It Takes Two to Tango: Combining Conventional Culture With Molecular Diagnostics Enhances Accuracy of Streptococcus pneumoniae Detection and Pneumococcal Serogroup/Serotype Determination in Carriage

Willem R. Miellet1,2, Janieke van Veldhuizen2, David Litt3, Rob Mariman2, Aliene J. Wijmenga-Monsuur2, Paul Badoux4, Tessa Nieuwenhuijsen1, Rebecca Thombre3, Sanaa Mayet1, Seyi Eletu3, Carmen Sheppard4, Marianne Alice van Houten5, Nynke Y. Rots2, Elizabeth Miller4, Norman K. Fry5,6, Elisabeth A. M. Sanders1,2 and Krzysztof Trzciński1*

1 Department of Pediatric Immunology and Infectious Diseases, Wilhelmina Children’s Hospital, University Medical Center Utrecht (UMCU), Utrecht, Netherlands, 2 Centre for Infectious Disease Control, National Institute for Public Health and the Environment (RIVM), Bilthoven, Netherlands, 3 Respiratory and Vaccine Preventable Bacterial Reference Unit (RVPBRU), Public Health England – National Infection Service, London, United Kingdom, 4 Regional Laboratory of Public Health (Streeklab) Haarlem, Haarlem, Netherlands, 5 Spaarne Gasthuis, Haarlem, Netherlands, 6 Immunisation and Countermeasures Division, Public Health England (PHE) – National Infection Service, London, United Kingdom

Background: The specificity of molecular methods for the detection of Streptococcus pneumoniae carriage is under debate. We propose a procedure for carriage surveillance and vaccine impact studies that increases the accuracy of molecular detection of live pneumococci in polymicrobial respiratory samples.

Methods: Culture and qPCR methods were applied to detect pneumococcus and pneumococcal serotypes in 1,549 nasopharyngeal samples collected in the Netherlands (n = 972) and England (n = 577) from 946 toddlers and 603 adults, and in paired oropharyngeal samples collected exclusively from 319 Dutch adults. Samples with no live pneumococci isolated at primary diagnostic culture yet generating signal specific for pneumococcus in qPCRs were re-examined with a second, qPCR-guided culture. Optimal Cq cut-offs for positivity in qPCRs were determined via receiver operating characteristic (ROC) curve analysis using isolation of live pneumococci from the primary and qPCR-guided cultures as reference.

Results: Detection of pneumococcus and pneumococcal serotypes with qPCRs in cultured (culture-enriched) nasopharyngeal samples exhibited near-perfect agreement with conventional culture (Cohen’s kappa: 0.95). Molecular methods displayed increased sensitivity of detection for multiple serotype carriage, and implementation of qPCR-guided culturing significantly increased the proportion of nasopharyngeal and
Miellet et al. Two to Tango

INTRODUCTION

Streptococcus pneumoniae (pneumococcus) is the most common etiological agent of invasive bacterial disease (O’Brien et al., 2009) and of community-acquired pneumonia of bacterial etiology (Welte et al., 2012). Despite being vaccine-preventable, pneumococcal disease remains among the leading causes of death in childhood (World Health Organization, 2019) as available vaccines target only a various subsets of only 24 from ca. 100 of the known serotypes. Due to high pneumococcal carriage rates (Bogaert et al., 2004; Satzke et al., 2013) children are considered the primary reservoir of pneumococcus and the main drivers of transmission and infections in any population (Wyllie et al., 2016a, 2020; Flasche et al., 2020). Children are also the primary group targeted with pneumococcal vaccination (Berical et al., 2016).

With pneumococcal vaccines protecting not only against disease but also against colonization, carriage is now an accepted endpoint in vaccination studies (Dagan et al., 1997; van Gils et al., 2009; Flasche et al., 2011; Auranen et al., 2013). Following the introduction of pneumococcal conjugate vaccines (PCV), of which 10-valent (PHiD-CV, GSK), 13-valent (Prevnar 13, Pfizer), 15-valent (Vaxneuvance, MSD), and 20-valent (Prevnar 20, Pfizer) are currently marketed, epidemiological surveillance of carriage became an essential tool for monitoring the direct and indirect (herd protection) effects of vaccination. Surveillance of carriage is also used to detect the emergence and track expansion of non-vaccine serotypes, the phenomenon described as vaccine-induced serotype replacement (Weinberger et al., 2011). Finally, carriage studies are instrumental in monitoring serotype-associated invasiveness (Sleeman et al., 2006).

To understand the trends in pneumococcal epidemiology and guide strategies for PCV use, it is critical to establish methodology for S. pneumoniae carriage detection that is sufficiently sensitive across all ages (Slotved, 2016; Weinberger et al., 2016). However, conventional culture, which is considered to be the gold standard method in carriage detection, is not suitable for detection of multiple serotype carriage (Valente et al., 2013) and lacks the sensitivity when used to detect S. pneumoniae in age groups other than children (Krone et al., 2014). There is also evidence that testing solely a single site within the upper airways reduces sensitivity of carriage detection (Korona-Glowniak et al., 2011; Trzcinski et al., 2013; Almeida et al., 2020).

Molecular methods have largely improved the sensitivity of S. pneumoniae detection and have made multiple serotype carriage detection feasible (Turner et al., 2011; Satzke et al., 2013). Using assays developed by our groups and others, we have tested the serotype composition of respiratory samples from children and adults and have demonstrated an under-detection of S. pneumoniae and of individual pneumococcal serotypes by culture when compared with molecular methods (Trzcinski et al., 2013; Wyllie et al., 2014, 2016a,b; Krone et al., 2015; Miellet et al., 2020). However, some caution that molecular methods exhibit poor specificity due to the presence of pneumococcal genes among commensal streptococci (Carvalho Mda et al., 2012, 2013; Boelsen et al., 2020). It could also be argued that molecular detection is unable to discriminate between live bacteria and presence of relic DNA (Lennon et al., 2018).

Here, we outline the protocol that combines conventional culturing with pneumococcus-specific qPCRs and employs statistical procedures to interpret the molecular results and enhances the specificity of the molecular methods. We show that molecular methods applied to nasopharyngeal samples demonstrate near-perfect agreement with primary culture and yet the sensitivity of S. pneumoniae carriage surveillance can be greatly enhanced by complementing conventional culture with qPCRs and vice versa. We also show that testing nasopharyngeal....
samples alone leads to underestimation of pneumococcal carriage in adults.

MATERIALS AND METHODS

Study Design and Ethics Statement

Pneumococcal carriage was investigated in two cross-sectional prospective observational studies conducted in 2015/2016 in community-dwelling individuals in the Netherlands (Vissers et al., 2018) and in England (Southern et al., 2018). The study conducted in the Netherlands was approved by the Medical Ethics Committee Noord Holland (NTR5405 on). The study conducted in England was approved by the NHS Health Research Authority and the London Fulham Research Ethics Committee (reference 15/LO/0458) and was registered on clinicaltrials.gov (reference NCT02522546). In both studies written informed consent was obtained from the parent or guardian of every participating child and adults provided written consent for their own participation. Both studies were conducted in accordance with Good Clinical Practice and the Declaration of Helsinki.

Sample Collection and Laboratory Processing

Respiratory samples were collected in the Netherlands between October 2015 and March 2016 in the study on carriage of respiratory bacterial pathogens coordinated by National Institute of Public Health and the Environment. Nasopharyngeal samples were collected in children aged 24 months (±1 month) vaccinated with PHiD-CV according to “2 primary + 1 booster” (2p + 1b) dose schedule, children aged 44–49 months vaccinated with PHiD-CV in 3p + 1b schedule, and parents of the 24-month-old children (one parent per child). Oropharyngeal swabs were also collected from Dutch adults (Watt et al., 2004; Vissers et al., 2018). The English study took place between July 2015 and June 2016 and was conducted by the National Vaccine Evaluation Consortium which included Public Health England (Southern et al., 2018). In England, nasopharyngeal samples were collected in children aged 1–5 years vaccinated with PCV13 according to a 2p + 1b schedule and in their household contacts (adults aged > 20 years).

