A comparison between longitudinal shedding patterns of Salmonella Typhimurium and Salmonella Dublin on dairy farms.

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Abstract

*Salmonella* in cattle herds may behave as epidemic or endemic infections. An intensive longitudinal sampling study across all management groups and ages on six dairy farms in the UK was used to examine patterns of *Salmonella* shedding, following the prior identification of either *Salmonella* Dublin (three farms) or *Salmonella* Typhimurium (three farms) on the premises in the context of clinical salmonellosis. Individual faeces, pooled faeces and environmental samples (total 5711 samples), taken approximately every six weeks for 15 to 24 weeks, were cultured for *Salmonella*. *S*. Dublin was detected at low frequency (on any visit 0.5 to 18.3 per cent of samples positive) and most consistently in calves. By contrast, *S*. Typhimurium was isolated at higher frequency (on any visit 6.8 to 75 percent of samples positive), and in higher numbers, up to 10⁷ cfu.g⁻¹ faeces. Significantly more samples from calves were positive for *S*. Typhimurium than were positive for *S*. Dublin (50.6% versus 3.1%; p < 0.001), which was also true for milking cows (46.3% versus 4.4%; p < 0.001). The differences could help to explain the different patterns of bovine infection classically associated with these two serovars in the UK. No consistent effect upon shedding was seen among the *S*. Typhimurium-infected herds following vaccination.

Keywords: Cattle, *Salmonella*, epidemiology, zoonoses

INTRODUCTION

Clinical *Salmonella enterica* subspecies *enterica* infections in cattle in the UK have for decades been associated principally with the two serovars *Salmonella* Dublin (SD) and *Salmonella* Typhimurium (ST) (Jones and others 2004, Wray and Davies 2004, Davison and others 2005). In passive surveillance of cattle in Great Britain, *Salmonella* incidents have fluctuated between 620 and 1196 annually in the last decade. SD and ST accounted for 66% and 10%, respectively, of 887 incidents in 2010 (AHVLA 2011). Epidemic and endemic patterns of herd infection have been described for cattle *Salmonella* infections; the former involving rapid spread, commonly with eventual disappearance, and the latter showing a persistent, fluctuating incidence (Nielsen and others 2004, Sternberg and others 2008, Pradhan and others 2009). Certain serovars or strains have historically been associated with one or other of these patterns, and in the UK, ST and SD have been regarded respectively as the classic ‘epidemic’ and ‘endemic’ serovars (Wray and others 1989, Wray and Davies 2004).

SD is considered to be host-adapted, showing a preference for cattle and occurring less frequently in other hosts. In British cattle there is a steady, fluctuating incidence (Wray and Davies 2004, Carrique-Mas and others 2010). A distinctive feature is the apparently common phenomenon of long-term carriage of SD by infected adult cattle with continuous or intermittent shedding, including at times conducive to transmission, such as calving (Lawson and others 1974, Richardson 1975, Nielsen and others 2004, Wray and Davies 2004).

ST infections in cattle have undergone a series of shifts in definitive phage-type (DT) isolated over the last forty years, with DT29 being superseded by DT204 complex, which was in turn superseded by DT104 (Rabsch and others 2001, 2002, Threlfall 2005). ST DT104 now seems to be in decline in cattle, pigs, poultry and humans in Great Britain (Davison and others 2005,
Counter to the classical epidemic and endemic patterns, prolonged herd infections with ST and short-lived infections with SD may also be observed. Environmental sampling on UK dairy farms over 16 months repeatedly yielded ST on one of the seven premises that were consistently *Salmonella*-positive (Davison and others 2005). Shedding of ST by cattle has been demonstrated for up to 18 months after initial clinical incidents on several farms, and shedding has been observed to reoccur on premises after a period when it was undetectable (Davies 1997).

Many aspects of herd management have been identified as relevant to the presence or persistence of bovine *Salmonella* infection; these principally emphasise larger herd size and poor biosecurity as major risk factors (Evans and Davies 1996, Evans 1996, Trueman and others 1996, Vaessen and others 1998, Davies 2001, Huston and others 2002, Boqvist and Vagsholm 2005, Fossler and others 2005, Davison and others 2006, Nielsen and others 2007, Vanselow and others 2007a). Another theme has been the potential for the recycling or acquisition of infection through the farm environment and/or via manure or effluent. Housing of cattle and exposure of calving cows to accommodation previously occupied by sick animals were risk factors in a UK study (Evans and Davies 1996). Sick pens, calving pens and manure stores may frequently be contaminated (Fossler and others 2005), and *Salmonella* may be imported with manure (Veling and others 2002, Vanselow and others 2007b). Poor control of rodents, wild birds and cats have also been identified as risks (Evans 1996, Veling and others 2002, Boqvist and Vagsholm 2005).

Patterns of infection appear to vary considerably between herds (Fossler and others 2004, Davison and others 2006). Attempts to model non-transient *Salmonella* herd infections need to take account of factors that contribute to forming epidemic or endemic patterns. It may be helpful to our understanding of such factors to trace both endemic and epidemic infections over time. Such monitoring would ideally include the prevalence and intensity of shedding among the various age and production groups on premises, plus the survival and dissemination of the organism in the environment and in wildlife.

