Salmonella nomenclature in the genomic era: a time for change

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Salmonella enterica nomenclature has evolved over the past one hundred years into a highly sophisticated naming convention based on the recognition of antigens by specific antibodies. This serotyping scheme has led to the definition of over 2500 serovars which are well understood, have standing in nomenclature and, for the majority, biological relevance. Therefore, it is highly desirable for any change in naming convention to maintain backwards compatibility with the information linked to these serovars. The routine use of whole genome sequencing and the well-established link between sequence types and serovars presents an opportunity to update the scheme by incorporating the phylogenetically relevant sequence data whilst preserving the best of serotyping nomenclature. Advantages include: overcoming the variability in antibody preparations; removing the need to use laboratory animals and implementing a truly universal system. However, the issue of trying to reproduce the phenotyping gold standard needs to be relaxed if we are to fully embrace the genomic era. We have used whole genome sequence data from over 46,000 isolates of Salmonella enterica subspecies enterica to define clusters in two stages: Multi Locus Sequence Typing followed by antigen prediction. Sequence type—serotype discrepancies were resolved using core SNP clustering to determine the phylogenetic groups and this was confirmed by overlaying the antigenic prediction onto the core SNP clusters and testing the separation of clusters using cgMLST Hierarchical Clustering. This allowed us to define any major antigenic clusters within an ST—here called the MAC type and written as ST-serovar. Using this method, 99.96% of Salmonella isolates reported in the UK were assigned a MAC type and linked to a serovar name taken from the Kauffmann and White scheme. We propose a change for reporting of Salmonella enterica sub-types using the ST followed by serovar.

We identify and name pathogenic organisms because in many cases this informs clinical and public health management of the diseases they cause. For Salmonella, the second most common cause of bacterial food poisoning, specific antibodies are used which recognise cell wall (O) and phase 1 and phase 2 flagella (H) antigens. The combination of O:H1:H2 is known as the antigenic formula and each unique combination is given a serovar (or serotype) name. Clinically the most critical differentiation for Salmonella is between the enteric fever-causing typhoidal serovars and the non-typhoidal (NTS) serovars that most commonly cause gastroenteritis. Enteric fever requires antibiotic therapy and if a typhoidal serovar: Salmonella enterica Typhi or Paratyphi, is identified then contact tracing is considered. For infection with NTS, specific therapy is not required in the immunocompetent host but because of the impact on society, outbreak investigations are often initiated where there is an exceedance in the population of a clonal strain. In the UK, if an isolate is identified as Salmonella then under the Public Health (Control of Disease) Act 1984 this must, by law, be notified to Public Health England and so typing carries both a legal and a public health responsibility. The local Health Protection Team is informed and after confirmation and typing from the reference laboratory the case is included in the national databases for infection surveillance; outbreak investigations are initiated if appropriate.

Salmonella is currently classified into two species, Salmonella bongori (originally classified as Salmonella enterica subspecies V), rarely associated with human infection, and Salmonella enterica. S. enterica is a diverse species which infects and colonises many animals including humans. Historically S. enterica classification has been based on biochemistry, to define 6 subspecies (I enterica, II salamae, IIIa arizonae, IIIb diarizonae, IV houn tenae, VI indica)1. Most human infections involve serovars within subspecies I2, which are named according to the Kauffmann-White scheme3. The first published Kauffmann-White Scheme (1934) described 44 serovars4 and

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be designated a serovar based on ST and analysis of antigen encoding genes were defined as being problematic. Salmonella and propose a method for naming all antibody based serotyping.

