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Reduced antibacterial drug resistance and blaCTX-M β-lactamase gene carriage in cattle-associated Escherichia coli at low temperatures, at sites dominated by older animals and on pastureland: implications for surveillance

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Running title: Resistance in cattle-associated E. coli
Abstract

Little is known about the drivers of critically important antibacterial resistance in species with zoonotic potential present on farms (e.g. CTX-M β-lactamase-positive *Escherichia coli*). We collected samples – monthly, between January 2017 and December 2018 - on 53 dairy farms in South West England along with data for 610 variables concerning antibacterial usage, management practices and meteorological factors. We detected *E. coli* resistant to amoxicillin, ciprofloxacin, streptomycin and tetracycline, respectively, in 2754/4145 (66%), 263/4145 (6%), 1475/4145 (36%) and 2874/4145 (69%) of all samples from faecally contaminated on-farm and near-farm sites. *E. coli* positive for *bla*CTX-M were detected in 224/4145 (5.4%) of samples. Multilevel, multivariable logistic regression showed antibacterial dry cow therapeutic choice (including use of cefquinome or framycetin) to be associated with higher odds of *bla*CTX-M positivity. Low average monthly ambient temperature was associated with lower odds of *bla*CTX-M *E. coli* positivity in samples and with lower odds of finding *E. coli* resistant to each of the four test antibacterials. This was additional to the effect of temperature on total *E. coli* density. Furthermore, samples collected close to calves had higher odds of having *E. coli* resistant to each antibacterial as well as positive for *bla*CTX-M. Samples collected on pastureland had lower odds of having *E. coli* resistant to amoxicillin or tetracycline as well as lower odds of being positive for *bla*CTX-M.

Importance

Antibacterial resistance poses a significant threat to human and animal health and global food security. Surveillance for resistance on farms is important for many reasons, including to track the impacts of interventions aimed at reducing the
prevalence of resistance. In this longitudinal survey of dairy farm antibacterial resistance, we showed that local temperature - as it changes over the course of a year - was associated with the prevalence of antibacterial-resistant *E. coli*. We also showed that prevalence of resistant *E. coli* was lower on pastureland and higher in environments inhabited by young animals. These findings have profound implications for routine surveillance and for surveys carried out for research. They provide important evidence that sampling at a single time-point and/or single location on a farm is unlikely to be adequate to accurately determine the status of the farm regarding the presence of samples containing resistant *E. coli*. 
Introduction

Antimicrobial resistance - and particularly antibacterial resistance (ABR) - is a significant global challenge. Many countries are implementing plans to reduce the use of antibacterial drugs (ABs) in food-producing animals. For example, the most recent UK five-year National Action Plan includes a target to reduce AB use (ABU) in the treatment of food-producing animals by 25% (1). In Europe, AB sales for food-producing animals fell by 20% from 2011 to 2016 (2). In the UK dairy industry, overall ABU dropped from 24 mg/kg in 2015 to 17 mg/kg in 2018 (3, 4). In 2018, additional industry-led policies were enforced in the UK that aimed to almost eliminate the use of highest-priority critically important antimicrobials (HP-CIAs) such as third- and fourth-generation cephalosporins (3GCs and 4GCs) as well as fluoroquinolones on dairy farms. One reason for reducing ABU in farming is to reduce the prevalence of ABR bacteria carried by farm animals. However, there is a need for better data on drivers of ABR in farming. More granularity of understanding is required concerning the risks of using individual ABs and other management practices. This is especially important in terms of drivers of HP-CIA resistance. A focus within HP-CIAs is on 3GC and fluoroquinolone resistance in Escherichia coli, a species commonly found in animal faeces and considered one of the most significant potential zoonotic ABR threats to humans (5).