The present study aimed to examine some of these parameters using intensive, longitudinal sampling of livestock management groups within a small number of previously-identified ST- and SD-affected farms. This complements previous UK longitudinal studies that were broader surveys across many farms or intensive observations of outbreaks associated with a single serovar. By focusing on ST and SD the intention was to include both endemic and epidemic infections.

**METHODS**

**Farm selection**

Dairy farms with recent incidents of clinical disease involving either ST DT104 or SD in cattle and which were accessible for intensive sampling from the Animal Health and Veterinary Laboratories Agency (AHVLA) site at Weybridge (Southeast England) were recruited. Once farmers had agreed to participate, visit dates were agreed. As it was the intention to sample herds where *Salmonella* infections were non-transient and not in a final phase, if the target serovar was not isolated at the first visit, the farm was removed from the
study and an alternative farm was sought, using the above criteria. A questionnaire on herd size, management and recent medication was completed during the initial visit.

**Farm sampling**

Each farm was sampled approximately every six weeks (tables 2 and 3) for a period of 15 to 24 weeks (four to five visits in total). A stratified sampling approach was adopted, taking on each occasion up to 300 individual voided faecal pats that provided a representative spread across all management groups and ages of animals (i.e. dairy herd, calves, dry cows, heifers and bulls) present on the farm. In addition, naturally pooled faeces from collecting yards and slurry storage areas were collected. Faeces sampling in this fashion allowed repeated and non-invasive monitoring of each group of animals.

For each sample, a minimum of 100 g faeces was collected using a new disposable plastic glove and placed in a new plastic jar. Additional environmental samples taken were: dust, debris scraped from walls, bedding, feed, water from drinkers, pooled water, run-off and drainage water. Also, faeces from wild birds, rodents, other wildlife pests or animals in close association with the cattle were collected.

**Salmonella detection**

Samples were transported to the laboratory at ambient temperature, stored at 4 °C and cultured for *Salmonella* on the day of collection. Pre-enrichment in buffered peptone water (BPW) and culture on modified semi-solid Rappaport-Vassiliadis agar (MSRV; Difco 218681) then Rambach agar (Merck 107500) was performed, as described previously (Wales and others 2009).

If *Salmonella* was confirmed in the initial culture, the original samples (stored at 4 °C in their original containers) were subjected to enumeration of the pathogen using a dilution-enrichment (modified most-probable number) method when sufficient sample remained for testing. Up to 27 samples were enumerated per visit, this comprising all positive samples when fewer than 20 were available, or otherwise a subset representing all *Salmonella*-positive areas of the farm. Sample material (25 g) was mixed with 225 ml BPW (preparation ‘0’), and a 10 ml aliquot of this (preparation ‘1’) was separated and used to start a decimal dilution series in BPW. All preparations were incubated at 37 °C for 16-20 h. Preparations ‘0’ and ‘1’ were cultured using MSRV/Rambach as described above. All higher dilutions were stored at 4 °C, then cultured if either of preparations ‘0’ or ‘1’ yielded *Salmonella*. The *Salmonella* count in the original sample was then estimated from the highest dilution in which *Salmonella* was isolated.

**Salmonella typing**

A proportion of isolates were serotyped at the AHVLA *Salmonella* Reference section. When the numbers of positive samples exceeded 15 per visit, a subset of isolates was chosen, representing each age group and area of the farm. Two ST strains isolated from farms which had been recruited as ‘SD’ farms were also phage-typed.
**Statistical analyses**

Chi-square tests were used to compare unweighted proportions of *Salmonella*-positive samples, in various groups. Positive proportion ratio comparisons of *Salmonella* frequency between the major sampling strata were made within the SD and ST farm groups. Using only milking cow samples from both SD and ST farm groups, (this being the group with the largest dataset) a proportion of positive samples ratio comparison for isolating *Salmonella* from pooled versus individual samples was calculated. Confidence intervals were calculated and proportion ratio comparisons were performed using the EpiInfo statistical software package (http://www.cdc.gov/epiinfo/).

**RESULTS**

**Recruitment of farms**

Six suitable farms were visited initially, but from two of these the original serovar of interest (one each of SD and ST) could not be re-isolated. Two additional ST-positive farms were subsequently recruited, making a total of six study farms, three each with SD and ST infections, all within 180 miles from AHVLA Weybridge and separated from each other by a minimum of 18 miles. Details are given in Table 1.

**S. Dublin-infected farms**

The numbers of *Salmonella*-positive samples obtained are detailed in Table 2. Between 98 and 288 (median 170) individual bovine faeces samples and zero to 70 (median 20) pooled bovine faeces samples were taken per visit.

On Farm 1, the initial isolation of SD was made following a problem with abortions. SD was isolated from calves’ faeces at visit 1 and was also associated mainly with calves at visit 2. It was not isolated from the main dairy herd, and was not detected after visit 2. Other *Salmonella* serovars were isolated after visit 1, collectively at a higher proportion than SD, These were: Agama, Ajiobo, Goldcoast, Newport, Saintpaul and Typhimurium (non-phage typable).

On Farm 2, the initial SD isolation had been from a sick cow, and at visit 1 the sick pen where this cow had been housed was still positive for SD in the lower layers of litter. At the same visit, SD was associated mainly with the calves, and was also found in rat faeces in the calf barn. *Salmonella* was infrequently isolated from this farm, and other serovars (Ajiobo, Binza and Kimuenza) individually were isolated at similar frequencies to SD.