From England and Wales to the Salmonella enterica analysis of the sequence data generated at Public Health England from all isolates referred is to reach a consensus for the definition of strains at the SNP and allele level for outbreak detection, the naming of serovars by referral to ST10. Another approach is to reproduce the entire Kauffmann-White scheme using the genetic sequence of every antigen encoding gene to predict the antigen encoded—genoserotyping. Programmes such as SeqSero11 have been developed with this aim in mind and are very successful; 98% concordance with serotyping reported from routine use on 520 isolates (20 serotypes)12 but the genetic basis for some antigens, particularly the cell wall (O) antigens remains elusive. The combination of the two sequence based approaches, phylogenetic methods with genoserotyping, for example the Salmonella in Silico Typing Resource (SISTR)13, reports 94.6% concordance on a dataset comprised of 4188 Salmonella genomes. However, the gold standard of the Kauffmann-White scheme is not perfect and remains a controversial topic14. Where WGS reports 94.6% concordance on a dataset comprised of 4188 Salmonella genomes. However, comparison with the gold standard then serotyping with antibodies raised in rabbits will remain necessary for the foreseeable future14. The solution may be with international groups such as PulseNet International, a global network dedicated to laboratory-based surveillance for food-borne diseases which is the most widely accepted process for reviewing Salmonella typing. Their vision is the implementation of WGS surveillance15 however, as the focus of PulseNet typing is in transition, the Kauffmann-White Scheme has not been updated since 2007, and there is no yet agreement on what should replace it—genomic approaches are generally considered to be the most promising but consensus is needed to ensure a standardisation of approach. One of the main issues is with isolates for which the link between serovar and DNA sequence data is not clear. These problematic isolates fall into three groups: (1) unknown genetics of antigen production—genoserotyping has not been defined; (2) lack of concordance with gold standard—commonly serotyping differentiates two isolates which have the same ST; and (3) novel STs—new STs which have not been formally approved. If we continue to follow the formally recognised gold standard then serotyping with antibodies raised in rabbits will remain necessary for the foreseeable future14.

Methods

Salmonella enterica subspecies enterica sequenced isolates14 reported between January 2014 and 21st June 2019 selected for this study included 46,268 strains. Data was analysed as follows:

Serovar inference using MLST. Analysis was undertaken on the 46,268 strains to understand the number of isolates in which ST alone could be used for identification without the need for any further testing.

Assigning isolates to novel STs. Since routine implementation in 2015, sequences that did not map to any known sequence type (i.e. novel ST) were assigned a new sequence type via PubMLST (https://pubmlst.org/Salmonella/) and validated by antibody-based testing. In brief, antibody-based testing was undertaken using in house sera antibodies against the isolate antigen using slide agglutination, microtitre dilution and Craigie motility agar methods according to the Kauffmann-White scheme. At the time of this study, antibody-based serotyping was performed on three independent isolates before an ST was validated and then used to define a serovar14. Novel STs were also compared against SeqSero11 for identification and unresolved serovars were then assessed using predicted ST serovars in Enterobase as previously described16.

Defining major antigen clusters (MACs) within problematic groups. The groups that could not be designated a serovar based on ST and analysis of antigen encoding genes were defined as being problematic. Usually this was because one ST contained more than one serovar, or the serovar prediction programmes were
not able to assign a serovar. A random subset of strains from each group also had antibody-based testing as described above. Problematic groups were analysed further to determine if the different serovars with the same ST came from distinct phylogenetic groups or were in fact from a single population—we term this the MAC type which was achieved as follows.

**Somatic antigen clusters.** At the time of writing, the available serovar prediction programmes were not able to resolve all O antigens from the genome sequence. Isolates from STs containing multiple serovars (defined by serological based typing of O antigens) were tested for phylogenetic separation using core genome SNP clustering.

**Flagella clusters.** We defined H antigens using serovar prediction programmes and reproduced the names in the Kauffmann-White scheme for most of the problematic STs using SeqSero—we checked three for concordance with antibody-based serotyping. The necessity of using *fliC* sequences as a differentiating factor within a ST has been questioned, therefore variation at the genomic level for serovars within a single ST that differed only by an H antigen was investigated using core genome SNP clustering.

