Development and evaluation of a multiple-locus variable-number tandem-repeats analysis assay for subtyping Salmonella Typhi strains from sub-Saharan Africa

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Abstract

Purpose. Molecular epidemiological investigations of the highly clonal Salmonella enterica subspecies enterica serovar Typhi (S. Typhi) are important in outbreak detection and in tracking disease transmission. In this study, we developed and evaluated a multiple-locus variable-number tandem-repeats (VNTR) analysis (MLVA) assay for characterization of S. Typhi isolates from sub-Saharan Africa.

Methodology. Twelve previously reported VNTR loci were evaluated and an MLVA assay consisting of five polymorphic loci was adopted. The MLVA assay was developed for use on capillary electrophoresis systems by testing a collection of 50 S. Typhi isolates. This S. Typhi strain panel consisted of six outbreak related isolates and 44 epidemiologically unlinked isolates. Amongst these were nine S.Typhi haplotype H58 isolates.

Results. The MLVA assay characterized the 50 isolates into 47 MLVA profiles while PFGE analysis of the same isolates revealed 34 pulsotypes. MLVA displayed higher discriminatory power (Simpson’s index of diversity [DI] 0.998 [95% confidence interval (CI) 0.995–1.000]) as compared to pulsed-field gel electrophoresis [Simpson’s DI 0.984 (95% CI 0.974–0.994)].

Conclusion. The MLVA assay presented in this study is a simple, rapid and more accessible tool that serves as a good alternative to other molecular subtyping methods for S. Typhi.

INTRODUCTION

Typhoid fever, a systemic infection caused by Salmonella enterica subspecies enterica serovar Typhi (S. Typhi) remains a significant public health problem worldwide [1, 2]. Recent global estimates indicate that typhoid fever causes approximately 26.9 million illnesses annually [1]. This disease is one of the most important causes of morbidity and mortality in Asia and Latin America where it is endemic. The spread of typhoid fever is aggravated by poor living conditions, substandard hygiene practices and unsafe drinking water [3–8]. Although S. Typhi has proven to be a significant public health problem in Africa [9], little is known about the continental-level molecular epidemiology and strain relatedness of this pathogen. Highly discriminatory molecular subtyping methods, which are accessible to strategic African laboratories, are essential to elucidate the epidemiology of S. Typhi, which, in turn, would allow the implementation of appropriate control strategies in the sub-Saharan Africa (SSA) region; for many of these countries, the true burden of typhoid fever is unknown [1, 10].

Molecular subtyping techniques have enhanced the ability to discriminate bacterial strains [11, 12]. These methods allow for the examination of bacterial strain relatedness at the DNA level and as a result provide a powerful tool for surveillance and outbreak investigations [10–12]. The increasing importance of S. Typhi as well as the emergence and
dissemination of the multidrug-resistant *S. Typhi* haplotype H58 (*S. Typhi* H58) across SSA has made it imperative to develop new molecular subtyping methods that allow for sensitive strain discrimination [9, 13–15]. Current methodologies used for characterization of *S. Typhi* include multi-locus sequence-typing (MLST), pulsed-field gel electrophoresis (PFGE) and whole-genome sequencing (WGS) [15–17]; amongst these, only PFGE and WGS can discriminate *S. Typhi* strains [15, 18].

PFGE is widely used for subtyping of *S. Typhi* [10, 17]. This involves macro-restriction analysis of bacterial chromosomal DNA and discrimination of isolates based on the resulting banding patterns. The major drawbacks of PFGE are that it is time consuming, which can delay foodborne outbreak identification, and that subjective interpretation of PFGE patterns still remains even with the availability of standardized PulseNet protocols [10]. Furthermore, the discriminatory capacity of this methodology in strain typing *S. Typhi* is not absolute [10, 19]. *S. Typhi* is prone to chromosomal rearrangements, which involve the reshuffling of DNA throughout the genome from one location to another. This may result in alteration of PFGE patterns, which may lead to identification of multiple PFGE patterns from a single outbreak strain [19–21].

