Identification of appropriate reference genes for qPCR studies in *Staphylococcus pseudintermedius* and preliminary assessment of *icaA* gene expression in biofilm-embedded bacteria

Evan C Crawford1*, Ameet Singh1, Devon Metcalf2, Thomas WG Gibson1 and Scott J Weese2

**Abstract**

**Background:** Quantitative PCR is rapidly becoming the standard method for analyzing gene expression in a wide variety of biological samples however it can suffer from significant error if stably expressed reference genes are not identified on which to base the analysis. Suitable reference genes for qPCR experiments on *Staphylococcus pseudintermedius* have yet to be identified.

**Results:** Three reference genes in *S. pseudintermedius* were identified and validated from a set of eight potential genes (*proC*, *gyrB*, *rplD*, *rho*, *rpoA*, *ftsZ*, *recA*, *sodA*). Two strains of *S. pseudintermedius* were used, and primer specificity and efficiency were confirmed and measured. Ranking of the genes with respect to expression stability revealed *gyrB*, *rho* and *recA* as the best reference genes. This combination was used to quantify expression of a single biofilm associated gene, *icaA*, in logarithmic, stationary and biofilm growth phases, revealing that expression was significantly upregulated in the biofilm growth phase in both strains.

**Conclusion:** Three reference genes, *gyrB*, *rho* and *recA*, were identified and validated for use as reference genes for quantitative PCR experiments in *S. pseudintermedius*. Also, the biofilm associated gene *icaA* was shown to be significantly upregulated in biofilm samples, consistent with its role in biofilm production.

**Keywords:** qPCR, Canine, MRSP, Biofilm, Reference gene

**Background**

*Staphylococcus pseudintermedius* is a common commensal organism of canines, but is also one of the most common causes of opportunistic infections [1-3]. Recently, methicillin resistant *S. pseudintermedius* (MRSP) has emerged and disseminated internationally [4,5], with 2 major sequence types (ST68 and ST71) representing the variety of clinical infections in most regions [3]. One area of concern is the ability of this bacterium to produce biofilm something that might be an important virulence factor and complicate elimination of infections [6,7]. Expression of genes pertaining to initial bacterial surface adherence and intercellular adhesion following biofilm formation, such as microbial surface components recognizing adhesive matrix molecule (MSCRAMMs) which mediate cellular adhesion, and the intracellular adhesion (*icaADBC*) operon [8], reported to be at least partially responsible for biofilm formation, likely affect the *in vivo* behavior of this organism, including resistance to therapy [9,10]. While there is significant postulation regarding these factors, understanding of the expression of antimicrobial resistance and biofilm associated genes in *S. pseudintermedius* and the subsequent clinical implications is still poor.

Quantitative real-time PCR (qPCR) is increasingly employed to quantify gene expression. While it can be very sensitive and specific, there are numerous pitfalls in its application that can easily result in misleading and incorrect conclusions. On of the most frequent errors is a failure to confirm the constitutive expression of the reference genes
used to measure the relative expression of genes of interest. Normalization of results in qPCR is vital to limit variability introduced by experimental conditions, sample preparation and analysis, and is one of the main underlying tenets of qPCR analysis. Selection of inappropriate reference genes can result in grossly incorrect conclusions owing to the miscalculation of gene expression. These and other reasons have prompted the development of minimum information for publication of qPCR experiments (MIQE) guidelines to ensure integrity, consistency and transparency of qPCR experiments, including standards for all aspects of experimental design, analysis and reporting [11].

Current recommendations suggest a minimum of three reference genes (ideally with M values below 1 for heterogenous samples), and the inclusion of additional genes as necessary to obtain a pairwise variation value < 0.15 [12]. This requires specific validation of candidate reference genes in the bacterium (and ideally strains) to be studied, something that is lacking for *S. pseudintermedius*. While multiple qPCR studies have been performed in various *Staphylococcus* spp [13,14], study and validation of reference genes in *S. pseudintermedius* is lacking, and it cannot be assumed that data from other staphylococci apply to this species.

The objectives of this study were to evaluate several potential reference genes in *S. pseudintermedius*, to identify the optimum gene or gene combinations for future qPCR expression studies and to evaluate expression of *icaA* using validated reference genes.

### Results and discussion

#### RNA isolation

The modified protocol afforded good to excellent yields (mean yield 53.5 ± 23.4 μg, range 16.9 – 92.2 μg, concentration 63.9 ± 94 μg/μL), as well as good RNA purity and integrity (RNA Integrity Number mean 9.1 ± 0.4) (Table 1), though there was evidence of some mild contamination from purification reagents (260/230 < 2). qPCR was performed on all RNA preparations to confirm DNA elimination; quantification cycle numbers were at least 20 cycles lower for RNA samples than equivalent amounts of DNA.