3GC resistance is increasingly prevalent in E. coli causing infections in humans (6) and is also found in farmed and domestic animals around the world (7). The production of CTX-M (an extended-spectrum β-lactamase) is the most common mechanism of 3GC resistance in E. coli in humans in the UK; for example, in a recent study of urinary E. coli from humans in South West England, 82.2% of 3GC-resistant isolates carried bla_{CTX-M} (8).
The objective of this study was to describe the prevalence of 3GC-resistant *E. coli* carrying *bla*\textsubscript{CTX-M} as well as *E. coli* resistant to amoxicillin, tetracycline, streptomycin and the fluoroquinolone ciprofloxacin found in faecally contaminated environments of dairy cattle in a geographically restricted population of dairy farms in South West England. These are, or represent, ABs widely used on dairy farms in the UK (3, 4). Furthermore, this study investigated environmental, ABU and management practice risk factors for the presence of such *E. coli*.

**Results**

**Prevalence and PCR characterisation of 3GC-resistant *E. coli* from dairy farms**

4581 samples were collected from faecally contaminated sites on 53 dairy farms. Samples were collected on each farm monthly between January 2017 and December 2018. 4145 samples were positive for growth of *E. coli* on non-selective agar. Of these, 384/4145 (9.3%) samples representing 47/53 (88.7%) of farms were positive for growth of *E. coli* on agar containing the 3GC cefotaxime. From these, 1226 3GC-resistant isolates were taken forward for PCR testing for possible cephalosporinase genes of interest (GOIs): *bla*\textsubscript{CTX-M} (groups 1, 2, 8, 9 and 25), *bla*\textsubscript{CMY}, *bla*\textsubscript{DHA} and *bla*\textsubscript{SHV}. Over half (648/1226; 52.7%) of all isolates tested were found to harbour *bla*\textsubscript{CTX-M} genes. Of these, 547/648 (84.4%) were of group 1, 99/648 (15.3%) were of group 9, and, in one case, both gene groups were identified. Twelve isolates harboured a *bla*\textsubscript{CMY} gene – one alongside *bla*\textsubscript{CTX-M} group 1 – and one isolate was *bla*\textsubscript{DHA-1}-positive. No isolates were positive for *bla*\textsubscript{SHV} and the remaining 566/1226 (46.2%) isolates were PCR-negative for all GOIs. These isolates were hypothesised to hyper-produce the
chromosomally encoded AmpC β-lactamase; some of these isolates have been characterised in detail in a separate study (9).

Farm- and sample-level risk factors for bla<sub>CTX-M</sub> E. coli positivity

Based on PCR, carriage of bla<sub>CTX-M</sub> was the most common mechanism of 3GC-resistance in E. coli from dairy farms in this study. Identifying management practice- and AMU-associated risk factors for bla<sub>CTX-M</sub> E. coli positivity was therefore considered to be an important objective. Overall, 5.4% (224/4145) of samples representing 42/53 (79.2%) of farms contained 3GC-resistant E. coli confirmed to carry bla<sub>CTX-M</sub> using PCR. Positivity for bla<sub>CTX-M</sub> E. coli was three times higher in samples collected from the environments of calves (Calf samples; 98/631 [15.5%] of samples) than overall (Table 1).

Given the high positivity rate for bla<sub>CTX-M</sub> E. coli in Calf samples, a separate risk factor analysis using only Calf data was performed. One farm-level fixed effect and three sample-level fixed effects were retained in the final multilevel, multivariable logistic regression model (Table S1, Table 2). The use of cefquinome or framycetin dry cow therapies were both associated with higher odds of bla<sub>CTX-M</sub> E. coli positivity, as was higher average monthly temperature. Plotting sample-level positivity for E. coli carrying bla<sub>CTX-M</sub> versus average monthly temperature revealed that the relationship between positivity and temperature was primarily driven by low bla<sub>CTX-M</sub> E. coli positivity rates in months where the average temperature was below 10°C (Figure 1A).

Risk factor analysis was also performed for the full dataset. One farm-level fixed effect and three sample-level fixed effects were retained in the final model (Table
Interestingly, this model revealed that bla<sub>CTX-M</sub> E. coli was less likely to be found in samples obtained from pastureland, which included publicly accessible farmland (Footpaths) compared with other sample types. Analysis of the full dataset confirmed what was seen with the Calf dataset: higher average monthly temperature was associated with a higher odds of bla<sub>CTX-M</sub> E. coli positivity. Again, visualisation of the data confirmed that this was primarily driven by a reduction in bla<sub>CTX-M</sub> E. coli positivity rate in months with an average temperature below 10°C (Figure 1B).

