occur.^{17,18} Additionally, when using silica filters or magnetic beads for DNA extraction, a high total DNA amount can cause overloading, decreasing the efficiency.^{19,20} Another challenge are biochemical reagents that inhibit the polymerase and decrease the qPCR efficiency.²¹⁻²⁴ All these factors, can cause unreliable results and since the cultivation approach achieves often false negative results due overgrowth, the results cannot be verified. In our study we show that agricultural exhaust air purification systems from pig fattening farms are one example where very strong matrices in air and process water occur. Due to their use of an active microbiome to degrade nitrogen and a circulating pH neutral wash water, they have a very high microbial load.

Quantification by qPCR is based on calibration with DNA standards provided by the manufacturers, expressed in genomic units. The cycle threshold (C_t) value obtained for the sample is then calculated, considering the dilution factors and sample volume used, and expressed as GU per mL. Since GU standards do not undergo sample preparation steps such as filtration, cell lysis, and DNA extraction, the quantitative results may lack trueness. Loss of DNA during extraction is inevitable. The extraction efficiency is around 20 to 90% and depends on the method used, sample type, and person who performs it.²⁵⁻²⁷ Simpler extraction methods can increase the DNA yield and assay sensitivity but also increase matrix effects.^{28,29} Even, when the extraction efficiency is known in a certain matrix such as mineral water, the transfer to other matrix is not satisfactory and the measured quantity value is not in the agreement of a true quantity value. Furthermore, microbiome-containing matrices of environmental samples are influenced by various factors such as sampling site, temperature, weather, and transportation-(time) until measurement, making the matrix effects never identical.³⁰⁻³² Additionally, often no blank samples are available since ubiquitous bacteria like Legionella spp. are always present, and removing a single bacterium without influencing other microorganisms is impossible. All these factors lead to the conclusion that a GU calibration or a single calibration in a simulated or even real matrix always entails a certain degree of inaccuracy. Also the addition of DNA^{33,34} or non-target bacteria^{35,36} into the sample to estimate the efficiency of the qPCR is a known strategy, but cannot be

directly compared with the efficiency of the target quantification. To account for all of the mentioned factors, we conclude that the inclusion of the entire sample preparation procedure can be favorable for calibration in environmental samples, treating the whole process as one combined bioanalytical method (Fig. 1). The addition of the viable target L. pneumophila to calibrate the total bioanalytical workflow is derived from a standard addition approach well known from instrumental analysis such as gas chromatography,³⁷⁻³⁹ atomic absorption spectroscopy,⁴⁰ and isotope analysis.^{41,42} Standard addition is performed by spiking defined levels of the analyte of interest. Using linear regression, analyte concentrations in the original sample can be back-calculated. This requires a linear response behavior of the analytical method and the exact dosability of the analyte. For the quantification of Legionella spp. in environmental samples with qPCR, defined living L. pneumophila cryopreserved standards are added directly into it. The semilogarithmic calibration of qPCR shows linearity between Ct value and bacterial concentration only on a logarithmic scale and therefore adjustments need to be made to obtain quantitative results through the standard addition approach used. Therefore, a standard addition algorithm designed for immunoassays is applied to overcome this limitation.43

In the present work, for the first time to our knowledge, this standard addition approach for qPCR was used to overcome matrix effects in complex air and water samples of a biotrickling filter allowing a more reliable quantification than possible with commonly used GU standards. Since cultivation lead to false negative results, the IMS-FCM was used as complementary cultivation-independent method to confirm the results. While calibration with GU standards is faster and needs less laboratory work, the standard addition approach can be utilized to quantify pathogens in complex water or other environmental matrices, such as those with a high microbial load if quantification with GU standards leads to unreliable results. It also enables the use of simpler DNA extraction methods, thereby enhancing the sensitivity, accuracy and reliability of qPCR.



Fig. 1 Workflow of a (GU-)calibration and measurement of a biotrickling filter sample. With a commonly performed GU calibration, GU standards are directly measured without pretreatment. When analyzing real samples, concentration and DNA extraction are important factors not included by this calibration strategy. The standard addition method using cryopreserved *L. pneumophila* includes all these factors, resulting in a more accurate quantification.