All samples were obtained according to World Health Organization standard procedures (Satzke et al., 2013). Immediately after sampling, swabs in the Netherlands were placed in liquid Amies transport medium (ESwab 482C, Copan, Brescia, Italy) and within 8 h transported to the diagnostic laboratory. In England swabs were placed in skim-milk, tryptone, glucose and glycerol (STGG) broth and delivered to a diagnostic laboratory within 48 h (Table 1).

Conventional Culture

Thirty-five microliters (Netherlands) or 50 µl (England) of specimen was used as an inoculum in detection of pneumococci by conventional culture (primary culture). In the Netherlands samples were cultured on SB7-Gent agar selective for streptococci (BA-GENT, Oxoid, Badhoeve Dorp, The Netherlands) and on non-selective Columbia blood agar (CBA, Oxoid). In England samples were cultured on Streptococcus-selective Blood Agar (COBA, Oxoid, Basinstoke, United Kingdom) and CBA (Oxoid). Remaining transport media were stored frozen at −70°C. In the Netherlands, samples were aliquoted into 200 µl volumes, and a single aliquot was supplemented with glycerol (10% v/v final concentration) prior to freezing.

After overnight incubation at 37°C and 5% CO₂, cultures were screened for pneumococcus-like colonies to be re-cultured. Once screened, in the Netherlands all colony growth was harvested from BA-GENT plate into 10% glycerol in brain heart infusion broth (Oxoid) (Miellet et al., 2020). In England, all colony growth was harvested from any plates containing alpha-hemolytic colonies into PBS, centrifuged, the supernatant removed, and any growth stored as a pellet (Southern et al., 2018). These samples were considered culture-enriched (CE) for pneumococci and stored at −80°C. Cultured strains (one per sample, but more if distinct pneumococcal morphotypes were apparent) were serotyped by Quellung method in the Netherlands (Vissers et al., 2018) and with the previously described PneumoCaT bioinformatic pipeline in England that was supplemented with slide agglutination when required (Kapatai et al., 2016; Southern et al., 2018). Since non-typeable thus unencapsulated pneumococci are not the target of pneumococcal vaccines and are also avirulent, cultured non-typeable strains were not considered in the analysis. Non-typeable pneumococci are negative for piaB (Whalan et al., 2006; Trzcinski et al., 2013; Wyllie et al., 2017) and their inclusion would complicate a comparison of detection methods.

Molecular Detection of Streptococcus pneumoniae

Molecular detection of S. pneumoniae in respiratory samples was conducted as described previously (Miellet et al., 2020). For this, Amies medium, STGG medium, and culture-enriched samples were shipped on dry ice from primary diagnostic sites to the study central laboratory in the Netherlands. There, pellets of culture-enriched samples collected in England were reconstituted to the 200 µl that matched original volume of an aliquot. Next, nucleic acids were extracted from 100 µl of the transport media using the DNeasy Blood and Tissue Kit (Qiagen) and eluted into 200 µl. These extractions represented minimally processed samples. For culture-enriched samples, 100 µl of a bacterial growth harvest was centrifuged for 2 min at 14,000 × g, the pellet was resuspended with 90 µl of TE buffer [20 mM Tris-HCl (pH 8.0), 2 mM EDTA] and incubated for 15 min at 95°C. Next, 90 µl of lysis buffer [20 mM Tris-HCl (pH 8.0), 2 mM EDTA, 2.4% Triton X-100 and 40 mg/ml lysozyme] was added, and the samples were processed as above. Pneumococcal DNA was detected via singleplex qPCR (Supplementary Table 1) using primers and probe targeting sequences (Supplementary Table 2) within genes coding for the pneumococcal iron uptake ABC transporter lipoprotein PiaB (Trzcinski et al., 2013), and for the major pneumococcal autolysin LytA (Carvalho Mda et al., 2007).
by testing 5.5 μl DNA from minimally processed or 1.0 μl of culture-enriched samples within a total qPCR reaction volume of 12.5 μl.

**Molecular Detection of Pneumococcal Serotypes**

DNA extracted from culture-enriched harvests was used to determine the serotype composition of nasopharyngeal and oropharyngeal samples using a panel of primers and probes (Azzari et al., 2010, 2012; Pimenta et al., 2013), targeting all serotypes covered by pneumococcal vaccines available either in the Netherlands or in England at the time of sample collection (PHID-CV, PCV13 and 23-valent polysaccharide vaccine, PPV23, Merck Sharp Dohme) except for serotype 2. The panel also covered selected non-vaccine serotypes (serotypes 6C, 6D, 7A, 10B, 12A, 12B, 15A, 15C, 15F, 16F, 21, 22A, 23A, 23B, 33A, 34, 35B, 35F, 37, and 38). However, with certain assays targeting more than one serotype it was not possible to distinguish between 35B, 35F, 37, and 38. For a particular serotype/serogroup were tested (Wyllie et al., 2016a) and the remaining samples were individually tested in qPCR for piaB and lytA and confirmed to be S. pneumoniae based on susceptibility to optochin.

**Statistical Analysis**

Data analysis was performed in GraphPad Prism software version 9.1.0 and R version 4.1.0. Receiver operating characteristic (ROC) curve analysis was performed using the “cutpoints” R package to validate qPCR results with culture results (primary culture plus qPCR-guided culture). Maximum Youden index values, the sum of sensitivity and specificity minus one, were estimated via bootstrapping (n = 1,000) on piaB and lytA qPCR data from respiratory samples to determine optimal cut-off values for qPCR detection (Nutz et al., 2011). Two-way mixed effects intraclass correlations (ICCs) (Koo and Li, 2016) and Bland-Altman plots (Bland and Altman, 1986) were used to evaluate agreement for quantitative results using the “irr” and “bland” R packages, respectively. Carriage rates were compared with McNemar’s test unless otherwise stated. Estimates for accuracy of diagnostic tests between methods (subgroups) were compared with a test of interaction (Altman and Bland, 2003). A p-value of < 0.05 was considered significant. Diagnostic test parameters (predictive values, sensitivity, and specificity) were calculated using an in-house made R function, and 95% confidence intervals were calculated with Wilson-Brown score. Cohen’s kappa, a measure of chance-corrected agreement, and its 95% confidence interval was calculated as described by McHugh (2012) and ratios of ≤ 0, 0.01–0.20, 0.21–0.40, 0.41–0.60, 0.61–0.80, and > 0.81 were interpreted as exhibiting poor, slight, fair, moderate, substantial, and near-perfect agreement, respectively (Landis and Koch, 1977). For comparison between serotyping by culture and by qPCR, analysis was limited to qPCR-targeted serotypes. For serogroup-specific qPCR assays a result was considered congruent when a serogroup was detected in qPCR that matched the serogroup of the serotype detected by

---

**TABLE 1** | Methodological differences between Streptococcus pneumoniae carriage studies conducted in the Netherlands and in England.

| Processing step                  | The Netherlands                                      | England                                      |
|----------------------------------|------------------------------------------------------|----------------------------------------------|
| Transport medium                 | 1 ml Amies                                           | 1 ml STGG                                   |
| Transport temperature            | Ambient                                              | 2–8°C                                       |
| Duration of transport            | < 8 h                                                | < 48 h                                      |
| Arrival to the diagnostic lab    | Culturing on receipt                                 | Storage at −80°C                             |
| Culture medium                   | BA-GENT and CBA                                      | COBA and CBA                                |
| Inoculum volume                  | 35 μl                                                | 50 μl                                       |
| Identification of S. pneumoniae  | Optochin-susceptibility and bile solubility tests    | Bioinformatics-based method applied to bacterial DNA |
| Minimally processed (MP) sample  | An aliquot of Amies stored at −80°C                  | An aliquot of skim milk tryptone glucose and glycerine broth medium (STGG) stored at −80°C |
| Culture-enriched (CE) sample     | Whole BA-GENT plate colony growth washed with 2.1 ml of BHI with 10% glycerol | Swab of a-hemolytic colonies from COBA (first choice) or CBA, and into 1 ml of STGG |

STGG, skim milk tryptone glucose and glycerine broth medium; CBA, Columbia blood agar plate; BA-GENT, blood agar plate gentamycin; COBA, blood agar plate with Streptococcus selective medium.