Strikingly, farm-level positivity for bla<sub>CTX-M</sub> E. coli at the sequential monthly sampling visits was higher in warmer months and lower in the coldest month (Figure 2A).

A Bayesian logistic regression model was also constructed in which the effect of total farm ABU and specifically total 3GC and 4GC use were tested as predictors for bla<sub>CTX-M</sub> E. coli positivity in the total dataset, with 102 potential predictors included. The impact of temperature (odds ratio 1.71 [1.42, 2.05]) on bla<sub>CTX-M</sub> E. coli positivity was also retained in this alternative model (Table S3).

Defining sample-level positivity for bla<sub>CTX-M</sub> E. coli is dependent upon finding bla<sub>CTX-M</sub> using PCR in E. coli colonies that have grown on agar containing cefotaxime. If bla<sub>CTX-M</sub> E. coli in a sample exist at such a low density that they are not detected using selective agar, the sample will be falsely identified as negative for bla<sub>CTX-M</sub> E. coli. This impact of bacterial density on assay sensitivity is an important consideration in the context of the finding that bla<sub>CTX-M</sub> positivity is low at low temperatures. To account for this, the logistic link function was adjusted (see Supplementary). This only modestly altered the effect sizes and the p-values for the risk factors (Figure S1), confirming that the effect of low temperature on bla<sub>CTX-M</sub> E. coli positivity was additional to its effect on E. coli prevalence. All values in Tables 2 and 3 come from models with this adjusted logistic link function applied.
Prevalence and risk factor analysis for *E. coli* resistant to other antibacterial classes

All 4145 samples positive for growth of *E. coli* on non-selective agar were also tested for detectable numbers of *E. coli* resistant to four non-cephalosporins: amoxicillin, tetracycline, streptomycin, and ciprofloxacin, the last being representative of the HP-CIA class, the fluoroquinolones. Resistance to amoxicillin and tetracycline were the most prevalent types of resistance found, with ciprofloxacin resistance being the least commonly detected (Table 4).

Using a Bayesian logistic regression method, factors associated with the risk of a sample being positive for *E. coli* resistant to each of the test ABs were identified from the total dataset. As seen for *bla*\textsubscript{CTX-M} *E. coli*, where and when the samples were collected were more consistently associated with the odds of finding resistant *E. coli* in a sample than farm-level management practices or ABU, with all four models showing a positive association between average monthly temperature and the odds of finding resistant *E. coli* in a sample (Table 5). Also consistent across all models was the significance of sampling different areas of the farm. Again, as with *bla*\textsubscript{CTX-M} *E. coli*, samples from the environments of calves were more likely to harbour *E. coli* resistant to all four ABs than samples collected elsewhere on the farm. Samples collected from pastureland were, like *bla*\textsubscript{CTX-M} *E. coli*, negatively associated with the presence of amoxicillin and tetracycline resistance (Table 5).

Full results for all variables tested can be found in Table S4. Re-running the models with sceptical priors did not affect the results; only very small differences in the model coefficients were observed (Table S5).
Discussion

Prevalence of \textit{bla}_{CTX-M} positive \textit{E. coli}

This study is unique in its scale: extensive management practice and ABU data along with multiple samples from multiple farms were collected monthly over a two-year period. Overall, 224/4145 (5.4\%) of samples were positive for \textit{E. coli} carrying \textit{bla}_{CTX-M}. This is similar to previously calculated \textit{bla}_{CTX-M} \textit{E. coli} carriage of approximately 7\% in Danish slaughter pigs (10) and 3.6\% in UK broiler chickens and turkeys (11). Various studies have identified much higher prevalence in chicken meat, but this could be due to cross-contamination at slaughter and in the food chain (12, 13).