On Farm 3, SD was the most prevalent serovar. It was isolated at each visit, mainly from the milking herd. Other serovars isolated were: Anatum, Agama, Enteritidis, Montevideo, Typhimurium DT104b, 3,10:−:1,6, O-rough and O-rough:g,p:−.

**S. Typhimurium-infected farms**

Initial isolates were all DT104. Representative isolates of ST obtained during the study were also DT104. The numbers of *Salmonella*-positive samples obtained from the visits to farms 4 to 6 are detailed in Table 3. Between 103 and 207 (median 152) individual bovine faeces samples and 14 to 67 (median 27) pooled bovine faeces samples were taken per visit.
On Farm 4, ST was isolated from most areas, with the number of positive samples fluctuating substantially between visits. Only one non-ST isolate was found: S. Ajiobo from pet faeces, on the fifth visit.

On Farm 5, ST was first isolated from dry cows, but was found in most areas of the farm on all visits. The proportion of Salmonella-positive samples was high at each visit. At the first visit the only animal groups where Salmonella was not found were 40 bulling heifers and 20 calves, kept in a neighbour’s field. The heifers were found to be positive for Salmonella at the third visit, after they had been moved back to the main farm. At the same visit the calving pen was found to be positive (included under ‘in-calf heifers’ in Table 3).

On Farm 6, ST was first isolated after the farmer noted a drop in milk yield. All groups of animals on the farm were infected during the study period. The proportion of positive samples declined over time, most markedly in the milking herd.

Enumeration data

Most (71%) of the enumeration cultures were set up by one week after sample collection, and 90% and 100% were set up within 10 and 18 days, respectively. Following tabulation, (not shown), there was no general pattern discernable of decreasing maximum counts with sample storage time, and two of eight samples yielding $>10^4 \text{ cfu.g}^{-1}$ had been stored for 18 days before processing. Results from the 302 samples identified as positive for the target serovar on the farm (SD or ST) are summarised in the Figure. Many of these enumeration cultures did not yield Salmonella, and these were allocated to the <1 to 10 cfu.g$^{-1}$ category, as they had previously been shown to be positive.

On each of farms 1 and 2, fewer than 10 enumerated samples proved to be SD-positive. Calves were the largest source of these, and they also produced the only samples exceeding 10 cfu.g$^{-1}$. On Farm 3, 50 enumerated samples were positive for SD. These were mostly pooled or individual faecal samples from the milking herd.

On Farm 4, some of the individual samples from calves, milkers and dry cows had high counts of Salmonella, including up to $10^7 \text{ cfu.g}^{-1}$ in two individual samples from dairy and dry cows. Farm 5 had a particularly large proportion of high-count samples on all visits, with some containing up to $10^6 \text{ cfu.g}^{-1}$. The highest counts were in individual faecal samples from the milking herd. The counts of ST were lower on Farm 6 than on farms 4 and 5, as were the number of positive samples. The highest counts seen were in samples taken from calves.

Overall, the distribution of Salmonella counts was similar for both individual and pooled faeces samples, although the highest counts were only seen in individual samples. A similar fraction of individual and pooled faeces samples (80% and 75%, respectively) fell into the <1 to 10 cfu.g$^{-1}$ category. Likewise, 11% of individual and 13% of pooled faeces samples were in the >10 to 100 cfu.g$^{-1}$ category. However, no pooled samples exceeded $10^4 \text{ cfu.g}^{-1}$, whereas 2.4% of individual samples did, up to $10^7 \text{ cfu.g}^{-1}$. On farms where maximum individual faeces counts did not exceed $10^4 \text{ cfu.g}^{-1}$ (1, 2, 3 and 6), pooled sample counts were heavily distributed towards the minimum of <1 to 10 cfu.g$^{-1}$, whereas on the two farms with higher individual faeces counts, the pooled sample counts were more widely distributed.
Fifty-three of the 302 samples were not pooled or individual bovine faeces, but only three of these (effluent and manure from Farm 4 and pooled water from Farm 5) yielded counts exceeding 10 cfu.g$^{-1}$. Four samples of faeces from badger latrines on farms 1 and 2 (not included in the main dataset illustrated in the Figure) yielded non-SD and non-ST serovars of *Salmonella* at levels between 10$^4$ and 10$^6$ cfu.g$^{-1}$.

**Serovars other than Dublin or Typhimurium**

The non-SD serovars isolated from farms 1 to 3 were, in order of their frequency of identification: Ajiobo (30 isolates), Agama (22 isolates), Anatum (17 isolates), Binza (12 isolates) and Newport (eight isolates), with serovars Goldcoast, Kimuenza, Enteritidis, Montevideo, Saintpaul, Typhimurium, O rough, O rough:g:p:- and 3,10:-:1,6 being isolated between one and three times. By comparison, SD was identified 90 times from these farms (Table 5). It is likely that the O rough:g:p:- strains identified were SD, but insufficient serotyping information was available to demonstrate a definite identity. All *S. Newport* strains isolated were fully susceptible to the 16 antibiotics used in the VLA panel (VLA 2006). On two farms, faeces were collected from badger latrines: isolates of *S. Agama*, *S. Ajiobo* and *S. Saintpaul* were obtained from latrines on Farm 1 and *S. Binza* similarly from Farm 2.