---

2 https://github.com/wmiellet/test-comparison-R
culture. Expected frequencies of multiple serotype carriage were calculated squaring observed pneumococcal prevalence rates and expected and observed multiple serotype frequencies were compared using a one-proportion Z-test.

**Assessment of Method’s Inter-Laboratory Reproducibility**

One-hundred seventy-six nasopharyngeal samples collected in England and \( n = 59 \) oropharyngeal samples collected in the Netherlands have been selected to evaluate the inter-laboratory reproducibility of molecular methods. For this, an aliquot of minimally processed STGG nasopharyngeal sample, a fresh preparation of culture enriched nasopharyngeal sample from a subset of the English study samples and culture-enriched oropharyngeal samples from a subset of the Dutch study samples were tested at the study site in England, as described above. Results for paired samples were compared between centers by calculating the percent agreement and Cohen's kappa. Quantitative results of both laboratories were also compared by calculating an intraclass correlation coefficient (ICC) and by comparing results in Bland-Altman plots. Carriage rates between both laboratories were compared using Cohen’s kappa.

**RESULTS**

We evaluated the performance of conventional and molecular methods in a new protocol (Figure 1) and applied it to detect *S. pneumoniae* and pneumococcal serotypes in 1,549 nasopharyngeal samples collected from 946 children aged 1–5 years (\( n = 653 \) in the Netherlands and \( n = 293 \) in England) and from 603 adults (\( n = 319 \) in the Netherlands and \( n = 284 \) in England), and on oropharyngeal samples from 319 adults in the Netherlands.

Detection of *Streptococcus pneumoniae*

When analyzing culturing results without the addition of qPCR-guided culturing, live *S. pneumoniae* was isolated from 29% (445/1,549) of primary cultures of nasopharyngeal swabs (Figure 2A), with significantly higher fraction of samples from children (45% or 422/946) compared with adults (4% or 23/603) positive for pneumococcus (Pearson’s chi-square, \( p < 0.0001 \)). Among 319 adults sampled in the Netherlands (Figure 2B), live pneumococcus was significantly more often cultured from nasopharyngeal (\( n = 15 \) or 5%) compared with oropharyngeal (\( n = 3 \) or 1%) samples (\( p < 0.01 \)).

Next, we conducted qPCRs on minimally processed and culture-enriched samples. To enhance the specificity of detection we used a “two-to-tango” approach by quantifying *piaB* and *lytA* genes and considering a sample to be positive for *S. pneumoniae* when both targets were detected. When applying an arbitrary quantification cycle (\( \Delta \Delta C_q \)) of \( < 40 \) \( C_q \) as criteria for positivity altogether 583 nasopharyngeal swabs (38% of 1,549) were identified as positive for pneumococcus (Figures 3A,B). The fraction of positive samples was significantly higher among minimally processed compared with culture-enriched swabs (561 or 36% vs. 479 or 31% of 1,549; \( p < 0.0001 \)). In line with...
FIGURE 2 | (A) Matrix layout for observed intersections of the *Streptococcus pneumoniae* detection procedures applied to \( n = 1,549 \) nasopharyngeal (NP) samples, sorted by size. (B) Matrix layout for observed intersections of the *S. pneumoniae* detection procedures applied to nasopharyngeal and oropharyngeal (OP) samples from \( n = 319 \) adults sampled in the Netherlands, sorted by size. Circles in the matrix indicate sets that are part of the intersection, with nasopharyngeal and oropharyngeal samples colored in green and blue, respectively. The bar diagram displaying the intersection size is colored using the same scheme, with orange indicating an intersection that represents positivity in both sample types. MP stands for minimally processed and CE for culture-enriched.
results of primary diagnostic culture, here too the proportion of positive samples was higher in children compared with adults (Pearson’s chi-square, \( p < 0.0001 \)) whenever nasopharyngeal swabs were tested minimally processed (56% or 528/946 vs. 5% or 33/603) or culture-enriched (48% or 454/946 vs. 4% or 25/603). However, contrary to results of primary diagnostic culture, among adults significantly larger proportions of oropharyngeal compared with nasopharyngeal samples \( ( p < 0.0001 ) \) have been classified as positive for pneumococcus with qPCR whenever tested minimally processed (26% or 83/319 vs. 6% or 19/319) or culture-enriched (18% or 58/319 vs. 6% or 18/319).

Next, samples negative for pneumococcus in primary diagnostic cultures yet generating signal in qPCRs for \( piaB \) and \( lytA \) were cultured again. This qPCR-guided culture effort increased the number of samples from which live pneumococci was cultured, in relatively small albeit significant increase of 3% (from 445 vs. 460 of 1,549; \( p < 0.0001 \)) for nasopharyngeal (Figures 2A, 3A,B) and 666% increase (from 3 to 23 of 319; \( p < 0.0001 \) ) for oropharyngeal samples (Figures 2B, 3E,F).

To validate qPCR results using culture and to further enhance the specificity of qPCR methods for the detection of live pneumococci we performed receiver operating characteristic (ROC) curve analysis and identified \( C_q \) cut-off values that yielded maximum Youden indices (Table 2). Reliability of positive qPCR results was assessed with Bland-Altman plots (Figures 3C,D,G,H and Supplementary Table 3) and by calculating the intraclass correlation coefficient between \( piaB \) and \( lytA \). Samples with \( C_q \) values below the ROC-derived \( C_q \) criterium \( (\text{ROC}_d C_q) \) demonstrated excellent agreement between \( piaB \) and \( lytA \) while samples with a \( C_q \) value above the \( \text{ROC}_d C_q \) threshold displayed poor agreement (Supplementary Table 3). Using the \( \text{ROC}_d C_q \) criterium we classified 33% (512/1,549) of minimally processed nasopharyngeal samples as positive for pneumococcus (Figure 2A). It included 52% (489/946) samples from children and 4% (23/603) from adults. For culture-enriched samples application of \( \text{ROC}_d C_q \) resulted in 30% (469/1,549) classified as positive including 47% (447/946) samples from children and 4% (22/603) from adults (Figure 2A). With \( \text{ROC}_d C_q \), the adults in the Netherlands were again significantly more often positive for pneumococcus in oropharyngeal compared to nasopharyngeal samples whenever tested minimally processed (19% or 59/319 vs. 5% or 15/319; \( p < 0.0001 \)) or culture-enriched (12% or 39/319 vs. 5% or 15/319; \( p < 0.01 \) ) (Figure 2B).

For culture-enriched nasopharyngeal and oropharyngeal samples of molecular detection in culture-enriched oropharyngeal samples \( ( p < 0.05 ) \) and also when compared with molecular detection with either culture-enriched sample of either type \( ( p < 0.0001 ) \). To this extent, molecular detection in culture-enriched oropharyngeal plus nasopharyngeal samples displayed the highest agreement to the reference out of all other evaluated approaches (Table 4).

Comparison of Serotype Carriage Detection Methods

With conventional culture (without qPCR-guided additional culture), 29% (445/1,549) of nasopharyngeal samples including 45% (422/946) from children and 4% (23/603) from adults were positive for a serotype (non-typeable pneumococci excluded), as already described above. Next, we assessed the accuracy of molecular methods when applied to detect carriage of pneumococcal serotypes. Supplementary Figure 1 depicts results of serotype detection in culture-enriched samples tested with serotype-specific qPCR assays. None of the samples generated any signal in qPCRs targeting serotype 1, and 23F and serogroup 18, nor was positive for any of these serotypes by culture. The assays targeting serotypes 4, 5 showed a lack of specificity and the assay targeting serotype 17F showed lack of sensitivity when applied to both, nasopharyngeal and to oropharyngeal samples. Results of these three assays were excluded from analysis.