In recent years, several approaches have been made in an effort to improve molecular subtyping of *S. Typhi* using multiple-locus variable-number tandem-repeats (VNTR) analysis (MLVA) [22–25]. Even so, till today, a standardized set of VNTR loci for the typing of homologous *S. Typhi* strains has not been established. In this study, we evaluated 12 previously reported VNTR markers for epidemiological investigation of *S. Typhi* strains from SSA. Our aim was to develop an MLVA assay suitable for use on a capillary electrophoresis system that targeted five VNTR markers.

**METHODS**

**Bacterial isolates**

The Centre for Enteric Diseases (CED) of the National Institute for Communicable Diseases, a division of the National Health Laboratory Services, serves as a reference centre for human enteric pathogens. *S. Typhi* cultures isolated from all body sites (i.e. normally sterile body sites as well as gastrointestinal sites) in both in-patients and out-patients are submitted to the CED from across the country for national surveillance through the Group for Enteric, Respiratory and Meningeal Disease Surveillance network in South Africa (GERMS-SA). A total of 50 *S. Typhi* isolates from the CED database were used to evaluate the MLVA assay. The selected isolates gave a good representation of diverse collection dates and good representation of geographic areas in South Africa. Amongst the strain collection were four isolates from known outbreaks and 9 *S. Typhi* H58 strains. *S. Typhi* isolates collected from Zimbabwe (*n*=2) and the Ivory Coast (*n*=1) were also included.

**Pulsed-field gel electrophoresis**

PFGE was performed as part of routine surveillance using the standardized PulseNet protocol for *Salmonella*, *Escherichia coli* and *Shigella sonnei* incorporating *XbaI* restriction enzyme for genomic digestion [26]. PFGE fingerprint patterns were analysed and compared using the BioNumerics software (Applied Maths, Belgium) with dendrograms of the fingerprint patterns created using the unweighted pair group method with arithmetic averages (UPGMA), and with analysis of banding patterns incorporating the Dice coefficient at an optimization setting of 1.5% and a position tolerance setting of 1.5%.

**Bacterial DNA preparation**

Bacterial DNA was prepared by inoculating half a loopful of bacterial culture in autoclaved 400 µl Tris-EDTA buffer (10 mM Tris: 1 mM EDTA, pH 8.0) and incubated for 25 min at 95°C. The supernatant containing DNA was used as a PCR template.

**PCR identification of *S. Typhi* H58**

*S. Typhi* H58 strains in the validation panel were identified using previously described PCR [27]. *S. Typhi* NCTC 8385 was used as a positive control. PCR reactions were carried out in 25 µl volume with the forward primer (H58F: 5′-GCAGG-CAAAATCGAAATCGA-3′) and reverse primer (H58R: 5′-CAAACCGTTGAATCGGAAAT-3′) at final concentrations of 0.4 µM as well as MgCl₂ and deoxynucleotide triphosphate at a final concentration of 1.5 mM and 200 µM, respectively. PCR thermal cycling included an initial denaturation at 94°C for 2 min, followed by 30 cycles at 94°C for 30 s, 60°C for 30 s, 70°C for 1 min, and a final extension at 72°C for 5 min. PCR amplicons were analysed on a 1% electrophoresis gel containing ethidiun bromide solution.

**MLVA**

**Selection of VNTR loci and PCR primers**

Twelve previously reported VNTR were selected and used as markers to explore their potential in determining the strain relatedness of *S. Typhi* isolates from SSA [22–25].

**Screening for length polymorphism of VNTR using simplex PCR**

Evaluation of the presence of allelic variation at each VNTR locus in the 50 *S. Typhi* strain panel began with simplex PCRs. *S. Typhi* NCTC 8385 was used as a positive control. All PCR assays were performed using the Qiagen Multiplex PCR Kit. Each 25 µl reaction contained 12.5 µl of the Qiagen Master Mix, 2.5 µl Qiagen Q-solution, 1 µM each of the forward [labelled with a 6-FAM or NED fluorophore (Table 1)] and reverse primers. For PCR amplification, the initial denaturation at 95°C for 15 min was followed by 25 cycles of a three-step cycle protocol: 94°C for 60 s, 55°C for 90 s and 72°C for 90 s, and a final extension at 72°C for 10 min. PCR amplicons were diluted 1:35 in sterile distilled water and 2 µl aliquots of the dilutions were mixed with 0.2 µl GeneScan 600 LIZ Size Standard v2.0 (Applied Biosystems) and 11 µl Hi-Di formamide solution (Applied Biosystems).
The samples were evaluated by capillary electrophoresis on the Applied Biosystems 3500 Genetic Analyzer (Applied Biosystems) and fragment sizes were analysed using the Gene-Mapper Software (Applied Biosystems). DNA fragments were automatically allocated to length bins and alleles were assigned based on the bin fragment sizes. The VNTR allele numbers were entered into the BioNumerics 6.5 software as character values and a dendrogram was constructed using a categorical coefficient with a 1.5 tolerance and UPGMA.

