RNA-based qPCR as a tool to quantify and to characterize dual-species biofilms

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While considerable research has focused on studying individual species, we now face the challenge of determining how interspecies interactions alter bacterial behaviours and pathogenesis. *Pseudomonas aeruginosa* and *Staphylococcus aureus* are often found to co-infect cystic-fibrosis patients. Curiously, their interaction is reported as competitive under laboratory conditions. Selecting appropriate methodologies is therefore critical to analyse multi-species communities. Herein, we demonstrated the major biases associated with qPCR quantification of bacterial populations and optimized a RNA-based qPCR able not only to quantify but also to characterize microbial interactions within dual-species biofilms composed by *P. aeruginosa* and *S. aureus*, as assessed by gene expression quantification. qPCR quantification was compared with flow-cytometry and culture-based quantification. Discrepancies between culture independent and culture dependent methods could be the result of the presence of viable but not-cultivable bacteria within the biofilm. Fluorescence microscopy confirmed this. A higher sensitivity to detect viable cells further highlights the potentialities of qPCR approach to quantify biofilm communities. By using bacterial RNA and an exogenous mRNA control, it was also possible to characterize bacterial transcriptomic profile, being this a major advantage of this method.

The recognition that biofilm communities, which typically comprise multiple species, with a spatiotemporal heterogeneous chemical, physiological and genetic composition poses a serious healthcare concern regarding the synergies that arise from the co-infecting species. Hence, selecting appropriate methodologies that provides reliable quantitative measures of individual populations within the polymicrobial biofilms is crucial to predict the behaviour and dynamic of these communities.

Culture-based methods have long been used to detect microbial pathogens; however, since these methodologies require the growth of the cultures, it takes a long period of time to obtain results. Furthermore, culturing is not always an option since some bacterial species are difficult to growth in the laboratory and many species develop a viable but non-cultivable state. Other limitations are the presence of cell-aggregates that strongly influence the outcome of bacterial culture quantification. Alternatively, culture-independent assessments of microbial communities, particularly through quantitative real-time PCR (qPCR) or sequencing, have allowed a rapid screening and/or quantification of the specific microorganisms within the consortia. The most frequently used qPCR method is based in the amplification of gDNA molecules. Unfortunately, DNA-based methodologies cannot unequivocally differentiate between live (including viable but non-cultivable cells (VBNC)) and dead cells and, therefore, are likely to overestimate the microbial communities. Other specialized methods, such as propidium-monoazide (PMA)-qPCR, are becoming more popular as the function and organization of the entire community depends on which members are alive or dead. Nevertheless, PMA-based approaches have reportedly important limitations, including changes in cell membrane permeability (influencing the uptake of PMA dye) and false-positive signals.

A less explored alternative is the utilization of RNA-based qPCR approaches for the detection of live members of a polymicrobial consortia. This principle was recent demonstrated by several independent studies which demonstrated that RNA-based quantification, was superior to DNA-based methods, in detecting live bacterial cells, providing a more relevant insight into the microbial community composition. Of utmost importance, RNA-based qPCR approach has the significant potential of performing not only a quantification analysis, but also addressing specific microbial interactions, as detected by the expression of key virulence genes of interest.
In fact, infections caused by the presence of multispecies are often more virulent or recalcitrant to treatment than those caused by each independent species. However, relatively little is known about the interspecies interactions that modulate the community dynamics in terms of composition and virulence; ultimately contributing to increased infection severity and chronicity.

P. aeruginosa and S. aureus are versatile bacterial pathogens and common etiological agents in several polymicrobial infections, including wounds, otitis media and oral infections, and CF lung disease. Of particular interest is the ecological interactions between P. aeruginosa and S. aureus, since microbial communities containing both of these pathogens can display enhanced virulence. Despite routine administration of antibiotics, these infections are often highly resilient and tolerant to treatment. This might explain why aggressive antibiotics treatment, patients with CF will eventually succumb to the chronic persistent infections. Robust measures of these community changes (composition and/or abundance) are therefore urgent, since misinterpretation of these communities can impair antimicrobial treatment.