Four-hundred and twenty-three nasopharyngeal samples were positive for one or more serotypes targeted in qPCRs either by culture \( ( n = 393 \) samples) or with molecular methods \( ( n = 411 \) samples) and applying \( \text{ROC}_d C_q \) criterium for positivity for a
Miellet et al. Two to Tango

FIGURE 3 | Scatter plots (A,B,E,F) displaying correlation between cycle threshold (Cq) from qPCR assays targeting the Streptococcus pneumoniae genes piaB and lytA for minimally processed (A,E) and culture-enriched (B,F) nasopharyngeal (A,B) and oropharyngeal (E,F) samples followed by the Bland-Altman plots (C,D,G,H) displaying agreement between piaB and lytA for samples positive with minimally processed (C,G) and culture-enriched (D,H) NASOPR criteria in nasopharyngeal (C,D) and oropharyngeal (G,H) samples. The shaded gray areas in (G,H) mark the 95% confidence interval of the upper limit of agreement, bias and lower limit of agreement. The Bland-Altman plots show the absence of systematic bias between piaB and lytA because the line of equality (solid line) is within the limits of agreement. Bias between piaB and lytA is more often observed in high Cq samples. Primary culture-positive and qPCR-guided culture-positive samples are indicated with red and green dots, respectively. Culture-negative samples are indicated with blue dots. In (A,B,E,F) black dashed lines indicate the arbitrary Cq criterium (<40 Cq), green dashed lines indicate ROCR-Q criteria derived by validating qPCR results with primary and qPCR-guided cultures results. Red dashed lines in (B) indicate ROCR-Q criteria derived by validating qPCR results with primary culture alone. In (A) red lines overlap with green lines and only the latter are shown.
TABLE 2 | Optimal qPCR cycle threshold \( C_q \) and corresponding parameters for Streptococcus pneumoniae carriage detection in nasopharyngeal \((n = 1,549)\) and oropharyngeal \((n = 319)\) samples.

| Parameter                  | Sample Reference | Optimal threshold \( piaB \) \((95\% CI)\) | Youden index \( piaB \) | Sensitivity \( piaB \) | Specificity \( piaB \) | Optimal threshold \( lytA \) \((95\% CI)\) | Youden index \( lytA \) | Sensitivity \( lytA \) | Specificity \( lytA \) |
|----------------------------|------------------|---------------------------------------------|--------------------------|--------------------------|--------------------------|---------------------------------------------|--------------------------|--------------------------|--------------------------|
| Minimally processed        | NP Primary cultures | 37.13 \((36.49–37.76)\) | 0.85                      | 0.95                      | 0.90                      | 37.47 \((36.44–38.39)\) | 0.82                      | 0.94                      | 0.87                      |
|                           | NP Primary culture and reculture | 37.21 \((36.51–38.46)\) | 0.85                      | 0.95                      | 0.90                      | 37.29 \((36.52–38.61)\) | 0.83                      | 0.94                      | 0.89                      |
|                           | OP Primary culture and reculture | 38.73 \((37.29–39.39)\) | 0.69                      | 0.91                      | 0.78                      | 37.47 \((36.52–37.66)\) | 0.69                      | 0.87                      | 0.82                      |
| Culture-enriched           | NP Primary cultures | 28.71 \((26.59–29.69)\) | 0.94                      | 0.98                      | 0.97                      | 29.19 \((27.41–30.25)\) | 0.93                      | 0.98                      | 0.95                      |
|                           | NP Primary culture and reculture | 30.26 \((29.91–30.47)\) | 0.96                      | 0.98                      | 0.98                      | 29.99 \((28.91–30.47)\) | 0.94                      | 0.98                      | 0.96                      |
|                           | OP Primary culture and reculture | 29.78 \((28.84–29.93)\) | 0.91                      | 0.96                      | 0.95                      | 30.73 \((29.42–30.81)\) | 0.85                      | 0.91                      | 0.94                      |

Results from qPCR were validated in a receiver operating characteristic curve analysis with culture as reference. With S. pneumoniae isolated from only three samples there was insufficient statistical power to perform the receiver operating characteristic curve analysis on oropharyngeal samples with primary cultures as a reference.

TABLE 3 | The accuracy of Streptococcus pneumoniae detection in \(n = 1,549\) nasopharyngeal samples from children \((n = 948)\) and adults \((n = 603)\) tested using conventional diagnostic culture and molecular methods applied to DNA extracted from minimally processed and culture-enriched samples and applying ROC\(_{C_q}\) thresholds for a sample positivity in qPCRs.

| Method                     | Study population  | Percent \((n)\) of positive samples \(95\% CI\) | PPV \%(95\% CI) | NPV \%(95\% CI) | Sensitivity \%(95\% CI) | Specificity \%(95\% CI) | Concordance \%(95\% CI) | \(\kappa\) \((95\% CI)\) |
|----------------------------|-------------------|-----------------------------------------------|----------------|----------------|--------------------------|--------------------------|------------------------|------------------|
| Primary culture            | All               | 28.7 \((445)\) \((26.5–31.0)\) | 100            | 98.6           | 96.7                      | 100                      | 99.0                   | 0.98              |
|                           | Children          | 44.6 \((422)\) \((41.5–47.8)\) | 100            | 97.1           | 96.6                      | 100                      | 98.4                   | 0.97              |
|                           | Adults            | 3.8 \((23)\) \((2.6–5.7)\) | 100            | 99.3           | 100                      | 100                      | 100                   | 1                |
| qPCRs on minimally         | All               | 33.1 \((512)\) \((30.8–35.4)\) | 83.2           | 96.7           | 92.6                      | 92.1                     | 92.3                   | 0.82              |
| processed samples          | Children          | 51.7 \((489)\) \((48.5–54.9)\) | 83.6           | 93.9           | 93.6                      | 84.3                     | 88.6                   | 0.77              |
|                           | Adults            | 3.8 \((23)\) \((2.6–5.7)\) | 73.9           | 99.0           | 99.0                      | 98.0                     | 0.73                   | 1                |
| qPCRs on culture-enriched  | All               | 30.3 \((469)\) \((28.0–32.6)\) | 95.7           | 99.0           | 97.6                      | 98.2                     | 98.0                   | 0.95              |
| samples                    | Children          | 47.3 \((447)\) \((44.1–50.4)\) | 96.0           | 98.4           | 98.2                      | 96.5                     | 97.3                   | 0.94              |
|                           | Adults            | 3.7 \((22)\) \((2.4–5.5)\) | 90.9           | 99.5           | 87.0                      | 99.7                     | 99.2                   | 0.88              |

Measures of diagnostic accuracy were calculated by comparing the number of detected samples positive per method with the number of \(n = 460\) for all individuals, and \(n = 437\) and \(n = 23\) for nasopharyngeal samples from children and adults, respectively. Individuals positive for S. pneumoniae based on isolation of live pneumococcus either from the primary diagnostic or qPCR-guided culture. PPV, positive predictive value; NPV, negative predictive value; 95\%CI, 95\% confidence interval; \(\kappa\), Cohen’s kappa where \(\leq 0\), 0.01–0.20, 0.21–0.40, 0.41–0.60, 0.61–0.80, > 0.81 are interpreted as poor agreement, slight, fair, moderate, substantial, and almost perfect agreement, respectively.

serotype. It included 42% \((395/946)\) samples from children and 3% \((16/603)\) from adults. Altogether, there were \(n = 479\) serotypes carriage events detected by testing nasopharyngeal samples with either conventional culture or molecular methods \((Supplementary Table 5)\). For serotypes targeted by qPCRs detected in nasopharyngeal samples the results of molecular detection displayed excellent agreement \(ICC 0.93, 95\% CI 0.92–0.94\) with \(piaB\) and \(lytA\) \(C_q\) \((Figure 4)\) and almost perfect agreement \(Cohen’s kappa > 0.81\) with isolation of live strain of a particular serotype from nasopharyngeal swab \((Figure 5)\). Also, results of serotype detection in oropharyngeal samples from adults displayed excellent agreement with \(piaB\) and \(lytA\) \(C_q\) \((ICC 0.96, 95\% CI 0.92–0.98)\) \((Supplementary Figure 2)\). Finally, there was near-perfect agreement between overall serotypes carriage...
events detected by qPCR compared with detected by culture (Table 5). Multiple-serotype carriage events were significantly more often detected using molecular methods compared with culture (p < 0.0001). Furthermore, despite a limited number of different serotypes tested by qPCR, observed multiple-serotype carriage frequencies were not significantly different from expected frequencies based on molecular detection on culture-enriched NP samples (11% vs. 9%, respectively; one proportion Z-test, p = 0.2669) unlike detection of multiple-serotype carriage by culture which significantly underestimated expected frequencies (2% vs. 9%, respectively; one-proportion Z-test, p < 0.0001).

