Serotyping of sub-Saharan Africa 
Salmonella strains isolated from poultry feces using multiplex PCR and whole genome sequencing

Assèta Kagambèga1,2,3*, Lari M. Hiott1, David S. Boyle4, Elizabeth A. McMillan1, Poonam Sharma1, Sushim K. Gupta1, Hazem Ramadan1,5, Sohyun Cho1, Shaheen B. Humayoun1, Tiffanie A. Woodley1, Nicolas Barro2, Charlene R. Jackson1 and Jonathan G. Frye1

Abstract

Background: Salmonella enterica remains a leading cause of food-borne diseases worldwide. Serotype information is important in food safety and public health activities to reduce the burden of salmonellosis. In the current study, two methods were used to determine serotypes of 111 strains of Salmonella isolated from poultry feces in Burkina Faso. First, Salmonella Multiplex Assay for Rapid Typing (SMART) Polymerase Chain Reaction (PCR) was used to determine the serovars of the S. enterica isolates. Second, serovar prediction based on whole genome sequencing (WGS) data was performed using SeqSero 2.0.

Results: Among the 111 Salmonella isolates, serotypes for 17 (15.31%) isolates were identified based on comparison to a panel of representative SMART codes previously determined for the 50 most common serovars in the United States. Forty-four (44) new SMART codes were developed for common and uncommon serotypes. A total of 105 (94.59%) isolates were serotyped using SeqSero 2.0 for serovar prediction based on WGS data.

Conclusion: We determined that SeqSero 2.0 was more comprehensive for identifying Salmonella serotypes from Burkina Faso than SMART PCR.

Keywords: Salmonella, Serotyping, Molecular method

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Background
The discovery of *Salmonella* was made by Theobald Smith in 1855 from the intestines of a pig suffering from swine fever [1, 2]. *Salmonella* is a genus of gram-negative bacteria in the family of Enterobacteriaceae with two species: *Salmonella bongori* and *Salmonella enterica*. The species *S. enterica* includes more than 2579 serovars and is a major cause of food-borne illness in humans [3, 4]. The subspecies of *S. enterica* are *enterica*, *salamae*, *arizonae*, *diarizonae*, *houtenae*, and *indica* [5]. *Salmonella enterica* subsp. *enterica* includes over 1400 serotypes and causes approximately 99% of *Salmonella* infections in humans and warm-blooded animals [6]. *Salmonella* serovars Typhi, Paratyphi A, and B cause enteric fever, a systemic febrile illness that only occurs in humans. Non-typhoidal *Salmonella* (NTS) infect a variety of hosts including warm blood animals. NTS are one of the leading causes of bacterial diarrhea worldwide, but the majority of cases occur in Sub-Saharan Africa [7]. Human disease can result from exposure to many sources such as infected animals, contaminated foodstuffs, contaminated water, and direct contact with infected environment or directly between humans.

In Sub-Saharan Africa, the socio-economic burden of NTS is difficult to quantify due to the lack of a standard method of assessment, which is compounded by under-reporting in many cases [8–10]. NTS serotype identification is important for the control of foodborne disease incidence. The traditional method for *Salmonella* serotyping is the slide and tube agglutination tests using the O and H antigen according to Kauffman-White scheme [11, 12]. However, this method is not always accessible in low- and middle-income countries (LMICs) because the anti-sera used for agglutination is very expensive and some stocks are often not available. In many LMICs like Burkina Faso, only antisera for the agglutination of Vi antigen is available for the detection of *Salmonella* Typhi and Paratyphi A or B in hospitals. A few studies have investigated *Salmonella* serotyping using the Braumann-White scheme in Burkina Faso by collaborating with laboratories in industrialized countries [13–16]. Studies in other Sub-Saharan African countries, such as Rwanda and The Central African Republic, and have also utilized traditional serotyping through collaboration to investigate NTS from human clinical sources [17, 18].

To establish the true incidence of NTS diseases in developing countries, development of new, cheaper methods is needed for *Salmonella* serotyping.

Molecular methods are well suited for this because their specificity is comparable with traditional Kauffman-White serotyping. However, many of these modern detection methods for *Salmonella* are still absent in some developing countries, particularly in Burkina Faso. Countries with limited financial resources cannot implement well-established yet complex nucleic acid analysis systems or laboratory-developed tests through a network of centralized laboratories. These molecular methods require specific and complex equipment, sensitive reagents, dedicated infrastructure, and deep technical knowledge, which are not available in many LMICs. Therefore, it is often a necessity for researchers from LMICs to collaborate with high income countries and test available modern techniques for *Salmonella* serotyping to determine if the method can be adapted for their country.

In this study, the high-throughput molecular determination of *Salmonella enterica* serovars by use of *Salmonella* Multiplex Assay for Rapid Typing (SMART) PCR using capillary electrophoresis and whole genome sequencing (WGS) were compared to determine their accuracy in identifying serotypes of NTS isolated from Burkina Faso. The SMART method was developed by Leader et al. [12] for discrimination of most common serotypes that are reported in the United States based upon their genetic differences. The genotypic serotype prediction from WGS data was done using SeqSero 1 and 2 [19, 20].

Methods

Bacterial strains
The 111 isolates used in this study were obtained from the Laboratoire de Biologie Moléculaire, d’épidémiologie et de surveillance des bactéries et virus transmissible par les aliments (LaBESTA)/Université Joseph KI-ZERBO, Burkina Faso. The strains were isolated from poultry feces and the serotype of each confirmed following the methodologies described in the International Organization for Standardization 6579–2017 [21].

