Analysis of multidrug resistant group B streptococci with reduced penicillin susceptibility forming small, less hemolytic colonies

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Abstract

Group B streptococci (GBS; Streptococcus agalactiae) are the leading cause of neonatal invasive diseases and are also important pathogens for elderly adults. Until now, nearly all GBS with reduced penicillin susceptibility (PRGBS) have shown β-hemolytic activity and grow on sheep blood agar. However, we have previously reported three PRGBS clinical isolates harboring a CylK deletion that form small less hemolytic colonies. In this study, we examined the causes of small, less hemolytic colony formation in these clinical isolates. Iso- genic strains were sequenced to identify the mutation related to a small colony size. We identified a 276_277insG nucleic acid insertion in the thiamin pyrophosphokinase (tpk) gene, resulting in premature termination at amino acid 103 in TPK, as a candidate mutation responsible for small colony formation. The recombinant strain Δtpk, which harbored the 276_277insG insertion in the tpk gene, showed small colony formation. The recombinant strain ΔcylK, which harbored the G379T substitution in cylK, showed a reduction in hemolytic activity. The phenotypes of both recombinant strains were complemented by the expression of intact TPK or CylK, respectively. Moreover, the use of Rapid ID 32 API and VITEK MS to identify strains as GBS was evaluated clinical isolates and recombinant strains. VITEK MS, but not Rapid ID 32 API, was able to accurately identify the strains as GBS. In conclusion, we determined that mutations in tpk and cylK caused small colonies and reduced hemolytic activity, respectively, and characterized the clinical isolates in detail.
Introduction

Streptococcus agalactiae (GBS) is the leading cause of neonatal sepsis and meningitis and is responsible for high mortality and morbidity, particularly in neonates and those suffering from underlying medical conditions, such as diabetes [1–3]. β-Lactams are first-line antimicrobial agents for intrapartum antibiotic prophylaxis and the treatment of GBS infections [4, 5]. However, GBS clinical isolates with reduced penicillin susceptibility (PRGBS) have emerged via the acquisition of substitutions, including V405A and/or Q557E, in penicillin-binding protein 2X [6–10]. PRGBS clinical isolates tend to be non-susceptible or resistant to fluoroquinolones and macrolides [11–13]. Most PRGBS clinical isolates show β-hemolytic activity and grow on sheep blood agar. However, we previously reported three multidrug-resistant PRGBS clinical isolates (MRY11-004, MRY11-005, and NUBL-2449) that form atypical, small less-β-hemolytic colonies on sheep blood agar [14]. These clinical isolates harbor a G379T nucleic acid substitution in the cylK gene, resulting in premature termination at amino acid 127 in CylK, which is required for full hemolytic activity of GBS [14].

Small colony variants (SCVs) are characterized by reduced growth, small colony size, and atypical colony morphology. Additional features, such as decreased respiration, increased resistance to aminoglycosides, reduced fermentation of sugars, and an unstable phenotype are common. SCVs are often linked to a deficiency in electron transport or thymidine biosynthesis [15], but the precise causes of small colony formation in other cases are unclear [16–20]. Although studies of the morphological and biochemical characteristics of SCVs have been most extensively studied in staphylococci [21–23], SCVs are found in various genera and species, e.g., enterococci [17, 18, 24], Streptococcus pneumoniae [19, 20], Streptococcus tigurinus [16], Escherichia coli [25], and Pseudomonas aeruginosa [15].

To our knowledge, there is one report of GBS opacity variants obtained from an unknown number of passages of a clinical isolate [26]. However, there are no previous reports of clinical GBS small colony variants. In this report, we elucidated the causes of reduced hemolytic activity and small colony formation in three clinical isolates of PRGBS.

Materials and methods

Bacterial strains and culture conditions

The three clinical isolates (MRY11-004, MRY11-005, and NUBL-2449) were recovered from two patients in one hospital in 2011. The first patient was an 88-year-old man who had underlying diseases, including diabetic peripheral neuropathy. The isolates were recovered in January of 2011. MRY11-004 and MRY11-005 were isolated from his blood and sputum, respectively. The second patient was an 83-year-old man. NUBL-2449 was isolated from his sputum in November of 2011. All three clinical isolates were classified as ST1. Moreover, all three clinical isolates showed an identical pulsotype according to PFGE. The details of these clinical isolates are described in [14].