Interlaboratory Reproducibility of Molecular Methods

To evaluate the reproducibility of molecular methods and to assess the agreement in laboratory results between the Netherlands and England we processed a subset of culture-enriched samples from both countries in both laboratories. Results for piaB and lytA qPCRs demonstrated good reliability between both laboratories (Supplementary Table 6). We observed near-perfect agreement identifying culture-enriched samples as positive for pneumococcus with molecular methods, and substantial agreement for minimally processed nasopharyngeal samples (Supplementary Table 7). For culture-enriched nasopharyngeal samples we evaluated agreement between both laboratories for serotype carriage detection by qPCR. Overall, near-perfect agreement was observed (Cohen’s kappa 0.82, 95% CI 0.74–0.90).

DISCUSSION

In the current study we have demonstrated that molecular methods exhibit near-perfect agreement to conventional culture in the detection and serotyping of *S. pneumoniae* in children and adults when a nasopharyngeal swab is the only sample tested. Furthermore, we have observed increased sensitivity of *S. pneumoniae* carriage detection among adults by testing oropharyngeal samples with molecular methods and conducting qPCR-guided culturing. We highlight several statistical procedures that can be used to evaluate the reliability of molecular results and enhance the specificity of molecular methods for the detection of live *S. pneumoniae*.

The current gold standard method for carriage detection is the isolation of live pneumococci from cultures of deep trans-nasal nasopharyngeal swab, in adults complemented with culture of a swab collected trans-orally (OP) (Satzke et al., 2013). However, the gold standard lacks sensitivity in case of low-density carriage or when applied to poly-microbial samples in which *S. pneumoniae* is not a dominant bacterium and it does not allow the detection of co-carriage of pneumococcal strains (Heffron, 1939; Trzcinski et al., 2013; Wyllie et al., 2014, 2016a; Almeida et al., 2020). Since the density of pneumococcal carriage episodes in adults is much lower...
FIGURE 4 | Scatter plots (A–C) displaying correlation between cycle threshold ($C_q$) from real-time PCR (qPCR) assays targeting the Streptococcus pneumoniae piaB and serotype/serogroup specific signal detected with qPCR for nasopharyngeal samples classified positive for S. pneumoniae according to ROC$_Cq$ criterium (green dashed lines). Bland-Altman plot (D) displaying agreement between piaB and dominant serotype detected by qPCR. Each dot represents an individual serotype carriage event detected by qPCR in culture-enriched nasopharyngeal sample (A–C). Dots in (A) depict serotypes detected with qPCR. Dots are color-coded according to serotypes/serogroup targeted in an assay (see legend). Color dots in (B) depict subdominant serotypes detected qPCR while gray dots mark dominant serotypes. Red dots in (A,B) depict samples not generating any signal in serotype/serogroup-specific qPCRs or with the signal of $C_q$ higher than the ROC$_Cq$ threshold, hence classified as negative for a serotype. In (C,D) green dots mark samples with congruent serotype between culture and molecular methods and orange dots mark samples with non-congruent result. In (D) shaded gray areas mark the 95% confidence interval of the upper limit of agreement, bias and lower limit of agreement. The continuous line marks the line of equality.

FIGURE 5 | Forest plot displaying the point estimate and 95% confidence of intervals sensitivity (A), specificity (B), concordance (C), and Cohen’s kappa (D) for molecular diagnostic tests applied to culture-enriched samples when compared to isolation of S. pneumoniae strains of particular serotype/serogroup from nasopharyngeal swabs collected in the study. Graphs displayed results for serotypes/serogroups that have been cultured from > 5 nasopharyngeal samples.

than in children (Trzcinski et al., 2013), sampling multiple sites increases sensitivity of carriage detection (Trzcinski et al., 2013; Krone et al., 2014; Wyllie et al., 2016a). However, carriage surveillance based exclusively on primary diagnostic cultures of nasopharyngeal and oropharyngeal samples often provides low quality data in adults. This limitation of the gold standard method is of particular concern for surveillance of carriage in older adults (Almeida et al., 2020; Miellet et al., 2020), the age group with the
largest incidence and burden of pneumococcal pneumonia and invasive pneumococcal disease (Jansen et al., 2009; Welte et al., 2012) as carriage is often reported to be virtually absent when the gold standard method is the only applied (Krone et al., 2014; Arguedas et al., 2020).

To overcome these limitations, we complement conventional culture with molecular methods to improve the overall sensitivity of S. pneumoniae carriage detection (Trzcinski et al., 2013; Wyllie et al., 2014, 2016a,b; Krone et al., 2015; Miellet et al., 2020). This approach is particularly effective when applied to highly poly-microbial samples from the oral niche. To validate this method, we have compared the performance of molecular methods to the gold standard. While demonstrating near-perfect agreement with primary nasopharyngeal cultures from children, application of qPCR-based methods still significantly increased the number of carriers detected. Near perfect agreement was also observed in adults when analysis focused explicitly on nasopharyngeal swabs. In adults, application of qPCR-based methods to nasopharyngeal samples did not increase sensitivity of carriage detection.

Importantly, in adults testing oropharyngeal samples changed the results dramatically. By revisiting samples negative in primary culture yet positive by qPCR with qPCR-guided culturing we have significantly increased the number of adult carriers from whom viable pneumococci were isolated. It demonstrates that molecular methods can improve the overall sensitivity of S. pneumoniae detection, a result in line with previous reports by us and others (Trzcinski et al., 2013; Krone et al., 2014, 2015; Wyllie et al., 2016a; Almeida et al., 2021). It also highlights that the gold standard method applied to oropharyngeal samples severely underestimates presence of S. pneumoniae in adults as qPCR-guided culturing increased by 7.7-fold the number of oropharyngeal samples from which viable pneumococci were isolated. Interestingly, there is evidence that testing oropharyngeal in addition to nasopharyngeal samples substantially enhances sensitivity of carriage detection also in children (Korona-Glowniak et al., 2011).

Concerns have been raised that the molecular methods are overly sensitive and lack specificity for the detection of live bacteria as “relic DNA” (DNA from non-intact cells) could be detected as well (Lennon et al., 2018). Indeed, live pathogens are less likely to be cultured from samples displaying weak positivity by qPCR (Wyllie et al., 2014; Miellet et al., 2021). While this may reflect limitations in current culturing techniques, it could also indicate presence of relic DNA that may reflect recent exposure to pneumococcus (e.g., prior to antibiotic therapy) instead of colonization proper (Lennon et al., 2018). Alternatively, suboptimal sample transport and storage conditions can also greatly limit the success rate of culturing in a carriage study. The addition of the culture-enrichment procedure prior to molecular detection reduces the risk of misclassifying these events as carriage, thus improving the specificity of detection for live pneumococci. Culture-enrichment also increases the sensitivity of carriage surveillance, in particular for poly-microbial samples (Trzcinski et al., 2013; Wyllie et al., 2014, 2016a,b; Krone et al., 2015; Rodrigues et al., 2019; Miellet et al., 2020, 2021; Almeida et al., 2021). Indeed, we observed that S. pneumoniae detection in culture-enriched samples displayed significantly increased, sensitivity, specificity, and agreement with culture and qPCR-guided culture when compared with detection in minimally processed samples.

