Whole-genome sequencing is rapidly replacing current molecular typing methods for surveillance purposes. Our study evaluates core-genome single-nucleotide polymorphism analysis for outbreak detection and linking of sources of \textit{Salmonella enterica} serovar Typhimurium and its monophasic variants during a 7-month surveillance period in Denmark. We reanalyzed and defined 8 previously characterized outbreaks from the phylogenetic relatedness of the isolates, epidemiologic data, and food traceback investigations. All outbreaks were identified, and we were able to exclude unrelated and include additional related human cases. We were furthermore able to link possible food and veterinary sources to the outbreaks. Isolates clustered according to sequence types (STs) 19, 34, and 36. Our study shows that core-genome single-nucleotide polymorphism analysis is suitable for surveillance and outbreak investigation for \textit{Salmonella} Typhimurium (ST19 and ST36), but whole genome–wide analysis may be required for the tight genetic clone of monophasic variants (ST34).

The foodborne pathogen \textit{Salmonella} is responsible for tens of millions of human infections worldwide each year \cite{1}. It constitutes a substantial health and economic burden, especially in developing countries \cite{1}. Fast, accurate, and highly discriminatory typing methods are crucial for detecting outbreaks, identifying sources of the outbreaks, and preventing further spread of the bacteria as part of effective surveillance.

In Denmark, 1,122 cases of human \textit{Salmonella} infections were registered in 2014. \textit{Salmonella enterica} serovar Typhimurium accounted for 17.6\% of cases, and its monophasic variants accounted for 20.5\%. Cases are often associated with consumption of swine and poultry products \cite{2}.

In Denmark, as in many countries worldwide, the monophasic \textit{Salmonella} Typhimurium variants have emerged in the past decades \cite{3–8}. The monophasic variants are circulating in multiple clonal lineages, and owing to the relatively rapid emergence of the clones that often also exhibit multidrug resistance, these types of monophasic variants are considered an important epidemic health risk \cite{3,9–11}.

Whole-genome sequencing (WGS) is a widely used technique for molecular subtyping of bacteria, and it is replacing the more laborious current molecular typing methods. The vast amount of data provided by this method not only enables high-resolution typing for surveillance but also provides valuable additional data regarding further characterization of emerging clones based on genetic differences and evolutionary studies. Several studies have proven WGS-based typing to have an enhanced discriminatory power in comparison to current molecular typing methods used for \textit{Salmonella} \cite{12–19}, although few studies have evaluated WGS analysis in real-time surveillance. A retrospective study of \textit{Salmonella} Enteritidis showed that single-nucleotide polymorphism (SNP)–based WGS analysis was suitable for surveillance purposes \cite{15}. However, the authors also emphasized the importance of evaluation and interpretation of the SNP-based analysis within serovars or even lineages before applying the method in real-time surveillance.

In Denmark, surveillance of \textit{Salmonella} Typhimurium and its monophasic variants is conducted at Statens Serum Institut (human clinical isolates) and the National Food Institute (food and veterinary isolates). Surveillance is based on serotyping, drug-susceptibility testing, and multilocus variable-number tandem-repeat analysis (MLVA). The aim of our study was to evaluate WGS as a typing method for routine surveillance of \textit{Salmonella} Typhimurium and its monophasic variants. To do so, we selected an already typed collection of strains from 2013 and 2014 and reanalyzed them to redefine outbreaks and detect outbreak sources based on core-genome SNP analysis.
Material and Methods
We selected 372 isolates of *Salmonella Typhimurium* and its monophasic variants for this study (online Technical Appendix 1, https://wwwnc.cdc.gov/EID/article/23/10/16-1248-Techapp1.xlsx). The collection included 292 human clinical isolates from the national surveillance system in Denmark (Statens Serum Institut, Copenhagen) collected during January 2013—April 2013 (previously sequenced isolates) and June 2014—October 2014 (isolates sequenced during this study). During the 7 months of surveillance, 8 outbreaks were previously defined based on epidemiologic data, serotyping, drug-susceptibility testing, and MLVA (Table 1). Outbreak investigations during that period were initiated when 5 isolates with an indistinguishable MLVA profile were collected within a 4-week period. The investigations included patient interviews, typing of food and veterinary isolates, and examination of isolates with closely related MLVA (1 locus difference) and resistance profiles.

In addition, we selected 80 food and veterinary isolates linked or possibly linked to the outbreaks from the National Food Institute collection in Denmark (DTU Food, Technical University of Denmark, Lyngby, Denmark). The food and veterinary isolates were isolated from swine, poultry, cattle, and feed in 2010, 2013, and 2014.