The GBS type V strain ATCC BAA-611 (2603 V/R) and type Ia ATCC BAA-1138 (A909) were used as the parent strains to generate recombinants. The GBS recombinant strain ΔcylK was based on ATCC BAA-611, harboring the G379T substitution in cylK, resulting in premature termination at amino acid 127 in CylK. The GBS recombinant strain Δtpk was based on ATCC BAA-1138, harboring the 276_277insG insertion in tpk, resulting in premature termination at amino acid 103 in thiamin pyrophosphokinase (TPK), which catalyzes the direct phosphorylation of thiamin via ATP to form thiamin pyrophosphate (TPP).

GBS was cultivated in Todd–Hewitt broth (THB) (BD, Franklin Lakes, NJ, USA) and Todd–Hewitt agar (THA) at 37˚C in 5% CO₂. GBS strains carrying recombinant pG+host6 or...
pDL278 derivatives were grown in the presence of erythromycin (5 μg/ml), chloramphenicol (10 μg/ml), and/or spectinomycin (300 μg/ml). Cultivation of Δtpk was performed at 37˚C in 5% CO₂ in THB and on THA containing TPP (500 μg/L). E. coli DH10B was grown at 37˚C in LB broth and strains carrying recombinant pG+host6 or pDL278 derivatives were selected in the presence of ampicillin (200 μg/ml) and spectinomycin (200 μg/ml).

Next-generation sequencing and analysis

Bacteria were cultured overnight at 37˚C in 5% CO₂, and chromosomal DNA was extracted using the QIAamp DNA Mini Kit (Qiagen, Hilden, Germany) following the manufacturer’s protocol, with slight modifications in order to completely remove RNA. The overnight culture (1 mL) was centrifuged for 10 min at 3,000 × g, and the supernatant was removed. The pellet was resuspended in 200 μL of the suspension solution and then incubated at 37˚C for 90 min. Then, 20 μL of proteinase K was added, and the solution was incubated for 60 min at 57˚C, until it became clear. RNase A (4 μL) was added, and the solution was incubated for 5 min at 25˚C. DNA was then purified following the QIAamp DNA Kit protocol. Concentrations of extracted DNA were measured using both NanoDrop and QuBit fluorometric quantitation. Extracted DNA was then prepared for next-generation sequencing using the Nextera XT DNA Preparation Kit. Sequencing was performed using a MiSeq sequencer (Illumina, San Diego, CA, USA) with the MiSeq Reagent Kit v3 (2 × 300 mer). Paired-end sequencing data from the MiSeq reporter software were further analyzed using CLC genomics Workbench ver. 9 (CLC Bio).

Construction of ΔcylK and a complementation plasmid to express intact CylK in ΔcylK

pG+host6-ΔcylK was constructed to introduce the G379T substitution into cylK. The plasmid was comprised of a thermosensitive plasmid pG+host6 backbone, with a fragment of the cylK gene containing the G379T substitution (nucleotide positions 659180 to 659657 in the GBS ATCC BAA-611 genome), the chloramphenicol acetyltransferase gene, conferring resistance to chloramphenicol, and a fragment of DNA from the region directly downstream of the cylK gene (nucleotide positions 659073 to 659648 in the GBS ATCC BAA-611 genome). These fragments were amplified by PCR using primers listed in S1 Table. Then, pG+host6-ΔcylK was transformed into E. coli DH10B for amplification. The purified plasmid was transformed into GBS BAA-611 and the transformants were selected on THA containing 0.5 μg/ml erythromycin at 30˚C. Successful integrant strains were then cultivated for 3 days in THB at 30˚C without erythromycin selection to facilitate the excision of the vector pG+host6-ΔcylK [27].

The pCyIK plasmid was constructed to express full-length CylK in ΔcylK. pCyIK was comprised of the E. coli-GBS shuttle vector plasmid pDL278 backbone, with a fragment of the promoter region from the bca gene (nucleotide positions 459015 to 459255 in the GBS ATCC BAA-1138 genome) and the full-length cylK gene (nucleotide positions 459015 to 459255 in the GBS ATCC BAA-611 genome) [28, 29]. The pCyIK plasmid was transformed into E. coli DH10B for amplification. The purified plasmid was transformed into ΔcylK and the transformants were selected on THA containing 300 μg/ml spectinomycin.