To further improve the specificity of molecular methods for live pneumococci we performed ROC curve analysis to estimate with the Youden index Cq cut-off values that display an optimal combination of sensitivity and specificity for samples positive for S. pneumoniae by culture in primary cultures or qPCR-guided cultures (Nutz et al., 2011). This approach excluded samples that exhibited minimal positivity for targeted genes by qPCR or displayed poor agreement between pIA and lytA genes as shown in Bland-Altman graphs and with the intra-class correlation coefficient. The use of the Youden index in a ROC curve analysis to identify optimal Cq cut-off values enhances the specificity of molecular methods for detection of live pneumococcus, and the ROCq criteria can be used to direct qPCR-guided culturing. The success of this approach is dependent on culturing methods, therefore care should be taken to employ sensitive culturing techniques, such as the use of selective culture plates (Satzke et al., 2013). Furthermore, for molecular methods to be informative for culturing efforts DNA extraction should be performed irrespective of the identification of pneumococcal growth in cultures. ROC curve analysis with the Youden index may yield overly stringent Cq cut-off values.

### Table 5: The accuracy of Streptococcus pneumoniae serotype detection in n = 1,549 nasopharyngeal samples from children (n = 946) and adults (n = 603) tested using molecular methods applied to DNA extracted culture-enriched samples.

| Study population | PPV % (95% CI) | NPV % (95% CI) | Sensitivity % (95% CI) | Specificity % (95% CI) | Concordance % (95% CI) | $\kappa$ (95% CI) |
|------------------|----------------|----------------|------------------------|------------------------|------------------------|-----------------|
| All (n = 1,549)  | 100 (99.0–100) | 97.8 (97.9–99.2) | 96.2 (93.8–97.7) | 100 (99.7–100) | 99.0 (99.7–100) | 0.97 (0.96–0.99) |
| Children (n = 946) | 98.1 (96.6–98.9) | 98.1 (96.6–98.8) | 97.1 (94.8–98.4) | 100 (99.3–100) | 98.8 (97.9–99.3) | 0.98 (0.96–0.99) |
| Adults (n = 603) | 100 (78.5–100) | 99.3 (98.3–99.7) | 77.8 (54.8–91.0) | 100 (99.3–100) | 99.3 (98.3–99.7) | 0.87 (0.75–1) |

Measures of diagnostic accuracy were calculated by comparing the numbers of serotype carriage events detected with molecular methods with $n = 393$ samples from which serotypes were cultured that were targeted by serotype-specific qPCR assays. PPV, positive predictive value; NPV, negative predictive value; 95%CI, 95% confidence interval; $\kappa$, Cohen’s Kappa where $\leq 0$, 0.01–0.20, 0.21–0.40, 0.41–0.60, 0.61–0.80, > 0.81 are interpreted as poor agreement, slight, fair, moderate, substantial, and almost perfect agreement, respectively.
if qPCR-guided culturing efforts are not conducted, which could result in classifying samples containing live pneumococci as negative by qPCR. Recent advances in culturing techniques may further enhance the specificity of molecular methods for the detection of live *S. pneumoniae* (York et al., 2021).

Some studies have cautioned against the use of molecular methods due to the presence of pneumococcal genes among commensal streptococci (Carvalho Mda et al., 2012, 2013; Boelsen et al., 2020; Ganaie et al., 2021). This phenomenon is likely to be common among bacterial species co-existing in a shared niche (Kroll et al., 1998; Kilian et al., 2014; Price et al., 2015) and of particular concern for highly polymicrobial samples, hence careful selection of targeted genes is important (Greve and Moller, 2012). As previously described by our group (Trzcinski et al., 2013; Krone et al., 2015; Wyllie et al., 2016b) and others (Tavares et al., 2019) a “two-to-tango” approach, quantifying both *piaB* and *lytA* genes with molecular methods enhances the specificity of *S. pneumoniae* detection in poly-microbial samples than either target alone. This approach allows for measurements to be evaluated in a reliability analysis on a per sample basis (Bland and Altman, 1986) and for bias between measurements to be identified. While systemic biases between quantified targets can reflect dissimilarities between qPCR assay efficiencies, non-systemic biases can arise due to the presence of a gene or closely related sequence in DNA from other bacterial species. In case of multiple serotype carriage, serotype-specific abundance of the dominant serotype should display high agreement to targeted genes used to detect pneumococcus (e.g., *piaB* and *lytA*) while serotype-specific abundances of non-dominant serotypes may display reduced agreement. Serotypes that exhibit greater abundances than *piaB* or *lytA*, or appear to be present in samples classified as negative for *S. pneumoniae* are likely to be due to commensal streptococci harboring genes involved in the biosynthesis of the pneumococcal polysaccharide capsule (Lessa et al., 2018; Pimenta et al., 2019). As such, certain serotype-specific assays may be non-reliable in poly-microbial sample types. Importantly, this also concerns serotype-specific assays targeting vaccine, such as serotypes 4 and 5 (Carvalho Mda et al., 2012, 2013; Wyllie et al., 2014, 2016a, 2017; Lessa et al., 2018; Pimenta et al., 2019; Ganaie et al., 2021).

Insights into the co-occurrence of multiple serotypes in carriage is critical for understanding the dynamics of the serotypes during colonization, host-to-host transmission, and carriage progression into disease. Detection of secondary strains present in co-carriage is also important when distinguishing between unmasking and serotype replacement in assessment of pneumococcal vaccines impact (Huebner et al., 2000). However, as demonstrated in our study, and described previously the gold standard method does not readily allow detection of multiple serotypes, an event that is likely to occur often in carriage (Huebner et al., 2000). Furthermore, using our methodology we have observed no significant difference in observed and expected frequencies of multiple-serotype carriage despite what has been reported previously by others who exclusively used data from conventional culture as method of detection (Numminen et al., 2013).

An important strength of our study is the application of the procedure to two different carriage studies conducted in two different countries, and the evaluation of reproducibility in an interlaboratory comparison. Another strength of our study is that the procedure described is flexible and can be readily adapted to the carriage surveillance of other bacterial pathogens, such as *Neisseria meningitidis* (Miellet et al., 2021), for which detection of live bacteria is important or for which conventional culture may display insufficient sensitivity.

Our study had a number of limitations, the impact of testing oropharyngeal samples was only evaluated for Dutch adults and not adults sampled in England nor for any children. Since oropharyngeal samples tested were collected explicitly from Dutch adults, our findings could be unique for that demographic group and geographic location. In the protocol described we only used *piaB* and *lytA* to detect *S. pneumoniae* with molecular methods and we did not evaluate the procedure with alternative targets described by others (Shafeeq et al., 2013; Tavares et al., 2019). Furthermore, we tested for a limited number of serotypes by qPCR. Finally, not all serotypes were shown to be equally reliably detected with molecular methods and for a number of qPCR assays we were not able to identify the serotype within a serogroup.

In summary, we argue that accurate detection of pneumococcal carriage using qPCR requires concordant quantification of two genes ("two-targets-to-tango") to classify a sample as positive for pneumococcus. Similarly, qPCR-based detection requires concordance between pneumococcal and serotype-specific quantification to assure specificity of the method ("two-targets-to-tango"). We provide evidence that accurate detection of pneumococcal carriage in adults requires at least testing of both, nasopharyngeal and oropharyngeal samples ("two-samples-to-tango") and requires molecular detection to be intertwined with culture ("two methods-to-tango"). Finally, we advise revisiting samples for qPCR-guided culturing ("two-cultures-to-tango") when positive by qPCR but negative at primary diagnostic culture. The use of qPCR-guided culturing is of utmost importance for oropharyngeal swabs.

We have outlined the procedure that enhances the specificity of molecular methods for the surveillance of pneumococci and of pneumococcal serotypes in nasopharyngeal and oropharyngeal samples. Our results demonstrate near-perfect agreement between conventional culture and molecular methods when applied to nasopharyngeal samples from children. We have shown that the sensitivity of *S. pneumoniae* carriage surveillance can be greatly enhanced by complementing conventional culture with qPCRs. In adults, testing oropharyngeal on the top of nasopharyngeal samples was of paramount importance for accuracy of pneumococcal carriage detection. Studies investigating impact of testing oropharyngeal samples on detection of pneumococcal carriage in children are needed.