### Diversity and discriminatory power

In order to determine the measure of diversity and the degree of polymorphism at each VNTR locus, Simpson’s index of diversity (DI) and 95% CIs were calculated using an online tool available at the Public Health England (PHE) website (www.hpa-bioinformatics.org.uk/cgi-bin/DICI/DICI.pl).

We used an online tool (www.comparingpartitions.info/) to determine Simpson’s DI as well as the Wallace coefficient for the MLVA assay and PFGE analysis. Statistical comparisons of Simpson’s DIs and the Wallace coefficients were also performed online.

### Nucleotide sequencing

Nucleotide sequencing was performed in order to determine the size of the flanking region at each of the chosen VNTR loci and to also confirm that the variations in the length of amplicons were a result of variation in copy number of tandem repeats. For each VNTR locus, PCR amplicons of ten isolates representing various VNTR amplicon sizes served as template DNA in a PCR cycle sequencing reaction using the BigDye Terminator Cycle Sequencing Kit (Applied Biosystems, Foster City, CA, USA) and an Applied Biosystems 3500 Genetic Analyzer. DNA sequences were analysed using DNASTAR Lasergene (version 8.0) software (DNASTAR, Madison, WI, USA).

### Multiplex PCR assay

VNTR loci that had the highest degree of polymorphism were selected for the development of the multiplex PCR assay. Each 25 µl reaction contained 12.5 µl of the Qiagen master mix, 2.5 µl Qiagen Q-solution, 1 µl crude DNA and primer concentrations as shown in Table 2. Forward primers for TR1, TR2, Sal02, Sal20 and TR4699 were labelled with PET, 6-FAM, 6-FAM, VIC and NED fluorophores, respectively.

### RESULTS

#### Evaluation of VNTR loci polymorphism

We evaluated a total of 12 VNTR loci using a panel of 50 S. Typhi isolates from SSA. The primer sets for all 12 VNTR loci were able to produce PCR amplicons in all isolates at an annealing temperature of 55°C. Of the 12

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**Table 1. Simpson’s diversity indices and primers selected for the amplification of 12 S. Typhi VNTR**