WGS and Sequence Analysis
We analyzed all human isolates by using WGS at Statens Serum Institut’s Department of Microbiology and Infection Control and sequenced food and veterinary isolates at the Technical University of Denmark’s National Food Institute. We sequenced isolates by using an Illumina Miseq (Illumina, San Diego, CA, USA). All sequences were de novo assembled and sequence type (ST) determined. We identified core-genome SNPs by using an in-house SNP pipeline, and then analyzed a selected subgroup of isolates by using the SNP pipelines NASP (20) and CSI Phylogeny 1.2 (21). A description of sequencing procedures and sequence analysis is provided (online Technical Appendix 2, https://wwwnc.cdc.gov/EID/article/23/10/16-1248-Techapp2.pdf). We assessed quality of the sequences and excluded 6 isolates from the study because of poor quality. Additional information on sequences also is provided (online Technical Appendix 1).

Sequence reads were deposited in the European Nucleotide Archive (study accession no. PRJEB14853).

Results
We detected core-genome SNPs in the entire isolate collection by using the complete genome of *Salmonella Typhimurium* 14028S (ST19) as the reference genome. The SNP analysis resulted in 14,326 SNPs. We constructed a maximum-parsimony tree from the core-genome SNPs and observed 3 ST-specific groups (Figure 1); 1 group mainly consisted of ST19 strains, 1 solely consisted of ST34 stains, and 1 solely consisted of ST36 strains. A long branch separated all 11 Typhimurium ST36 isolates from the remaining isolates with 3,707 SNPs. Furthermore, we observed a distinct cluster of 242 isolates of ST34. The close genetic cluster included isolates of both serovar Typhimurium and monophasic variants, with the monophasic variants being most prevalent. The remaining 113 isolates clustering together were ST19, ST376, ST568, and ST2212, all identified as serovar Typhimurium.

Outbreak Investigation
We analyzed the 3 observed ST groups separately. Core-genome SNPs were detected by using an internal de novo assembled reference genome for each ST group. We examined the 8 previously defined outbreaks (outbreaks A–H; Table 1) on the basis of the genetic relatedness of the isolates, the epidemiologic data, and the food traceback investigations. We identified and redefined all 8 outbreaks on the basis of the SNP analysis (Table 2). We also plotted the distribution of the outbreaks over time (online Technical Appendix 2, Figure).

The ST36 Group
We analyzed 11 ST36 isolates and detected 3,146 core-genome SNPs with SNP distances between isolates ranging from 0 to 1,694. The previous definition of outbreak A included 6 human cases and no suspected food and veterinary isolates (Table 1). The SNP analysis clustered all 6 isolates with a SNP distance between the isolates of 0 to 7 SNPs. Nearest neighbor isolate was separated from the cluster with 143 SNPs, clearly differentiating the outbreak cluster from the remaining isolates (Figure 2; Table 2).

Table 1. Previously defined outbreaks included in the data collection in an outbreak investigation of *Salmonella enterica* serovar Typhimurium and its monophasic variants, Denmark*

<table>
<thead>
<tr>
<th>Outbreak</th>
<th>No. sequenced strains</th>
<th>Year</th>
<th>Outbreak characteristics</th>
<th>Serovar variant</th>
<th>Food/veterinary isolates linked to outbreak</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>6</td>
<td>2014</td>
<td>Same MLVA and resistance profile</td>
<td>Typhimurium</td>
<td>No</td>
</tr>
<tr>
<td>B</td>
<td>35</td>
<td>2013</td>
<td>Same MLVA, various resistance profiles</td>
<td>Typhimurium</td>
<td>Yes</td>
</tr>
<tr>
<td>C</td>
<td>7</td>
<td>2013</td>
<td>Same MLVA and resistance profile</td>
<td>Typhimurium</td>
<td>No</td>
</tr>
<tr>
<td>D</td>
<td>8</td>
<td>2013</td>
<td>Same MLVA, various resistance profiles</td>
<td>Typhimurium</td>
<td>No</td>
</tr>
<tr>
<td>E</td>
<td>24</td>
<td>2014</td>
<td>Same MLVA, various resistance profiles</td>
<td>Monophasic</td>
<td>Yes</td>
</tr>
<tr>
<td>F</td>
<td>19</td>
<td>2014</td>
<td>Various MLVA, same resistance profile</td>
<td>Monophasic</td>
<td>Yes</td>
</tr>
<tr>
<td>G</td>
<td>14</td>
<td>2014</td>
<td>Same MLVA, various resistance profiles</td>
<td>Monophasic</td>
<td>No</td>
</tr>
<tr>
<td>H</td>
<td>5</td>
<td>2014</td>
<td>Same MLVA and resistance profile</td>
<td>Monophasic</td>
<td>Yes</td>
</tr>
</tbody>
</table>

*MLVA, multilocus variable-number tandem-repeat analysis.*
The ST19 Group

The ST19 group comprised 98 human isolates and 5 food and veterinary isolates. Within the ST19 group, we detected 6,549 SNPs with distances between isolates ranging from 0 to 982 SNPs (Figure 3). Two outbreaks (B and C) were detected in 2013; B comprised 35 human isolates, and C comprised 7 human isolates (Table 1).