Experimental

Bacteria cryopreserved standards

Cryopreserved standards of viable L. pneumophila Sg 1 subtype Bellingham were prepared according to standardized method developed by rqmicro¹⁵ to ensure a stable and well-defined cell count in the spiking experiments. Two quality controls via flow cytometry, one after growing in media and the second after conditioning in mineral water, were performed. In each quality control, the Total Cell Count (TCC), the Total Legionella Count (TLC), and the Intact Legionella Count (ILC) were measured via flow cytometry on a rqmicro.COUNT device (rqmicro AG, Schlieren, Switzerland). The cryopreserved standards were only used if they were of sufficient purity (TLC/TCC > 90%) and the viability was sufficiently high (ILC/TLC > 60%). The cryo buffer consisted of 2× PBS (5.4 mM KCl, 20 mM Na₂HPO₄, 3.3 mM KH₂PO₄, 0.27 M NaCl, pH 7.0), 10 g L⁻¹ BSA (Sigma Aldrich, St. Louis, USA) and 120 g L⁻¹ dextran 40 (Sigma Aldrich, St. Louis, USA). The standards were stored at -80 °C and directly used after thawing at room temperature. Each day, a new cryopreserved standard was used from the same batch. The long-term storage and thawing stability of the cryopreserved standards was shown elsewhere by Streich et al.44 The initial concentration after 8 months was slightly reduced by $7.7 \pm 1.2\%$. The TLC of the used cryopreserved standard was 6.7×10^7 bacteria per mL and the ILC was 3.4×10^7 bacteria per ml.

Sample preparation

For calibration in mineral water, a dilution series of cryopreserved bacteria standards were performed in Evian water (1.5 L-bottles, Danone, Paris, France). 1 mL of sample was spiked with 10 μ L of diluted standards. For extraction of environmental samples, 1 mL of the sample was directly used and spiked with 10 μ L of standard solution. Samples and samples spiked with cryopreserved standards were centrifuged at 8000*g* for 5 min, followed by DNA extraction of a formed pellet using foodproof® StarPrep Two Kit Procedure B (Biotecon Diagnostics, Potsdam, Germany). Extraction works by a combination of mechanical lysis and heat. After lysis, samples were centrifuged, and supernatants (100 μ L) were used for analysis. The DNA extracts were stored at 4 °C and used the next day.

Quantitative polymerase amplification reaction

The qPCR Kit microproof® *Legionella* Quantification LyoKit (Biotecon, Potsdam, Germany) was used for all experiments. Measurements were performed on the qTOWER³G (Analytik Jena, Jena, Germany) instrumentation. 25 μ L of the DNA extract was added to the qPCR reaction mix according to the manual delivered by Biotecon. A negative control (PCR-H₂O) was added for every run. Thermal cycling conditions were chosen according to the given manual. First, a pre-incubation was performed (1 cycle) at 37 °C for 4 minutes and 95 °C for 5 minutes. Followed by 50 amplification cycles containing 3 steps. Step 1: 95 °C for 5 s, step 2: 60 °C for 60 s and step 3: 72 °C for 60 s. The fluorescence measurement was performed

in step 2. Four different fluorescence channels were measured, Cy5 (internal control), HEX (*Legionella* spp.), FAM (*L. pneumophila* Sg 1–15), and ROX (*L. pneumophila* Sg 1). If one identical DNA extract was measured in replicates, it is indicated as a technical replicate. If, for each measurement, a different replicate of a DNA extract was measured, it is stated as a biological replicate. A negative control (PCR-grade H_20) was always added to a qPCR run.