The non-ST serovars isolated from farms 4 to 6 were: Ajiobo (one isolate), Durham (two isolates), O rough:i:1,2 (three isolates), and 4,5,12:-:1,2 (two isolates). The last two are likely to have been *S. Typhimurium*; however insufficient serotyping information was available to demonstrate a definite identity.

**Statistical analyses**

**Proportion of positive samples.** There was a highly significant ($\chi^2 = 942, P < 0.001$) difference between the unweighted proportions of *Salmonella* positive samples from SD (7.5%) versus ST (42%) farms. There were similar differences between the two groups of farms when considering just the principal serovars, i.e. SD or ST (Table 4). For calves, SD was isolated from 3.1% of samples from ‘SD’ farms and ST from 50.6% of samples from ‘ST’ farms ($\chi^2 = 252, P < 0.001$). For milking cows the equivalent statistics were 4.4% for SD and 46.3% for ST ($\chi^2 = 739, P < 0.001$).

On SD-infected farms only, the proportion of SD-positive samples was highest among the milking herd at 4.4%. Comparing 95% confidence intervals of ratios of the proportion of positive samples, and using the milking herd as the reference, the proportion in the milking herd was significantly higher than in other groups except calves (Table 4). The principal reason for this concentration in the milking cows was shedding of SD in this production group on Farm 3 (Table 2). For all serovars other than SD the proportion of positive samples ratio was similar among milking cows, calves and all other samples (data not shown). On ST-infected farms the positive proportion ratios were not significantly different between milking cows and calves, but they were significantly lower among other cattle and the environment, compared with milking cows (Table 4). This is a similar pattern to that seen with the SD data, although for SD the difference between milking cows and other adult stock (i.e. not calves) was far more marked than with ST.
Pooled versus individual samples. The proportion of positive samples ratio for detecting *Salmonella* in pooled compared with individual samples was 2.6 (95% confidence interval 2.4 - 2.9).

DISCUSSION

The ST isolation frequencies among important production groups (calves and milking cows) on ‘ST’ farms were significantly higher than the equivalent SD isolation frequencies on ‘SD’ farms. The highest proportion of SD shedding was observed amongst calves, plus amongst milkers on Farm 3, whereas in ST-positive farms shedding was more evenly spread across all production groups. There was a particularly low frequency of isolation of SD from adult cattle not in the milking herd. Farm 3 was visited especially soon after identification of infection, which may account for the greater shedding of SD by milking cattle than on the other two SD farms, especially if the milking herd was the origin of the infection. Alternatively, management and/or hygiene factors may have predisposed that milking herd to frequent *Salmonella* shedding.

ST was dominant on farms 4 to 6 (in respect of proportion of positive samples, maximum counts and other serovar isolates) by contrast with SD on farms 1 to 3. There may have been some ‘masking’ by ST of smaller numbers of other salmonellas through the enrichment-detection process. Nonetheless, SD exhibits long-term latency in some animals and adaptation to periparturient transmission to calves and it is possible that these characteristics may allow endemic persistence but show relatively light environmental contamination alongside more transient infections by other serovars.

The *Salmonella* density in positive samples was typically (>65% of samples on all farms) below 10 cfu.g⁻¹, regardless of serovar. However, occasional samples contained much higher numbers of *Salmonella*, all from ‘ST’ herds. It may be that substantial amplification of the organism occurs in only a small proportion of animals. Such individuals may be epidemiologically highly significant, for example if the establishment of *Salmonella* infection within a group requires a certain threshold challenge dose. There is evidence, from other longitudinal studies of other serovars, of especially prolonged or heavy shedding of *Salmonella* by a minority of individuals (Jones and others 1983, Roy and others 2001, Van Kessel and others 2007), and such individuals can substantially affect outcomes in mathematical models of bovine *Salmonella* epidemiology (Lanzas and others 2008). It is unclear in the present study whether the heavily-shedding individuals were doing so transiently or persistently.

The high prevalence, and sometimes high density, of ST shedding seems likely to have promoted the rapid spread of infection within farms 4 to 6, and to have increased the risk of the infection being carried off the premises. An ‘epidemic’ disease pattern would be expected in these circumstances. Conversely an ‘endemic’ pattern of infection may result if spread is limited and consequent immunity is patchy. This may happen if the level of shedding does not result in an infectious dose for much of the herd, or if groups are housed in different areas of the farm or on separate holdings. As a tentative hypothesis, this endemic pattern could be typical among adults with a less frequently-shed serovar such as SD. It may exceptionally be seen with a more heavily-shed serovar (such as ST) in the context of frequent introductions of immunologically naïve stock into the infected area. Mathematical models of *Salmonella* epidemiology in dairy herds (Xiao and others 2006, Lanzas and others 2008) lend support to
the significance of such factors as the duration and intensity of shedding and the quality of immune response in the development of epidemic versus endemic infection patterns.

It may be that factors such as biosecurity, feeding, management and movement of young stock and parasite control influenced the various farms’ *Salmonella* risk and subsequent epidemiology. Comprehensive risk factor modelling was outside the scope of the present study, but the suggested interplay between shedding levels, immunity, and the extent and duration of herd infections does not exclude other factors that may influence which serovar is likely to be present and how it may behave.