Studies examining the prevalence of \textit{bla}_{CTX-M} \textit{E. coli} in human populations have shown mixed results. A prevalence of 65.7\% was found amongst commensal isolates in Thailand (14). In the UK, a study across four regions reported commensal faecal carriage of \textit{bla}_{CTX-M} \textit{E. coli} to be approximately 7\% (15). A recent analysis of human urinary samples from the same region as the farms surveyed in this study gave a sample-level prevalence of \textit{bla}_{CTX-M} \textit{E. coli} of approximately 5\% (8). It should be noted that all farm samples in the present study were from faecally contaminated sites, not individual animals, and so it is possible that the number of animals carrying \textit{bla}_{CTX-M} \textit{E. coli} was much lower than the reported sample-level prevalence. Direct comparison with human and other farm animal carriage studies should therefore be made with caution.
Impact of temperature on the odds of finding resistant and \( \text{bla}_{\text{CTX-M}} \)-positive \( E. \) \( coli \)

This study found 42/53 (79\%) of farms to be positive for \( \text{bla}_{\text{CTX-M}} \) \( E. \) \( coli \), based on phenotypic analysis and PCR. This was higher than seen in other studies using similar methodology; for instance, 17/48 (35\%) of randomly selected UK dairy farms (16) and 5/25 (20\%) of farms in Ohio (17) have been previously shown to be positive. In the present study, samples were collected each month over two years, hence the chances of finding a positive sample on each farm may have been greater than in these earlier point-prevalence studies (16,17). When farm-level positivity for \( \text{bla}_{\text{CTX-M}} \) \( E. \) \( coli \) was plotted on a month-by-month basis (Figure 2A), the highest prevalence for a single monthly survey was 22.5\%, which fits more closely with these other studies.

Sample-level prevalence of \( \text{bla}_{\text{CTX-M}} \) \( E. \) \( coli \) was low overall (5.4\%). This contrasts with >90\% (18) or 50\% (19) of \( \text{bla}_{\text{CTX-M}} \) \( E. \) \( coli \) in samples taken from bovine faecal pats. This difference could be because these earlier studies used enrichment culture prior to testing for resistance at sample level, which increases the chances of finding resistance at sample level. Another explanation for the difference, is the large number of samples collected in the present study, particularly over winter, given low temperature was associated with low \( \text{bla}_{\text{CTX-M}} \) \( E. \) \( coli \) positivity (Fig. 1A; 1B). Indeed, the observation that average monthly temperature had a significant effect on \( \text{bla}_{\text{CTX-M}} \) \( E. \) \( coli \) positivity (Table 3), as well as positivity for resistance to amoxicillin, ciprofloxacin, streptomycin, and tetracycline (Table 5) highlights problems with studies where a single time-point or sampling season is used. Figure 2 shows the stark impact of this in real terms: \( \text{bla}_{\text{CTX-M}} \) \( E. \) \( coli \) positivity and positivity for
Ciprofloxacin-resistant *E. coli* at farm level was zero in February, the coldest month of the year (based on average temperature; **Fig 2A, B**).

Whilst average annual temperature found at locations across an entire continent has previously been shown to impact average ABR levels at those locations (20), the finding that periods of low temperatures were associated with lower prevalence of a dominant cause of ABR - and particularly HP-CIA resistance at a given location during the course of a year - is particularly important. This observation also leads to concern about the impact of climate change - and especially increasing temperatures - on attempts to reduce ABR. Whilst temperature was associated with the total number of *E. coli* found in each sample, this was accounted for using a measurement error method incorporated into the model; as such, the effect of temperature on ABR or *bla* _CTX-M_-positive *E. coli*, whilst in part driven by the effect on total *E. coli* number, also had an independent association suggestive of a temperature-dependent fitness burden of resistance.

**High levels of ABR and *bla* _CTX-M_ positive *E. coli* in farm locations dominated by young animals and low levels on pastureland**

There were clear differences in the risk of encountering *bla* _CTX-M_ *E. coli* at different sites on a farm (e.g., 15.5% in Calf samples, 4.1% in Adult samples). The Calf environment was also much more likely to have *E. coli* resistant to amoxicillin, tetracycline, streptomycin, and ciprofloxacin, so this seems to be a universal effect. Other studies have also generally found high levels of resistance in samples collected from or in the environment of younger calves (21-24). There may also be an association with temperature here since calves are generally kept in warmer
environments than are adult cows but may also be due to some physiological change with age.