Construction of Δtpk and a complementation plasmid to express intact TPK in Δtpk

Δtpk was constructed using a similar method to that used for the construction of ΔcylK, with minor modifications. pG+host6-Δtpk was constructed in order to introduce the 276_277insG
insertion in \( tpk \). The plasmid was comprised of a thermosensitive plasmid pG+host6 backbone, with a fragment of the \( tpk \) gene containing the 276_277\( \text{ins} \)G insertion (nucleotide positions 1790161 to 1790826 in the GBS ATCC BAA-1138 genome), the chloramphenicol acetyltransferase gene, and a fragment of DNA from the region directly downstream of the \( tpk \) gene (nucleotide positions 1789494 to 1790190 in the GBS ATCC BAA-1138 genome). Integron strains were successively cultivated for 3 days in THB containing TPP (500 \( \mu \text{g/L} \) at 30°C without erythromycin selection to facilitate the excision of vector pG+host6-\( \Delta \)tpk.

The pTPK plasmid was constructed to express full-length TPK in \( \Delta \)tpk. pTPK was comprised of the \( E. \text{coli} \)-GBS shuttle vector plasmid pDL278 backbone, with a fragment of the promoter region from the \( bca \) gene (nucleotide positions 459015 to 459255 in the GBS ATCC BAA-1138 genome), and the full-length \( tpk \) gene (nucleotide positions 1790174 to 1790806 in the GBS ATCC BAA-1138 genome). pTPK was transformed into \( E. \text{coli} \) DH10B for amplification. The purified plasmid was transformed into \( \Delta \)tpk and MRY11-004. Thereafter, transformants were selected on THA containing 300 \( \mu \text{g/ml} \) spectinomycin at 37°C.

Growth curve

Bacterial growth in THB at 37°C in ambient air was monitored using an OD monitor (ODBox-C; TAITEC, Koshigaya, Japan). Overnight culture (60 \( \mu \text{L} \)) was diluted in 6 mL of fresh THB. The cultures were shaken at 160 rpm and the optical density (600 nm) was measured every 5 min for 24 h. The experiment was performed five times.

Transmission electron microscopy

For transmission electron microscopy (TEM) analyses, BAA-1138 and \( \Delta \)tpk cultures were incubated for 12 h in THB and THB containing TPP (500 \( \mu \text{g/L} \)), respectively. MYR11-004 and \( \Delta \)tpk cultures were also incubated for 12 h in THB containing no TPP. After washing with PBS (0.1 M Phosphate Buffer Solution, pH 7.4) three times, bacterial cells were washed in PBS containing 2% glutaraldehyde and then stored at 4°C overnight. Bacterial cell pellets were washed with PBS four times, resuspended in PBS containing 2% osmium tetroxide, and incubated at 4°C for 2 h. Samples were cut using an ultramicrotome. TEM was performed using the JEM-1400PLUS (JEOL, Tokyo, Japan).

Rapid ID 32 API and VITEK MS assay

Clinical isolates, \( \Delta \)tpk, and BAA-1138 were analyzed using Rapid ID 32 API (bioMérieux, Marcy-l’Étoile, France) and VITEK MS (Sysmex bioMérieux) according to the manufacturer’s instructions. The three clinical isolates and \( \Delta \)tpk were cultured on sheep blood agar (Nissui, Tokyo, Japan) for 3 days and on chocolate II agar (BD) in 5% \( \text{CO}_2 \) overnight. BAA-1138 cultured on sheep blood agar and on chocolate II agar in 5% \( \text{CO}_2 \) overnight was also analyzed.

Nucleotide sequence accession numbers

Raw sequence data from Miseq sequencing are deposited as fastq files at EMBL/GenBank under the accession numbers SAMD00077745, SAMD00077746, SAMD00077747, SAMD00077757, SAMD00077758 and SAMD00077759 for MRY11-004, MRY11-005, NUBL-2449, MRY11-004L, MRY11-005L and NUBL-2449L, respectively. The raw sequencing data have been deposited in the DDBJ Sequence Read Archive (DRA) under accession numbers DRA005682, DRA005683, DRA005684, DRA005686, DRA005687, and DRA005688 for MRY11-004, MRY11-005, NUBL-2449, MRY11-004L, MRY11-005L, and NUBL-2449L, respectively [30].
Statistical analysis

Growth curve data are expressed as means ± standard deviation. The statistical significance (P-values) of differences in mean values for two-sample comparisons was determined using the Student’s t-test implemented in Microsoft Excel. P < 0.05 indicated statistical significance [31].