<table>
<thead>
<tr>
<th>VNTR locus</th>
<th>Number of alleles</th>
<th>DI</th>
<th>Confidence interval</th>
<th>Repeat sequence</th>
<th>VNTR primer sequences*</th>
<th>VNTR primer references</th>
</tr>
</thead>
<tbody>
<tr>
<td>TR2</td>
<td>26</td>
<td>0.940</td>
<td>0.916–0.964</td>
<td>CCAGTTCC</td>
<td>Forward: CCGTGGTTTTTGGCTGATACG</td>
<td>[22]</td>
</tr>
<tr>
<td>TR4699</td>
<td>22</td>
<td>0.921</td>
<td>0.892–0.950</td>
<td>TGTTGG</td>
<td>Reverse: CAGAGGATATCCGCAATACTCCG</td>
<td>[23]</td>
</tr>
<tr>
<td>Sal02</td>
<td>16</td>
<td>0.916</td>
<td>0.896–0.936</td>
<td>TACCCAG</td>
<td>Forward: CGATAGACAGCAGACAGCAGA</td>
<td>[24]</td>
</tr>
<tr>
<td>TR1</td>
<td>11</td>
<td>0.868</td>
<td>0.836–0.900</td>
<td>AGAAGAAA</td>
<td>Reverse: CACAGATGGCTGACTACTACCC</td>
<td>[23, 25]</td>
</tr>
<tr>
<td>Sal16</td>
<td>10</td>
<td>0.839</td>
<td>0.805–0.874</td>
<td>ACCATG</td>
<td>Forward: TGGCTGTAATCTGGCTGATCTCA</td>
<td>[24]</td>
</tr>
<tr>
<td>Sal20</td>
<td>6</td>
<td>0.730</td>
<td>0.657–0.804</td>
<td>CAG</td>
<td>Reverse: AACTTCGGGATGTATGTCGCC</td>
<td>[23]</td>
</tr>
<tr>
<td>TR4500</td>
<td>3</td>
<td>0.607</td>
<td>0.530–0.684</td>
<td>GGACTC</td>
<td>Forward: CACGCTGCTGCGACTATCC</td>
<td>[23]</td>
</tr>
<tr>
<td>Sal06</td>
<td>3</td>
<td>0.339</td>
<td>0.185–0.494</td>
<td>CTCAAT</td>
<td>Reverse: CTTCTGCTGATGGCCACTCC</td>
<td>[25]</td>
</tr>
<tr>
<td>TR5</td>
<td>2</td>
<td>0.039</td>
<td>0.000–0.114</td>
<td>CGTCACG</td>
<td>Forward: GTAAAAAGCGGTCGTAAGCTGTTGAA</td>
<td>[22]</td>
</tr>
<tr>
<td>Sal15</td>
<td>1</td>
<td>0.000</td>
<td>0.000–0.132</td>
<td>No data published</td>
<td>Reverse: TCGAGCGGCTGCTGACGACTG</td>
<td>[24]</td>
</tr>
<tr>
<td>Sal10</td>
<td>1</td>
<td>0.000</td>
<td>0.000–0.132</td>
<td>ACGGCCGCTGCCG</td>
<td>Forward: AAGGGAAGCGGCTGACGACTG</td>
<td>[25]</td>
</tr>
<tr>
<td>TR4</td>
<td>1</td>
<td>0.000</td>
<td>0.000–0.132</td>
<td>GAAATAAAAAATG</td>
<td>Reverse: AAAAGCCGCTGCTGACGACTG</td>
<td>[24]</td>
</tr>
</tbody>
</table>

*For simplex PCRs forward primers for VNTR locus TR2, TR4699, TR4500, Sal06, Sal15 and Sal10 were labelled with NED fluorophore, while VNTR locus Sal02, TR1, Sal16, Sal20, TR5 and TR4 were labelled with 6-FAM fluorophore. For the multiplex PCR, forward primers for VNTR locus TR1, TR2, Sal02, Sal20 and TR4699 were labelled with PET, 6-FAM, 6-FAM, VIC and NED fluorophores, respectively.
potential VNTR loci evaluated, seven (TR1, TR2, TR4500, TR4699, Sal02, Sal16 and Sal20) demonstrated the ability to discriminate S. Typhi strains. Simpson’s DI for these loci ranged from 0.607 to 0.940 (Table 1). For the remaining five VNTR loci, two (TR5 and Sal06) demonstrated decreased ability to discriminate S. Typhi strains with lower diversity indices of 0.039 and 0.339 respectively, while the other three VNTR loci (Sal15, Sal10 and TR4) were homogenous and showed no variation amongst the strains.

**MLVA validation by nucleotide sequencing**

The seven most variable VNTR loci with high Simpson’s DI were selected for nucleotide sequencing. VNTR loci with 100 % conserved flanking sequences were selected for inclusion in our MLVA assay. The flanking sequences in all the five VNTR loci, including TR1, TR2, Sal02, Sal20 and TR4699, were conserved and polymorphism was due to variation in the number of tandem-repeat units within these loci (Table 2). Nucleotide sequencing revealed that the flanking sequences at VNTR locus Sal16 were not conserved. Aside from the six-base-pair tandem-repeat sequence ‘ACCATG’ at VNTR locus Sal16, an additional 12-base-pair repeat sequence ‘ACCA CAC TAC AG’ was identified. Therefore, polymorphism at this VNTR locus was due to variation at both tandem-repeat regions. Nucleotide sequencing also revealed that the flanking sequences at VNTR locus TR4500 were not conserved. A seven-base-pair insertion sequence ‘TTGCCCAC’ was identified in six of the ten S. Typhi isolates that were subjected to nucleotide sequencing. The correct number of repeat units for each strain could not be accurately determined as not all strains harboured the seven-base-pair sequence. Redesigning primers would not have resolved the problem as the insertion sequence is located adjacent to the tandem-repeat region. For this reason, VNTR loci Sal16 and TR4500 were excluded from the MLVA assay.