The modified protocol for RNA and DNA extraction, namely the inclusion of lysostaphin (and Dispersin B for biofilm samples) in the initial lysis buffer resulted in substantial and reproducible increases in RNA and DNA yield during harvesting (results not shown). Dispersin B catalyzes the hydrolysis of polysaccharide intercellular adhesin (PIA), a major constituent of the extracellular matrix of *Staphylococcus* spp. biofilms. Incorporating this enzyme in the solution to recover the biofilm from a surface, and in the lysis buffer is thought to increase recovery by enzymatically degrading the extracellular matrix of biofilm, releasing adherent bacteria for recovery, and exposing them to the lysis solution. The ability of this simple method to yield an adequate quantity and quality of DNA from biofilm-embedded bacteria was an important finding and will facilitate future studies of gene expression in biofilms.

### Amplification specificity and determination of PCR efficiency of reference genes

Quantification cycle (Cq) for each reaction was plotted against the log of DNA concentration, and the slopes of the curves were used to calculate the PCR efficiency values, which ranged from 1.79 to 1.87, with excellent regression coefficients (Table 2). Despite the range in melting points of the qPCR products (Table 2), all primers amplified well under the conditions listed. Melt curve analysis showed a single melt curve for each target gene, and DNA agarose gel electrophoresis revealed a single peak for each product. Sequencing of the qPCR products matched the sequence under the conditions listed. Melt curve analysis showed a single melt curve for each target gene, and DNA agarose gel electrophoresis revealed a single peak for each product. Sequencing of the qPCR products matched the sequence.

### Comparison of qPCR reaction products

Melt curve analyses and gel electrophoresis were also performed on and compared between samples recovered from qPCR reactions performed on cDNA prepared using random hexamers, single gene specific primer and combination gene specific primer reverse transcript reactions; there were no differences in any of the products obtained from those obtained from qPCR or genomic DNA.

---

**Table 1 RNA quality and recovery**

| Sample ID | Concentration (ng/μL) | 260/280 | 260/230 | Total RNA recovery (μg) | RIN |
|-----------|-----------------------|---------|---------|-------------------------|-----|
| A42 log A | 616.2                 | 2.15    | 1.09    | 61.6                    | 9.6 |
| A42 log B | 771.4                 | 2.15    | 2.27    | 77.1                    | n.t.|
| A42 log C | 805.5                 | 2.15    | 2.27    | 80.5                    | 9.0 |
| A54 log A | 823.1                 | 2.14    | 2.20    | 82.3                    | 9.4 |
| A54 log B | 919.4                 | 2.13    | 1.67    | 91.9                    | n.t.|
| A54 log C | 921.6                 | 2.14    | 2.03    | 92.2                    | 9.4 |
| A42 stat A| 535.4                 | 2.10    | 2.00    | 53.5                    | 9.2 |
| A42 stat B| 449.4                 | 2.02    | 1.95    | 44.9                    | 8.8 |
| A42 stat C| 493.0                 | 2.05    | 1.66    | 49.3                    | 8.8 |
| A54 stat A| 403.3                 | 2.05    | 2.05    | 40.3                    | 8.5 |
| A54 stat B| 630.1                 | 2.13    | 1.89    | 63.0                    | n.t.|
| A54 stat C| 397.5                 | 2.05    | 1.89    | 39.7                    | 8.8 |
| A42 BF A  | 732.4                 | 2.12    | 2.17    | 36.6                    | 9.4 |
| A42 BF B  | 738.6                 | 2.11    | 2.18    | 36.9                    | n.t |
| A42 BF C  | 887.3                 | 2.12    | 2.19    | 44.4                    | 8.4 |
| A54 BF A  | 606.9                 | 2.11    | 1.97    | 30.3                    | 9.5 |
| A54 BF B  | 337.9                 | 2.06    | 1.74    | 16.9                    | n.t.|
| A54 BF C  | 421.9                 | 2.03    | 1.99    | 21.1                    | 8.9 |

(n.t: not tested, log: logarithmic, stat: stationary, BF: biofilm samples).
Stability assessment and validation of reference genes

Relative expression levels (quantification cycle numbers) were entered into the Microsoft Excel (Microsoft Canada, Mississauga, ON) visual basic application geNorm [12], which calculated stability values (M values) for each gene (Table 3). GeNorm was also used to calculate normalization factors using combinations of the most stably expressed genes, and calculated the pairwise variation between these factors to identify the optimum number of reference genes to use (Table 4). The three genes with the lowest individual M values were used in combination (rho, recA, gyrB) for subsequent expression analysis.