**Core genome SNP clustering.** All data were analysed in Galaxy. Raw Illumina sequence data (previously generated by PHE) was downloaded from the NCBI sequence read archive (SRA, [https://www.ncbi.nlm.nih.gov/sra](https://www.ncbi.nlm.nih.gov/sra)). Strains with associated laboratory serovar identifications were assembled using SPAdes with default parameters and assessed with QUAST (v5.0.2). Per group, a high quality reference was selected that had the fewest contigs > 1000 bp and had the largest single contig. Reference assemblies were reduced to contigs > 1000 bp using seqtk (v1.3.3) with the -L 1000 flag. All sequenced isolates per group were compared to the reference using snippy (v3.2) and snippy-core was used to generate a core SNP alignment. Maximum likelihood phylogenetic trees were generated using IQ-TREE (Galaxy version 1.6.12) and visualised with associated metadata in iTOL. Differences in *fliC* and *fljB* nucleotide sequences were assessed by sequence comparison in Seaview (v4.7) after annotation of the reference sequence using prokka (Galaxy version 1.13). As ST226 (Carrau/Gatow) only comprised 9 isolates in the PHE sequences, this group was supplemented with 129 isolates obtained from Enterobase with the same ST and whole genome sequence available for download. Isolates assigned serovar Bredeney came from either ST241 or ST897. To assess whether these truly represented different phylogenetic groups, a maximum likelihood tree containing all isolates from ST241 and ST897 was generated and visualised as before.

**Hierarchical clustering.** Hierarchical Clustering of the groups was assessed using tools in Enterobase, an open access public database, as previously described to define the number of alleles (3002 cgMLST genes) at which the MAC types differentiate. This enables readers to differentiate the MAC types who may not have access to bespoke phylogenetic methods. Major Antigenic Cluster (MAC) types were designated to both serovars if serovars were genetically distinct and differed by a minimum 100 alleles using cgMLST Hierarchical Clustering, otherwise the most common serovar was assigned as the MAC type.

**Results**

DNA sequence data was available from 46,268 subspecies I *Salmonella* isolates sequenced at PHE over the period January 2014 to June 2019 comprising 550 named serovars and 1704 STs. In this study, a total of 11,605/46,268 (25%) isolates had antibody-based testing undertaken. Using traditional serological techniques during this time period, there were 970 (2.1%) isolates reported as unnamed serovars. Using a genomic approach and utilising the ST to associate sequence to serovar, the naming of 43,657 (94.37%) isolates was possible, leaving 2601 (5.63%) requiring further analysis. These isolates fell into two main categories, 1774 (3.85%) isolates were associated with ‘problematic groups’ and 827 (1.78%) isolates were novel STs. SeqSero could differentiate 1,607 isolates from problematic groups in accordance with MAC typing and 806 isolates with novel STs. This left a total of just 172 isolates (0.37%) from problematic groups that couldn’t be fully resolved using ST and SeqSero according to the Kaufmann and White Scheme (see Supplementary: Fig S1), further details described in below.

**Problematic groups (1774 isolates).** Each problematic group was investigated (Table 1) and the differences in the somatic (O) antigen or flagella (H) antigen were recorded. Clustering on core SNP similarity was performed and the serovar name was overlaid to visualise the distribution of serovars on the SNP tree—where the serovars clustered independently, a major antigen cluster or MAC type was defined within the ST. See Fig. 1A,B for examples and Supplementary Figs. S2–S14 for details.