Temporal, weather, management and geographical effects also should be considered as explanatory factors. Despite sampling of SD farms starting, on average, earlier in the year and later after diagnosis than ST farms, no trend for a decreasing or increasing proportion of positive samples with time after initial isolation is evident, neither are peaks or troughs of positive samples seen in concert with hotter (summer) or colder (late autumn) seasons. There were no marked differences in climate or geography within the study area. Thus, gross influences of the above factors are not evident, although fuller evaluation of such potential effects would require a larger study.

Spatial and temporal clustering of dairy herd infections with SD, and potentially with ST, has been reported in the UK (Fenton and others 2009). It is therefore possible that strain relatedness within the same serovars on different farms partly accounts for the differences observed between the serovars. i.e. there was a selection bias towards closely-related strains within each serovar. Certain strains of SD have proved difficult to culture from the faeces of certain cattle(Baggesen and others 2007), and it is certainly possible that related SD strains might be responsible for the infrequent isolations of SD on the three different premises. However, the higher proportion of positive samples seen in specific management groups suggest otherwise.

*Salmonella* vaccination immediately preceded a dramatic reduction in shedding in one instance: the milking herd of Farm 6, vaccinated between visits 1 and 2. However, the dry cows on the same farm and the adult stock on farms 4 and 5 did not show a similar drop in shedding following vaccination. This suggests that vaccination had, at best, a variable effect on shedding. This contrasts with a previous report, where a rapid reduction in the shedding of ST DT104 and its long-term cessation on several closed dairy farms was associated with the use of a killed SD/ST vaccine (Davies 1997).

Besides the target serovars, all named serovars isolated in the present study have been detected in diagnostic or environmental samples from British cattle in recent years (Davison and others 2006, VLA 2008, Carrique-Mas and others 2010). On some of the six farms there were serovars common to cattle and wildlife, some of which are not often cattle-associated. Some of these have been associated in the past with badgers, including Binza, Agama, Anatum, Durham and Ajobo (Wray and others 1977, Wilson and others 2003, R. H. Davies, unpublished). Serovars Goldecast, Kimuencia, Newport and Anatum have also previously been isolated from wildlife, while Montevideo has been associated with sheep and animal feed (VLA 2008). The repeated isolation of ST DT104 from wildlife is consistent with previous investigations (Davies 1997), where ST DT104 persisted in swallows and mice after disappearing from the herd and its direct environment.
The high density of *Salmonella* found in some badger faecal samples in the current study is evidence of the reinfection risk that badgers may pose. Fortunately the badger-associated serovars are not currently major public health threats, but earlier VLA investigations on highly-contaminated premises (Davies 1997, VLA unpublished data) have shown that badgers can be infected with ST DT104, and with *S. Goldcoast* that had been associated with human disease.

The present MSRV *Salmonella* culture method appears, unusually, to be generally sensitive and specific when used with environmental samples and with cattle faeces, including those containing SD (Baggesen and others 2007, Eriksson and Aspan 2007, Carrique-Mas and Davies 2008). However even the MSRV method shows unpredictably low sensitivity with certain combinations of faeces and SD (Baggesen and others 2007), which should be borne in mind when interpreting the present data. The comparison made between individual and pooled samples supports the value of pooling when sensitive *Salmonella* detection is used, as previously reported with cattle and pig faeces (Kivela and others 1999, Arnold and Cook 2009).

In summary, the present study examined two *Salmonella* serovars (SD and ST), classically associated with different patterns of bovine infection, on a small number of farms. Distinct patterns were associated with each serovar, both in the intensity of faecal shedding and in the livestock groups most affected. Among the study herds, ST DT104 affected all parts of a farm including wildlife, and was shed by a higher proportion of cattle, and in higher numbers than was SD. This, and the development of herd immunity, could help to partially explain the epidemic potential of ST as well as the subsequent clearance of infection, in an industry where movements of potential short- or long-term carrier animals are frequent and biosecurity measures are relatively poor or difficult to apply.

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### Table 1: Details of the six dairy farms studied