This study also identified a lower odds of detecting \( \text{bla}_{\text{CTX-M}} \) \( E. \ coli \) in samples collected on pastureland compared with those collected elsewhere on the farm. This relationship also held for \( E. \ coli \) resistant to amoxicillin and tetracycline. Because pastureland may be more affected by the elements, this finding may be partly linked with the association between temperature and ABR.

**AB contamination of colostrum as a possible driver of \( \text{bla}_{\text{CTX-M}} \) \( E. \ coli \) positivity in dairy calves - evidence of direct and co-selection**

Our analysis identified a small number of specific risk factors. There was an association between calf water trough cleaning and lower odds of Calf samples having \( E. \ coli \) with \( \text{bla}_{\text{CTX-M}} \) (Table 2). Whilst there are many reasons for providing the cleanest possible drinking water, this was not seen for other ABR phenotypes and it is unclear why this association was identified. Furthermore, feeding maize silage was associated with a higher odds of finding \( \text{bla}_{\text{CTX-M}} \)-positive \( E. \ coli \) across the whole dataset (Table 3). It would be interesting to take samples of silage and to test if resistant \( E. \ coli \) survive better in this type of medium, but again this association was not seen for other resistance phenotypes. There were also associations between the odds of finding tetracycline-resistant \( E. \ coli \) and ABU, calving and rainfall (Table 5).

The most interesting AB-related association found was specifically for Calf samples. It has been shown experimentally that feeding waste (AB-contaminated) milk to calves increases faecal excretion of ABR bacteria (25). This practice is reducing on UK dairy farms and, in the analysis presented here, waste milk feeding was not
associated with an increased risk of finding ABR or $\text{bl}_\text{CTX-M}$ positive $E. \ coli$. In contrast, the choice of dry cow therapy (an AB preparation inserted into a cow’s udder between lactations to help treat or prevent mastitis) was associated with $\text{bl}_\text{CTX-M}$ $E. \ coli$ positivity in Calf samples (Table 2). It has previously been shown that colostrum from cows given cefquinome dry cow therapy is heavily contaminated with cefquinome (26), and colostrum management is a hugely important part of early life for most farmed mammals and is universally encouraged in dairy farming. In this study, cefquinome (a 4GC) dry cow therapy was most significantly associated with $\text{bl}_\text{CTX-M}$ $E. \ coli$ in Calf samples (Table 2). This can be explained by direct selection because production of CTX-M confers 4GC resistance in $E. \ coli$ (27). There was also a clear positive association between the usage of framycetin as part of a dry cow therapy combination and the odds of finding $\text{bl}_\text{CTX-M}$ $E. \ coli$ in Calf samples (Table 2). Whilst no work has been published on the contamination of colostrum with framycetin, its use as a mastitis therapy for milking cows leads to identifiable residues in milk (28), so it is highly likely to also contaminate colostrum. It is possible, therefore, that feeding of colostrum - which can be contaminated with AB used for dry cow therapy - is a driver of $\text{bl}_\text{CTX-M}$ $E. \ coli$ in calves. An alternative (or indeed an additional) explanation for this observed association is that $E. \ coli$ (a species known to be found in the udders of dairy cows [29]) that carry $\text{bl}_\text{CTX-M}$ are selected within the udder during AB dry cow therapy and contaminate colostrum alongside the AB used. Others (16) have also identified overall use of 3/4GCs as a risk factor for $\text{bl}_\text{CTX-M}$ $E. \ coli$ presence on dairy farms but have not made a link between the usage of framycetin and prevalence of $\text{bl}_\text{CTX-M}$ $E. \ coli$. However, it is not always clear whether other studies have separated out different dry cow therapies since they have tended to focus on systemic AB use. Clearly, an
aminoglycoside like framycetin cannot directly select for blaCTX-M E. coli, but aminoglycoside resistance genes are common on plasmids (30), as is blaCTX-M (27). Hence, this may be an example of co-selection, where selection of resistance to one antibacterial class increases resistance to another.