**Somatic clusters.** There were three STs containing multiple serovars differentiated by O antigens. Comparison between antibody-based serotyping and the SeqSero predictor programmes showed that SeqSero could not predict all antigenic expressions for the somatic antigen. The common antigens involved in the sequence-based naming issues were O22 and O23 (ST1985, Fig. 1B), O6,7 and O6,14 (ST226, see Supplementary Fig. S9) and O6,8 and O8,20 (ST2256, see Supplementary Fig. S1). Phylogenetic clustering showed that within a single ST, two serovars, if differentiated only by O antigen, did not separate into distinct clusters (Table 1). The exception to this was the serovar Carrau or Gatow (ST226, See supplementary Fig. S9).
| ST  | Serotypinga | Differential antigenb | Phylogeny grouping | No. Isolates | %     | Proposed MAC type/ namec | Main hierarchical clustering level to differentiate | Figure |
|-----|-------------|-----------------------|--------------------|-------------|-------|---------------------------|---------------------------------------------------|--------|
| 20  | Brandenburg | H:lv                  | Distinct           | 24          | 0.05  | ST20-Brandenburg HC400_11346 | Figure 1A                                        |        |
| 20  | Sandiego    | H:e,h                 | Distinct           | 26          | 0.06  | ST20-Sandiego HC400_2225   | Figure 1A                                        |        |
| 1985| Bahati or Durham | O:22 or O:23 | Mixed              | 66          | 0.14  | ST1985-Durham N/A—interspersed | Figure 1B                                        |        |
| 49  | Saintpaul   | H:e,h                 | Distinct           | 75          | 0.16  | ST49-Saintpaul HC200_8     | Figure S2                                        |        |
| 49  | Haifa       | H:z10                 | Distinct           | 130         | 0.28  | ST49-Haifa HC200_1433      | Figure S3                                        |        |
| 582 | Kottbus     | H:e                    | Distinct           | 42          | 0.09  | ST582-Kottbus HC900_7054   | Figure S3                                        |        |
| 582 | Chailey     | H:z23                 | Distinct           | 6           | 0.01  | ST582-Chailey HC900_17     | Figure S3                                        |        |
| 22  | Braenderup  | H:e,n,z15             | Distinct           | 607         | 1.31  | ST22-Braenderup HC100_185  | Figure S4                                        |        |
| 22  | Larochelle  | H:1,2                 | Distinct           | 19          | 0.04  | ST22-Larochelle HC100_1136, 26664, 28707 | Figure S4                                        |        |
| 241 | Bredeney    | H:lv                  | Distinct           | 45          | 0.10  | ST241-Bredeney HC200_2494, HC200_1335 | Figure S5, S14 |        |
| 241 | Schwarzengrund | H: d                 | Distinct           | 12          | 0.03  | ST241-Schwarzengrund HC200_17392 | Figure S5                                        |        |
| 897 | Bredeney    | H:1,7                 | Distinct           | 8           | 0.02  | ST897-Bredeney HC400_31544 | Figures S6, S14 |        |
| 897 | Kimmuenza   | H:e,n,x               | Distinct           | 9           | 0.02  | ST897-Kimmuenza HC400_24937 | Figure S6                                        |        |
| 48  | Panama      | H:lv                  | Distinct           | 197         | 0.43  | ST48-Panama HC400_369      | Figure S7                                        |        |
| 48  | Miami       | H:a                   | Distinct           | 4           | 0.01  | ST48-Miami HC400_2307, HC400_67476 | Figure S7                                        |        |
| 2019| Napoli or Zaiman | H:z13 or H:l,v   | Mixed              | 21          | 0.05  | ST2019-Napoli N/A—interspersed | Figure S8                                        |        |
| 226 | Carrau      | O:6,14                | Distinct           | 4           | 0.01  | ST226-Carrau HC400_363     | Figure S9                                        |        |
| 226 | Gatow       | O:6,7                 | Distinct           | 3           | 0.01  | ST226-Gatow HC400_59526   | Figure S9                                        |        |
| 684 | Uganda or Sinstorf | H:1,or H:1,z13 | Mixed              | 75          | 0.16  | ST684-Uganda N/A—interspersed | Figure S10                                       |        |
| 909 | Bareilly    | H:1,5                 | Distinct           | 299         | 0.65  | ST909-Bareilly HC200_899   | Figure S11                                       |        |
| 909 | Richmond    | H:1,2                 | Distinct           | 86          | 0.19  | ST909-Richmond HC200_101  | Figure S11                                       |        |
| 2256| Brunel or tananarive | O:8, 20 or O:6:8 | Mixed              | 7           | 0.02  | ST2256-Brunel N/A—interspersed | Figure S12                                       |        |
| 101 | Bochum      | H:r                   | Distinct           | 4           | 0.01  | ST101-Bochum HC900_491    | Figure S13                                       |        |
| 101 | Wien        | H:b                   | Distinct           | 5           | 0.01  | ST101-Wien HC900_95        | Figure S13                                       |        |