<table>
<thead>
<tr>
<th></th>
<th>Farm 1</th>
<th>Farm 2</th>
<th>Farm 3</th>
<th>Farm 4</th>
<th>Farm 5</th>
<th>Farm 6</th>
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<tbody>
<tr>
<td><strong>Original isolate (date)</strong></td>
<td>S. Dublin (03/03/05)</td>
<td>S. Dublin (20/05/05)</td>
<td>S. Dublin (01/03/05)</td>
<td>S. Typhimurium (01/06/05)</td>
<td>S. Typhimurium (29/06/05)</td>
<td>S. Typhimurium (23/08/05)</td>
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<td><strong>Days to first sampling</strong></td>
<td>98</td>
<td>28</td>
<td>105</td>
<td>20</td>
<td>56</td>
<td>9</td>
</tr>
<tr>
<td><strong>Farm size</strong> *</td>
<td>Dairy herd 270</td>
<td>Dairy herd 171</td>
<td>Dairy herd 289</td>
<td>Milking 300</td>
<td>Dairy herd 150</td>
<td>Dairy herd 270</td>
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<tr>
<td></td>
<td>Calves &lt;6 mths† 80</td>
<td>Calves &lt;11 mths† 100</td>
<td>Calves &gt;6 mths† 60</td>
<td>Calves 80-100</td>
<td>Calves 100</td>
<td>Calves/young stock 190</td>
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<td></td>
<td>Bulling heifers 54</td>
<td>Bullocks 11</td>
<td>Bulls 2</td>
<td></td>
<td></td>
<td></td>
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<td><strong>Closed herd</strong></td>
<td>No</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
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<td>No</td>
</tr>
<tr>
<td><strong>Rear own replacements</strong></td>
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<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td><strong>Brought in stock</strong></td>
<td>Calves</td>
<td>Virgin bulls</td>
<td>Young stock</td>
<td>Bulls</td>
<td>Young stock</td>
<td>3% brought in (Heifers)</td>
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<td></td>
<td>In-calf heifers</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td><strong>Treatment of new stock</strong></td>
<td>New calves mixed immediately. In-calf heifers separate.</td>
<td>Mixed after 1 week</td>
<td>Mixed immediately</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td><strong>Salmonella vaccination</strong></td>
<td>Calves (Bovivac S²)</td>
<td>No</td>
<td>Last 100 cows calved plus young stock (Bovivac S³)</td>
<td>50 dry cows: 2 doses 20 days apart. All stock by 2nd visit (Bovivac S³)</td>
<td>All existing stock in June &amp; July. New calves not vaccinated. (Bovivac S³)</td>
<td>Dairy herd, dry cows &amp; in-calf heifers, 1st dose early Sept (Bovivac S³)</td>
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<tr>
<td><strong>Antibiotic treatment in last 3 months</strong></td>
<td>Injections (calves &amp; adults), intramammary and oral (adults).</td>
<td>Marbofloxacin for sick animals</td>
<td>Any sick adults or young stock treated.</td>
<td>Oxytetracycline given</td>
<td>Marbofloxacin given to Salmonella cases</td>
<td>Marbofloxacin given to Salmonella cases</td>
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<td><strong>Other animals on farm</strong></td>
<td>Dog, Pig</td>
<td>Dog, Horse, Pig</td>
<td>Dog, Pig</td>
<td>Dog</td>
<td>Cat, Dog</td>
<td>Chicken, Dog, Horse, Sheep, Guinea Fowl</td>
</tr>
<tr>
<td><strong>Wildlife problems</strong></td>
<td>Badger, Rat</td>
<td>Badger, Rat, Rabbit</td>
<td>Bird, Fox, Rat</td>
<td>Bird</td>
<td>Bird</td>
<td>Bird</td>
</tr>
</tbody>
</table>

* Numbers of animals in various management categories are given. †Months. ‡Inactivated, adjuvanted combined Salmonella Dublin and Salmonella Typhimurium vaccine, Intervet Schering-Plough Animal Health. Administration reportedly as per data sheet (two doses 3 weeks apart for adults and 2-3 weeks apart for calves).
Table 2: All *Salmonella* isolates from farms with original isolates of *Salmonella* Dublin

<table>
<thead>
<tr>
<th>Sample category</th>
<th>Farm 1</th>
<th></th>
<th></th>
<th></th>
<th>Farm 2</th>
<th></th>
<th></th>
<th>Farm 3</th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Visit 1</td>
<td>Visit 2</td>
<td>Visit 3</td>
<td>Visit 4</td>
<td>Visit 1</td>
<td>Visit 2</td>
<td>Visit 3</td>
<td>Visit 4</td>
<td>Visit 1</td>
<td>Visit 2</td>
<td>Visit 3</td>
<td>Visit 4</td>
<td>Visit 1</td>
<td>Visit 2</td>
</tr>
<tr>
<td>Milkers</td>
<td>09/06/05</td>
<td>05/09/05</td>
<td>18/10/05</td>
<td>10/08/05</td>
<td>20/09/05</td>
<td>25/10/05</td>
<td>17/06/05</td>
<td>10/08/05</td>
<td>20/09/05</td>
<td>25/10/05</td>
<td>14/06/05</td>
<td>02/08/05</td>
<td>19/09/05</td>
<td>01/11/05</td>
</tr>
<tr>
<td>(pooled samples)*</td>
<td>3$^D$/120</td>
<td>6$^D$/59</td>
<td>13$^D$/86</td>
<td>29$^D$/134</td>
<td>5$^D$/88</td>
<td>1$^D$/80</td>
<td>0 /125</td>
<td>1$^D$/100</td>
<td>8$^D$/217</td>
<td>22$^D$/129</td>
<td>11$^D$/169</td>
<td>36$^D$/158</td>
<td>2$^D$/133</td>
<td>0 /19</td>
</tr>
<tr>
<td>Dry cows</td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>In-calf heifers (pooled samples)*</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>6$^D$/22</td>
<td>0 /37</td>
<td>0 /20</td>
<td>0 /19</td>
<td>0 /19</td>
<td>0 /20</td>
<td>0 /19</td>
<td>0 /19</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bulling heifers (pooled samples)*</td>
<td></td>
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<td></td>
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</tr>
<tr>
<td>Heifers</td>
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<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Calves</td>
<td>9$^D$/78</td>
<td>6$^D$/41</td>
<td>10$^D$/37</td>
<td>2$^D$/26</td>
<td>1$^D$/49</td>
<td>0 /30</td>
<td>0 /58</td>
<td>1$^D$/40</td>
<td>0 /34</td>
<td>1$^D$/45</td>
<td>0 /41</td>
<td></td>
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</tr>
<tr>
<td>Bulls / bullocks</td>
<td>0 /6</td>
<td>1 /3</td>
<td>0 /10</td>
<td>0 /10</td>
<td>0 /10</td>
<td>0 /10</td>
<td>0 /10</td>
<td>0 /10</td>
<td>0 /10</td>
<td>0 /10</td>
<td>0 /10</td>
<td>0 /10</td>
<td>0 /10</td>
<td>0 /10</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Non-bovine stock</td>
<td>0 /3</td>
<td>0 /1</td>
<td>0 /1</td>
<td>0 /1</td>
<td>0 /1</td>
<td>0 /1</td>
<td>0 /1</td>
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<td>0 /1</td>
<td>0 /1</td>
<td>0 /1</td>
<td>0 /1</td>
<td>0 /1</td>
</tr>
<tr>
<td>Environmental</td>
<td>3 /32</td>
<td>2$^D$/17</td>
<td>1 /15</td>
<td>1 /15</td>
<td>1$^D$/23</td>
<td>0 /17</td>
<td>1 /9</td>
<td>0 /6</td>
<td>1$^D$/14</td>
<td>2 /27</td>
<td>0 /11</td>
<td>8$^D$/30</td>
<td>2 /23</td>
<td></td>
</tr>
<tr>
<td>Wildlife</td>
<td>0 /2</td>
<td>0 /4</td>
<td>3 /8</td>
<td>2 /5</td>
<td>3$^D$/9</td>
<td>1$^D$/9</td>
<td>1 /11</td>
<td>0 /2</td>
<td>0 /1</td>
<td>0 /3</td>
<td>0 /4</td>
<td>0 /4</td>
<td>0 /3</td>
<td></td>
</tr>
<tr>
<td>Pets (incl. horses)</td>
<td>2 /2</td>
<td>0 /2</td>
<td>1 /3</td>
<td>0 /3</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Feed in storage</td>
<td>0 /1</td>
<td>0 /1</td>
<td>0 /1</td>
<td>0 /1</td>
<td>0 /6</td>
<td>0 /2</td>
<td>0 /3</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>% positive</td>
<td>6.5</td>
<td>12.7</td>
<td>17.5</td>
<td>15.3</td>
<td>9.0</td>
<td>1.4</td>
<td>1.7</td>
<td>0.5</td>
<td>3.1</td>
<td>9.7</td>
<td>5.0</td>
<td>18.3</td>
<td>1.8</td>
<td></td>
</tr>
</tbody>
</table>
Table 3: All *Salmonella* isolates from farms with original isolates of *Salmonella* Typhimurium