**Optimization of the MLVA multiplex PCR assay consisting of five VNTR loci**

We developed an MLVA assay consisting of five highly polymorphic VNTR loci including VNTR loci TR1, TR2, Sal02, Sal20 and TR4699. Three of these VNTR loci (TR4699, Sal02 and Sal20) were located in gene regions and the other two VNTR loci (TR1 and TR2) were intergenic (Table 2). The five polymorphic VNTR loci were pooled into one PCR reaction for the development of the MLVA multiplex PCR. Capillary electrophoresis was used for the analysis of the MLVA assay in order to correctly determine the VNTR loci amplicon sizes. For easy identification of PCR amplicons, forward primers for each VNTR locus were labelled with fluorophores. The forward primers for VNTR loci TR1, TR2, Sal02, Sal20 and TR4699 were labelled with PET, 6-FAM, 6-FAM, VIC and NED, respectively (Fig. 1). Due to only four fluorophores being available for selection, this resulted in labelling of two VNTR loci (namely TR2 and Sal02) with the same fluorophore (6-FAM). PCR amplicons from these two VNTR loci could still be easily identified as their amplicon sizes did not overlap.

**Discriminatory power of MLVA and PFGE analysis of S. Typhi isolates from SSA**

Overall, PFGE distinguished the 50 S. Typhi isolates into 34 unique pulsotypes (Fig. 2). PFGE clusters were defined at 80 % similarity threshold. PFGE analysis revealed four clusters, with the largest cluster (cluster b) consisting of 34 isolates (Fig. 2). PFGE showed the ability to discriminate S. Typhi H58 isolates. The nine S. Typhi H58 strains were differentiated into seven pulsotypes. Unique pulsotypes were identified for isolates belonging to the Delmas, Mpumalanga 2005 outbreak and the Pretoria, Gauteng 2010 outbreak.

The MLVA assay characterized the 50 isolates into 47 unique MLVA profiles (Fig. 3). Only three clusters, consisting of two or more isolates with indistinguishable MLVA profiles, were identified. These clusters consisted of isolates from recent outbreaks in South Africa namely, the Delmas, Mpumalanga 2005 outbreak and the Pretoria, Gauteng 2010 outbreak. The MLVA assay was able to discriminate amongst S. Typhi H58 strains. The nine S. Typhi H58 strains in the validation panel were differentiated into nine MLVA profiles. MLVA concurd with PFGE analysis in clustering of outbreak isolates.

The discriminatory power of the MLVA assay was calculated using Simpson’s DI applied to the S. Typhi strain panel. Simpson’s DI does not only depend on the number of

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**Table 2. Features of seven highly polymorphic S. Typhi VNTR loci**