A tight genetic cluster with 0 to 4 SNP differences between the isolates comprised all 35 isolates previously defined in outbreak B. Two additional human isolates from 2013 with closely related MLVA profiles were located in this cluster and regarded as part of the outbreak, as defined by the SNP analysis. Likewise, 1 isolate from 2014 clustered with the outbreak cases but was not included in the new outbreak definition because of the difference in time. Patient interviews pointed to consumption of pork as the likely contamination source. Two suspected food isolates from pork with the same MLVA profile were, at the time, collected from 2 different meat-distributing companies. However, we could not confirm a clear connection between the food isolates and the human cases. Our SNP analysis showed that the 2 suspected isolates were located in the outbreak cluster and therefore provided additional evidence that pork was the likely source of the outbreak. The cluster was separated from the nearest neighbor isolate with 34 SNPs (Figure 3; Table 2).

All 7 isolates previously defined in outbreak C had identical core-genome SNPs. No food or veterinary isolates were linked to the cases, and the outbreak cluster was separated from the nearest neighbor with 64 SNPs (Figure 3; Table 2).

The ST34 Group

Most of the isolates observed in this study were ST34, and this ST group was dominated by the monophasic variants. The ST34 group included 169 human isolates and 73 food and veterinary isolates. Within the ST34 group, we detected 5,763 SNPs with distances between isolates ranging from 0 to 811 SNPs (Figure 3). Two outbreaks (D and E) were detected in 2014; D comprised 22 human isolates, and E comprised 9 human isolates (Table 1).

A tight genetic cluster with 0 to 4 SNP differences between the isolates comprised all 22 isolates previously defined in outbreak D. Two additional human isolates from 2014 with closely related MLVA profiles were located in this cluster and regarded as part of the outbreak, as defined by the SNP analysis. Likewise, 1 isolate from 2015 clustered with the outbreak cases but was not included in the new outbreak definition because of the difference in time. Patient interviews pointed to consumption of cattle as the likely contamination source. Two suspected food isolates from cattle with the same MLVA profile were, at the time, collected from 2 different meat-distributing companies. However, we could not confirm a clear connection between the food isolates and the human cases. Our SNP analysis showed that the 2 suspected isolates were located in the outbreak cluster and therefore provided additional evidence that cattle was the likely source of the outbreak. The cluster was separated from the nearest neighbor isolate with 20 SNPs (Figure 3; Table 2).

All 9 isolates previously defined in outbreak E had identical core-genome SNPs. No food or veterinary isolates were linked to the cases, and the outbreak cluster was separated from the nearest neighbor with 31 SNPs (Figure 3; Table 2).

Table 2. New definitions of previously defined outbreaks based on core-genome SNP analysis in an outbreak investigation of *Salmonella enterica* serovar Typhimurium and its monophasic variants, Denmark*

<table>
<thead>
<tr>
<th>Outbreak</th>
<th>No. human cases</th>
<th>Included/excluded compared with previously defined</th>
<th>Food/veterinary isolates</th>
<th>Sources</th>
<th>No. SNPs</th>
<th>Maximum SNP distance</th>
<th>SNP distance from nearest neighbor</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>6</td>
<td>–</td>
<td>0</td>
<td>Swine</td>
<td>11</td>
<td>8</td>
<td>7</td>
</tr>
<tr>
<td>B</td>
<td>37</td>
<td>+2</td>
<td>2</td>
<td>Swine</td>
<td>11</td>
<td>4</td>
<td>34</td>
</tr>
<tr>
<td>C</td>
<td>7</td>
<td>–</td>
<td>0</td>
<td>–</td>
<td>0</td>
<td>0</td>
<td>64</td>
</tr>
<tr>
<td>D</td>
<td>7</td>
<td>+1/–2</td>
<td>4</td>
<td>Swine</td>
<td>2 (6)</td>
<td>2 (6)</td>
<td>3 (7)</td>
</tr>
<tr>
<td>E</td>
<td>20</td>
<td>+1/–5</td>
<td>13</td>
<td>Swine/cattle</td>
<td>6 (7)</td>
<td>3 (4)</td>
<td>3 (4)</td>
</tr>
<tr>
<td>F</td>
<td>22</td>
<td>+3</td>
<td>2</td>
<td>Cattle</td>
<td>4</td>
<td>3</td>
<td>20</td>
</tr>
<tr>
<td>G</td>
<td>9</td>
<td>–</td>
<td>0</td>
<td>Swine</td>
<td>0</td>
<td>0</td>
<td>4 (7)</td>
</tr>
<tr>
<td>H</td>
<td>5</td>
<td>–</td>
<td>1</td>
<td>Swine</td>
<td>0</td>
<td>0</td>
<td>4 (7)</td>
</tr>
</tbody>
</table>