High-throughput molecular determination of *Salmonella enterica* serovars
We used the SMART method developed by Leader et al. [12] with slight modification. *Salmonella* strains were streaked onto blood agar and incubated for 18–20 h at 36 °C. Then, one colony from each plate was cultured in 5 mL of Luria Bertani (LB) broth, (Difco™, Becton Dickinson and Company, Sparks, MD) and incubated for 18 h at 37 °C with shaking. The genomic DNA was then isolated from the overnight culture using the GenElute bacterial genomic DNA kit (Sigma-Aldrich, St. Louis, MO, USA) and following the kit instructions for use. Once extractions were completed, the DNA was analyzed on the Nanodrop 2000 for DNA quality measuring the 260/280 nm. All DNA were then stored at −20 °C until ready for PCR and library preparation.

PCR amplification
Each PCR mixture contained 12.5 μL of Immolase DNA polymerase 2X master mix (Bioline, Inc., Randolph, MA, USA), 2.5 μL of 10X primer master mix, 3 mM MgCl₂,
and 1 μL of extracted DNA with addition of nuclease free water to a final reaction volume of 25 μL. The cycling conditions used in the thermal cycler were 94 °C for 10 min; 25 cycles of 94 °C for 30 s, 57 °C for 90 s, and 72 °C for 30 s; 72 °C for 5 min; 15 cycles of 94 °C for 30 s, 68 °C for 90 s, and 72 °C for 30 s; and 72 °C for 5 min. For each run, the negative control was sterile water and positive controls were genomic DNA from Salmonella Typhimurium LT2, S. Typhi CT18, S. Enteritidis strains 21,027 and 98,104. The primers used in this study were previously described by Leader et al. [12]. The amplicon samples were then diluted and analyzed on an ABI 3130XL Genetic Analyzer using capillary electrophoresis.

Genemapper software v3.5 (Applied Biosystems, Foster City, CA, USA) was used to analyze the sizes of resulting PCR products according to the protocol developed by Leader et al. [12]. Scoring was based upon the presence of a PCR product that corresponded to the predicted amplicon size, as detected in control reactions with DNA from S. Typhimurium, S. Typhi, and S. Enteritidis. Each PCR product detected was given a number (1 through 16) according to the size of the amplicon [12]. The amplicons detected for each isolate were combined to create a SMART code that corresponds to serotypes previously screened by this method.

Whole genome sequencing of Salmonella strains

Extracted DNA was quantified using the Qubit double-strandedDNA high-sensitivity assay kit according to the manufacturer’s instructions (Life Technologies Corp., Carlsbad, CA, USA). The Illumina libraries were prepared using the Nextera XT DNA library preparation kit and Nextera XT index primers (Illumina, san Diego, CA, USA). The library fragment size distribution was checked using the Bioanalyzer 2100 with an Agilent HS DNA kit (Agilent Technologies, Santa Clara, CA, USA) and quantified using a Qubit DNA HS assay kit in a Qubit fluorometer (Thermo Fisher Scientific, Waltham, MA, USA). The generated libraries were then sequenced using a MiSeq version 2 reagent kit (Illumina) with 500 and 300 cycles. The paired-end read length of 2 X 250 bp was used for 500 cycles and 2 X 150 bp for 300 cycles on the MiSeq platform (Illumina).

The quality metrics of the reads were performed by FastQC (http://www.bioinformatics.babraham.ac.uk/projects/fastqc/). The sequences were then assembled using the A5-miseq assembler [22], and deposited into NCBI under BioProject no. PRJNA679582 (https://www.ncbi.nlm.nih.gov/bioproject/PRJNA679582). The genome sequence was annotated via the NCBI Prokaryotic Genome Annotation Pipeline [23].

SeqSero method for determination of Salmonella serovars using whole genome sequencing data

We performed WGS on the 111 Salmonella isolates. Using SeqSero we determined the serotype of 105 (94.59%) Salmonella isolates. Twenty-six different serotypes were identified: Enteritidis, Typhimurium, Kentucky, Agona, Virchow, Anatum, Derby, Hato, Chester, Jedburgh, Schwarzengrund, Tennessee, Albany, Duesseldorf, Poona, Eastbourne, Gaminara, Drac, Alexanderplatz, Brancaster, Bredeney, Amoutive, Telekебir, Liverpool, Munster, Monschau. Among the serotypes predicted, S. Hato was the most frequently identified serotype with 28 of the 111 isolates (25.22%), followed by Derby (20.72%), Typhimurium (6.30%), Munster (6.30%), Tennessee (5.40%), and Kentucky (4.50%) (Table 1).

SMART PCR assigned some isolates with new codes without serotype predictions and SeqSero assigned them a serotype. For example, SMART PCR predicted serotype Dusseldorf for one isolate and SeqSero predicted a serotype. For example, SMART PCR predicted serotype Dusseldorf for one isolate and SeqSero predicted a serotype. For example, SMART PCR predicted serotype Dusseldorf for one isolate and SeqSero predicted a serotype. For example, SMART PCR predicted serotype Dusseldorf for one isolate and SeqSero predicted a serotype. For example, SMART PCR predicted serotype Dusseldorf for one isolate and SeqSero predicted a serotype. For example, SMART PCR predicted serotype Dusseldorf for one isolate and SeqSero predicted a serotype.