Herein, the objective of this study was to evaluate an optimal RNA-based qPCR method to investigate the community dynamics and microbial interactions between P. aeruginosa and S. aureus. Thus, microbial composition of 24- and 48-h-old dual-species biofilms was assessed by RNA-based qPCR as well as changes in the transcriptomic profile of key virulence-related genes. The qPCR bacterial quantification was first fully optimized to polymicrobial samples and further compared with conventional culture and flow-cytometry techniques.

Results and Discussion

Quantification of dual-species biofilm communities: plate count versus flow cytometry. Dual-species biofilms were first characterized over the course of 24- and 48-h by enumerating P. aeruginosa and S. aureus through CFU assessment on selective media (PIA and MSA, respectively) and also by using flow-cytometry (Fig. 1). At both time points, P. aeruginosa was the prevalent species detected, being the differences between P. aeruginosa and S. aureus more relevant at 48-h of growth. Interestingly, these differences were more dramatic (~6 log_{10} CFU/mL) when considering culture quantification only, but significantly lower (~2 log_{10} cells/mL) when quantified by flow-cytometry. These results highlight that culture methods are not reliable to analyse biofilm populations in dual-species biofilms by plate count versus flow cytometry.

In order to analyze and compare the biofilm structures, 24- and 48-h dual-species biofilms were directly examined using scanning electron microscopy (Supplementary Figure S1). The dual-species consortium showed, at 24-h, a non-contiguous layer of cells, that evolved, at 48-h, into mature biofilm producing a thick layer of co-aggregated cells surrounded by extracellular matrix. Both P. aeruginosa (road-shape) and S. aureus (coccos-shape) were distinguishable at 24-h but at 48-h, the dense structure of the biofilm did now allow to distinguish the bacterial species.

The impact of RNA extraction variability in the utilization of 16S rRNA as a direct quantification tool. Molecular-based methodologies have greatly increased our understanding regarding ecological interactions occurring among members of polymicrobial communities, when compared to culture-based methods. However, those techniques are not void of limitations and, especially when using RNA-based approaches, several controls and normalizations need to be considered.

Figure 1. Quantification of P. aeruginosa and S. aureus populations in dual-species biofilms by plate count (CFU) and flow-cytometry (FC), following 24- and 48-h of growth. Biofilm counts are expressed as means ± SD of CFU per mL or cells per mL, respectively. For each condition three independent experiments were performed. Statistical significance was determined by performing an ANOVA followed by a Tukey’s multiple comparison test to compare: (**) significantly different P. aeruginosa counts versus S. aureus counts for each method; ***P < 0.001; ****P < 0.0001. (,) significantly different P. aeruginosa counts between methods and significantly different S. aureus counts between methods; γγγγP < 0.0001. Abbreviations: PA = P. aeruginosa, SA = S. aureus.
A key aspect in RNA-based qPCR quantification is the need to normalize RNA extraction efficiencies, since the quality and quantity of extracted RNA can affect downstream applications. While often underreported, RNA extraction efficiency varies. To demonstrate this, we first assessed the variability of RNA extraction processes, followed an optimized protocol previously described. Briefly, we split planktonic suspensions of either high or low cell concentrations of *P. aeruginosa*, in six aliquots from the same biological sample, to perform total RNA extraction. As observed in Fig. 2, a high variability of total RNA was obtained, especially when using high density populations. Total RNA concentrations ranged from 26 to 220 ng/μL in the 10⁹ CFU/mL aliquots and 7 to 77 ng/μL in the 10⁵ CFU/mL aliquots. Noteworthy, aliquots of the same biological sample yielded distinct RNA concentrations, with a higher inter-assay variability in high biomass samples (coefficient of variation of 0.66) compared to the low biomass (0.23) (Supplementary Table S1).

Next, we quantified 16S rRNA for each aliquot. Not surprising, the cycle threshold (Ct) value detection was highly variable, in a range of 3 to 5 cycles (Fig. 3). These results confirmed that quantifying 16S rRNA-target, by itself, is not a good strategy to quantify bacterial communities.