*Previous outbreaks shown in Table 1. SNPs in parentheses are derived from reanalysis of closely related clusters with an internal de novo assembled reference genome. SNP, single-nucleotide polymorphism.
possibly linked food and veterinary isolates, and 5 outbreaks were detected (outbreaks D–H; Table 1).

The SNP analysis of the 242 isolates resulted in 1,488 core-genome SNPs. SNP distances ranged from 0 to 95. Based on core-genome SNPs, the genetic relation between isolates was distinctly more close in comparison to ST36 and ST19. For some clusters, few SNPs separated the isolates, so defining outbreaks based on the analysis was complicated. We recalculated 2 close clusters, which included outbreaks D, E, and H, separately with an internal de novo assembled reference genome to obtain a higher resolution (Figure 4; Table 2). The recalculations added a few extra SNPs; however, conclusions were still not clear cut. Analyzing statistics from our SNP pipeline revealed that ≈20% of the reference genome was discarded when all sequences were analyzed using the closed reference genome (Table 3). Likewise, in some cases, 10% of the reference genome was not used when analyzing an apparently closely related cluster separately. To rule out whether the disregarded data were attributable to the SNP pipeline used, we additionally analyzed the cluster including outbreaks E and H by using the 2 alternative core-genome SNP pipelines NASP (20) and CSI Phylogeny 1.2 (21). From our in-house pipeline, we identified 374 core-genome SNPs within this cluster. The NASP pipeline identified 404 SNPs, and CSI Phylogeny identified 361 SNPs. No further obvious changes were observed in the overall phylogeny in this cluster or for the resolution within the outbreaks, supporting the robustness of our pipeline.

Outbreak D, detected in 2013, previously comprised 8 human isolates. From our analysis, we could include 1 additional human case and excluding 2, on the basis of the genetic relatedness of the isolates (Figure 4). Patient interviews revealed a likely source being consumption of pork from a specific butcher. No relevant food samples from the butcher, patient households, or the companies distributing meat to the butcher were available. Isolates with the same MLVA profile were, at the time, isolated from different slaughterhouses. The SNP analysis linked 3 swine isolates collected from 3 different slaughterhouses with 2–4 SNP differences with the human cases. Another food isolate from pork separated with 6 SNPs was also likely connected to the outbreak. Four food and veterinary isolates, also separated with few SNPs to the outbreak cases, were collected in 2014 and therefore not considered as part of the outbreak. The SNP analysis provided additional evidence for the connection with consumption of pork, further indicating multiple possible sources and the presence of the strain in the food production in 2014.

A large outbreak (outbreak E) was detected in 2014; our collection included 24 isolates from that outbreak. Of the 24 isolates, 23 clustered with few SNP differences (Figure 4). Because the outbreak isolates were located in a closely related cluster in the ST34 group, a clear outbreak definition based on SNP analysis was difficult to make. One human isolate was clearly separated from the cluster and could be excluded as an outbreak case. One additional isolate with a different MLVA profile clustered with 0 SNPs to cases and was included. The probable source of the outbreak was connected to consumption of pork. Samples from swine with the same MLVA profile were collected from a slaughterhouse and suspected as the primary source. Traceback investigations pointed to a specific swine herd with high *Salmonella* carriage rate supplying swine to the slaughterhouse during the same period. We collected additional samples from a company cutting and distributing meat from the slaughterhouse and from products from companies receiving meat from the primary sources. Isolates from the suspected primary and secondary sources clustered with 0 SNPs to outbreak cases, confirming the connection. Further analysis of the accessory genome identified the presence of 1 region unique for 20 of the human outbreak isolates and the 13 confirmed
food and veterinary isolates. The identified region had an approximate size of 3,600 bp and contained a ColRNA1-like (92% sequence identity) compatibility gene related to plasmids, 2 genes associated with plasmid regulation (copG and rop), and 1 hypothetical protein (99% protein similarity with predicted plasmid protein identified in an enteropathogenic Escherichia coli 0119:H6 strain [GenBank accession no. AP014807.1]). The region was located on an entire single contig with a higher average read coverage and ≈10% lower GC-content than the average of the genome. The nearest neighbor isolates considered for inclusion in the outbreak did not harbor the plasmid-related region and were not included in the final definition (Table 2).