Discussion
Serotyping is an important tool for monitoring for foodborne outbreaks and in understanding the diversity and distribution of serotypes within populations, flocks, and herds. However, serotyping by traditional methods remains inaccessible for many LMICs. In this study we investigated two molecular serotyping method to identifying serotypes for 111 isolates from Burkina Faso. The goal was to serotype Salmonella enterica using rapid and accessible molecular methods as opposed to immunologic approaches to antigen characterization. The traditional method of serotyping using the Kauffman-White...
| Sample ID | Serogroup | SMART PCR code | Antigenic serotype by SeqSero2 | Serotype by SeqSero2 |
|-----------|-----------|----------------|-------------------------------|---------------------|
| S38       | C2-C3     | 1-2-5-6-11     | 8:e4,z24:-                   | Albany or Duesseldorf |
| S39       | B         | 1-5-7-9-11-16  | 4,e,he,n,x                   | Chester             |
| S40       | E1        | 1-2-5-7-11-13  | 3,10e29:-                    | Jedburgh            |
| S47       | B         | 1-2-5-6-10-11  | 4,g,m,s:-                    | Hato                |
| S52       | B         | 1-5-7-9-11-16  | 4:i,1,2                      | Typhimurium         |
| S53       | B         | 1-2-5-6-11     | 4,g,m,s:-                    | Hato                |
| S58       | G         | 1-5-6-7-11-16  | 13,d,e,n,z15                 | Teikelkebir         |
| S59       | B         | 1-2-5-6-7-8-11-12-13-14-16 | 4:i,1,2 | Typhimurium |
| S60       | G         | 1-5-6-7-9-11-16 | 13,d,e,n,z15             | Teikelkebir         |
| S61       | D1        | 1-2-3-5-6-11-14-15 | 9,g,m,-                   | Enteritidis        |
| S62       | D1        | 1-2-3-5-6-11-14-15 | 9,g,m,-                   | Enteritidis        |
| S63       | B         | 1-2-5-6-11     | 4,g,m,s:-                    | Hato                |
| S64       | B         | 1-2-5-11-13    | 4,f,g,-                      | Agona               |
| S65       | B         | 1-2-5-6-11-13  | 4,f,g:-                      | Derby               |
| S66       | E4        | 1-2-5-6-11-13-16 | 1,3,19,f,g,1,5          | N/A                 |
| S67       | B         | 1-5-7-9-11-16  | 4,e,he,n,x                   | Chester             |
| S68       | B         | 1-2-5-6-10-11  | 4,g,m,s:-                    | Hato                |
| S69       | C2-C3     | 1-2-5-10-11-13-16 | 4:i,1,2                 | Typhimurium        |
| S70       | B         | 1-2-5-6-11     | 4,f,g:-                      | Derby               |
| S71       | C1        | 1-5-7-11-13-16  | 7:r,1,2                      | Virchow             |
| S72       | M         | 1-2-5-11-13-16  | new code                     | Amoutive            |
| S74       | C2-C3     | 1-2-5-10-11-13-16 | new code                   | Kentucky            |
| S75       | B         | 1-2-5-6-7-8-11-12-13-14-16 | 4:i,1,2                 | Typhimurium        |
| S76       | B         | 1-2-5-6-10-11  | 4,f,g:-                      | Derby               |
| S77       | B         | 1-5-7-9-11-16  | 4,e,he,n,x                   | Chester             |
| S78       | B         | 1-2-5-6-7-8-11-12-13-14-16 | 4:i,1,2                 | Typhimurium        |
| S79       | B         | 1-2-5-6-7-10-11-13 | new code                   | Brancaster          |
| S80       | I         | 1-5-6-10-11-16  | 16:d,1,7                     | Gaminara            |
| S81       | B         | 1-2-5-6-7-10-11-13 | new code                   | Gaminara            |
| S82       | B         | 1-2-5-6-7-10-11-13 | new code                   | Gaminara            |
| S83       | B         | 1-2-5-6-7-10-11-13 | new code                   | Derby               |
| S84       | B         | 1-2-5-7-11-13-16  | 4,d,1,7                      | Schwarzengrund      |
| S85       | B         | 1-2-5-6-7-11-16  | 4,f,g:-                      | Derby               |
| S86       | B         | 1-2-5-6-7-10-11-13 | new code                   | Brancaster          |
| S87       | E        | 1-5-7-11-16     | 4,f,g:-                      | Derby               |
| S88       | PolyE     | 1-2-5-6-9-11    | 47z38:-                      | Alexanderplatz      |
| S89       | B         | 1-2-5-6-7-10-11-13 | new code                   | Derby               |
| S90       | B         | 1-2-5-7-11-16-17-19 | 8:iz6                     | Kentucky            |
| S91       | B         | 1-2-5-6-11-13-16-17-19 | new code                   | Derby               |
| S92       | B         | 1-2-5-6-7-11-16  | 4,f,g:-                      | Derby               |
| S93       | B         | 1-5-6-7-11-16-17-19 | new code                   | Schwarzengrund      |
| S94       | B         | 1-2-5-6-11-13-16-17-19 | new code                   | Derby               |
| S95       | Poly D    | 1-5-6-7-11-13-16-17-19 | new code                   | Brancaster          |
| S96       | B         | 1-2-5-6-7-11-13-16-17-19 | new code                   | Derby               |
| S97       | E1        | 1-2-5-6-7-11-13-16-17-19 | new code                   | Derby               |
| S98       | B         | 1-5-6-7-10-11-16  | 4,l,v,1,7                    | Bredeney            |
| S99       | E4        | 1-5-7-11-16     | 4,f,g:-                      | Derby               |
| S100      | PolyE     | 1-2-5-6-9-11    | 47z38:-                      | Alexanderplatz      |
| S101      | B         | 1-2-5-6-7-10-11-13 | new code                   | Derby               |
| S102      | C2-C3     | 1-2-5-10-11-13-16 | new code                   | Kentucky            |
| S103      | B         | 1-2-5-6-7-10-11-13 | new code                   | Derby               |
| S104      | C1        | 1-2-5-11-13     | 7:z29:-                      | Tennessee           |
| S105      | B         | 1-2-5-6-7-10-11-13 | new code                   | Derby               |
| S106      | D1        | 1-3-5-7-10-11-16  | 9:e,he,1,5                  | Eastbourne          |
Table 1  SMART-PCR and SeqSero results (Continued)