**Conclusions**

Overall, we provide important evidence that sampling at single time-points and/or limited locations on a farm are unlikely to be adequate to accurately determine the status of the farm concerning the prevalence of ABR E. coli. This makes comparisons between surveillance studies on farms - designed for research or for regulatory purposes - extremely difficult, and we urge for a standardised, multi-sample framework accounting for the differential risk factors identified here to be used in the design of future studies.
Materials and Methods

Farm recruitment and ethical approval

A convenience sample of 53 dairy farms was recruited through personal contacts, local veterinary practices, and milk processors. These represented a variety of dairy management systems, ranging from seasonally calving extensively managed herds to zero-grazed intensive systems. Recruited dairy farms were comparable to farms throughout the UK, with a median herd size of 193 compared to a UK median of 178, a median 305-day milk yield of 7488 L compared to a UK median of 8967 L, and a median somatic cell count (SCC) of 167,000 cells/mL of milk compared to a UK median of 178,000 cells/mL of milk. Antibiotic purchasing in 2016 was 26 milligrams per population corrected unit (mg/PCU) for the UK dairy industry (3) and 21 mg/PCU for represented farms.

Of the 53 farms recruited, 43 study farms were in a 50 x 50 km area defined based on the locations of 146 general practices that referred routine urine samples from human patients to the microbiology reference lab at Severn Pathology, Southmead Hospital (8). A further 10 study farms were clustered in a separate region of South West England.

All farmers gave fully informed consent to participate in the study. Ethical approval was obtained from the University of Bristol’s Faculty of Health Sciences Research Ethics Committee (ref 41562).

Farm sampling, sample characteristics and sample processing

Farms were visited monthly between January 2017 and December 2018. Samples were collected using sterile overshoes (over-boot socks) traversing farm areas.
Where access was restricted (e.g., for pens containing single or pairs of calves), samples were collected directly from the ground using gloved hands.

Samples were collected of the following six types: “Adult” samples, the faecally contaminated environment of the milking cow population - either the collecting yard, housing shed or field, collected every month from 52 farms (one farm was a youngstock-only unit). “Dry Cow” samples, the faecally contaminated environment of the dry cow population, collected from 46 farms. “Heifer” and “Calf” samples, the faecally contaminated environment of replacement dairy heifers. These categories were not mutually exclusive and reflected the animals present when the samples were collected. In addition, 209 (5%) samples were collected where no relevant animals were present. On 41 farms, approximately 10 heifers were followed per farm from birth until 18 months of age, with samples collected monthly from their environment, whether housed in a shed or out at pasture. Furthermore, 10 farms had samples collected from additional groups of pre-weaned replacement heifer calves (calves still being fed milk) to increase the number of samples available of this type. For analysis, because of the differing management practices at different life stages, replacement heifer samples were divided into those associated with pre-weaned calves (Calf samples) and post-weaned heifers (Heifer samples). “Pasture” samples from the faecally contaminated environments around the above animals whilst grazing on pastureland on 47 farms were also separated from a subset of “Footpath” samples which were taken from public footpaths or other rights of way that crossed the farm, where relevant, on 41 farms.

Characteristics for each sample were recorded on a datasheet at the time of collection. Different information was recorded depending on the sample type.
For all samples: sample number; textual description of sample location; whether animals were housed or in a field; Ordinance Survey reference for outdoor samples. For Calf samples: sample number; eartag numbers; textual description of the sample location; total number of animals in the group; presence or absence of beef calves in the group; date of birth for each calf; dry cow therapy used on the dam of each calf in the dry period before the birth. For Footpath samples: presence of livestock at time of sampling.

Samples were refrigerated (4-8°C) from collection to processing, were transferred into individual labelled sterile stomacher bags, and suspended in 10 ml.g\(^{-1}\) of phosphate buffered saline (PBS Dulbecco A; Oxoid, Basingstoke, UK). Samples were then mixed for one min in a stomacher (Stomacher 400, Seward, Worthing, UK). Samples were mixed 50:50 with sterile glycerol and aliquots stored at -80°C.