Our results showed that all 19 isolates defined in 2014 as part of outbreak F clustered with 0 to 3 SNPs differences. Three additional human isolates (all with different MLVA profiles) clustered with 0 SNPs to cases and were included in the new cluster definition. The nearest neighbor isolate was located 20 SNPs away, and the outbreak could clearly be defined. Outbreak and food traceback investigations identified beef as the most likely source. We confirmed 2 samples from companies with 0 to 3 SNPs differences with human cases (Figure 4; Table 2).

Likewise, outbreak G was clearly defined by the SNP analysis. Nine human isolates clustered with 0 to 2 SNPs and separated from nearest neighbor with 31 SNPs. Five isolates previously included were distantly related to the cluster and not included in the new outbreak definition (Figure 4; Table 2). No linked food and veterinary isolates were available; however, we did observe a geographic connection to northern Denmark. Three of 5 excluded patients were interviewed, and no travel to northern Denmark was reported, further indicating the exclusion of the patients from the outbreak.

Last, all 5 isolates previously defined in outbreak H clustered with 0 SNPs. Consumption of a specific swine product was reported by 2 case-patients, and a food sample from the specific product clustered with 0 SNPs (Figure 4; Table 2).
Influence of Reference Genome

We examined the influence of the choice of reference genome used in the SNP analysis on the cluster formation. As mentioned previously, we analyzed each ST group and each outbreak separately by using an internal de novo assembled reference genome for each ST group and outbreak. We evaluated the size of core-genome used, percentage of reference genome used, and number of called SNPs (Table 3). Our results showed that using an internal reference genome for ST36 yielded an extra 259 SNPs compared with using the complete ST19 genome 14028S. Using an internal reference for ST19 resulted in 47 fewer SNPs and using an internal reference for ST34 resulted in 2 fewer SNPs compared with using the 14028S genome. No extra SNPs were called within the outbreaks for ST19 and ST36, regardless of reference used or group of isolates analyzed. A few extra SNPs were added when analyzing the outbreaks in ST34 separately with an internal reference. Extra SNP resolution was mostly added on the longer branches and not within tight clusters. We also evaluated including poor-quality genomes in the analysis; inclusion resulted in a considerable loss of data (Table 3).

Discussion

In this retrospective study, we showed that core-genome SNP analysis could be applied for surveillance of Salmonella Typhimurium and its monophasic variants. We were able to recover the 8 previously defined outbreaks based on the SNP analysis, epidemiologic data, and food traceback investigations. With the analysis, we could exclude unrelated human isolates and include related isolates not previously defined in the outbreaks based on MLVA. Furthermore, we were able to link possible sources to the outbreaks and reject previously suspected food and veterinary sources. In 4 out of the 8 outbreaks, we could identify the likely source of the outbreak as related to swine. In 1 outbreak, consumption of beef product was confirmed. The remaining 3 outbreaks were not linked to any known sources. In Denmark, Salmonella Typhimurium and its monophasic variants are commonly isolated from swine or pork (2), and pork meat is considered the main source of infection as observed and reported in many other countries (9,22–24).

The overall phylogeny of all isolates showed 3 groups of isolates corresponding to the ST. The most commonly isolated ST19 and ST34 isolates clustered together, with ST34 isolates defined in a distinct tight cluster. ST36 isolates were separated from ST19 and ST34 isolates with a long branch indicating a distant relation to ST19 and ST34. The distant relation is further confirmed by ST19 and ST34 belonging to e-BurstGroup 1 (eBG1), whereas
ST36 is located in eBG138, having 3 out of 7 alleles identical with eBG1 (25).

For isolates with ST19 and ST36, the outbreak clusters were well-delimited and separated, as was the case for 2 outbreaks in the ST34 group. For 3 outbreaks with ST34 isolates, the low diversity of the core-genome complicated clear conclusions based on SNP differences between isolates. With further analysis of accessory genes and information from the outbreak investigation, we could more clearly define the outbreaks. Results from the SNP analysis showed that 20% of the reference genome was discarded when analyzing the entire ST34 group, indicating a large amount of accessory data not being used in the analysis. The close core-genome correlates well with the fact that the ST34 is considered a newly expanding clone (3,10,11,24). Additionally, the large variation detected in the accessory genome correlates well with the findings of Petrovska et al. (26), which also revealed a high amount of microevolution within a clonal expansion of ST34 in the United Kingdom.