Results

Phenotypes of the ΔcylK strain

To determine if ΔcylK causes reduced hemolytic activity and/or small colony formation, we constructed ΔcylK based on BAA-611 harboring the G379T substitution in cylK gene. The recombinant strain showed less β-hemolytic activity than that of the wild-type strain on sheep blood agar plates. However, ΔcylK did not show small colony formation (Fig 1). The mutant

![Image of colony size and hemolytic activity comparison](https://doi.org/10.1371/journal.pone.0183453.g001)

Fig 1. Comparison of hemolytic activity and colony formation to elucidate influence of deletion CylK. Colony size and hemolytic activity of a GBS ATCC strain (BAA-611), recombinant strains (ΔcylK), and complemented strains (ΔcylK+ pDL278 and ΔcylK+pCylK) on 5% sheep blood agar. The strains were grown at 37˚C in 5% CO2 for 16 h. ΔcylK is a recombinant strain based on ATCC BAA-611, harboring the G379T substitution in cylK, resulting in premature termination at amino acid 127 in CylK. pDL278 is a gram-positive and gram-negative shuttle vector. pCylK is a complementation plasmid used to express intact TPK in the ΔcylK strain.

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was then complemented with a plasmid harboring cyk and the resulting strain showed similar hemolytic activity to that of the wild type on sheep blood agar plates (S1 Fig).

Identification of the cause of small colony formation and phenotypes of the Δtpk strain

After several in vitro passages on 5% sheep blood agar, three clinical isolates reverted to a fast-growing phenotype and large colony size, equal to that of the GBS ATCC strains (approximately 1 mm in diameter). Large colonies were confirmed to be GBS using the agglutination method with anti-Lancefield B antigen serum and serotype VIII with anti-GBS serotype-specific serum, and these results were identical to those obtained for clinical isolates. To identify a causal genetic factor for the observed differences, the whole genomes of the clinical isolates and the derivative strains were sequenced. Genomes were compared using CLC Workbench ver. 9 to detect single nucleotide polymorphisms (SNPs). Although several SNPs were found in comparisons between the clinical isolates and derivative strains (Table 1), only the 276_277insG insertion in the tpk gene was found in all three clinical isolates. This insertion was confirmed using Sanger sequencing, and caused premature termination at amino acid 103 in TPK. To determine the effects of this insertion, we constructed the Δtpk recombinant strain based on S. agalactiae ATCC BAA-1138 harboring the 276_277insG insertion in the tpk gene. The Δtpk strain showed similar hemolytic activity on sheep blood agar to that of the wild-type strain (S1 Fig) and smaller colonies (less than 1 mm in diameter) than those of the wild-type strain (approximately 1 mm in diameter) on sheep blood agar (Fig 2). Moreover, when a plasmid expressing full-length TPK was introduced into Δtpk and MRY11-004, the resulting strain showed similar colony formation to that of the wild-type strain on sheep blood agar (Fig 2).

Table 1. Nucleic acid differences between MRY11-004 and MRY11-004L, MRY11-005 and MRY11-005L, NUBL2449 and NUBL2449L using CLC workbench ver. 9 (>80%; Frequency).

| Nucleic acid difference | Position | Effect (length with the insertion or deletion/full length of the gene) | Gene description (Locus tag) |
|-------------------------|----------|-------------------------------------------------------------------|-----------------------------|
| MRY11-004               |          |                                                                  |                             |
| MRY11-004L              |          |                                                                  |                             |
| G                       | -        | 276_277                                                          | thiamine pyrophosphokinase (SAG1775) |
| A                       | G        | 78 bases upstream of SAG2072 (78 bases upstream of SAG2073)       | uridine phosphorylase (SAG2072) (GmR family transcriptional regulator (SAG2073)) |
| -                       | C        | 226_227                                                          | Phosphoribosylaminomimidolecarbonamide formyltransferase/IMP cyclohydrolase (SAG0030) |
| -                       | C        | 341_342                                                          | Mannosyl-glycoprotein endo-β-N-acetylglucosaminidase (EN73_03515) |
| MRY11-005               |          |                                                                  |                             |
| MRY11-005L              |          |                                                                  |                             |
| G                       | -        | 276_277                                                          | thiamine pyrophosphokinase (SAG1775) |
| NUBL2449               | NUBL2449L|                                                                  |                             |
| G                       | G        | 276_277                                                          | thiamine pyrophosphokinase (SAG1775) |
| C                       | -        | 279                                                              | thiamine pyrophosphokinase (SAG1775) |
| -                       | C        | 1442_1443                                                        | elongation factor G (SAG1769) |