While stability of expression of reference genes is important, reference genes ideally also have a high PCR efficiency. The primers we identified had efficiency ranging from 1.79 to 1.84 (with a value of 2.0 representing 100% efficiency), and while the final three genes used for normalization did not have the highest efficiencies, the values are accounted for during normalization of qPCR expression data.

With respect to the identification of reference genes, these results are specifically only applicable to the two strains studied under the three growth phases sampled. However, the two strains that were studied comprised the two main international MRSP clones [3], suggesting that these genes will be suitable for broad studies of MRSP gene expression. The excellent stability of expression of these genes over the broad range of conditions studied also indicate that these targets will likely function very well as reference genes over a generally broad range of conditions.

icaA expression

Expression of icaA was significantly higher in the biofilm compared to logarithmic and stationary phases (p = 0.0093 (A42), p < 0.0001 (A54), Figure 1). Individual Tukey’s post-hoc p-values for individual comparisons were for logarithmic vs biofilm p = 0.015 (A42); p = 0.0001 (A54) and for stationary vs biofilm p = 0.015 (A42); p = 0.0002 (A54). There was no difference in expression level between logarithmic and stationary phases for either strain, individual Tukey post-hoc p > 0.9999 (A42), p = 0.8258 (A54).

These findings are unsurprising given the role of this gene in formation of polysaccharide intracellular adhesin, but it is noteworthy that such a profound alteration in expression was detectable. Further investigation of expression of this gene under other conditions and on a variety of surfaces, as well as studies of expression of other biofilm associated genes such as MSCRAMMs.

---

### Table 2 Reference and target gene efficiency determination and amplicon melting points, r² values are regression coefficients for the curves

| Candidate reference genes | Slope of the curve | r² on the slope | PCR efficiency | Melting temperature (°C) |
|---------------------------|--------------------|----------------|----------------|--------------------------|
| proC                      | −3.78              | 0.997          | 1.84           | 80.0                     |
| gyrB                      | −3.85              | 0.998          | 1.82           | 79.5                     |
| rplD                      | −3.86              | 0.998          | 1.82           | 75.5                     |
| rho                       | −3.68              | 0.997          | 1.87           | 81.0                     |
| rpoA                      | −3.89              | 0.996          | 1.81           | 76.5                     |
| ftsZ                      | −3.85              | 0.998          | 1.82           | 79.0                     |
| recA                      | −3.83              | 0.998          | 1.82           | 81.0                     |
| sodA                      | −3.97              | 0.996          | 1.79           | 79.5                     |
| Target gene               | −3.92              | 0.995          | 1.80           | 80.5                     |

### Table 3 GeNorm gene stability (M) values

| Candidate reference genes | geNorm M value |
|---------------------------|----------------|
| proC                      | 0.701          |
| gyrB                      | 0.604          |
| rplD                      | 0.769          |
| rho                       | 0.595          |
| rpoA                      | 0.663          |
| ftsZ                      | 0.848          |
| recA                      | 0.599          |
| sodA                      | 1.150          |

### Table 4 Pairwise variation in normalization factor for combinations of reference genes

| Number of genes used | Pairwise variation of normalization factors |
|----------------------|--------------------------------------------|
| 2 vs 3               | 0.119                                      |
| 3 vs 4               | 0.096                                      |
| 4 vs 5               | 0.082                                      |
| 5 vs 6               | 0.112                                      |
| 6 vs 7               | 0.107                                      |
| 7 vs 8               | 0.135                                      |

---

Crawford et al. BMC Research Notes 2014, 7:451
http://www.biomedcentral.com/1756-0500/7/451
Conclusion
Proper development of qPCR assays, including reference gene assessment, is a critical quality control step in the application of this tool. This study has identified a group of *S. pseudintermedius* reference genes, and used those reference genes to demonstrate a significant expression change in a biofilm associated gene. This information provides a vital background for the performance of gene expression studies in this increasingly important veterinary pathogen.

Methods
Bacterial strains and culture conditions
Two canine *S. pseudintermedius* strains were chosen, representing the two main international MRSP clones, sequence type (ST) 68 (strain A42) and ST71 (strain A54). These two strains were isolated from clinical infections, and were both previously classified as moderate biofilm formers using a polystyrene plate assay (unpublished data). Isolates were grown in tryptic soy broth supplemented with 1% (w/v) dextrose for all growth conditions. Samples from 1% (w/v) dextrose for all growth conditions. Samples were isolated as single colonies from streak plates prepared using a polystyrene plate assay (unpublished data).