**Impact of utilizing an external ref mRNA in bacterial quantification using 16 sRNA.** When quantifying gene expression by qPCR, a normalization strategy is required and often it includes the quantification of a reference gene, also known as housekeeping gene. Due to the variability of the experimental method the same principle needs to be applied when quantifying bacterial populations by qPCR. The most common alternative is to normalize the levels of the target mRNA to that of an exogenous mRNA spike-in transcript. The exogenous mRNA control is characterized by: (i) not being originally found in the sample, (ii) its addition should be previous to the RNA extraction procedure and (iii) its initial concentration is known. After laboratorial
The effects of the *ref* mRNA normalization on the 16S rRNA expression levels. Relative gene expression of the 16S rRNA normalized to the *ref* mRNA, using the delta Ct method, in 10⁹ CFU/mL and 10⁵ CFU/mL samples adjusted to the initial copies of *ref* mRNA used. For each bacterial concentration five technical replicates were spiked with standard concentrations of *ref* mRNA prior to RNA extraction and subsequent reverse transcription. For each condition two independent biological samples were used. Statistical significance within each samples’ set was determined by performing one-way ANOVA analyses followed by a Tukey’s multiple comparison test (** **P < 0.0001).

In this study, commercial luciferase mRNA (*ref* mRNA) was preferred as the exogenous reference gene since the specific sequence is not found in any of the bacterial species tested, as assessed via NCBI GenBank database (www.ncbi.nlm.nih.gov). To quantify the *ref* mRNA, a standard curve was constructed, by plotting the Ct values against the known initial *ref* mRNA copy number (Supplementary Figure S2A). The standard curve generated covers a linear range along the selected five orders of magnitude with a correlation coefficient (R²) of 0.977, suggesting that all of the tested concentrations can be used in the subsequent analysis. Nevertheless, considering that the initial concentration of *ref* mRNA spiked into the samples may interfere with the mRNA quantification due to competition for qPCR reagents⁴¹, five aliquots derived from the same *P. aeruginosa* biological culture were amended with 1 × 10⁶ to 1 × 10¹ⁱ copies of *ref* mRNA, immediately after cell lysis (Supplementary Figure S2B). The significance of correlation between the total copies of *ref* mRNA added and total copies of *ref* mRNA extracted from each sample was calculated based on Pearson’s coefficient (r = 0.99, P < 0.05 for 10⁹ CFU/mL samples and r = 0.90, P < 0.05 for 10⁵ CFU/mL samples). Next, in order to test the sensitivity and accuracy of the qPCR assay, the 16S rRNA gene was normalized to the exogenous control. This was done either in the 10⁹ and 10⁵ CFU/mL samples. Figure 4 demonstrates that in almost all conditions tested, no significant variations were found in the relative gene expression determined. The exception was in the low biomass sample amended with 1 × 10⁵ copies of *ref* mRNA. This suggests that at low amounts of genomic material, the efficiency drops below the linear range of the method⁴².

For further analysis, we selected the *ref* mRNA concentration of 1 × 10⁶ copies of *ref* mRNA/μL, since under these conditions the best mRNA recovery was found, for either the low or high cell concentrations (Supplementary Table S2). Interestingly, the low mRNA recovery estimated confirms that somewhat random physical losses during RNA isolation will impair some quantification approaches⁴⁴,⁴⁷. Overall, these results support the importance of a *ref* mRNA normalization for proper quantification of bacterial populations using the RNA-based method.

Validation of the *ref* mRNA normalization in controlled dual-species communities. To assess the robustness of this approach, the relative expression of the 16S rRNA gene was analysed in three well defined dual-species planktonic samples, (*P. aeruginosa/S. aureus* ratio equal to 10, 1 and 0.1). The C<sub>r</sub> of each 16S rRNA target was normalized to the *ref* mRNA in order to determine the experimental ratios, calculated by applying the equation described in the section Quantitative Real Time-PCR of the Experimental procedures (Supplementary Table S3). As shown in Fig. 5, the experimental ratios obtained by applying the *ref* mRNA strategy were consistent with the theoretical ratios for all *P. aeruginosa* and *S. aureus* populations. Statistical analysis showed no significant differences between the theoretical and the experimental PA/SA ratios (P > 0.05).