The strains MRY11-004L, MRY11-005L, and NUBL2449L are large colony strains derived from their parental clinical small colony strains MRY11-004, MRY11-005, and NUBL2449, respectively, after several passages on sheep blood agar. – indicates a deletion. Position is defined as the distance from the start codon. Frequency = Count (detected specific nucleic acids at a nucleic acid position)/Coverage (all nucleic acids at a certain nucleic acid position).

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Comparative analysis of growth characteristics

We monitored the growth of the clinical isolates GBS ATCC strain BAA-1138, $\Delta tpk$, $\Delta tpk+pDL278$, and $\Delta tpk+pTPK$. The GBS ATCC BAA-1138, $\Delta tpk+pTPK$, and $\Delta tpk$ supplemented with TPP (500 $\mu$g/L) had similar growth profiles. However, the clinical isolates $\Delta tpk$ and $\Delta tpk+pDL278$ displayed significantly reduced growth compared with that of the GBS ATCC strain BAA-1138, $\Delta tpk$ supplemented with TPP (500 $\mu$g/L), and $\Delta tpk+pTPK$ (Fig 3).

Auxotrophic testing

The clinical isolates and $\Delta tpk$ showed similar colony sizes to those of GBS ATCC strains on sheep blood agar, MHA, and THA containing greater than 50 $\mu$g/L TPP. Additionally, similar to A909, these isolates grew on sheep blood agar around discs containing 20 $\mu$g of TPP. However, on MHA with 5% sheep blood and sheep blood agar around discs with 20 $\mu$g of TPP.
Fig 3. Comparison of growth rate of GBS ATCC strain, recombinant strain and complemented strains.
Growth curves (OD$_{600nm}$) of the clinical isolates (MRY11-004, MRY11-005, and NUBL-2449), GBS ATCC strain (BAA-1138), recombinant strain ($\Delta$tpk), and complemented strains ($\Delta$tpk+pDL278 and $\Delta$tpk+pTPK) at 37˚C in ambient air (A,B). OD$_{600nm}$ = 0–2.5 (A), OD$_{600nm}$ = 0–0.2 (B). Data are presented as averages of five independent experiments. Data are the mean ± standard deviation OD$_{600nm}$.*,**Statistically significant difference: *P < 0.05; **P < 0.01. $\Delta$tpk indicates a recombinant strain based on ATCC BAA-1138, harboring the 276_277insG insertion in tpk, resulting in premature termination at amino acid 103 in thiamin pyrophosphokinase (TPK). pDL278 is a gram-positive and gram-negative shuttle vector. pTPK is a complementation plasmid used to express intact TPK in the $\Delta$tpk strain. $\Delta$tpk+TPP indicates $\Delta$tpk grown in Todd–Hewitt broth containing thiamine pyrophosphate (500 μg/L).

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thymidine, Δtpk showed medium colony sizes (1 mm in diameter), which were smaller than those of GBS ATCC strains (Fig 4). Moreover, THA containing more than 50 μg/ml thymidine showed medium-sized colonies. Although previous reports have shown that the addition of NAD⁺, menadione, thiamine, or hemin results in a normal colony morphology in S. aureus SCVs, the clinical isolates and Δtpk did not show visible growth around discs containing 20 μg of these compounds[15] [21] [22].

While the clinical isolates and Δtpk were able to grow on chocolate II agar, they showed medium colony sizes on BY chocolate agar. A comparison of the ingredients in these agars revealed that chocolate II agar contained 10 mg/L hemoglobin, while BY chocolate agar contained 5% horse blood. Accordingly, the hemoglobin content may explain the growth of the clinical isolates and Δtpk. Supplementation of MHA and THA with 10 mg/mL hemoglobin confirmed this hypothesis; the clinical isolates and Δtpk showed colony sizes similar to those of the ATCC strain.