<table>
<thead>
<tr>
<th>Sample category</th>
<th>Farm 4</th>
<th></th>
<th>Farm 5</th>
<th></th>
<th>Farm 6</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No. of positives / total samples</td>
<td></td>
<td>No. of positives / total samples</td>
<td></td>
<td>No. of positives / total samples</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Visit 1 21/06/05</td>
<td>Visit 2 16/08/05</td>
<td>Visit 3 27/09/05</td>
<td>Visit 4 09/11/05</td>
<td>Visit 5 05/12/05</td>
<td>Visit 1 24/08/05</td>
</tr>
<tr>
<td>Milkers</td>
<td>112T /179 (40)</td>
<td>27T /154 (37)</td>
<td>94T /145 (52)</td>
<td>45T /121 (20)</td>
<td>54T /133 (20)</td>
<td>66T /107 (27)</td>
</tr>
<tr>
<td>(pooled samples)</td>
<td>11T /17</td>
<td>5T /8 (14)</td>
<td>4T /36 (20)</td>
<td>9T /36 (20)</td>
<td></td>
<td>21T /39 (14)</td>
</tr>
<tr>
<td>Dry cows</td>
<td>3T /12</td>
<td></td>
<td>0 /5 (14)</td>
<td>3T /5 (14)</td>
<td>0 /2 (14)</td>
<td>1T /40 (14)</td>
</tr>
<tr>
<td>(pooled samples)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>In-calf heifers</td>
<td>11T /17</td>
<td>5T /8 (14)</td>
<td>4T /36 (20)</td>
<td>9T /36 (20)</td>
<td></td>
<td>21T /39 (14)</td>
</tr>
<tr>
<td>Bulling heifers</td>
<td>0 /19</td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Heifers</td>
<td>1T /38</td>
<td></td>
<td>0 /40 (14)</td>
<td>11T /30 (14)</td>
<td>19T /22 (14)</td>
<td>4T /11 (14)</td>
</tr>
<tr>
<td>(pooled samples)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Calves</td>
<td>26T /30</td>
<td>20T /40 (20)</td>
<td>26T /29 (20)</td>
<td>10T /31 (20)</td>
<td>7T /31 (20)</td>
<td>8T /32 (20)</td>
</tr>
<tr>
<td>(pooled samples)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bulls / bullocks</td>
<td>8T /8</td>
<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Mixed bovine areas</td>
<td>4T /40</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-bovine stock</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Environmental</td>
<td>19T /30</td>
<td>1T /12 (14)</td>
<td>8T /16 (14)</td>
<td>9T /28 (14)</td>
<td>4T /20 (14)</td>
<td>7T /12 (14)</td>
</tr>
<tr>
<td>Wildlife</td>
<td>1T /3</td>
<td>0 /2 (14)</td>
<td>0 /1 (14)</td>
<td>0 /2 (14)</td>
<td>2T /3 (14)</td>
<td>2T /2 (14)</td>
</tr>
<tr>
<td>Pets (incl. horses)</td>
<td>2T /2</td>
<td>1T /3 (14)</td>
<td>3T /6 (14)</td>
<td>0 /2 (14)</td>
<td>1T /2 (14)</td>
<td>2T /2 (14)</td>
</tr>
<tr>
<td>Feed in storage</td>
<td>0 /1</td>
<td>0 /2 (14)</td>
<td>0 /3 (14)</td>
<td>0 /2 (14)</td>
<td>0 /1 (14)</td>
<td>0 /2 (14)</td>
</tr>
<tr>
<td>Totals: fraction</td>
<td>171 /280</td>
<td>53 /259 (20)</td>
<td>136 /206 (20)</td>
<td>68 /220 (20)</td>
<td>75 /226 (20)</td>
<td>106 /237 (20)</td>
</tr>
<tr>
<td>% positive</td>
<td>61.1</td>
<td>20.5 (20)</td>
<td>66.0 (20)</td>
<td>30.9 (20)</td>
<td>33.2 (20)</td>
<td>44.7 (20)</td>
</tr>
</tbody>
</table>