| Sample ID | Serogroup | SMART PCR code | Serotype by SMART | Antigenic serotype by SeqSero2 | Serotype by SeqSero2 |
|------------|-----------|----------------|------------------|-------------------------------|---------------------|
| S111       | C1        | 1–2–5–11–13    | new code         | 7z29:-                        | Tennessee           |
| S112       | B         | 1–2–5–6–8–11–13| new code         | 4,g,m,s:-                     | Hato                |
| S114       | B         | 1–2–5–6–8–11–13| new code         | 4,g,m,s:-                     | Hato                |
| S115       | B         | 1–2–5–6–11–13  | new code         | 4,f,g:-                       | Derby               |
| S118       | G         | 1–6–7–9–11–16  | Poona            | 13z:1,6                       | Poona               |
| S120       | B         | 1–2–5–6–7–10–11–13| new code     | 4,f,g:-                       | Derby               |
| S121       | G         | 1–6–7–9–11–16  | Poona            | 13z:1,6                       | Poona               |
| S123       | B         | 1–5–6–7–10–11–16| Bredeney         | 4,lv:1,7                      | Bredeney            |
| S124       | B         | 1–2–5–6–11–13  | new code         | 4,f,g:-                       | Derby               |
| S125       | B         | 1–2–5–6–8–11–13| new code         | 4,g,m,s:-                     | Hato                |
| S126       | C2-C3     | 1–2–5–10–11–16| new code         | 8iz6                          | Kentucky            |
| S132       | B         | 1–2–5–6–7–10–11–13| new code     | 4,f,g:-                       | Derby               |
| S133       | B         | 1–2–5–6–7–10–11–13| new code     | 4,f,g:-                       | Derby               |
| S140       | PolyE     | 1–2–5–6–9–11   | new code         | 47z38:-                       | Alexanderplatz      |
| S143       | PolyE     | 1–2–5–6–9–11   | new code         | 47z38:-                       | Alexanderplatz      |
| S145       | C1        | 1–2–5–11–13    | new code         | 7z29:-                        | Tennessee           |
| S146       | C1        | 1–2–5–6–10–11–13| Tennessee       | 7z29:-                        | Tennessee           |
| S147       | PolyE     | 1–5–6–7–9–11–16| new code         | 47lyve,n,x                    | Drac                |
| S148       | E1        | 1–5–6–7–10–11–16| new code         | 3,10e,h:1,5                  | Muenster            |
| S149       | E1        | 1–5–6–7–10–11–16| new code         | 3,10e,h:1,6                  | Muenster            |
| S150       | E1        | 1–5–6–7–9–11–16| Muenster         | 3,10e,h:1,5                  | Muenster            |
| S151       | E1        | 1–5–6–7–9–11–16| Muenster         | 3,10e,h:1,5                  | Muenster            |
| S152       | D1        | 1–5–6–7–9–11–16| Muenster         | 3,10e,h:1,5                  | Muenster            |
| S153       | E1        | 1–5–6–7–9–11–16| Muenster         | 3,10e,h:1,5                  | Muenster            |
| S154       | E1        | 1–5–6–7–10–11–16| new code         | 3,10e,h:1,5                  | Muenster            |
| S155       | G         | 1–7–9–10–11–16  | new code         | 39f,ge,n,z15                  | N/A                 |
| S156       | G         | 1–7–9–10–11–16  | new code         | 13z:1,6                       | Poona               |
| S162       | E4        | 1–5–6–7–11–13  | new code         | 1,3,19b:-                     | N/A                 |
| S163       | B         | 1–2–5–6–11–13  | new code         | 4,f,g:-                       | Derby               |
| S164       | E4        | 1–5–6–7–11–13  | new code         | 1,3,19b:-                     | N/A                 |
| S165       | B         | 1–2–5–6–11–13  | new code         | 4,f,g:-                       | Derby               |
| S167       | E4        | 1–5–6–7–11–13  | new code         | 1,3,19b:-                     | N/A                 |
| S168       | B         | 1–2–5–6–7–8–11–12–13-14-16 | Typhimurium     | 4:i,1,2                       | Typhimurium         |
| S169       | B         | 1–2–5–6–7–10–11 | new code         | 4,g,m,s:-                     | Hato                |
| S170       | B         | 1–2–5–6–11–13  | new code         | 4,f,g:-                       | Derby               |
| S171       | B         | 1–2–5–6–10–11–13| new code     | 7z29:-                        | Tennessee           |
| S172       | C2-C3     | 1–2–5–10–11–13–16| new code     | 8iz6                          | Kentucky            |
| S183       | B         | 1–2–5–6–11–13  | new code         | 4,f,g:-                       | Derby               |
| S184       | B         | 1–2–5–6–11–13  | new code         | 4,g,m,s:-                     | Hato                |
| S185       | B         | 1–2–5–6–11–13  | new code         | 4,g,m,s:-                     | Hato                |
| S186       | B         | 1–2–5–6–11–13  | new code         | 4,g,m,s:-                     | Hato                |
| S187       | B         | 1–2–5–6–11–13  | new code         | 4,g,m,s:-                     | Hato                |
| S188       | B         | 1–2–5–6–11–13  | new code         | 4,g,m,s:-                     | Hato                |
| S191       | B         | 1–2–5–6–11–13  | new code         | 4,g,m,s:-                     | Hato                |
Scheme is expensive, time consuming, and training is needed to accurately read results. The SMART PCR is faster and cheaper than traditional serotyping and can be automated to read results. For example, it is possible to test two 96-well plates of Salmonella strains in one SMART PCR run. The reagents required are also less expensive than antisera and available from many different vendors worldwide [12]. Both factors would benefit outbreak control in the developing world. In the present study, 44 isolates were assigned new codes not previously included in the SMART PCR database. This result will be benefit LMICs by extending the original SMART code database with serotypes more prevalent in other parts of the world, particularly from Sub-Saharan Africa. This could be used in future studies to analyze the diversity of Salmonella serotypes. Moreover, Salmonella infections can globally circulate and a serotype from another region can potentially emerge as a common serotype persistent in other places than from were first reported. For example, Wong et al. [24] demonstrated that a Multidrug Resistant (MDR) S. Typhi H58 emerged in South Asia was propagated to many locations around the world, including countries in Southeast Asia, Western Asia and East Africa.