TEM analyses
BAA-1138 and Δtpk containing TPP (500 μg/L) exhibited regular cell division with single cross walls in dividing cells. However, cell division in MRY11-004 and Δtpk appeared to terminate before the cells could fully separate and show agglutination. Moreover, only MRY11-004 showed heterogeneous cell sizes and cell clusters (Fig 5).

Identification by Rapid ID 32 Strep API and VITEK MS
Although the clinical isolates and BAA-1138 were identified as GBS using the Rapid ID 32 Strep API system, Δtpk cultured on sheep blood was not identified as GBS and was judged as SDSE or GBS. Concerning biological profiles, the clinical isolates showed different trehalose (TRE), pullulan (PUL), sodium pyruvate (VP), and maltose (MAL) phenotypes compared to those of BAA-1138. Moreover, in addition to PUL, VP, and MAL, Δtpk showed different
sucrose (SAC) phenotypes. Therefore, Δtpk was unable to be correctly confirmed as GBS. However, the clinical isolates, Δtpk, and BAA-1138 cultured on sheep blood and chocolate II agar were identified as GBS (99.9% identity) using VITEK MS.

Discussion

We characterized the first clinical PRGBS with less hemolytic activity and auxotrophy for TPP. The G379T substitution in cylK gene and 276_277insG insertion in tpk gene caused the less hemolytic activity and small colony formation, respectively. In previous reports, SCVs with electron transport and thymidine biosynthesis defects have shown both small colony formation and low hemolytic activity due to the deletion of a single gene. However, in this study, small colony formation and reduced hemolytic activity were caused by different genes. The deletion of CylK results in reduced hemolytic activity on horse blood agar and similar growth to that of the parental strain in THB supplemented with 5% yeast extract [32, 33]. This suggests that full-length CylK is required for full hemolytic activity of GBS, and may not be related to cell growth and colony size. The G379T substitution in cylK, resulting in premature termination at amino acid 127 in CylK, was associated with low hemolytic activity. The deletion of amino acids 22 to 115 of CylK have been reported to cause reduced hemolytic activity [32]. Therefore, full-length CylK might be required for full hemolytic activity in GBS. In the TEM analysis, a clinical isolate showed heterogeneous cell sizes and cell clusters; however, there are many candidate loci associated with this heterogeneity based on next-generation sequence data for the three clinical isolates and therefore the underlying causes are unknown.

The Δtpk strain, which exhibited premature termination at amino acid 103 in TPK, showed small colony formation. Moreover, complementation of the strain with a plasmid harboring full-length TPK resulted in similar colony sizes to those of GBS ATCC strains. Therefore, the function of this domain is related to growth. The structure of TPK in yeast bound with TPP
has revealed the locations of the thiamin-binding site and probable catalytic residue [34]. Analysis of GBS TPK using Pfam showed that these sites were conserved. Therefore, because amino acid positions 103–210 in TPK correspond to the thiamine-binding domain, Δtpk and clinical isolates lacking these amino acids might not bind thiamine and accordingly might be unable to catalyze the production of TPP. Moreover, because TPP, a coenzyme form of vitamin B1, is important for the formation of a coenzyme required for central metabolic functions (pyruvate decarboxylase, pyruvate dehydrogenase, α-ketoglutarate dehydrogenase, and transketolase), TPP might be essential for cell growth in bacteria and Δtpk might exhibit growth defects [35]. Moreover, in Schizosaccharomyces pombe, trx3-recombinant strains (low-expressed TPK) grow slowly and show aberrant morphology [36]. The TPK deletion has not been detected in bacterial clinical isolates; therefore, these are the first known clinical isolates with the 276_277insG insertion in the tpd gene.