Data evaluation

The evaluation sheet, based on the ISO 12869:2019, included in the qPCR Kit microproof® Legionella Quantification LyoKit (Biotecon Diagnostics, Potsdam, Germany) was used for GU calibration. According to this standard a limit of quantification of 130 cells per mL was achieved for Legionella spp., L. pneumophila and L. pneumophila Sg 1. A limit of detection of 15 cells per mL for Legionella spp. and 25 cells per mL for L. pneumophila and L. pneumophila Sg 1 was obtained. The published Excel-Solver sheet from Pang et al.⁴³ developed for standard addition approaches for immunoassays was used to quantify L. pneumophila by qPCR. For adaption to qPCR, instead of signal values, medium cycle threshold (C_t) values of each spiking concentration were inserted in the sheet and an estimated concentration of the unspiked concentration was inserted as a starting point for the algorithmic-based calculation. Afterwards, the Excel-Solver algorithm was able to calculate the exact value.

Measurements with IMS-FCM

L. pneumophila were counted by combining immunomagnetic separation with flow cytometry (IMS-FCM). All measurements were performed on the device rqmicro.COUNT (rqmicro, Schlieren, Switzerland). Intact Legionella counts (ILC) were measured by the Legionella kit L.p. SG1-15 DETECT Kit (31102) (rqmicro, Schlieren, Switzerland). The assay was performed as described by the manufacturer with the following changes. Instead of the enrichment of samples with filtration of 100 mL sample, 3 mL of the clean gas samples were filtrated through a prefilter (5 µm diameter pore size, 39002, rqmicro, Schlieren, Switzerland) and then caught on a sterile filter (0.22 µm diameter pore size, 39001, rqmicro, Schlieren, Switzerland). Only a prefiltration (5 µm diameter pore size) was performed for the process water samples. Without any further enrichment, the analysis was proceeded as described in the kit. To measure the total microbial load, the total cell count (TCC)-Kit (30010) (rqmicro, Schlieren, Switzerland) was used and performed according to the kit manual.

Sampling at the biotrickling filter

At a fattening pig house with 2000 animals, the clean gas, and process water of an installed biotrickling filter were sampled. The cleaned gas was sampled at 3 different timepoints directly at the outlet of the unit. To protect the bioaerosol collector from wind, a tube (1 m deep, 1 m in diameter) was installed for the sampling. A Coriolis® μ device (Bertin Technologies SAS, Montigny-le-Bretonneux, France) was used as collection

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system. The collection vessels were filled with 15 ml sterile 0.9% NaCl solution. The sampling duration consisted of 3 rounds, each lasting 10 minutes, with a flow rate of 300 L min⁻¹. Three individual samples were combined into one composite sample, corresponding to a total air volume of 9 m³ per sample. The collected total volume was measured and divided by the initial volume of 45 mL to get the evaporation factor of liquid during measurement. This factor was used when calculating the raw data from the measurements to the unit cells per m³. Two process water samples at 2 different timepoints were taken from the middle of the basin using a 500 mL PE bottle. All samples were transported in sealed bottles, kept in the dark, without active cooling, to the laboratory of the Thünen Institute. They were then directly sent via express delivery (arriving around 9 a.m. the following day) to our institute at TUM, where the measurements were conducted.

Results and discussion

Spiking strategy for standard addition calibration in qPCR

For standard addition, samples with unknown concentrations of *Legionella* spp. or *L. pneumophila* must be spiked with a defined concentration of intact bacteria cells. This necessitates



in the regression plot is unknown, too. The used algorithm by Pang et al.43 for the quantitative analysis utilizes a linear behavior on a logarithmic scale within the linear working range of the qPCR assay when U is close to the true value. U is changed by a numeric algorithm until the highest linearity is found. This means the residual variation from the calculated values of the regression line and the observed values is the smallest, R^2 is the closest to 1. At least four different spiking concentrations in the sample are needed as data points for standard addition.⁴³ To minimize sample volume, time and costs, the amount of data points should be limited as much as possible while still generating quantitative results. For that, the spiking levels must be chosen so that the total concentration (sample concentration + spiking concentration) is within the linear range of the method to generate a proper concentration dependency. Both too high (Fig. 3A) and too low (Fig. 3B) spike concentrations need to be avoided, since either the signal of the actual concentration in the sample is dwarfed by the standard addition, or no concentration dependency is visible due to the high initial concentration in the non-spiked sample (Fig. 3B). This is achieved by properly estimating the





Fig. 2 Workflow of the standard addition approach. 1 mL of the sample is spiked with different concentrations of a *Legionella pneumophila* Sg 1 subtype Bellingham bacteria standard to adjust the spiking concentrations. Afterwards, unspiked sample plus the spiked dilution series are extracted and measured identically according to the kit manual. For the quantification, an algorithm⁴³ is used that is alternating the unknown concentration U until the residual sum of squares is minimal, resulting in the maximum linearity of the data points.