However, a limitation of SMART PCR is the identification of new codes without an assigned serotype. Previously pulsed-field gel electrophoresis patterns were used to compliment SMART codes. The SMART assay was initially developed to identify the 50 most common serotypes from clinical isolates found in the Northwestern USA [12]. Our study identified many new SMART codes associated with uncommon serotypes and there is an urgent need to extend the database to include more Salmonella serotypes as classified by the Kauffmann-White scheme [25]. This will greatly increase the usability of the SMART PCR around the world. The original SMART codes should be renewed every five years because Salmonella infections are in constant flux and any serotype can emerge as a top serotype at any time. Moreover, the capillary electrophoresis machine is very sensitive to power surges and needs to be protected appropriately, which is a challenge for laboratories in developing countries. Furthermore, the widespread application of NGS tools will at some point render capillary electrophoresis redundant.

| Sample ID | Serogroup | SMART PCR code | Serotype by SMART | Antigenic serotype by SeqSero2 | Serotype by SeqSero2 |
|-----------|-----------|----------------|-------------------|------------------------------|-------------------|
| S194      | B         | 1–2–5–6–11     | new code          | 4g,m,s:-                     | Hato              |
| S196      | B         | 1–2–5–6–11     | new code          | 4g,m,s:-                     | Hato              |
| S197      | B         | 1–2–5–6–11     | new code          | 4g,m,s:-                     | Hato              |
| S198      | B         | 1–2–5–6–11–13  | new code          | 4f,g:-                       | Derby             |
| S199      | B         | 1–2–5–6–11–13  | new code          | 4f,g:-                       | Derby             |
| S200      | B         | 1–2–5–6–11     | new code          | 4g,m,s:-                     | Hato              |
| S201      | B         | 1–2–5–6–11     | new code          | 4g,m,s:-                     | Hato              |
| S202      | B         | 1–2–5–6–11     | new code          | 4g,m,s:-                     | Hato              |
| S203      | B         | 1–2–5–6–11     | new code          | 4g,m,s:-                     | Hato              |
| S204      | B         | 1–2–5–6–11     | new code          | 4g,m,s:-                     | Hato              |
| S205      | B         | 1–2–5–6–11     | new code          | 4g,m,s:-                     | Hato              |
| S206      | B         | 1–2–5–6–11     | new code          | 4g,m,s:-                     | Hato              |
| S207      | B         | 1–2–5–6–11     | new code          | 4g,m,s:-                     | Hato              |
| S208      | B         | 1–2–5–6–11     | new code          | 4g,m,s:-                     | Hato              |
| S209      | B         | 1–2–5–6–11     | new code          | 4g,m,s:-                     | Hato              |
| S212      | B         | 1–2–5–6–11     | new code          | 4g,m,s:-                     | Hato              |
| S216      | B         | 1–2–5–6–11     | new code          | 4g,m,s:-                     | Hato              |
| S219      | B         | 1–2–5–6–11     | new code          | 4g,m,s:-                     | Hato              |
| S248      | B         | 1–2–5–6–10–11  | new code          | 4f,g:-                       | Derby             |
| S249      | B         | 1–2–5–6–10–11  | new code          | 4f,g:-                       | Derby             |
| S251      | B         | 1–2–5–6–10–11–13 | new code  | 4f,g:-                       | Derby             |
| S252      | C1        | 1–2–5–11–13    | new code          | 7z29:-                       | Tennessee         |
| S253      | E4        | 1–5–6–7–11–13  | new code          | 1,3,19b:-                    | N/A               |
| S254      | B         | 1–2–5–10–11–13–16 | new code  | 8iz6                         | Kentucky          |
| S255      | C2–C3     | 1–2–5–10–11–13–16 | new code  | 4i:1,2                       | Typhimurium       |
Table 2: New SMART codes generated by Sub-Saharan African salmonella serotypes as compared to existing serotype specific codes