Table 1. Proposed reporting guidelines for *Salmonella* using genome sequence data to define the major antigenic clusters (MACs). aNot used in MAC typing but presented for comparison and explanation. bAny serotype prediction programme can be used though some antigens cannot be distinguished. cFigure shows core SNP cluster for each antigenic type. dHierarchical Clustering Level at which the serovars can be differentiated genetically. No. = number of MAC type, % of the 46,268 Salmonella reported.

Flagella clusters. There were fourteen STs containing multiple serovars with an H antigen difference (Table 1). Comparison between antibody-based serotyping and the SeqSero predictor programmes showed that SeqSero could predict all antigens from the *flfC* and *flfB* gene sequences. The most common antigens involved in the sequence-based naming issues were the H antigens: *flfC*: l,v; e,h; z10; z23; z13; a; b; d; e and *flfB*: 1,2; 1,5; 1,7; z-; e,n=x. Phylogenetic clustering showed that the majority of serovars found in the same ST, which differed by H antigen, formed distinct clades (ST20-Fig. 1A, ST49-Fig. S2, ST582-Fig. S3, ST22-Fig. S4, ST241-Fig. S5, ST897-Fig. S6, ST48-Fig. S7, ST909-Fig. S11 and ST101-Fig. S13) and so were considered as separate entities (Table 1). There was also an example of a polyphyletic serovar found in two problematic groups defined as MAC type ST241-Bredeney and ST897-Bredeney but still genetically distinct (Table 1, Fig. S14). The exceptions were two groups that contained mixed clades (Table 1, ST2019-Fig. S8 and ST684-Fig. S10) which were differed by the l,v and l,v,z13 antigen (Table 1).

Major antigenic cluster (MAC) typing. We then used core SNP clustering to test if the different serovars, defined by traditional serotyping, separated into distinct phylogenetic clusters or were distributed together as a mixed cluster; this we named MAC typing and the groupings were confirmed by cgMLST Hierachial Clustering. Where the phylogenetic clusters contained mixed antigenic types the most common serovar seen by the laboratory was used to name all isolates in that cluster. Phylogenetic analysis and cgMLST Hierachial Clustering showed that distinct MAC types varied in their genetic relatedness and split from a range of 100 alleles level (ST22, see Supplementary Fig. S4) to 900 alleles level (ST582, see Supplementary Fig. S3) (Table 1).

Novel STs. Analysis in this study showed that novel STs (n = 827) make up a small percentage of *Salmonella enterica subspecies enterica* but the number of new STs arising continues to be stable accounting for 1.8–2.4% of *Salmonella enterica* reported each year (Table 2). The novel bioinformatic pipeline was not fully implemented in 2014 and novel STs were unrepresented in that year. In summary, of the 46,268 Salmonella analysed, using genomic testing and a polyphasic approach of ST, SeqSero and MAC typing enabled 99.96% of isolates to be reported as a named serovar. For 21 (0.04%) novel ST isolates, only a partial antigenic profile, according to the Kauffman-White Scheme could be characterised (see Supplementary Fig. S1).
Discussion

Based on the analysis reported here, we have shown that MLST and SeqSero can differentiate the majority of Salmonella previously defined isolates into major antigenic clusters—MAC types, which map to serovars in the Kauffmann-White scheme. The assignment of a MAC type to the 0.37% of problematic and novel isolates resulted in the total genomic identification of 99.96% of isolates (see Supplementary Fig. S1). The 0.04% of isolates which could not be fully resolved in terms of the full antigen prediction was still an improvement to 2.1% of isolates that could not be resolved via antibody-based testing which were reported as ‘unnamed Salmonella’. Although MAC typing of problematic groups depended initially upon core genome SNP typing, this will not need to be repeated for the STs described here. Assignment of a serovar name to an isolate with a sequence type can be achieved through many programmes freely available online. Laboratories using WGS can use this publication to ascribe serovar names to the 14 problematic STs described here. This publication also provides a road map for assigning serovar names to any new problematic STs seen in other regions of the world.