**DATA AVAILABILITY STATEMENT**

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.
ETHICS STATEMENT

The studies involving human participants were reviewed and approved by the Medical Ethics Committee Noord Holland (NTR5405 on http://www.trialregister.nl) and by the NHS Health Research Authority and the London Fulham Research Ethics Committee (reference 15/LO/0458; on clinicaltrials.gov reference NCT02522546). Written informed consent to participate in this study was provided by the participants’ legal guardian/next of kin.

AUTHOR CONTRIBUTIONS

ES and KT had an idea and initiated the study. NR, EM, NF, ES, and KT secured financial support. NF and KT led the project. AW-M, PB, CS, MH, NR, EM, and NF conducted carriage studies, collected the data, and provided study materials. WM, JV, and KT developed, validated laboratory methods, and wrote the laboratory protocol. WM, JV, DL, PB, TN, SM, RT, SE, and CS analyzed samples and collected the data. WM, RM, and KT contributed analytical tools. WM, JV, and DL curated the data. WM, DL, NF, and KT managed the study. WM and KT performed formal analysis of study data, visualized presentation of the results, and drafted the manuscript. All authors amended, critically reviewed, and commented on the final manuscript.

FUNDING

Funding for this study was provided to UMCU and PHE by GlaxoSmithKline Biologicals SA. GlaxoSmithKline Biologicals SA. GlaxoSmithKline Biologicals provided the opportunity to review a preliminary version of this manuscript for factual accuracy, but the authors are solely responsible for final content and interpretation. The authors received no financial support or other form of compensation related to the development of the manuscript. The collection of the samples in the UK was funded by the National Institute for Health Research Policy Research Programme (“Vaccine Evaluation Consortium Phase II,” 039/0031).

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2022.859736/full#supplementary-material

Supplementary Figure 1 | Results of molecular method-based quantification of serogroups/serotypes-specific sequences in DNA from culture-enriched nasopharyngeal and oropharyngeal samples. Individual scatter plots depict results of a single serotype-specific or serogroup-specific qPCR assay as labeled above the panel. Each symbol represents an individual sample: pale-red dots represent nasopharyngeal samples from children; blue triangles and green triangles represent nasopharyngeal and oropharyngeal samples from adults, respectively. Sample was classified as positive for a serotype/serogroup with molecular method when the signal detected by qPCR for piaB (X-axis) and serotype/serogroup (Y-axis) were both below the piaB Cq cut-off threshold of 30 Cq. Only samples that generated a signal of Cq < 40 in a particular serogroup/serotype-specific qPCR are depicted. Symbols of Cq > 40 for piaB depict individual samples or pools of samples negative for pneumococcus by qPCR. In the panel depicting qPCR results for serogroup 6 all symbols represent samples identified as positive for serotypes 6C or 6D except for red circles depicting two samples identified as positive for serotype 6A or 6B, both positive for serotype 6A by culture. In the panel depicting qPCR results for serogroup 22 all symbols represent positivity for serotype 22F. None of the samples generated any signal in qPCRs targeting serotype 1, serotype 23F and serogroup 18. In addition, no samples were identified as positive for serogroup 12 according to study criteria.

Supplementary Figure 2 | Scatter plot displaying correlation between cycle threshold (Cq) from real-time PCR (qPCR) assays targeting the Streptococcus pneumoniae piaB and serotype/serogroup specific signal detected with qPCR for culture-enriched (CE) oropharyngeal samples from adults classified positive for S. pneumoniae according to piaB Cq criterium (green dashed lines). Dots depicts serotypes detected with qPCR. Dots are color-coded according to serotypes/serogroup targeted in an assay (see legend).

REFERENCES

Almeida, S. T., Paulo, A. C., Froes, F., de Lencastre, H., and Sa-Leao, R. (2021). Dynamics of pneumococcal carriage in adults: a new look at an old paradigm. J. Infect. Dis. 223, 1590–1600. doi: 10.1093/infdis/jiaa558

Almeida, S. T., Pedro, T., Paulo, A. C., de Lencastre, H., and Sa-Leao, R. (2020). Re-evaluation of Streptococcus pneumoniae carriage in Portuguese elderly by qPCR increases carriage estimates and unveils an expanded pool of serotypes. Sci. Rep. 10:8373. doi: 10.1038/s41598-020-65399-x

Altman, D. G., and Bland, J. M. (2003). Interaction revisited: the difference between two estimates. BMJ 326:219. doi: 10.1136/bmj.326.7382.219

Arguedas, A., Tracinski, K., O’Brien, K. L., Ferreira, D. M., Wylie, A. L., Weinberger, D., et al. (2020). Upper respiratory tract colonization with Streptococcus pneumoniae in adults. Expert Rev. Vaccines 19, 353–366. doi: 10.1080/14760584.2020.1750378

Auranen, K., Rinta-Kokko, H., Goldblatt, D., Nohynek, H., O’Brien, K. L., Satzke, C., et al. (2013). Colonisation endpoints in Streptococcus pneumoniae vaccine trials. Vaccine 32, 153–158. doi: 10.1016/j.vaccine.2013.08.061

Azzari, C., Moriondo, M., Cortimiglia, M., Vallarieni, C., Canessa, C., Indolfi, G., et al. (2012). Potential serotype coverage of three pneumococcal conjugate vaccines against invasive pneumococcal infection in Italian children. Vaccine 30, 2701–2705. doi: 10.1016/j.vaccine.2011.12.008

Azzari, C., Moriondo, M., Indolfi, G., Cortimiglia, M., Canessa, C., Becciolini, L., et al. (2010). Realtime PCR is more sensitive than multiplex PCR for diagnosis and serotyping in children with culture negative pneumococcal invasive disease. PLoS One 5:e9282. doi: 10.1371/journal.pone.0009282

Bercial, A. C., Harris, D., Dela Cruz, C. S., and Possick, J. (2016). Pneumococcal vaccination strategies. An update and perspective. Ann. Am. Thorac. Soc. 13, 933–944. doi: 10.1513/AnnalsATS.201511-778FR

Bland, J. M., and Altman, D. G. (1986). Statistical methods for assessing agreement between two methods of clinical measurement. Lancet 1, 307–310. doi: 10.1016/S0140-6736(86)90837-8

Boelsens, L. K., Dunne, E. M., Gould, K. A., Ratu, F. T., Vidal, J. E., Russell, F. M., et al. (2020). The challenges of using oropharyngeal samples to measure pneumococcal carriage in adults. mSphere 5:e00478-20. doi: 10.1128/mSphere.00478-20

Bogaert, D., De Groot, R., and Hermans, P. W. (2004). Streptococcus pneumoniae colonisation: the key to pneumococcal disease. Lancet Infect. Dis. 4, 144–154. doi: 10.1016/S1473-3099(04)00938-7

Carvalho Mda, G., Bigogo, G. M., Junghae, M., Pimenta, F. C., Moura, I., Roundtree, A., et al. (2012). Potential nonpneumococcal confounding of PCR-based determination of serotype in carriage. J. Clin. Microbiol. 50, 3146–3147. doi: 10.1128/JCM.01505-12