<table>
<thead>
<tr>
<th>VNTR locus</th>
<th>Gene</th>
<th>Product</th>
<th>Repeat sequence</th>
<th>Unit length</th>
<th>Forward offset (bp)</th>
<th>Reverse offset (bp)</th>
<th>MLVA multiplex PCR primer concentrations</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>TR1</td>
<td></td>
<td>Intergenic region between yefD and yefE</td>
<td>AGAAGAA</td>
<td>7</td>
<td>39</td>
<td>116</td>
<td>1.2 μM</td>
<td>[22, 23]</td>
</tr>
<tr>
<td>TR2</td>
<td></td>
<td>Intergenic region between arcD and yffB</td>
<td>CCAGTTC</td>
<td>8</td>
<td>191</td>
<td>105</td>
<td>1.2 μM</td>
<td>[22, 23]</td>
</tr>
<tr>
<td>TR4699</td>
<td>sefC</td>
<td>Outer membrane fimbral usher protein</td>
<td>TGGTTGG</td>
<td>6</td>
<td>38</td>
<td>137</td>
<td>0.8 μM</td>
<td>[23]</td>
</tr>
<tr>
<td>Sal02</td>
<td>citT</td>
<td>Citrate carrier</td>
<td>TACCAG</td>
<td>6</td>
<td>136</td>
<td>59</td>
<td>1 μM</td>
<td>[23, 25]</td>
</tr>
<tr>
<td>Sal16</td>
<td></td>
<td>Intergenic region between STY3169 (pseudogene) and STY3172</td>
<td>ACCATG</td>
<td>6</td>
<td>90</td>
<td>91, 103, 109, 115, 127</td>
<td>–</td>
<td>[23]</td>
</tr>
<tr>
<td>Sal20</td>
<td>ftsN</td>
<td>Cell division protein</td>
<td>CAG</td>
<td>3</td>
<td>83</td>
<td>80</td>
<td>0.5 μM</td>
<td>[23, 25]</td>
</tr>
<tr>
<td>TR4500</td>
<td>STY4635</td>
<td>Hypothetical protein</td>
<td>GGACTC</td>
<td>6</td>
<td>76</td>
<td>195, 202</td>
<td>–</td>
<td>[23]</td>
</tr>
</tbody>
</table>
alleles present at each locus but it also takes into consideration the equitability with which the alleles are distributed at each locus [28, 29]. VNTR loci with a Simpson’s DI value closer to 1 are better markers to differentiate the strains for epidemiological purposes. For the MLVA assay, Simpson’s DI was calculated at 0.998 (95 % CI 0.995–1.000). Simpson’s DI for PFGE analysis of these isolates was calculated at 0.984 (95 % CI 0.974–0.994). The difference in Simpson’s DI for MLVA and PFGE were statistically significant (P=0.010).

The congruence between the MLVA assay and PFGE analysis was determined by calculating the Wallace coefficient. This coefficient indicates the probability that two isolates that cluster together by one subtyping method could also be clustered together using another subtyping method [30]. The Wallace coefficient between the MLVA assay and PFGE pulsotypes was 67 %. In contrast, the probability that two isolates with the same MLVA profile could have the same pulsotypes was 9 %. The MLVA assay showed a higher discriminatory power than PFGE analysis.

**DISCUSSION**

Epidemiological investigations are important for the control of the dissemination of typhoid fever. The ability to study the incidence and spread of *S. Typhi* in SSA relies on the selection and use of suitable and rapid molecular methods that are accessible to the many laboratories in this region. In order to study the dissemination of *S. Typhi* in SSA, highly discriminatory molecular methods are required for characterization of this pathogen. PFGE has been widely used for subtyping *S. Typhi* isolates; however, the suboptimal discriminatory power of this molecular method coupled with the intensive labour involved makes PFGE unsuitable. WGS has become the most commonly used molecular subtyping tool for human pathogens; however, the cost involved in using this molecular method for routine surveillance of *S. Typhi* infections in the developing countries of the SSA region makes WGS unfeasible, at present.

In our study, we evaluated 12 VNTR loci and used them as molecular markers to discriminate amongst *S. Typhi* isolates from SSA. VNTR loci with shorter repeat sequences (≤8 bp in length) were included for selection, as these show more variation in copy number. In addition, VNTR loci and primers that harboured perfect homogenous repeat sequences and had 100 % conserved flanking sequences with no insertions or deletions in repeat sequences were selected. Five of the 12 VNTR loci evaluated in this study (TR1, TR2, Sal02, Sal20 and TR4699) proved to be polymorphic and showed the ability to discriminate *S. Typhi* isolates. VNTR loci TR1 and TR2 were first tested by Liu *et al.* [22] and were found to be highly