**Microbiology and PCR analysis**

Twenty microlitres of sample (diluted 1:10) were spread onto tryptone bile X-glucuronide agar (TBX; Scientific Laboratory Supplies); 20 µL of undiluted sample were spread onto TBX agar containing 16 mg.L\(^{-1}\) tetracycline, 8 mg.L\(^{-1}\) amoxicillin, 0.5 mg.L\(^{-1}\) ciprofloxacin, 16 mg.L\(^{-1}\) streptomycin or 16 mg.L\(^{-1}\) cephalxin. Plates were incubated at 37°C, and the number of blue colonies (\textit{E. coli}) counted. Samples yielding no \textit{E. coli} colonies on antibiotic-free agar were excluded from further analysis. Up to five \textit{E. coli} isolates from each cephalxin TBX agar plate were transferred onto cefotaxime (CTX, 2 mg.L\(^{-1}\)) TBX agar. All AB concentrations were chosen as those which define clinically relevant resistance in humans according to EUCAST (31). Two multiplex PCRs were performed to screen for β-lactamase genes in CTX-R \textit{E. coli}. The first was to detect \textit{bla}\textsubscript{CTX-M} groups as previously described (32) and the second
was to detect the following additional β-lactamase genes: $bla_{CMY}$, $bla_{DHA}$, $bla_{SHV}$, $bla_{TEM}$, $bla_{OXA-1}$ (8).

**Obtaining farm management information**

Four management practice questionnaires were developed. *Questionnaire 1* was completed by the researcher in the presence of the farmer at the time of consent and the first farm visit. *Questionnaire 2* was completed by the researcher in the presence of the farmer using Epicollect5 approximately six to nine months into the project. *Questionnaire 3* was completed by the researcher in the presence of the farmer using Epicollect5 approximately 12-16 months into the project. *Questionnaire 4* was completed by the researcher during a telephone call with the farmer within two months of the last visit to the farm. All questionnaires are presented in Table S6.

In total, there were 610 variables derived from the four questionnaires used in the analyses. Questions giving rise to these variables were validated and processed in the following way: Questions with a single response or zero variance were removed. Questions which had either been shown in the literature to be important risk factors or which were judged by veterinary experts to be of potential importance were selected. Categorical levels were collapsed to avoid small response counts. Some questions were combined as individually they provided detail irrelevant to this study. Repeat questions of demographic variables were averaged. Variables with missing values were removed; all variables removed for this reason were also considered likely to be of low importance. As a result, all the categorical variables were dichotomous.
Monitoring ABU

All dairy farmer participants gave permission for researchers to contact their veterinary practices and request AB prescription/sales data for a period of at least one year before the beginning of the project through the end of the project. All practices except one supplied records. This practice serviced two farms and on-farm records were used instead for these two farms. Data were assessed by a veterinary researcher to ensure consistent naming of products and quantification between practices, given a wide range of variations on product names and quantity denominators used. Usage metrics were produced using mg/PCU for the first 12 months of the project, from the date the farm enrolled on the project and had the first samples collected (range January 2017-July 2017) until 12 months after this date (range January 2018-July 2018).

Risk factor analysis

The risk factors examined fell into four categories: farm management, ABU, sample characteristics and meteorological. The first three are described above; for the last, local meteorological data were extracted from publicly available UK Met Office data (https://www.metoffice.gov.uk/pub/data/weather/uk/climate/stationdata/yeoviltnodata.txt).

Sample processing and data analysis workflows are illustrated in Figure S2. All data analysis was performed using R (https://www.r-project.org/). Two modelling approaches were used: 1) variable selection via univariable screening and stepwise model selection with a multilevel, multivariable logistic regression model and 2) a regularised Bayesian model. Both were used to analyse risk factors associated with
"bla\textsubscript{CTX-M} E. coli" positivity; the second was also used to analyse risk factors associated with positivity for *E. coli* resistant to amoxicillin, ciprofloxacin, streptomycin, and tetracycline. Sensitivity analyses were performed to test for measurement bias, to account for the fact that resistant *E. coli* were more likely to be found in a sample if there was a higher density of bacterial colony forming units. Further details of variable selection and development of the models and model checking are presented below. All code can be found at [https://github.com/HannahSchubert1/OH-STAR-modelling-code](https://github.com/HannahSchubert1/OH-STAR-modelling-code). Details of all model checking can be found in Figure S3.