DNA isolation
Cells were collected from a logarithmic phase sample as described above, and processed using the High Pure PCR Template Preparation Kit (Roche Applied Science, Indianapolis, IN) following the manufacturer’s instructions with the exception that lysostaphin (Sigma-Aldrich, Oakville, ON) was added at 0.1 mg/mL in the initial lysis step. Concentration and purity at 260/280 nm was measured using a Nanodrop ND-100 spectrophotometer (Nanodrop Technologies Inc., Wilmington, DE), and the samples were qualitatively examined for shearing by DNA gel electrophoresis.

RNA isolation
Each pellet of cells was resuspended in 2 mL PBS (pH 7.4), centrifuged to a pellet and the supernatant discarded. The
pelleted cells were subsequently processed using the Qiagen RNeasy Mini kit (Qiagen, Germantown, MD), following the manufacturer’s instructions, with the following changes. The cells were initially resuspended in tris-EDTA buffer (pH 7.5) containing 15 mg/ml lysozyme (Sigma-Aldrich), 2 mg/mL proteinase K (Sigma-Aldrich), 25 μg/mL Dispersin B (Kane Biotech Inc., Winnipeg, MB) and 0.1 mg/mL lysostaphin (Sigma-Aldrich). All optional steps for additional washes or spins during the procedure were performed. Logarithmic and stationary phase samples were eluted from the purification column with two volumes of 50 μL sterile nuclease free water, biofilm samples were eluted with one volume of 50 μL (volumes chosen so as to obtain final RNA concentrations above 400 ng/μL). All samples were treated with the DNAfree DNase kit (Ambion, Austin, TX) as per the manufacturer’s instructions using a total of 2 units per reaction, added in two aliquots. A Nanodrop ND-100 spectrophotometer (Nanodrop Technologies Inc.) was used to measure RNA concentration and purity at 260/280 nm. Three samples of each RNA sample were measured and the measured concentrations were averaged. Representative samples were submitted for Bioanalyzer analysis (Agilent Technologies, Santa Clara, CA). Real-time PCR was performed using the efficiency determination protocol on RNA samples to confirm the absence of DNA using gyrB primers. RNA samples were stored at −20 to −80°C until further use.

Reverse transcription/cDNA preparation
Two μg of total RNA was used in each 20 μL gene specific reverse transcription reaction using the Omniscript RT PCR kit (Qiagen), using 10 μM of each primer, 0.5 mM dNTP, 1 U reverse transcription enzyme, provided buffer diluted to 1x and the remainder as water. A mixture of forward and reverse primers for all nine genes was used. Reactions were incubated at 42°C for 60 minutes. Reverse transcription products were purified using the QIAquick PCR purification kit (Qiagen) as per the manufacturer’s instructions, eluted with 100 μL Qiagen EB buffer, and stored at −20°C until further use.

qPCR for gene stability and icaA
The same protocol for efficiency testing was used to examine stability of potential reference gene and icaA expression, with the only change being that only 30 cycles were performed. In triplicate, cDNA produced from all three growth phases were subjected to qPCR (as described for the efficiency evaluation) for each candidate gene. In each 20 μL reaction, 5 μL of purified RT reaction product (equivalent to 100 ng of RNA prior to reverse transcription) was used. No template (negative; water) and DNA (positive; 100 ng DNA) controls were included in each run. No amplification was identified in negative controls. Intra-plate normalization was performed using the measured level of icaA in the positive control on each plate.
icaA expression, statistical analysis
Expression of icaA was compared between the three growth phases for the two strains using the combination of three reference genes. Normalization factors, calibration factors and relative expression was calculated as was reported by Hellemans et al. [16]. A one-way ANOVA with Tukey’s post-hoc test was performed to compare expression between growth phases. P-values < 0.05 were considered significant.

Abbreviations
qPCR: Quantitative polymerase chain reaction; MRSP: Methicillin resistant Staphylococcus pseudintermedius; MSCRAMM: Microbial surface component recognizing adhesive matrix molecule; MIQE: Minimum information for quantitative PCR experiment; ST: Sequence type; OD: Optical density; w/v: Weight/volume; PBS: Phosphate buffered saline; cDNA: Complementary DNA; RIN: RNA integrity number; Cq: Quantification cycle.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
EC performed the majority of the laboratory work, assisted by DM. Primer design was completed by EC and DM. The study was conceived by SW and AS, who also both participated in the experimental design. EC drafted the initial manuscript, and all authors were involved in editing and production of the final version. All authors read and approved the final manuscript.

Acknowledgements
The authors wish to thank Joyce Rousseau for her tireless effort and support in the laboratory.