![Fig. 1. Electropherogram showing PCR fragments of all five VNTR loci incorporated in a multiplex PCR for MLVA analysis of *S. Typhi* isolates. The fragments were resolved by capillary electrophoresis. VNTR loci TR1, TR2, Sal02, Sal20 and TR4699 were labelled with PET (red), 6-FAM (blue), 6-FAM (blue), VIC (green) and NED (black) fluorophores, respectively.](image-url)
Fig. 2. Dendrogram analysis of PFGE patterns (XbaI digestion) showing genetic relationships amongst S. Typhi isolates from SSA. Percentage relatedness is indicated at nodes. Dendrogram analysis revealed 34 pulsotypes amongst 50 S. Typhi isolates. # denotes Mpumalanga, 2005 outbreak isolates and * denotes Gauteng, 2010 outbreak isolates. The blue blocks denote strains belonging to the same PFGE cluster at 80% similarity threshold (clusters a–d). MLVA profiles are provided for direct comparison.
Fig. 3. Dendrogram based on MLVA profiles of 50 S. Typhi isolates. Percentage relatedness is indicated at nodes. The dendrogram was generated using categorical coefficient with a 1.5 tolerance and UPGMA clustering. A total of 47 unique MLVA profiles were identified amongst 50 S. Typhi isolates. # denotes Mpumalanga, 2005 outbreak isolates and * denotes Gauteng, 2010 outbreak isolates.
This assay can assist in rapid identification of VNTR markers for analysis of S. Typhi isolates from Asia. These VNTR loci were also explored in several other MLVA assays and were found to be highly polymorphic, exhibiting diversity indices ranging from 0.87 to 0.90 [23–25]. Similarly, in our study, VNTR loci TR1 and TR2 were highly polymorphic and exhibited Simpson’s DI calculated at 0.87 and 0.94, respectively. VNTR loci Sal02 and Sal20 were first explored by Ramisse et al. [25]. In their study, they identified VNTR loci Sal02 and Sal20 as two of the most variable VNTR loci exhibiting diversity indices calculated at 0.87 and 0.81, respectively [25]. Other studies have also shown the ability of VNTR loci Sal02 and Sal20 in discriminating S. Typhi isolates [23, 24]. In our study, VNTR loci Sal02 and Sal20 were highly polymorphic and exhibited Simpson’s DI calculated at 0.92 and 0.73, respectively. VNTR locus TR4699 was first described by Octavia and Lan [23]. This VNTR locus was identified as one of the highly polymorphic loci in differentiating 73 global S. Typhi isolates and exhibited a diversity index calculated at 0.95. Tien et al. [24] also explored the use of VNTR locus TR4699 in an MLVA assay and found this locus to be highly polymorphic, exhibiting a diversity index of 0.92. The findings in our study concur with those of previous studies as VNTR locus TR4699 exhibited Simpson’s DI calculated at 0.92. The high Simpson’s DI values calculated at each locus indicated that the selected loci are of highly polymorphic nature and have greater discriminatory power sufficient to differentiate epidemiologically unrelated strains.

A multiplex PCR assay containing primers targeting the five highly polymorphic VNTR loci was developed and proved to be rapid and highly discriminatory in characterizing S. Typhi isolates from SSA. Simpson’s DI for the MLVA assay was calculated at 0.998, indicating high discriminatory abilities. The combination of the five VNTR loci showing high-diversity levels enabled differentiation of closely related and unrelated S. Typhi isolates from SSA. MLVA showed higher discrimination of S. Typhi (including S. Typhi H58) and would be an effective molecular subtyping tool for epidemiological investigation of S. Typhi outbreaks in SSA.

Conclusion

Typhoid fever surveillance is of crucial importance in the SSA region where there is a high burden of disease. MLVA has a great ability to differentiate S. Typhi strains from SSA. Being PCR-based, MLVA can be used in many laboratories in developing countries which may not have access to WGS.

Our study describes an MLVA assay consisting of five VNTR markers for analysis of S. Typhi strains from SSA. This assay can assist in rapid identification of S. Typhi strain relatedness as well as typhoid fever outbreak detection. To the best of our knowledge, capillary electrophoresis equipment is available in four other laboratories in the SSA region to which the MLVA assay can be adopted. These include Kenya, Gambia, the Ivory Coast and Uganda. The implementation of MLVA analysis in these laboratories could be helpful in monitoring the spread of typhoid fever across the continent and highlight the role of specific clones in disease causation. The MLVA assay could also assist in identifying the role of imported cases of typhoid fever into South Africa and their contribution to the burden of disease. This will promote effective and appropriate disease intervention strategies, including prevention and treatment.

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Conflicts of interest

The authors declare that there are no conflicts of interest.

References


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