There were fourteen polyserovar groups addressed in this paper which contains a comprehensive dataset of Salmonella reported in England and Wales, it is likely that there will be more groups in the future and in different regions of the world. Analysis of the groups showed that genoserotyping programmes can usually differentiate polyserovars that differ by the flagella antigen using current software programmes such as SeqSero and that these polyserovars, were genetically different. Therefore, those serovars which form distinct clades, should continue to be differentiated even though they are in the same ST. There are exceptions to this rule including those differentiated by the lv,iz13 antigen (Serovar Uganda or Sinstorf) or the H:z13 or H:v antigen (Serovar Napoli or Zaiman) which formed mixed clades. This is likely due to the quality or specific binding properties of the antibodies. Interpretation of results may also be a factor as shown by MAC types ST22-Braenderup (H:e,n,z15)
and ST22-Larochelle (H:1,2), a clearly distinct group, in which antibody-based results incorrectly identified some of the strains (see Supplementary Fig. S8), (Table 1, Figs. 1A, see Supplementary Figs. S2–S7, S10, S11, S13).

Polyserovars that differed by the somatic antigen could not be differentiated by WGS genoserotyping software programmes and the majority were not genetically different (Table 1, Fig. 1B, see Supplementary Fig. S12). With the exception of the distinct MAC Types such as ST226–Gatow and ST226–Carrau (see Supplementary Fig. S9), Even though genoserotyping cannot differentiate these groups, Hierarchical Clustering can be used, in this case at the 400-allele level (Table 1). The exceptions from the majority of strains where differing flagella antigens are genetically distinct but slightly flawed serotyping.

Biotypes. Essentially, SISTR and SeqSero databases are based on the K&W scheme which depends upon the different groups of Salmonella varied between HC100-HC900 for defining MAC types (Table 1). Therefore, these formatic skills. Having a strict cut off to define MAC types is not possible because the genetic variability within bioinformatic platforms, both methodologies are also available on Enterobase and don't require extensive bioin-

Salmonella national communication and the understanding of biological, clinical, transmission and outbreak association of Salmonella middle income countries who don't have access to molecular techniques. At the very least however, the concept the future. It is also useful for frontline laboratories where presumptive identification is required and in low to holds value in microbiology, retaining the skills in specific institutes (as with viral culture) will be important for

3–14 days; additional staff resources and expertise; and additional quality testing systems and cost. Serology still expresses but that there is a very low impact in clinical or public health management. It is unlikely that a full reform of the Kauffmann–White naming scheme based on genetic differences will occur, to differentiate polyphyletic or polyserovar groups but the use of MAC types will resolve expressions but that there is a very low impact in clinical or public health management. It is unlikely that a full

Definition of Salmonella in the genomic era. The MAC type can be inferred by validated genomic

approaches with validated databases, for example by ST9,10 prediction software programmes11,13,27 and MAC
typing (this study). It should be recognised that for novel STs, not all somatic antigens will be predicted and so an agreed approach to naming is required. It is therefore recommended that Salmonella strains are named first by Salmonella species and subspecies and then the MAC type (ST or provisional ST plus historical serovar name or Serogene). Examples of MAC types: (a) S. enterica ST34—S. Typhimurium-monoaphasic variant (b) S. enterica ST43—S. Java (c) S. enterica ST86—S. Paratyphi B (d) S. diarizonae ST1262—18l,vz, (e) S. bongori, ST398—60:z41:–. An.