Fig. 3 Schematic overview for finding the proper spiking levels. The *y*-axis represents the C_t value, the *x*-axis the sample (U) and the spiking levels (U + spiking concentration) on a logarithmic scale. (A) Pattern for a low sample concentration and overly high spiking levels. (B) Spiking low concentrations in a sample with a high initial analyte concentration results in a saturated pattern. (C) Use of proper spiking levels.

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sample concentration by preliminary measurements and the knowledge of the linear working range of the assay (see Fig. 3C).

Proof-of-principle study

The qPCR with standard addition was initially tested using a sterile Evian mineral water matrix. Drinking water does not show significant influences on qPCR applications. It is a welldefined water matrix ideal for quantitative calibration studies of pathogens. In this experiment, Evian water was spiked with L. pneumophila Sg 1 subtype Bellingham at varying concentrations. Biological triplicates were conducted, where each was measured as a triplicate using qPCR. The spiked concentrations were calculated by standard addition and in comparison, a GU calibration was conducted according to the manual of the qPCR kit. For GU calibration, a calibration curve was constructed using four different levels of GUs, ranging from 25 to 25 000 GU per reaction. The C_t values of the measurements were fitted for GU calibration using an evaluation Excel sheet based on the ISO 12869:2019 provided by the company Biotecon. 10²-10⁶ cells per mL were spiked into Evian water to

perform the standard addition approach. The algorithm described in the Excel-Solver program by Pang et al.43 was applied to determine the unknown concentration of L. pneumophila. The calibration results are shown in Fig. 4. The summary of the data is shown in Table 1. For the standard addition, a linear behavior within the logarithmic scale is needed. In all 3 fluorescence channels, a high linearity of the data points was observed. For the Legionella spp. channel, a R^2 of 0.998 was achieved. For the L. pneumophila and L. pneumophila Sg 1 channel a R^2 of 0.999 was determined. Blank measurements (sterile mineral water with no added pneumophila Sg 1) showed no C_t values for the L. L. pneumophila primer set and very high C_t values for Legionella spp. and L. pneumophila Sg 1 primer set, indicating a minor carryover of DNA during DNA extraction or qPCR runs in this experiment. On average, both quantification principles slightly overestimated the true value with a mean of 139% (GU approach) and 135% (standard addition approach).

Based on the data, this suggests that the standard addition approach yields comparable results in the drinking water matrix, where the GU calibration is considered as the gold



Fig. 4 Calibration with different concentrations of *L. pneumophila* Sg 1 subtype Bellingham in Evian water. Three primer sets are included, detecting *Legionella* spp., *L. pneumophila* Sg 1–15 and *L. pneumophila* Sg 1 (number of measurements m = 8, number of biological replicates n = 3). (A) qPCR raw data. C_t values corresponding to the spiked *L. pneumophila* concentrations. This represents the known "true value" of bacteria in the Evian water. The known values are treated as unknown and were calculated with the standard addition approach (B) and the kit GU-calibration (C). (B) Quantification of the data points of (A) using the standard addition algorithm and the Excel-Solver program from Pang *et al.*⁴³ As described previously, the algorithm is used to calculate U, resulting in the highest linearity of the plot, which gives the value of U. (C) Calibration of the data points of GU standard as described in the kit manual (m = 4, n = 2). The data points of A are fitted in the calibration to get the GU number of the data points and afterward transferred in cells per mL.