Our results show that the SNP analysis is a suitable typing method in relation to surveillance of *Salmonella* Typhimurium, with the possible exception of some lineages of the monophasic variants. Before applying the method in real time, parameters like isolates analyzed (e.g., ST and clusters), choice of reference genome, and sequence quality need to be addressed and taken into account when setting up a workflow. The reference genome used and the group of isolates analyzed had some, although mostly minor, effect on the SNPs called. However, in general, the choice of reference genome and the selection of isolates analyzed did not change our outbreak definitions. For ST19 and ST34, an overall reference genome, either ST19 or ST34, could be applied. A few extra SNPs were added within outbreaks when analyzing smaller clusters of isolates or outbreaks with an internal reference genome. However, within the tight ST34 group, the few extra SNPs within outbreaks might help in defining some outbreaks more clearly. We observed the largest differences when analyzing the ST36 isolates with a close ST36 reference genome compared with results using an ST19 reference genome. This analysis identified an extra 259 SNPs, and because the ST36 group is distantly related to ST19 and ST34, we recommend using an ST36 reference genome for this group of isolates.

The parameter that did affect the outcome considerably was the quality of the genomes used. We excluded 6 genomes out of 372 isolates because of poor quality. Including these in the analysis resulted in ≈29% fewer SNPs; therefore, we recommend quality assessment of the genomes before analysis. Last, SNP analysis does not give a static value easily communicated between institutions. Adding new isolates to the analysis results in new calculations and a new core-genome. Potential output differences might occur when different pipelines are used; however, the 3 SNP pipelines used in this study did not result in any major differences in phylogeny and had no influence on the outbreak investigations. A clear consensus of the workflow, quality criteria, and the bioinformatics tools used would resolve practical issues regarding the method. Alternatively, the widely used gene-by-gene approach is faster in real-time surveillance and more easily comparable between laboratories provided identical schemes are used. However, this approach requires expensive software (unless data are uploaded to public repositories) and agreement on schemes and curation of allele databases.

Studies on other *Salmonella* serovars using SNP analysis for outbreak investigation and detection revealed a variable number of SNP cutoff values for defining outbreak clusters, ranging from 0 to 30 SNPs (14,15,17–19,27,28). Likewise, our study shows that SNP cutoffs for outbreaks vary, even within a single serovar, lineage, or clone, and evaluation from outbreak to outbreak is needed. The nature of the outbreak must be taken into account when defining a single outbreak, given that many parameters are possibly affecting the results (e.g., the genetic makeup of the serovar, routes of infection, source types, and time). The

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**Table 3.** Statistical data from core-genome SNP analysis of different subgroups with complete genome of *Salmonella enterica* serovar Typhimurium 14028S or an internal de novo assembled genome as reference in an outbreak investigation of *Salmonella* Typhimurium and its monophasic variants, Denmark*

<table>
<thead>
<tr>
<th>Isolates within</th>
<th>Selection of isolates</th>
<th>Size of core-genome used, bp</th>
<th>% Reference genome used</th>
<th>Called SNPs</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>14028S Internal de novo</td>
<td>14028S Internal de novo</td>
<td>14028S Internal de novo</td>
</tr>
<tr>
<td>All</td>
<td>All</td>
<td>3606685</td>
<td>78.16</td>
<td>13,246 (10,163)</td>
</tr>
<tr>
<td>ST36</td>
<td>All</td>
<td>4527849</td>
<td>92.97</td>
<td>2,587</td>
</tr>
<tr>
<td></td>
<td>Outbreak A</td>
<td>4604894</td>
<td>94.53</td>
<td>8</td>
</tr>
<tr>
<td>ST19</td>
<td>All</td>
<td>4314989</td>
<td>88.60</td>
<td>6,596</td>
</tr>
<tr>
<td></td>
<td>Outbreak B</td>
<td>4648245</td>
<td>95.44</td>
<td>16</td>
</tr>
<tr>
<td></td>
<td>Outbreak C</td>
<td>4711565</td>
<td>96.74</td>
<td>16</td>
</tr>
<tr>
<td>ST34</td>
<td>All</td>
<td>4055543</td>
<td>83.27</td>
<td>1,490</td>
</tr>
<tr>
<td></td>
<td>Outbreak D</td>
<td>4665201</td>
<td>95.79</td>
<td>32</td>
</tr>
<tr>
<td></td>
<td>Outbreak E</td>
<td>4706765</td>
<td>96.64</td>
<td>7</td>
</tr>
<tr>
<td></td>
<td>Outbreak F</td>
<td>4705075</td>
<td>96.61</td>
<td>9</td>
</tr>
<tr>
<td></td>
<td>Outbreak G</td>
<td>4694550</td>
<td>96.39</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>Outbreak H</td>
<td>4734518</td>
<td>97.21</td>
<td>0</td>
</tr>
</tbody>
</table>