*Number of samples, included in total, made up of pooled faeces from communal areas.

T *Salmonella* Typhimurium isolated.
Table 4: Comparisons between sampling categories of the proportion of samples positive for *Salmonella* serovars Dublin and Typhimurium

<table>
<thead>
<tr>
<th></th>
<th>Total positives</th>
<th>Total samples</th>
<th>Per cent positive (95% CI†)</th>
<th>Positive sample ratio (95% CI†)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>S. Dublin</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(farms 1-3)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Milkers</td>
<td>71</td>
<td>1598</td>
<td>4.4 (3.5 - 5.6)</td>
<td>1.00 (reference)</td>
</tr>
<tr>
<td>Calves</td>
<td>15</td>
<td>479</td>
<td>3.1 (1.9 - 5.1)</td>
<td>0.70 (0.40 - 1.22)</td>
</tr>
<tr>
<td>Other cattle</td>
<td>1</td>
<td>494</td>
<td>0.2 (0.0 - 1.1)</td>
<td>0.05 (0.01 - 0.33)</td>
</tr>
<tr>
<td>Environment*</td>
<td>7</td>
<td>361</td>
<td>1.9 (0.9 - 4.0)</td>
<td>0.43 (0.20 - 0.92)</td>
</tr>
<tr>
<td><strong>S. Typhimurium</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(farms 4-6)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Milkers</td>
<td>748</td>
<td>1615</td>
<td>46.3 (43.9-48.8)</td>
<td>1.00 (reference)</td>
</tr>
<tr>
<td>Calves</td>
<td>174</td>
<td>344</td>
<td>50.6 (45.3-55.8)</td>
<td>1.09 (0.97-1.23)</td>
</tr>
<tr>
<td>Other cattle</td>
<td>139</td>
<td>508</td>
<td>27.4 (23.7-31.4)</td>
<td>0.59 (0.51-0.69)</td>
</tr>
<tr>
<td>Environment*</td>
<td>118</td>
<td>312</td>
<td>37.8 (32.6-43.3)</td>
<td>0.82 (0.70-0.95)</td>
</tr>
</tbody>
</table>

* The group ‘environment’ includes samples from the environment, pets and wildlife.
† Confidence interval

Table 5: Frequency of isolation of all serovars on *Salmonella* Dublin-contaminated farms

<table>
<thead>
<tr>
<th>Serovar</th>
<th>Farm 1</th>
<th>Farm 2</th>
<th>Farm 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dublin</td>
<td>2/4</td>
<td>3/4</td>
<td>5/5</td>
</tr>
<tr>
<td>Agama</td>
<td>4/4</td>
<td></td>
<td>1/5</td>
</tr>
<tr>
<td>Ajiobo</td>
<td>4/4</td>
<td>1/4</td>
<td></td>
</tr>
<tr>
<td>Anatum</td>
<td></td>
<td>4/5</td>
<td></td>
</tr>
<tr>
<td>Binza</td>
<td>2/4</td>
<td></td>
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</tr>
<tr>
<td>Enteritidis</td>
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<td>1/5</td>
<td></td>
</tr>
<tr>
<td>Goldcoast</td>
<td>2/4</td>
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</tr>
<tr>
<td>Kimuenza</td>
<td></td>
<td>1/4</td>
<td></td>
</tr>
<tr>
<td>Montevideo</td>
<td></td>
<td></td>
<td>1/5</td>
</tr>
<tr>
<td>Newport</td>
<td>3/4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Saintpaul</td>
<td>1/4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Typhimurium</td>
<td>1/4</td>
<td>1/5</td>
<td></td>
</tr>
<tr>
<td>O rough</td>
<td></td>
<td></td>
<td>1/5</td>
</tr>
<tr>
<td>O rough:g,p:-</td>
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<td>1/5</td>
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<tr>
<td>3,10:-:1,6</td>
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<td></td>
<td>1/5</td>
</tr>
</tbody>
</table>
Figure: Salmonella counts in representative samples positive for Salmonella Dublin (farms 1 to 3) or Salmonella Typhimurium (farms 4 to 6).