Spiked value in <i>L. pneumophila</i> Sg 1 cells per mL	Ca	lculated value with standard addition <i>Legionella</i> spp./mL (recovery)	Calculated value with GU-calibration in <i>Legionella</i> spp. cells per mL (recovery)
<i>Legionella</i> spp. 0 100 500 1000 5000	18 10 56 12 36	6 (105%) 7 (113%) 22 (122%) 59 (73%)	<15 <130 630 (126%) 1500 (150%) 7100 (142%)
Spiked value in <i>L. pneumophila</i> Sg 1 cells per mL	Calculated value with standard addition in <i>L. pneumophila</i> Sg 1–15 cells per mL (recovery)		Calculated value with GU-calibration in <i>L. pneumophila</i> Sg 1–15 cells per mL (recovery)
<i>L. pneumophila</i> Sg 1–15 0 100 500 1000 5000	128 (128%) 737 (147%) 1343 (1349) 6603 (1329))) 6)	<130 620 (124%) 1200 (120%) 5700 (114%)
Spiked value in <i>L. pneumophila</i> Sg 1 cells per mL		Calculated value with standard addition in <i>L. pneumophila</i> Sg 1 cells per mL (recovery	Calculated value with GU-calibration) in <i>L. pneumophila</i> Sg 1 cells per mL (recovery)
<i>L. pneumophila</i> Sg 1 0 100 500 1000 5000		11 138 (138%) 606 (121%) 1198 (120%) 3913 (78%)	<15 <130 620 (124%) 1200 (120%) 5700 (114%)

Table 1 Summary of the concentration for the Evian calibration in Fig. 4A calculated with the GU-calibration (Fig. 4C) or with the standard addition approach (Fig. 4B). The recovery shows the over- or underestimation compared to the true spiked value

standard. The approach demonstrates a recovery rate of 73%–134% within a concentration range of 100–5000 *Legionella* per mL (Table 1). All recovery values are within the acceptable range of $-0.6 \log_{10}$ unit and $+0.3 \log_{10}$ unit defined in the ISO12869:2019,¹⁶ calculated by the decimal logarithm of the measured value divided by the true value. Consequently, it can be concluded that the standard addition method has been effectively adapted from the immunoassay approach proposed by Pang *et al.*⁴³ to a qPCR method.

Quantification in complex matrices: a proof-of-principle study on a biotrickling filter

As an example for a challenging real matrix, environmental samples from agricultural biotrickling filters were used. Clean gas samples were collected with a cyclone sampler and the measurement with flow cytometry resulted in a high microbial load of 3.1×10^7 total cells per m³ (Fig. S1A[†]). For the analysis of Legionella using the qPCR standard addition method, spiking levels of 10³-10⁵ cells per mL *L. pneumophila* Sg 1 were chosen since a concentration in this range was expected in the sample. Fig. 5 presents the qPCR results of one gas sample (sample 3). 3 samples were analyzed and the results are summarized in Table 2. For all samples, L. species concentrations determined by the GU calibration are 292-380% higher than the results of the standard addition approach. Since no blank samples were available and the true concentration of Legionella spp. in the sample was not known, an orthogonal reference method was needed. Therefore, as a comparison to



Fig. 5 qPCR results for a clean gas sample (sample 3) from an agricultural biotrickling filter. The unspiked clean gas sample (U) is marked in green and the sample with defined spiking levels (U + concentration) marked in orange (number of measurements m = 4, number of technical replicates n = 3). The negative control (qPCR-H₂O) showed no C_t values.

the qPCR results, the concentration of *L. pneumophila* was also determined by IMS-FCM (Fig. S1B[†]). Combining immunomagnetic separation with subsequent flow cytometry allows counting *L. pneumophila* labeled by a panel of antibodies specific for Sg 1–15. This system generates reliable results on drinking