Quantification of a dual-species biofilm communities by RNA-based qPCR. Following the experimental validation of the *ref* mRNA normalization strategy in the well-defined planktonic cultures, 24- and 48-h-old dual-species biofilms of *P. aeruginosa* and *S. aureus* were formed and quantified by qPCR (Supplementary Table S4). The same biofilms were also characterized by culture dependent (plate count) and culture independent (flow cytometry) methods (Supplementary Table S5). As shown in Fig. 6, the ratios of *P. aeruginosa/S. aureus* varied significantly accordingly with the method used, especially in the 48-h-old biofilm (P < 0.0001). The differences between culture-dependent and culture-independent methods were more significant than between both culture-independent methods. Interestingly, both qPCR and flow-cytometry methods have ascertained *P. aeruginosa* as the predominant species in the biofilm, but a significant population of *S. aureus* was also detected by these two methods in the 48-h-old consortium, in striking contrast with the CFU quantification. It is conceivable that after 48-h of co-culture with *P. aeruginosa*, a remarkable portion of *S. aureus* cells...
acquire a VBNC state. Indeed, as biofilms mature, often portions of bacteria within the community enter a VBNC state, in response to stressful conditions, such as starvation or antimicrobial treatments, suggesting that this is an adaptive strategy for long-term survival under unfavourable conditions. Although a series of physiological changes occur during the transition from cultivable to VBNC state, VBNC cells were found to maintain high levels of rRNA, similar to cultivable cells, being qPCR RNA-based assays commonly used to evaluate gene expression profile in cultures containing VBNC cells. However, an important technical limitation of these studies is that transcription is evaluated as the global gene expression within the biofilm, not able to differentiate the physiological state of sub-populations. To determine the differences within such populations, the new generation of single-cells transcriptomic analysis should be used, but as far as we are aware, this has not been attempted in dual-species biofilms.

We also performed a Live/Dead (L/D) viability assay on disrupted 48-h-old dual-species consortia, to inspect the overall biofilm-cell viability. L/D staining showed that cocci-like bacterial cells were more abundant in the green than the red filter. Those results reinforce the previous conclusion that even though S. aureus lost its culture capability after 48-h of co-culture with P. aeruginosa, some bacteria remains in a viable state.

Applicability of the RNA-based qPCR quantification method to further study inter-species microbial interactions. A potential advantage of the RNA-based qPCR quantification approach to study dual-species biofilms, is its ability to not only discern between the relative proportions of each species, but also to determine if a specific gene is being induced or repressed. To achieve this, the gene of interest should be normalized to the reference gene (in this case, 16s RNA gene), while this reference gene is also normalized against the exogenous ref mRNA (in this case luciferase) in order to account for the relative quantification of each bacterial population.

As a demonstration of principle of this method potential, we quantified the differences in expression of key virulence genes of P. aeruginosa and S. aureus, between single- or dual-species biofilms. To inspect the impact of S. aureus on P. aeruginosa virulence potential, we analysed the expression of pqsA (Pseudomonas quinolone signal), rhlA (rhamnolipids) and mucA (alginate biosynthesis). Pseudomonas quinolone signal (PQS) is an important P. aeruginosa signalling molecule regulating the production of quorum-sensing (QS) dependent virulence factors, such as elastase, pyocyanin and rhamnolipids. Rhamnolipids are biosurfactants produced by P. aeruginosa and...
Figure 7. Gene expression profile of (A) *P. aeruginosa* and (B) *S. aureus* virulence-related genes in 24- and 48-h-old dual-species biofilms, as compared to single-species biofilms. Transcripts were normalized to 16S rRNA and results were calculated as fold change relative to gene expression values obtained for single-species biofilms. Dashed lines represent a 2-fold cut-off value. For each condition at least two independent experiments were performed with three technical replicates. Statistical significance was determined by performing an unpaired two-tailed *t* test (**P < 0.001).