Historical methods for naming new serotypes will also play a role, if the antigen had historically reacted with a known antibody, even if non-specific, it was recorded as being the same. We see evidence of this when looking at sequences of the flagella from 6,7:c:1,5 strains (S. Decatur and the S. Choleraesuis/S. Paratyphi C group, originally differentiated by biotyping) in which antibody-based testing is not efficient as distinguishing distinct sequences of fijB genes which has resulted in serological conflation of these genetically unrelated serovars9. We see the same issue where genoserotyping can't differentiate historical biotyped groups with S. Paratyphi B and S. Paratyphi B var Java, which can't be differentiated serologically. They are genetically distinct with the former causing invasive disease10 and fall into distinct ST groups9 and therefore should be clearly differentiated in name. The use of MAC typing could also resolve these groups without the need for biochemical testing to differentiate biotypes. Essentially, SISTR and SeqSero databases are based on the K&W scheme which depends upon the excellent but slightly flawed serotyping.

Currently for novel Salmonella, antibody-based serology is still undertaken to comply with the Kauffmann-White scheme which does not take the genotype into account. The impact of continuing to use antibody based serological methods includes: the continued use of animal model products; increase in turnaround times by 3–14 days; additional staff resources and expertise; and additional quality testing systems and cost. Serology still holds value in microbiology, retaining the skills in specific institutes (as with viral culture) will be important for the future. It is also useful for frontline laboratories where presumptive identification is required and in low to middle income countries who don't have access to molecular techniques. At the very least however, the concept of using ST, genotypic expression and MAC typing for defining Salmonella where genomic methods are available should be adopted. Particularly in reference laboratories and accepting that 0.04% may not predict all phenotypic expressions but that there is a very low impact in clinical or public health management. It is unlikely that a full reform of the Kauffmann and White scheme based on genetic differences will occur, to differentiate polyphyletic or polyserovar groups but the use of MAC types will resolve Salmonella nomenclature issues.

This study/opinion piece supports the continued use of historical names, they are valuable in terms of international communication and the understanding of biological, clinical, transmission and outbreak association of groups. It is recommended that Salmonella are named genetically as the gold standard where these practices are available. To define new MAC types, either SNP or cgMLST hierarchical clustering approaches, with a minimum of three of each serovar, could be used as this study shows. Although these methods can be performed via bespoke bioinformatic platforms, both methodologies are also available on Enterobase and don't require extensive bioin-

formatic skills. Having a strict cut off to define MAC types is not possible because the genetic variability within different groups of Salmonella varied between HC100-HC900 for defining MAC types (Table 1). Therefore, these distinctions would be required on a group by group basis.
Conclusion
A previous report from PHE, using the same dataset, stated that almost half of sequenced *Salmonella* isolates that were checked phenotypically were due to discrepancies with the findings from the serology of sending laboratory.\(^1\) We show here that this is not necessary as the serovar designation from genome sequence data alone can be trusted and used for reporting. Using WGS, 99.96% of *Salmonella* isolates reported in the UK can currently be assigned a MAC type name taken from the MLST\(^1\,\(^2\) and genetic antigenic profile in line with the Kaufmann and White Scheme.\(^1\) We recommend a change to using sequence data for the routine naming of all *Salmonella*.

Received: 13 December 2020; Accepted: 9 March 2021

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Acknowledgements
We would like to thank Amy Gentle, Martin Day, Andrew Levy, Clare Maguire and Tracey Dealey for undertaking antibody-based typing in the laboratory. This study was funded by Public Health England. The authors gratefully acknowledge the support of the BBSRC; G.C.L. and J.W were funded by the BBSRC Institute Strategic Programme Microbes in the Food Chain BB/R012504/1 and its constituent project BBS/E/F/000PR10352.
Author contributions
M.A.C. designed the study and prepared the tables and figure S1, M.A.C., G.C.L., J.W. wrote the main manuscript and performed analysis. G.L. performed phylogenetic analysis and prepared figures.

Competing interests
The authors declare no competing interests.

Additional information
Supplementary Information The online version contains supplementary material available at https://doi.org/10.1038/s41598-021-86243-w.

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