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water⁴⁵ and was shown to work also well on gas samples, sampled with the Coreolis μ^{15} as well as for process water of cooling towers.44 Prefiltration was conducted to remove larger particles and prevent clogging of the flow paths in the applied cartridge. Regarding our samples, the loss during prefiltration is expected to be minimal in clean gas samples and minor for process water samples, making it as a suitable reference method. The standard addition methods overestimate the FCM-IMS value (823 L. pneumophila cells per m³) for L. pneumophila Sg 1 by 131%. In contrast, the conventional GU calibration resulted in an overestimation of 729%. The standard addition approach is closer to the results of the IMS-FCM. The GU calibration has a relative overestimation when compared to both the standard addition method (557%) and IMS-FCM (729%). One plausible explanation is that the fluorescence channel, which measures the internal standard, is used as a correction factor for other fluorescence channels that detect the amplicons of different primer sets. If inhibitory effects exist in the internal standard channel, the measured value is corrected, leading to potential overestimation if the inhibition of the internal standard primer set differs from the analyte primer set. Another reason can be an increase in DNA extraction efficiency. While efficiency generally decreases with higher microbial load,⁴⁶ a larger and stronger bacterial pellet forms during the initial centrifugation step when there is a higher amount of bacteria. The presence of substantial amounts of other cells and DNA may lead to a higher recovery of Legionella spp. DNA in biotrickling filter samples compared to water matrices, where the system is calibrated by the provider. This difference can result in overestimation. Furthermore, the GU-calibration initially provides the value per reaction, which then needs to be calculated based on the volume of DNA extract used in the qPCR reaction relative to the total DNA extract volume and the sample volume before extraction. All these mathematical corrections increase the likelihood of error propagation. In contrast, our alternative approach - the described qPCR with standard addition - takes into account the whole sample preparation process for calibration. No presumed correction factor is required for DNA extraction and importantly, the unit can be expressed in bacteria per mL, making it more comparable to other methods, such as cultivation, immunoassays and flow cytometric approaches.

For a more in-depth analysis of the prevalence of *Legionella* spp. in biotrickling filters, process water samples were analyzed, posing an even more significant challenge. The process water, although initially fresh water, continuously accumulates

deposited dust, ammonia, and organic substances during the operation of the system. This results in a cloudy (139 ± 9 NTU (Nephelometric Turbidity Unit)) yellow-brownish sample with a very high microbial load (TCC = 2.2×10^7 cells per mL, Fig. S2†). Due to the anticipated high concentrations of *Legionella* spp. in process water, samples were diluted at a ratio of 1:100 before extraction. Moreover, higher concentrations were used for spiking levels to prevent saturation effects. The qPCR results of one process water sample (sample 2) are presented in Fig. 6.

The concentration values of two different samples are summarized in Table 3. Once again, the qPCR results indicate higher quantities of *L. pneumophila* than to IMS-FCM (20 740 *L. pneumophila* cells per mL). When employing the standard addition approach, a smaller overestimation is observed in contrast to the GU calibration method. It is worth noting that alternative quantification method IMS-FCM may slightly underestimate the presence of *L. pneumophila* in process water due to prefiltration using a 5 μ m pore-sized filter. This filtration step is implemented to prevent cartridge blockage and has been observed in previous experiments to retain some *Legionella* spp. that are bound to bigger particles or due to clump formation of the bacteria, resulting in a slight reduction



Fig. 6 qPCR results for a process water sample (sample 2) from an agricultural biotrickling filter. The unspiked process water sample is (U) marked in green and the sample with defined spiking levels (U + concentration) is marked in orange (m = 5, n = 2). The negative control (qPCR-H₂O) showed no C_t values.

Table 2 Summary of the environmental clean gas sample concentrations determined with qPCR (standard addition method and GU calibration)

Biotrickling filter clean gas sample	Calculated concentration according to standard addition	Calculated concentration according to GU-calibration
Sample 1 c(<i>Legionella</i> spp. cells per m ³)	2010	5875
Sample 2 c(<i>Legionella</i> spp. cells per m^3)	1305	4000
Sample 3 c(<i>Legionella</i> spp. cells per m^3)	9710	36 850
Sample 3 c(<i>L. pneumophila</i> cells per m ³)	1150	6410

 Table 3
 Summary of the process water concentrations of an agricultural biotrickling filter determined with qPCR (standard addition method and GU-calibration)