| Serotypes       | New SMART codes                          | Existing SMART code from Leader et al. 2009 |
|-----------------|------------------------------------------|--------------------------------------------|
| Hato            | 1–2–5–6–11                               | None                                       |
|                 | 1–2–5–6–8–11–13                          |                                            |
|                 | 1–2–5–6–10–11                            |                                            |
|                 | 1–2–5–6–7–10–11                          |                                            |
|                 | 1–2–4–5–6–11                            |                                            |
| Typhimurium     | 1–5–7–9–11–16                            | 1–2–5–6–7–8–11–12–13–14–16               |
|                 | 1–5–6–7–11–16                            |                                            |
|                 | 1–2–5–10–11–13–16                        |                                            |
| Kentucky        | 1–2–4–5–6–10–11–13–16                    | 1–5–6–9–11                                |
|                 | 1–2–5–11–16                              |                                            |
|                 | 1–2–5–10–11–13–16                        |                                            |
|                 | 1–2–5–10–11–16                           |                                            |
| Derby           | 1–2–5–6–11–13                            | 1–2–5–6–7–11–14                          |
|                 | 1–2–5–6–11                              |                                            |
|                 | 1–2–5–6–7–8–11–12–13–14–16              |                                            |
|                 | 1–2–5–6–10–11                            |                                            |
|                 | 1–2–5–6–7–10–11–13                       |                                            |
|                 | 1–2–5–6–11                               |                                            |
|                 | 1–2–5–6–10–11                            |                                            |
|                 | 1–2–5–11–13                              |                                            |
|                 | 1–2–5–13–16                              |                                            |
| Muenster        | 1–5–6–7–10–11–16                         | 1–5–6–7–9–11                              |
| Schwarzengrund  | 1–5–6–11–16                              | 1–5–6–7–11                                |
| Tennessee       | 1–2–5–11–13                              | None                                       |
| Teitelkebir     | 1–5–6–7–9–11–16                          | 1–5–6–7–11                                |
| Brancaster      | 1–2–5–6–11–13                            | None                                       |
|                 | 1–2–5–6–7–10–11                          |                                            |
|                 | 1–2–5–6–7–10–11–13                       |                                            |
|                 | 1–5–6–7–11–13–16                         |                                            |
| 1,3,19b:-       | 1–5–6–7–11–13                            | None                                       |
| 1,3,19f,g:1,5   | 1–2–5–6–11–13                            | None                                       |
| 39z10e,n2,15    | 1–2–4–5–10–11–13–16                     | None                                       |
|                 | 1–7–9–10–11–16                           |                                            |
| Albany or Duesseldorf | 1–2–5–6–11                        | None                                       |
| Chester         | 1–5–7–9–11–16                            | 1–5–7–11                                  |
| Jedburgh        | 1–2–5–7–11–13                            | None                                       |
| Amnoutive       | 1–2–5–7–11–13–16                         | None                                       |
| Gaminara        | 1–2–5–6–7–10–11–13                      | None                                       |
|                 | 1–2–5–6–11–13                            |                                            |
| Anatum          | 1–2–5–6–7–11–16                          | 1–2–5–6–7–11–12                          |
| Alexanderplatz  | 1–2–5–6–9–11                            | None                                       |
| Eastbourne      | 1–3–5–7–10–11–16                         | None                                       |
| Poona           | 1–7–9–10–11–16                           | 1–6–7–9–10–11                             |
| Drac            | 1–5–6–7–9–11–16                          | None                                       |
SeqSero 2
SeqSero 2.0 is a new software tool for Salmonella serotype determination from WGS data described by Zang et al. [20]. SeqSero 2.0 identifies Salmonella serotypes with more precision in comparison to the first edition of SeqSero [19]. We initially used SeqSero 1.0 and found many strains with unknown serotypes. When we used SeqSero 2.0 there was a significant improvement in serotyping results (data not shown). Therefore, we can say that SeqSero 2.0 is a very powerful tool for determining serotypes using WGS data. However, this tool must be constantly updated to consider new serotypes that are identified by the Kauffman-White scheme [25]. In the present study, SeqSero 2.0 was able to predict 94.59% of the serotypes from the submitted strains. This result agrees with the results found by Banerji et al. [26]; SeqSero 2.0 was able to accurately predict most serotypes but not all. As serotyping based on WGS becomes the new gold stand, it will be important to resolve serotypes to a single correct identification.

SeqSero 2.0 has many advantages for Salmonella serotyping and fewer limitations as compared to SMART PCR. SeqSero 2.0 uses assembled whole genomes and returns a result of an antigenic profile with the serotype name in a few minutes. In the present study, we were able to predict 105 Salmonella serotypes from 111 submitted (94.59%) using SeqSero 2.0. However, using SMART PCR, only 17 of 111 strains analyzed (15.31%) were assigned a serotype. These findings demonstrate the limitation of SMART PCR assays for Salmonella strains isolated outside the U.S. The original SMART codes were created using only clinical isolates from the North Western USA, in this current study there are differences not only in the geographic region but also in using predominantly veterinary and food related isolates which may have a bearing on the significant variation in SMART codes.

Whole genome sequencing is a powerful tool for understanding Salmonella epidemiology and distribution of disease. However, sequencing is very expensive, time consuming, and requires data storage capacities and staff with high technical and bioinformatic skills. SeqSero 2 analysis of some isolates provided two possible serotypes sharing the same antigenic profile (or formula) but with differing minor O antigenic factors or in other cases gave no serotype. These unpredicted serovars are either not included in the SeqSero database and could be shared as part of the iterative construction of the serotype database for use in the next version 3.0.

In this study we noticed that SeqSero and SMART PCR can be complementary for the determination of certain serotypes. For example, SMART PCR predicted some isolates as S. Duesseldorf, and these same isolates were predicted by SeqSero 2 as S. Albany or Duesseldorf.