Carvalho Mda, G., Pimenta, F. C., Moura, I., Roundtree, A., Gertz, R. E. Jr., Li, Z., et al. (2013). Non-pneumococcal mitis-group streptococci confound detection
of pneumococcal capsular serotype-specific loci in upper respiratory tract. PeerJ 1:e97. doi: 10.7717/peerj.97
Carvalho Mda, G., Tondella, M. L., McCaustland, K., Weidlich, L., McGee, L., Mayer, L. W., et al. (2007). Evaluation and improvement of real-time PCR assays targeting lytA, ply, and psaA genes for detection of pneumococcal DNA. J. Clin. Microbiol. 45, 4260–4266. doi: 10.1128/JCM.02896-06
Dagan, R., Mualem, M., Melamed, R., Leryo, O., and Yagupsky, P. (1997). Reduction of pneumococcal nasopharyngeal carriage in early infancy after immunization with tetravalent pneumococcal vaccines conjugated to either tetanus toxoid or diphtheria toxoid. Pediatr. Infect. Dis. J. 16, 1060–1064. doi: 10.1097/00006454-199711000-00011
Flasche, S., Lipsitch, M., Ojal, J., and Pinsent, A. (2020). Estimating the contribution of different age strata to vaccine serotype pneumococcal transmission in the pre vaccine era: a modelling study. BMC Med. 18:129. doi: 10.1186/s12916-020-01601-1
Flasche, S., Van Hoek, A. J., Sheasby, E., Waight, P., Andrews, N., Sheppard, C., et al. (2011). Effect of pneumococcal conjugate vaccination on serotype-specific carriage and invasive disease in England: a cross-sectional study. PLoS Med. 8:e1001017. doi: 10.1371/journal.pmed.1001017
Ganaie, F., Branche, A. R., Peasley, M., Rosch, J. W., and Nahm, M. H. (2021). Oral streptococci expressing pneumococci-like cross-reactive capsule types can affect WHO recommended pneumococcal carriage procedure. Clin. Infect. Dis. [Epub ahead of print]. doi: 10.1093/cid/ciab1003
Greve, T., and Moller, J. K. (2012). Accuracy of using the lytA gene to distinguish Streptococcus pneumoniae from related species. J. Med. Microbiol. 61(Pt 4), 478–482. doi: 10.1099/jmm.0.036574-0
Heffron, R. (1939). Pneumonia; with Special Reference to Pneumococcus Lobar Pneumonia. Oxford: Oxford University Press.
Huebner, R. E., Dagan, R., Porath, N., Wasan, D. A., and Langford, P. R. (1998). Natural history of carriage of Streptococcus pneumoniae in saliva and oropharyngeal samples from college students. Sci. Rep. 11:23138. doi: 10.1038/s41598-021-02555-x
Miellet, W. R., van Veldhuizen, J., Nicolae, M. A., Mariman, R., Bootsma, H. J., Bosch, T., et al. (2020). Influenza-like illness exacerbates pneumococcal carriage in older adults. Clin. Infect. Dis. 73, e2680–e2689. doi: 10.1093/cid/ciaa1551
Numminen, E., Cheng, L., Gyllenborg, M., and Corander, J. (2013). Estimating the transmission dynamics of Streptococcus pneumoniae from strain prevalence data. Biometrics 69, 748–757. doi: 10.1111/biom.12040
Nutz, S., Dall, K., and Karlovsky, P. (2011). Determination of the LOQ in real-time PCR by receiver operating characteristic curve analysis: application to qPCR assays for Fusarium verticillioides and F. proliferatum. Anal. Bioanal. Chem. 401, 717–726. doi: 10.1007/s00216-011-5089-x
O’Brien, K. L., Wolfson, L. J., Watt, J. P., Henkle, E., Deloria-Knoll, M., McCall, N., et al. (2009). Burden of disease caused by Streptococcus pneumoniae in children younger than 5 years: global estimates. Lancet 374, 893–902. doi: 10.1016/S0140-6736(09)61204-6
Pimenta, F., Gertz, R. E. Jr., Park, S. H., Kim, E., Moura, I., Milucky, J., et al. (2019). Streptococcus infantsis, Streptococcus mitis, and Streptococcus oralis strains with highly similar cps5 loci and antigenic relatedness to serotype 5 Pneumococcus. Front. Microbiol. 9:3199. doi: 10.3389/fmicb.2018.03199
Pimenta, F. C., Roundtree, A., Soysal, A., Bakir, M., du Plessis, M., Wolter, N., et al. (2013). Sequential triplex real-time PCR assay for detecting 21 pneumococcal capsular serotypes that account for a high global disease burden. J. Clin. Microbiol. 51, 647–652. doi: 10.1128/JCM.02927-12
Price, E. P., Sarovich, D. S., Nowosilsky, E., Beisbarth, J., Marsh, R. L., Pickering, J., et al. (2015). Haemophilus influenzae: using comparative genomics to accurately identify a highly recombinogenic human pathogen. BMC Genomics 16:641. doi: 10.1186/s12864-015-1857-x
Rodrigues, F., Christensen, H., Morales-Aza, B., Sikora, P., Oliver, E., Oliver, J., et al. (2019). Viable Neisseria meningitidis is commonly present in saliva in healthy young adults: non-invasive sampling and enhanced sensitivity of detection in a follow-up carriage study in Portuguese students. PLoS One 14:e0209905. doi: 10.1371/journal.pone.0209905
Satzke, C., Turner, P., Virolainen-Julkunen, A., Adrian, P. V., Antonio, M., Hare, K. M., et al. (2013). Superiority of trans-oral over trans-nasal sampling in detecting Neisseria meningitidis in saliva and oropharyngeal samples from college students. Sci. Rep. 3:2670–2674. doi: 10.1038/s41598-016-11977-1
Shafeeq, S., Kuipers, O. P., and Kloosterman, T. G. (2013). Cellobiose-mediated gene expression in Streptococcus pneumoniae: a repressor function of the novel GntR-type regulator BguR. PLoS One 8:e57586. doi: 10.1371/journal.pone.0057586
Slotved, H. C. (2016). Other age groups than children need to be considered as carriers of Streptococcus pneumoniae serotypes. Hum. Vaccin. Immunother. 12, 2670–2674. doi: 10.1080/21645515.2016.1197451
Southern, J., Andrews, N., Sandu, P., Sheppard, C. L., Waight, P. A., Fry, N. K., et al. (2018). Pneumococcal carriage in children and their household contacts six years after introduction of the 13-valent pneumococcal conjugate vaccine in England. PLoS One 13:e0195799. doi: 10.1371/journal.pone.0195799
Tavares, D. A., Handem, S., Carvalho, R. J., Paulo, A. C., de Lencastre, H., Hinds, J., et al. (2019). Identification of Streptococcus pneumoniae by a real-time PCR assay targeting SP2020. Sci. Rep. 9:3285. doi: 10.1038/s41598-019-39791-1
Trzcinski, K., Bogaert, D., Wylie, A., Chu, M. L., van der Ende, A., Bruin, J. P., et al. (2013). Superiority of trans-oral over trans-nasal sampling in detecting Streptococcus pneumoniae colonization in adults. PLoS One 8:e60520. doi: 10.1371/journal.pone.0060520
Two to Tango

Miellet et al. (2022). Two to Tango. Frontiers in Microbiology, 13, 859736. doi: 10.3389/fmicb.2022.859736

Conflict of Interest: Public Health England (PHE) has provided vaccine manufacturers (GSK, Pfizer, Sanofi) with post-marketing surveillance reports on pneumococcal infection which the companies are required to submit to the UK Licensing authority in compliance with their Risk Management Strategy. A cost recovery charge is made for these reports. PHE’s Respiratory and Vaccine Preventable Bacteria Reference Unit has received unrestricted research grants from Pfizer to participate in pneumococcal surveillance projects. KT received funds for an unrestricted research grant from GlaxoSmithKline Biologicals SA, consultation fees, fees for participation in advisory boards, speaking fees and funds for unrestricted research grants from Pfizer, and fees for participating in advisory boards from Merck Sharp and Dohme, all paid directly to his home institution. Except for the funds from GlaxoSmithKline Biologicals SA none was received in the relation to the work reported here.

The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Publisher’s Note: All claims expressed in this article are solely those of the authors and do not necessarily represent those of their affiliated organizations, or those of the publisher, the editors and the reviewers. Any product that may be evaluated in this article, or claim that may be made by its manufacturer, is not guaranteed or endorsed by the publisher.

Copyright © 2022 Miellet, van Veldhuizen, Litt, Mariman, Wijmenga-Monsuur, Badoux, Nieuwenhuijzen, Thombre, Mayet, Eletu, Sheppard, van Houten, Rots, Miller, Fry, Sanders and Trzcinski. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.