Biotrickling filter process water	Calculated concentration according to standard addition	Calculated concentration according to GU-calibration
Sample 1 c(<i>Legionella</i> spp. cells per mL)	160 000	1 900 000
Sample 2 c(<i>Legionella</i> spp. cells per mL)	257 000	1 000 000
Sample 2 c(<i>L. pneumophila</i> cells per mL)	57 900	220 000

in concentration within the filtered sample. Furthermore, also clump formation due a high bacteria concentration inside the cartridge during measurement can result in an underestimation. One bacteria aggregate containing more than one *L. pneumophila* can result in only one count. Nevertheless, the tendency that was shown in the gas samples, where the IMS-FCM is in better agreement with the SA-approach is also observed in process water. The GU calibration results in a factor of 3.9 to 11.9 higher *Legionella* spp. concentration than the SA approach. For *L. pneumophila* concentration, the GU calibration is 3.8 times higher than the SA approach.

In summary, the standard addition approach proved to generate more reliable results in the biotrickling filter process water than the GU calibration, thus enhancing the analysis of these complex samples.

Conclusion

In this study a novel standard addition approach for qPCR was implemented for the first time, encompassing the entire sample preparation process. Cryopreserved L. pneumophila standards of known concentrations were used for spiking, enabling the generation of quantitative results in mineral water comparable to those obtained with a GU calibration. In contrast, when applied to biotrickling filter samples with high microbial loads, the conventional GU calibration yielded unreliable results. Here, the standard addition approach proved to be more reliable in generating accurate results. It could be demonstrated, that biotrickling filters are a potential source of Legionella spp. and the developed method is a valuable tool for further investigation of these systems. Furthermore, the qPCR approach enables the quantification of both, Legionella spp. and L. pneumophila, allowing for a more comprehensive analysis compared to immunoassays. It is important to acknowledge that this method entails higher costs and increased workload, making it less suitable for routine measurements with a large number of samples to analyze on a daily basis. However, it exhibits significant potential for analyzing complex and poorly characterized samples where the impact of the matrix on traditional calibration methods is unknown. Particularly in cases where the sample composition is unstable and subject to change, the standard addition approach can outperform classical calibration methods, providing more dependable results. This reliable qPCR method becomes particularly crucial when cultivating the target bacteria, as observed in biotrickling filters, is not feasible. Quantitative results are then unattain-

able through culture-based techniques and subsequent analyses such as MALDI-TOF are not possible. Other methods like digital PCR (dPCR) using absolute quantification is due to the separation in small reaction chambers less affected to matrix effects. This results in a more robust assay in comparison to classical qPCR approaches.⁴⁷ Nevertheless, the availability of qPCR devices in most laboratories and accessibility of different kits on the market is still bigger than for dPCR. Additionally, the DNA extraction in dPCR has the same challenges as qPCR. The efficiency can highly depend on the matrix. While the quantification is different between qPCR and dPCR, our work can be adapted and be applied in the future for dPCR approaches to overcome matrix influences during DNA extraction by spiking living bacteria standards. For qPCR the SA addition approach can also enable the use of a simpler DNA extraction method resulting in higher yields and better sensitivity of the assay since it is less affected by matrix effects. Furthermore, if the cryopreserved standards are added directly after sampling, not only DNA extraction but also the transportation can be covered in the calibration. The strength of the standard addition approach lies in its versatility, allowing its application to a wide range of environmental analytical inquiries where the detection of microorganisms and consideration of matrix effects are essential factors.

Author contributions

The manuscript was written through contributions of all authors. Gerhard Schwaiger planed and performed the experiments of the quantification with the help of Marco Matt. Gerhard Schwaiger evaluated the data and wrote the manuscript with the input from the co-authors. Philipp Streich characterized the cryopreserved bacteria standards. Sarah Bromann and Marcus Clauß were taking the samples. M. Elsner and M. Seidel supervised the project and were responsible for funding acquisition and resources. All authors have given approval to the final version of the manuscript.

Notes

The AI tool "Grammarly" was used to improve the English written language.

Conflicts of interest

There are no conflicts to declare.

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