Conclusions
Salmonella epidemiology is a worldwide public health problem. In this study, the results highlight the accuracy of modern molecular methods and in doing so also the need for less expensive methods for rapid serotyping of Salmonella in developing countries. SeqSero 2.0 is very accurate for Salmonella serotyping, but WGS is very expensive, especially for LMICs and a role for NGS may be as a shared resource with other needs such as antimicrobial drug resistance, a growing risk to public health with grave consequences. Therefore, researchers should continue developing less expensive and accurate methods of Salmonella serotyping that can be accessible worldwide. However, both methods require a clonal culture isolate and so while these molecular tools offer great accuracy, the need for classical microbiology cannot be overlooked in first culturing and identifying the pathogen from its sample matrix.

Abbreviations
SMART: Salmonella Multiplex Assay for Rapid Typing; PCR: Polymerase Chain Reaction; WGS: Whole Genome Sequencing; LMICs: Low- and middle-income countries; NTS: Non-typhoidal Salmonella; ISO: International Organization for Standardization; CDC: Centers for Disease Control and Prevention; NGS: New Generation Sequencing; MDR: Multidrug Resistant; LaBESTA: Laboratoire de Biologie Moléculaire, d'épidémiologie et de surveillance des bactéries et virus transmissible par les aliments; U.S.: United States; USA: United States of America

Acknowledgements
We would like to thank the members of the Bacterial Epidemiology and Antimicrobial Resistance Research Unit, USDA, ARS, Athens, Georgia, USA for their valuable contributions for this research project.

Authors’ contributions
AK carried out the strain’s characterization. HR, SC, SBH, TAW participated in phenotypic characterization. LH, AK, SP, SKG, HR carried out the WGS analysis. AK drafted the manuscript. DSB, EAM, LH participated in writing the manuscript. CRJ, NB and JGF supervised the WGS and participated in writing the manuscript. All authors read, commented on, and approved of the final manuscript.

Funding
This study was supported by the United States Fulbright scholarship grant to AK and the Bacterial Epidemiology and Antimicrobial Resistance Research Unit, USDA, ARS, Athens, Georgia, USA. The funding sources had no role in the design of the study and collection, analysis, and interpretation of data and in writing the manuscript.

Availability of data and materials
The datasets and Accession numbers generated during the current study are available in the [NCBI] repository under BioProject no. PRJNA679982 [https://www.ncbi.nlm.nih.gov/bioproject/PRJNA679982], and available in the [NCBI] repository under BioSample no. PRJNA679582 [https://www.ncbi.nlm.nih.gov/biosample?Db=biosample&DbFrom=bioproject&Cmd=Link&LinkName=bioproject_biosample&LinkReadableName=BioSample&ordinalpos=1&IdsFromResult=679582].

Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

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

1. Bacterial Epidemiology and Antimicrobial Resistance Research Unit, U.S. National Poultry Research Center, USDA, ARS, Athens, GA, USA. 2. Laboratoire de Biologie Moléculaire, d’épidémiologie et de surveillance des bactéries et virus transmissibles par les aliments (LaBEST)/École Doctorale Sciences et Technologies (EDST)/Université Joseph Ki-ZERBO, Ouagadougou, Burkina Faso. 3. Institut des Sciences, Ministère des enseignement supérieur, de la recherche scientifique et de l’innovation, Ouagadougou, Burkina Faso. 4. PATH, Suite 200, Seattle, WA 98121, USA. 5. Hygiene and Zoonoses Department, Faculty of Veterinary medicine, Mansoura University, Mansoura 35516, Egypt.

Received: 18 July 2020 Accepted: 1 January 2021
Published online: 19 January 2021

References

1. Kim AY, Goldberg MB, Rubin RH. Salmonella infections. In: Gorbath SL, Bartlett JG, Blacklow NR, editors. Infectious diseases. 3rd ed. Philadelphia: Lippincott Williams and Wilkins; 2004. p. 68.

2. Eng SK, Pusparajaha P, Mutalib Ab NS, Sera HL, Chand KG, Leea LH. A study on the prevalence of Salmonella species in chicken meat in selected markets in Malaysia. Food Control. 2011;22(7):1101–5.

3. Brenner FW, Villar RG, Angulo FJ, Tauxe R, Swaminathan B. Salmonella classification. In: Porwollik S, editor. Salmonella from genome to function. Norfolk: Academic Press; 2011. p. 1–23.

4. Malorny B, Hauser E, Dieckmann R. New approaches in subspecies-level Salmonella classification. In: Porwollik S, editor. Salmonella from genome to function. Norfolk: Academic Press; 2011. p. 2465–7.

5. Brenner FW, Villar RG, Angulo FJ, Tauxe R, Swaminathan B. Salmonella nomenclature. J Clin Microbiol. 2000;38:2465–7.

6. Bopp CA, Brenner FW, Fields PI, Wells JG, Stockbine NA. Escherichia, Shigella, Salmonella: A laboratory manual for microbiology. Washington, D.C.: ASM Press; 2003. p. 564–71.

7. Centers for Disease Control and Prevention. Travelers health, Salmonellosis. Nat Rev Immunol. 2015;15:452–7.

8. Olenkova I, Olojede J, Oyedapo A, Ajibade O, Oyeniyi O, Ekanem J. Prevalence of Salmonella from chicken meat in selected wholesale markets in Nigeria. J Food Prot. 2011;74:1547–51.

9. Leader BT, Frye JG, Hu J, Fedorka-Cray PJ, Boyle DS. High-throughput PCR and capillary electrophoresis analysis. J Clin Microbiol. 2009;47:1290–5.