Major conclusions and study limitations. Our results underline the importance of selecting appropriate methodologies to analyse dual-species biofilm communities, since some techniques, if used alone, may not provide accurate insights into biofilm composition, resulting in misleading conclusions. Both culture-independent techniques used in this study provided a more rapid and sensitive assessment of the individual species present in the dual-species biofilm consortia, comparing with conventional culture, which underestimated the real abundance of *S. aureus* population. As each technique has its particularity, Table 1 summarise the advantages, disadvantages, and purposes of use of all methodologies applied in this study.

Of relevance, our experiments demonstrated the advantage of using a RNA-based qPCR quantification and highlighted how important is the use of an appropriate mRNA control molecule in the qPCR assay, in order to address experimental biases. Overall, this experiment confirmed that this qPCR method is able to assess specific gene expression profiling together with quantification of bacterial populations in dual-species biofilms. Despite the advantages of the qPCR assay optimized through this study, it must be recognized that potential bias can also be introduced. For instance, when gene transcript abundance is calculated by normalization against the ref mRNA, the assumption is that it reflects the behaviour among the whole RNA pool over the entire sample’s processing steps. One potential bias in normalization against the ref mRNA is that it cannot compensate the differences of RNA extraction and reverse transcription efficiencies between the targeted species. While the occurrence of this bias cannot be excluded under the tested conditions, it is believed that is safe to assume that it is not very significant, given the results achieved. The qPCR sensitivity to quantify species with very low cell numbers in certain scenarios (e.g. after exposure of an antimicrobial therapy targeting specifically a member of the polymicrobial community) may also limit its usefulness on the quantification of less prevalent species within the community.

**Experimental Procedures**

**Bacterial strains and culture conditions.** *P. aeruginosa* (strain UCBPP-PA14) and *S. aureus* (strain ATCC 25923) were used throughout this work. Both strains were stored at −80 ± 2 °C in Tryptic Soy Broth (TSB, Liofilchem) supplemented with 20% glycerol. Prior to each assay, bacteria were subcultured from frozen stock preparations onto plates of TSB supplemented with 2% (W/V) agar (TSA) and incubated aerobically at 37 °C for 24-h.

**Planktonic growth.** All assays were carried out by using a standardized bacterial inoculum. Briefly, a few colonies of each species were collected from the TSA plates and grown overnight in batches of TSB (Liofilchem) at 37°C under agitation (120 rpm). Subsequently, cells were harvested by centrifugation (9000g, 5 min) and washed...
in sterile saline solution (0.9% NaCl). The concentration of cellular suspensions was then adjusted in TSB by spectrophotometric measurement at 640 nm (calibrations were performed for each bacterial strain to relate the absorbance at 640 nm with the number of colony forming units, CFU) to obtain a final concentration of 1 × 10^5 CFU/mL or 1 × 10^6 CFU/mL. Unless otherwise stated, two independent assays with five to six replicates for each condition tested were performed. For the preparation of the dual-species planktonic suspension, overnight grown cultures of each species were adjusted to a final concentration of 1 × 10^6 CFU/mL and then mixed in three samples of various compositions (P. aeruginosa/S. aureus ratios of 10, 1 and 0.1). Two independent assays for each condition with two replicates were used. To perform the RNA extraction and subsequent reverse transcription, single- and dual-species planktonic suspension were centrifuged (7000 rpm, 10 min, 4 °C) and the pellet resuspended in RNA protect (Qiagen), as described below. Planktonic samples preparation is schematically represented in Fig. 8.