The three clinical isolates analyzed in this study somewhat resemble previously reported SCVs. However, they also have different characteristics from previously reported SCVs. The three clinical isolates showed slow growth phenotypes on THA, sheep blood agar, and in THB. This phenotype resembles that of previously reported SCVs [15–18, 24]. However, the clinical isolates in this study did not show irregular cell shapes, as observed in other SCVs, when analyzed by TEM [17, 18], nor did they show auxotrophy for menadione, hemine, or thiamine, as did SCVs of S. aureus. The clinical isolates and Δtpk showed auxotrophy for TPP. Moreover, small colony formation was caused by a partial deletion in the TPK, which has not been reported in other SCVs to date. In general, electron transport deficient-SCVs and thymidine biosynthesis deficient-SCVs show an increase in gentamicin and sulfamethoxazole/trimethoprim MIC, respectively. Although the three clinical isolates showed resistance to sulfamethoxazole/trimethoprim (MIC >256 μg/ml), there was no difference observed in the MIC of gentamicin for the clinical isolate MRY11-004 (MIC 32 μg/ml) compared to those of MRY11-004+pTPK (MIC 32 μg/ml, expressing full-length TPK in MRY11-004), Δtpk (MIC 64 μg/ml), and Δtpk +pTPK (MIC 64 μg/ml). Furthermore, SCVs are typically isolated from patients undergoing long-term antibiotic therapy, and can cause latent or recurrent infections [15]. For the three clinical isolates, sulbactam/ampicillin therapy was documented from October 25 to October 31 and from November 28 to December 2 in 2010. Thereafter, two of the clinical isolates (MRY11-004 and MRY11-005) were isolated from sputum and blood on January 4, 2011. Moreover, piperacillin/tazobactam and levofloxacin therapy was documented from October 22 to October 25 and from November 27 to November 31 in 2010, respectively. The clinical isolate NUBL-2449 was isolated from sputum on November 14, 2012 [14]. Therefore, no cases of long-term antibiotic therapy were documented in the two patients, and this likely does not explain the small colony formation phenotype. On the other hand, these β-lactam and quinolone therapies might have selected for multidrug resistance and/or PRGBS. Selection for PRGBS by long-term β-lactam therapy has been observed [9]. Additionally, conventional identification systems (e.g., the API Rapid system) have failed to identify E. coli, E. faecium, and E. faecalis SCVs correctly [18, 24, 25]. Although the clinical isolates cultured on chocolate II agar and sheep blood agar could be detected using the Rapid ID 32 Strep API system, Δtpk on sheep blood agar yielded no definitive results owing to an inconsistent biological phenotype. Furthermore, several biochemical phenotypes were different from those of the parental strain and GBS ATCC strains. Therefore, clinical isolates of GBS with deletions in the tpd gene might be misidentified in the future. In contrast to conventional identification systems based on biochemical characteristics, the clinical isolates and Δtpk were correctly identified by VITEK MS (99.9% identity). Moreover, E. faecium SCVs were precisely identified by VITEK MS [18]. These findings suggest that MALDI-TOF MS is a powerful tool for the identification of strains with similar atypical phenotypes.
In conclusion, we characterized the first PRGBS clinical isolates with reduced hemolytic activity and auxotrophy for TPP. Moreover, we found that VITEK MS correctly identified GBS with a deletion in TPK. In this study, we revealed the causes of small colony formation with less hemolytic activity and characterized clinical isolates in detail. Because these clinical isolates may be misclassified using routine bacterial tests and are multidrug resistant, they represent a potential public health concern.

Supporting information

S1 Fig. Comparison of hemolytic activity of clinical isolates, recombinant strains and complemented strains. Clinical isolates (MRY11-004, MRY11-005 and NUBL-2449), recombinant strains ($\Delta$cyIK and $\Delta$tpk) and complemented strains ($\Delta$cyIK+pDL278, $\Delta$cyIK+pCylIK, $\Delta$tpk+pTPK and $\Delta$tpk+pTPK) on Mueller Hinton Agar with 5% Sheep Blood. $\Delta$cyIK indicates a recombinant strain based on ATCC BAA-611, harbouring the G379T substitution in cyIK, resulting in premature termination at amino acid 127 in CylK. pDL278 is a Gram-positive and gram-negative shuttle vector. $\Delta$cyIK+pCylIK indicates a complementation plasmid to express intact TPK in $\Delta$cyIK strain. $\Delta$tpk indicates a recombinant strain based on ATCC BAA-1138, harbouring the 276_277insG insertion in tpk, resulting in premature termination at amino acid 103 in thiamin pyrophosphokinase. $\Delta$tpk+pTPK indicates a complementation plasmid to express intact TPK in $\Delta$tpk strain. A: MRY11-004 B: MRY11-005 C: NUBL-2449 D: 2603 V/R E; $\Delta$cyIK F: $\Delta$cyIK+pDL278 G: $\Delta$cyIK+CylIK H: $\Delta$cyIK+pCylIK I: A909 J: $\Delta$tpk K: $\Delta$tpk+pDL278 L; $\Delta$tpk+pTPK.

(TIFF)

S1 Table. Primers used for PCR amplification in this study.

(DOCX)

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