10. Ao TT, Feasey NA, Gordon MA, Kuniholm P, Boot M, van Bo榆dijk D, de Kruijff B, Savelkoul PHF. The genetic susceptibility of Salmonella enterica serovar Typhimurium to quinolones and resistance determinants in the chromosome. J Antimicrob Chemother. 2010;65:2053–9.

11. Kagambéga A, Haukka K, Siitonen A, Traoré AS, Barro N. Serotyping and antimicrobial susceptibility of Salmonella from cattle, poultry, pigs, hedgehogs and humans in Burkina Faso. BMC Microbiol. 2013;13(1253).

12. Bonkoungou JO, Haukka K, Österblad M, Hakanen AJ, Traoré AS, Barro N, Sitonen A. Bacterial and viral aetiology of childhood diarrhea in Ouagadougou, Burkina Faso. BMC Pediatr. 2013;13:1–6.

13. Dembélé R, Konaté A, Bonkoungou UDJ, Kagambéga A, Konaté K, Bagné T, Traoré AS, Barro N. Serotyping and antimicrobial susceptibility of Salmonella isolated from children under five years of age with diarrhea in rural Burkina Faso. Afr J Microbiol Res. 2014;8:1517–63.

14. Post AS, Diallo SN, Guiraud I, Lombo P, Tahita MC, Maltha J, Puybelde SV, Matthieu W, Ley B, Thriemer K, Rouamba E, Diarra K, Debongrave S, Tinto H, Jacob J. Supporting evidence for a human reservoir of invasive non Typhoidal Salmonella from households in Burkina Faso. PLoS Negl Trop Dis. 2019;13:e0007782. https://doi.org/10.1371/journal.

15. Byukuwenge R, Mi L, Li L, Uwanyirigira M, Vepachedu VR, Kanyawasim S, Nzayiramahoro M, Kuchipudi SV, Jayarao BM. Complete genome sequences of 20 nontyphoidal Salmonella isolates from Rwanda. Microbiol Res Announc. 2019;8:e00016–9. https://doi.org/10.1128/MRA.00016-19.

16. Breurec S, Reynaud Y, Frank T, Farra A, Costilhes G, Well X-F, Le Hello S. Serotype distribution and antimicrobial resistance of human Salmonella enterica in Bangui, Central African Republic, from 2004 to 2013. PLoS Negl Trop Dis. 2019;13:e0007917. https://doi.org/10.1371/journal.

17. Zhang S, Yin Y, Jones MB, Zhang Z, Deatherage Kaiser BL, Dinsmore BA, Fitzgerald C, Fields PI, Deng X. Salmonella serotype determination utilizing high-throughput genome sequencing data. J Clin Microbiol. 2015;53:1685–92. https://doi.org/10.1128/JCM.00323-15.

18. Zhang S, de Bakker HC, Li S, Chen J, Dinsmore BA, Lane C, Lauer AC, Fields PI, Deng X. SeqSero2: rapid and improved salmonella serotype determination using whole-genome sequencing data. Appl Environ Microbiol. 2019. https://doi.org/10.1128/AEM.01746-19.

19. International Organization for Standardization (ISO) 6160-1:2017. Microbiology of the food chain—horizontal method for the detection, enumeration and serotyping of Salmonella. Part 1: detection of Salmonella spp.

20. Coll D, Jospin G, Darling AE. AS-miseq: an updated pipeline to assemble microbial genomes from Illumina MiSeq data. Bioinformatics. 2015;31:587–9.

21. Tatsouva T, DiCuccio M, Badreddin A, Chetverin V, Nawrocki EP, Zaslavsky L, Lomsadze A, Kudorov KD, Borodovsky M, Ostell J. NCBI prokaryotic genome annotation pipeline. Nucleic Acids Res. 2016;44(614–24. https://doi.org/10.1093/nar/gkw569.

22. Wong VK, Baker S, Pickard DJ, Parkhill J, Page AJ, Feasey NA, Kingsley BA, Thomson NR, Keane JA, Well FX, Edwards DJ, Hawkey J, Hedges SR, Mather AE, Cai A, Balfour J, Hill PJ, Theu NTV, Klemm EJ, Ginios DA, Breiman RF, Watson CH, Karukski S, Gordon MA, Heidelberg RS, Okoro C, Jacobs J, Lunguya O, Edmunds WJ, Mfiecula F, Chabalgoity JA, Kuma M, Jenkins K, Dutta S, Marks F, Campos J, Thompson C, Obaro S, Mannen CA, Dolecek C, Keddy KH, Smith AM, Pary C, Kariyawasam S, Mulholland EK, Campbell J, Dongol S, Baxtyn B, Dufour M, Bandaranayake D, Nagesia TT, Singh SP, Hatta M, Newton P, Onsare RS, Isara L, Dance D, Davong Y, Thwaites G, Wijedoru L, Crump JA, De Pinna E, Nair S, Niles EL, Thanh DP, Turner P, Soeng S, Valcanis M, Powelling J, Dimovski K, Hogg G, Farrar J, Holt KE, Dougan G. Phylogeographic analysis of the dominant multidrug-resistant H58 clade of Salmonella Typhi identifies inter- and intracontinental transmission events. Nat Genet. 2015;47(6):632–9. https://doi.org/10.1038/ng.3281.

23. Kaufmann F. Classification and nomenclature of the genus Salmonella. Acta Pathol Microbiol Scand B: Microbiol Immunol. 1971;79:421–2.

24. Banerji S, Simon S, Tille A, Fruth A, Flieger A. Genome-based Salmonella serotyping as the new gold standard. Nat Sci Rep. 2020;10:4333. https://doi.org/10.1038/s41598-020-61254-1.

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