*SNPs in parentheses are called SNPs with inclusion of poor-quality sequences. SNP, single-nucleotide polymorphism; ST, sequence type.*
SNP analysis and other WGS methods for typing provide a higher discrimination of isolates in comparison with other conventional typing methods (12–19) and gives the advantage of many additional analyses. However, new typing approaches also lead to many new questions on how to interpret data and, in a surveillance context, how to define outbreak cases and sources. Despite the high resolution of the new typing methods, detailed and extended information on epidemiology and food traceback are still crucial elements in effective surveillance. This study has not only provided valuable information on the core-genome SNP analysis for surveillance and outbreak and source detection but also has given insight into the phylogenetic relationships between isolates of Salmonella Typhimurium and its monophasic variants in Denmark.

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Ms. Gymoese is a PhD student at the University of Copenhagen, Denmark, and is employed at Statens Serum Institut’s Department of Microbiology and Infection Control. Her main research interests are genomics using whole-genome sequence analysis to investigate and optimize surveillance of foodborne bacterial infections.

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Salmonella Typhimurium, Denmark


Address for correspondence: Mia Torpdahl, Section for Foodborne Infection, Department of Microbiology and Infection Control, Statens Serum Institut, Arntlerivej 5, 2300 Copenhagen S, Denmark; email: mtd@ssi.dk

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• Hendra Virus Vaccine, a One Health Approach to Protecting Horse, Human, and Environmental Health
• Possible Role of Songbirds and Parakeets in Transmission of Influenza A(H7N9) Virus to Humans
• Hantavirus Infections among Overnight Visitors to Yosemite National Park, California, USA, 2012

• Comparison of Imported Plasmodium ovale curtisi and P. ovale wallikeri Infections among Patients in Spain, 2005–2011
• Coxiella burnetii Seroprevalence and Risk for Humans on Dairy Cattle Farms, the Netherlands, 2010–2011

EMERGING INFECTIOUS DISEASES
Investigation of Outbreaks of *Salmonella enterica* Serovar Typhimurium and Its Monophasic Variants Using Whole-Genome Sequencing, Denmark

Technical Appendix 2

Strain Collection

For this study, 372 isolates of *Salmonella* Typhimurium and its monophasic variants were selected, including 292 human clinical isolates and 80 food and veterinary isolates (Technical Appendix 1). Human isolates were collected from January 2013 until April 2013 and from June 2014 until October 2014, including 8 outbreaks. Food and veterinary isolates were selected based on connection or possible connection to the outbreaks. Further information of the isolates are listed in Technical Appendix 1 (https://wwwnc.cdc.gov/EID/article/23/10/16-1248-Techapp1.xlsx)

Whole-Genome Sequencing

Pure bacterial cultures were cultivated overnight at 37°C on SSI 5% blood agar plates (SSI Diagnostica, Hillerød, Denmark). Genomic DNA was purified at SSI using Qiagen DNeasy Blood and Tissue Kit (Qiagen, Valencia, USA) according to the kit protocol, and at DTU Food using Invitrogen Easy-DNA Kit (Thermo Fisher Scientific, Waltham, USA). Initial DNA concentration was measured and quantified using the Qubit Fluorometer and dsDNA BR/HR Assay Kit (Thermo Fisher Scientific). Sample and library preparation was performed using the Nextera XT v2 DNA Library Preparation kit (used at SSI) or Nextera XT v3 DNA Library Preparation kit (used at DTU Food) (Illumina, San Diego, USA). Libraries were finally purified by Agencourt AMPpure XP System (Beckman Coulter, Indianapolis, USA), and whole-genomes were sequenced using an Illumina Miseq with paired-end technology (250 basepair reads).
Average read coverage at 22 was set as a preliminary quality assessment. Sequences are available at the European Nucleotide Archive (http://www.ebi.ac.uk/ena) under study accession number PRJEB14853.

**Sequence Analysis**

Sequence reads were de novo assembled using CLC Genomic Workbench (Qiagen) with default settings and a minimum contig size of 500 bp. Statistics from the genome assembly were used as further quality filtering and N50 values higher than 50,000 were accepted. Sequence types (ST) based on the seven gene MLST scheme for *Salmonella enterica* (1) were determined from the de novo assembled genomes using the webtool MLST 1.8 (https://cge.cbs.dtu.dk//services/MLST) (2).