### Biofilm formation

Dual-species biofilms encompassing P. aeruginosa and S. aureus were grown as previously described, with minor modifications. Briefly, overnight cultures of each strain, grown in TSB at 37 °C and 120 rpm in aerobic conditions, were washed in sterile water and diluted in TSB to obtain 1 × 10^6 CFU/mL as final concentration. The dual-species cultures were then prepared by mixing the inoculum of each species in a 1:1 ratio. Each well of a 24-well polystyrene plate (Orange Scientific, Braine-l’Alleud, Belgium) was seeded with 1 mL of P. aeruginosa and S. aureus suspension and incubated at 37 °C on a horizontal shaker (120 rpm) for 24- and 48-h. Following biofilm growth, the planktonic fraction of the polystyrene plates was removed and the wells were washed twice with distilled sterile water. Biofilms cells were resuspended in 1 mL of 0.9% NaCl, detached by scraping and collected by centrifugation (7000 rpm, 10 min, 4 °C). Three biologic replicates of each condition were performed. To perform the RNA extraction and subsequent reverse transcription, a biofilm pooling (10 replicates of each condition tested were performed. For the preparation of the dual-species planktonic suspension, overnight grown cultures of each species were adjusted to a final concentration of 1 × 10^6 CFU/mL and then mixed in three samples of various compositions (P. aeruginosa/S. aureus ratios of 10, 1 and 0.1). Two independent assays for each condition with two replicates were used. To perform the RNA extraction and subsequent reverse transcription, single- and dual-species planktonic suspension were centrifuged (7000 rpm, 10 min, 4 °C) and the pellet resuspended in RNA protect (Qiagen), as described below. Planktonic samples preparation is schematically illustrated in Fig. 8.

### RNA extraction

RNA was extracted using the RNeasy mini kit (Qiagen), as previously optimized. Briefly, bacteria were first resuspended in 500 μL lysis buffer (provided in the kit). Then, 500 μL of phenol solution (AppliChem) was added, and the entire suspension was transferred to a safe lock tube (2 mL) with 0.4 g of acid-washed 150–212 mm silica beads (Sigma). The tube content was vortexed for 20 s previously to be placed in the FastPrep cell disruptor (BIO 101, Thermo Electron Corporation, Thermo Scientific) at 6.5 m/s for 35 s. The samples were then cooled on ice and the beat- heading step repeated twice. Afterwards, Afterwards, 700 μL of 70% ethanol (Fisher Scientific) was added to the supernatants, followed by 1 μL of 1 × 10^6 luciferase control RNA transcripts (ref mRNA) (Promega). To achieve the final RNA fraction, 30 μL of RNA-free water was added to the RNA columns. To digest possible contaminating genomic DNA, RNA was treated with DNase I (Fermentas, Ontario, Canada) following manufacturer instructions. Briefly, the RNA samples were incubated at 37 °C for 30 min after the addition of 2 μL of DNase I and 5 μL of reaction buffer. Then, 5 μL of 25 mM EDTA was added to the mixture and incubated at 65 °C for 10 min to inactivate the DNase I enzyme.

### RNA quality determination

RNA integrity and purity were assessed by gel electrophoresis and the total RNA concentration was spectrophotometrically determined using a NanoDrop 1000TM (Thermo Scientific). As indicator of protein contamination and polysaccharide, phenol, and/or chaotropic salt contamination, the absorbance ratio A_260/A_280 and A_260/A_230 were measured, respectively. The RNA integrity was assessed by 23S/16S banding pattern visualization. Electrophoresis was carried out at 80 V for 60 min using a 1.5% agarose gel. The gel was stained with GelRed (Thermo Fisher Scientific) and visualized using a GelDoc2000 (Bio-Rad, Hercules, CA, US) (Supplementary Figure S4). RNA was stored at −80 °C for further use.

### Determination of RNA recovery using an exogenous internal reference mRNA

Luciferase mRNA (Promega) was used as the exogenous internal reference (ref mRNA) to estimate the percentage of RNA

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### Table 1. Comparison of the methodologies described in this study, conventional culture, flow-cytometry and qPCR.