Funding
This project was supported by the AKC Canine Health Foundation (AS). The contents of this publication are solely the responsibility of the authors and do not necessarily represent the views of the Foundation.

Author details
1Department of Clinical Studies, Ontario Veterinary College, University of Guelph, 50 Stone Road, Ontario N1G 2 W1, Canada. 2Department of Pathobiology and Centre for Public Health and Zoonoses, Ontario Veterinary College, University of Guelph, 50 Stone Road, Ontario N1G 2 W1, Canada.

References
1. Weese JS: A review of multidrug resistant surgical site infections. Vet Comp Orthop Traumatol 2008, 21:1.
2. Griffith GC, Morris DO, Abraham JL, Shofer FS, Rankin SC: Screening for skin carriage of methicillin-resistant coagulase-positive staphylococci and Staphylococcus schleiferi in dogs with healthy and inflamed skin. Vet Dermatol 2008, 19:142–149.
3. Perreten V, Kadlec K, Schwarz S, Andersson UG, Finn M, Greko C, Moodley A, Kania SA, Frank LA, Bemis DA, Franco A, Iurescia M, Battisti A, Duim B, Wagenaar JA, van Duikken E, Weese JS, Fitzgerald JR, Rossano A, Guardabassi L: Clonal spread of methicillin-resistant Staphylococcus pseudintermedius in Europe and North America: an international multicentre study. J Antimicrob Chemother 2010, 65:1145–1154.
4. van Duikken E, Catry B, Greko C, Moreno MA, Pomba MC, Pyorala S, Ruzauskas M, Sanders P, Threlfall EJ, Torren-Edo J, Tompkins C: Review on methicillin-resistant Staphylococcus pseudintermedius. J Antimicrob Chemother 2011, 66:2705–2714.
5. Weese JS, van Duikken E: Methicillin-resistant Staphylococcus aureus and Staphylococcus pseudintermedius in veterinary medicine. Vet Microbiol 2010, 140:418–429.
6. Otto W: Staphylococcal Infections: Mechanisms of Biofilm Maturation and Detachment as Critical Determinants of Pathogenicity. Annu Rev Med 2012, 64:175–188.
7. Dicicco M, Neethirajan S, Singh A, Weese JS: Efficacy of clarithromycin on biofilm formation of methicillin-resistant Staphylococcus pseudintermedius. BMC Vet Res 2012, 8:225.
8. Fey PD, Olson ME: Current concepts in biofilm formation of Staphylococcus epidermidis. Future Microbiol 2010, 5:917–933.
9. Rohde H, Frankenberger S, Zähringer U, Mack D: Structure, function and contribution of polysaccharide intercellular adhesin (PIA) to Staphylococcus epidermidis biofilm formation and pathogenesis of biomaterial-associated infections. Eur J Cell Biol 2010, 89:103–111.
10. Römling U, Balsalobre C: Biofilm infections, their resilience to therapy and innovative treatment strategies. J Intern Med 2012, 272:541–561.
11. Bustin SA, Benes V, Garson JA, Hellemans J, Huggett J, Kubista M, Mueller R, Nolan T, Pfaffl MW, Shipley GL, Vandesompele J, Wittwer CT: The MIQE Guidelines: Minimum Information for Publication of Quantitative Real-Time PCR Experiments. Clin Chem 2009, 55:611–622.
12. Vandesompele J, De Preter K, Pattyn F, Poppe B, Van Roy N, De Paepe A, Speleman F: Accurate normalization of real-time quantitative RT-PCR data by geometric averaging of multiple internal control genes. Genome Biol 2002, 3:71–8.
13. Chung PY, Chung L, Navaratnam P: Identification of novel gene targets in Staphylococcus aureus treated with betulinic acid by gene expression profiling analysis. Res Microbiol, in press.
14. Thes T, Skurray RA, Brown MH: Identification of suitable internal controls to study expression of a Staphylococcus aureus multidrug resistance system by quantitative real-time PCR. J Microbiol Meth 2007, 70:355–362.
15. Singh A, Walker M, Rousseau J, Weese JS: Characterization of the biofilm forming ability of Staphylococcus pseudintermedius from dogs. BMC Vet Res 2013, 9:93.
16. Hellemans J, Mortier G, De Paepe A, Speleman F, Vandesompele J: qBase relative quantification framework and software for management and automated analysis of real-time quantitative PCR data. Genome Biol 2007, 8(2):R19.

Cite this article as: Crawford et al.: Identification of appropriate reference genes for qPCR studies in Staphylococcus pseudintermedius and preliminary assessment of icaA gene expression in biofilm-embedded bacteria. BMC Research Notes 2014 7:451.