Core-genome SNPs were detected using an in-house SNP-pipeline based on GATK and BWA-MEM. The core-genome was defined as positions present in all strains, including intergenic regions and with no phage masking, but filtered from recombination events. Reads were mapped and aligned against the complete genome of *Salmonella* Typhimurium strain 14028S (Accession NC_016856.1) (3) or against an internal de novo assembled reference genome using BWA v. 0.7.4 (4). Aligned reads were sorted, filtered and duplicate reads removed with Elprep v. 1.02 (5). GATK v. 2.5–2 (6) was used to call variants, and called variants were filtered by parsing the variant call files using an in-house python script. A minimum read support of 90% was required to make a variant call. Positions with less than ten times depth of coverage or with ambiguous calls in any genome sequence were excluded. Recombination regions were detected based on the base pattern for position of variants in all isolates, where each base pattern was considered a putative branch in a phylogenetic tree. Segments of identical base patterns were sorted by size and considering the frequency of the base pattern of the segments, the probability of observing a segment of a given or longer length was calculated and Bonferroni corrected. Segments with probability below a threshold of 0.05 were removed, and the frequency of the base pattern was reduced by the length of the segment.

Quality of the sequences analyzed in the SNP-pipeline was evaluated by parsing the discarded SNPs file and sorting discarded SNPs based on low depth, ambiguous calls and gaps. Four human and 2 veterinary isolates were excluded from this study based due to poor quality.
Some clusters were recalculated in the pipeline with the complete genome of strain 14028S or a closely related de novo assembled reference genome to evaluate the potential influence of reference genome on the SNP analysis and to evaluate the influence of analyzing clusters separately. Size of the core-genome and coverage of the reference genome used for SNP analysis was extracted from the pipeline output binary alignment/map files using Samtools v. 0.1.19 (7) and an in-house python script. The de novo assembled genomes used as close reference genomes in the SNP analysis were remapped against itself in the SNP pipeline, to check for false positive SNPs.

A selected subgroup of isolates was additionally analyzed using the SNP pipelines NASP (http://tgennorth.github.io/NASP) (8) and CSI-phylogeny (https://cge.cbs.dtu.dk/services/CSIPhylogeny) (9). Output data with missing SNPs (SNPs called but not present in all genomes) from the NASP pipeline were furthermore used for identification of unique regions in a selected cluster within the ST34 subgroup. Unique regions were BLAST searched against remaining isolates in the cluster using CLC Genomic Workbench. Annotation of the regions was done using PROKKA (10), and BLAST search in the NCBI database (https://blast.ncbi.nlm.nih.gov/Blast.cgi) were performed for closer identification of the regions. Possible plasmid regions were identified from de novo assembled genomes using PlasmidFinder 1.3 (https://cge.cbs.dtu.dk//services/PlasmidFinder) (11).

Construction of phylogeny of the isolates by multiple alignment of SNPs and calculation of maximum-parsimony (MP) trees were performed using Bionumerics 7.6 (Applied Maths, Sint-Martens-Latem, Belgium).

**Outbreaks as Defined by the SNP Analysis**

Distribution of the outbreaks A–H and sporadic isolates over time (Technical Appendix 2 Figure). Larger peaks in sporadic isolates were seen, especially in the 2014 period. The peaks were a result of summer season peak in general, and not due to accumulation of e.g., travel related isolates.
References


**Technical Appendix 2 Table.** Number of isolates selected and sequenced in this study, grouped according to source type, outbreak association and serovar variant

<table>
<thead>
<tr>
<th>Source</th>
<th>Total</th>
<th>Linked to outbreak*</th>
<th>Typhimurium</th>
<th>Monophasic variants</th>
</tr>
</thead>
<tbody>
<tr>
<td>Human (2013)</td>
<td>106</td>
<td>54</td>
<td>76</td>
<td>30</td>
</tr>
<tr>
<td>Human (2014)</td>
<td>186</td>
<td>68</td>
<td>70</td>
<td>116</td>
</tr>
<tr>
<td>Swine</td>
<td>64</td>
<td>14</td>
<td>21</td>
<td>43</td>
</tr>
<tr>
<td>Animal</td>
<td>3</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Food/fresh meat</td>
<td>58</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Environmental</td>
<td>3</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Poultry</td>
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<td>0</td>
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<td>9</td>
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<tr>
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<tr>
<td>Food/fresh meat</td>
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<tr>
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<td>3</td>
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<tr>
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<td></td>
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<tr>
<td>Feed</td>
<td>3</td>
<td>0</td>
<td>0</td>
<td>3</td>
</tr>
</tbody>
</table>

*Outbreak linked isolates as previously defined based on MLVA, antibiotic susceptibility testing and outbreak/food trace back investigations. Food and veterinary isolates are only marked as linked to outbreaks if the isolates were identified as sources or possible sources.

**Technical Appendix 2 Figure.** Distribution of 8 outbreaks (A–H) and sporadic isolates of *Salmonella Typhimurium* and its monophasic variants over time in weeks. The 8 outbreaks are defined based on core-genome SNP analysis.