| Advantages | Flow Cytometry | qPCR |
|------------|----------------|------|
| ♦ Standardized method that allows the identification of the most abundant members | ♦ Allows a rapid detection of multi-species samples | ♦ Detection of live cells |
| ♦ Time consuming analysis; requires selective media; not detection of cells in a viable but non-cultivable state | ♦ Require full optimization of the method to each species | ♦ Can be used for different applications: bacterial detection and to assess changes in the gene expression profile |
| ♦ Limit of cell detection | ♦ Optimization (primers specificity, RNA extraction) | |

| Disadvantages | Fast method for enumeration of a large number of samples. |
|--------------|----------------------------------------------------------|
| ♦ Limit of cell detection | ♦ Fast method for detection and quantification; can simultaneously analyse gene expression profile of key genes (e.g. virulence genes) in the tested samples. |

| Purposes of use |
|----------------|
| Screening method for bacterial enumeration. |
loss during extraction. After the centrifugation of the lysed cells, 1 μL of a range of 1 × 10^{11} to 1 × 10^{7} ref mRNA transcripts/μL were added to the aqueous phase. After RNA extraction, the amount of ref mRNA recovered from samples was measured by qPCR and the standard curve generated from serial dilutions of ref mRNA. The recovery fraction was calculated by dividing the determined levels by the initial number of ref mRNA transcripts added to the samples.

Quantitative Real Time-PCR (qPCR). For the quantification of the 16S rRNA by qPCR, total RNA was reversely transcribed to complementary DNA (cDNA) using GRS cDNA Synthesis Kit (Grisp). Specifically, 100 ng of total RNA was reversely transcribed in 10 μL of reaction volume by using random primers. The samples were incubated at 65 °C for 5 min, 37 °C for 60 min, and 70 °C for 10 min. Primers, specific for 16S rRNA, housekeeping gene, and virulence-related genes of P. aeruginosa and S. aureus were designed (Supplementary Table S5). The qPCR reaction was performed by mixing together 5 μL of master mix SYBR Xpert Fast SYBR (Grisp), 2 μL of 1:100 diluted cDNA, 0.5 μL of forward and reverse primes (5 μM), and water up to a total volume of 10 μL. No reverse transcriptase (NRT) and a no template control (NTC) were included to verify the reaction mixtures were DNA and other contaminants-free. The efficiency of the primers used was measured by the dilution method\(^3\). To perform the qPCR run, a CFX 96 (Bio-Rad) was used with the following cycle parameter: 95 °C for 3 minutes, 40 cycles of 95 °C for 5 s, and 60 °C for 20 s. qPCR products were analysed by melting curves for unspecific products or primer dimer formation. The qPCR was applied to independently quantify the 16S rRNA of P. aeruginosa and S. aureus and the ref mRNA in each cocktailed mixed sample. The normalized gene expression was determined by using the delta Ct method (2^{∆Ct}), a variation of the Livak method, where

\[ ∆Ct = Ct(ref\ mRNA) - Ct(16S\ rRNA). \]

After normalization of the 16S rRNA for both species in each cocktail sample, the PA/SA ratio was calculated by applying the following equation:

\[ \text{Ratio} \left( \frac{PA}{SA} \right) = \frac{\text{Normalized} \left[ \text{ref mRNA/16S rRNA (PA)} \right]}{\text{Normalized} \left[ \text{ref mRNA/16S rRNA (SA)} \right]} \times \text{Normalized} \left[ \frac{16S \ rRNA (PA)}{16S \ rRNA (SA)} \right] \]

Relative fold increase of \( \text{pqsA, rhlA, mucA, sodA, sarA, agrB} \) and \( \text{hld} \) genes, was determined by applying the Pfaffl equation\(^7\) using 16S rRNA as reference gene. At least two biologic replicates of each condition were used with three technical replicates.

Biofilm cultivability. The number of adhering bacteria was determined after biofilm cell detachment. To remove any aggregates, biofilm suspensions were first sonicated for 10 s at 30% amplitude\(^11\) (Cole Parmer Ultrasonic Processor, IL, USA) and then the culturable cell count was carried out. Briefly, the disrupted biofilms

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**Figure 8.** Experimental workflow of the planktonic and biofilm samples preparation.
were serially diluted (1:10) in 0.9% NaCl, streaked onto different selective agar media and incubated at 37 °C for 18–24 h, for CFU counting. Pseudomonas isolation agar (PIA) and Mannitol Salt Agar (MSA) were the selective media used to discriminate *P. aeruginosa* and *S. aureus*, respectively. Values of cultivable biofilm cells were expressed as CFU/mL and represented as the average of three independent experiments.

**Flow-cytometry.** The total number of *P. aeruginosa* and *S. aureus* cells in 24- and 48 h-old mixed biofilms was determined using flow-cytometry. In brief, biofilm cells were collected by centrifugation (7000 rpm, 10 min, 4 °C) and suspended in 100 μL of a solution with 25 μg/mL of wheat germ agglutinin (WGA) conjugated with FITC (Molecular probes, Thermofisher Scientific). This suspension was incubated at room temperature, in the dark, for 15 min. Thereafter, cells were washed twice with 0.9% NaCl, suspended in 1 mL of the same solution and sonicated for 10 s at 30% amplitude (Cole Parmer Ultrasonic Processor, IL, USA) to disrupt aggregates. The bacterial fluorescence analysis was carried out using an EC800 Sony flow cytometer (Sony Biotechnologies Inc., CA, USA) equipped with a 488 nm laser. Multiparametric analyses were performed on both scattering signals (FSC, SSC) and FL1 channel. All the detectors were set to logarithmic amplification. Samples were acquired with a flow rate of 10 μL/min and the analysis stopped when 80,000 events were detected. Three biologic replicates of each condition were used and each sample was analysed at least twice to ensure an accurate counting. Data analysis was performed using EC800 analysis software (Sony Biotechnologies).

**Assessment of cell viability by Live/Dead staining.** To analyse the viability of dual-species biofilm consortia 48-h-old biofilms were scraped, and the biofilm disrupted-cells were collected in 1 mL sterile distilled water. Afterwards, disrupted biofilm cells were stained in the dark, for 15 min, with SYTO® BC (2 μM) and Propidium Iodide (15 μM) (both from Invitrogen™, CA, USA) prepared in saline solution, in order to differentially label live and dead cells by microscopy. For microscopic observation, an Olympus BX51 microscope fitted with fluorescence illumination was used. The optical filter combination consisted of 470 to 490 nm and 530 nm excitation filters.

**Scanning electron microscopy.** Scanning electron microscopy (SEM) was used to examine the dual-species biofilm described before, with minor modifications. Briefly, dual-species biofilms were formed on Thermostar™ plastic coverslips (Thermo Scientific™; Rochester, NY, USA) placed in the bottom of the wells of 24-well microtiter plates (Orange Scientific, Belgium). Prior to SEM observation, discs were added to aluminum pin stubs with electrically conductive carbon adhesive tape (PELCO® Tabs™). Samples were coated with 2 nm of Au for improved conductivity. The aluminum pin stub was then placed inside a Phenom Standard Sample Holder or Phenom Charge Reduction Sample Holder. The analysis was conducted at 5 kV with intensity image. The samples were characterized using a desktop Scanning Electron Microscope (SEM) (Phenom ProX, Netherlands). All results were acquired using the ProSuite software.

**Statistical analysis.** All assays were carried out in triplicate and repeated at least twice, and the results are presented as means ± SDs. The Pearson's correlation coefficient was calculated to determine the association between two variables. All P-values were based on two-tailed tests of significance, and confidence interval of 95% was used in all the analysis performed. Statistical analysis was performed using Graph Pad Prism 7.0.

**Data Availability**

The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request.

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Acknowledgements
This study was supported by the Portuguese Foundation for Science and Technology (FCT) under the scope of the strategic funding of UID/BIO/04469/2019 unit and BioTecNorte operation (NORTE-01-0145-FEDER-000004) funded by the European Regional Development Fund under the scope of Norte2020 – Programa Operacional Regional do Norte. Grant of APM (SFRH/BD/132165/2017).

Author Contributions
N.C. and M.O.P. conceptualized and designed the study. A.P.M. and A.F. performed the experiments. All authors contributed in data interpretation. A.P.M. wrote the first draft of the paper. N.C., A.F. and M.O.P. reviewed the manuscript. All authors read and approved the final manuscript.

Additional Information
Supplementary information accompanies this paper at https://doi.org/10.1038/s41598-019-50094-3.

Competing Interests: The authors declare no competing interests.

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