**Modelling risk of \textit{bla}\textsubscript{CTX-M}  \(\beta\)-lactamase gene carriage**

Risk factor analysis was performed separately on Calf data and on the full dataset (all samples combined). Thirty-seven variables were selected for the Calf risk factor analysis and 110 for the full risk factor analysis as described below. Exploratory data analysis revealed the main source of clustering of observations was at the farm level. There were no noticeable longitudinal patterns or clustering due to the location within each farm. Random intercept logistic regression models with farm as a random effect were used throughout the analysis.

Two approaches to risk factor analysis were performed: a frequentist variable selection method (using univariable screening followed by step-down model selection and maximum likelihood estimation; Method 1) and a Bayesian method with regularization (Method 2, on the full dataset only).

**Method 1: Variable selection**

Variable selection was performed by first screening the variables for association with *E. coli* carrying *bla\textsubscript{CTX-M} using univariable, multilevel logistic regression (33) with
each variable entered as a fixed effect and with random intercepts for each farm. Continuous variables were first checked for linearity with the log odds of the outcome to assure they did not violate model assumptions. For the analysis of the full dataset, it was not possible to converge such a model for 27 of the variables (most likely due to the low number of samples that were positive for \(E. \ coli\) carrying \(bla_{CTX-M}\), so 83 variables were examined. See Tables S1 and S2 for full univariable results.

Variables with associations where \(p < 0.25\) after controlling for false discovery rate using the Benjamini-Hochberg procedure (chosen to be the most appropriate method for the exploratory nature of the project) were entered in to a multivariable, multilevel logistic regression model with random intercepts for each farm. A backwards stepwise procedure was used to further refine the model, selecting only those variables where the regression coefficient maintained \(p < 0.05\). Variables which survived this analysis were checked for multicollinearity by removing each variable in turn and checking that the confidence intervals for the estimate for each variable still overlapped. The predictive accuracy was checked using area under the Receiver Operating Characteristic Curve (0.84 for the full dataset; 0.80 for Calf data).

**Method 2: Bayesian model for predicting presence of \(bla_{CTX-M}\)-positive \(E. \ coli\)**

A farm-level random intercept model was fit using the R package BRMS (34) on the full dataset. All variables were used as predictors of \(bla_{CTX-M}\)-positive \(E. \ coli\) but were split into two groups. The first group comprised farm-level 1) total antibiotic use, 2) total third- and fourth-generation cephalosporin use, and 3) total first-generation cephalosporin use. The second group contained the remaining meteorological and farm management variables. For the first group, uninformative priors (normal distribution with a mean 0, standard deviation 5) were used; for the second group, a regularizing prior (horseshoe prior with a single degree of freedom [35]) was used.
The mean intercept was also given a diffuse prior (normal distribution with mean 0, standard deviation 5) while the standard deviation of the random effects was given a half-Student-T distribution with three degrees of freedom and a scale factor of 10 (the default in BRMS). Four independent Markov chains were sampled with 1000 warmup iterations and 1000 sampling iterations. The target acceptance criterion was increased from the default 0.8 to 0.95 to decrease the chance of sampling divergencies (although two remained, these were not deemed significant). The sampling was assumed to be well converged as the Gelman-Rubin statistic for each variable was 1.00. Results are shown in Table S3.

Accounting for measurement error

To account for measurement error of the bla_{CTX-M}-positive *E. coli* (whereby if more *E. coli* were found in a sample, the sensitivity of the test for finding bla_{CTX-M}-positive *E. coli* was higher) the logistic link function was altered to include the sensitivity and specificity of each sample, following the work by Coutinho *et al.* (36). The altered logistic link function was derived by equating the conditional probability of the true bla_{CTX-M} status \( P(Y_{\text{true}} = 1|X) \) to the conditional probability of the observed bla_{CTX-M} status, \( P(Y_{\text{obs}} = 1|X) \) for a given sensitivity (s) and specificity (e).

The specificity of the bla_{CTX-M} test was assumed to be 100% across samples whereas the sensitivity (s) was estimated separately for each observation using the number of *E. coli* colonies grown on antibiotic-free agar (k) and a minimum detectable prevalence of bla_{CTX-M}-positive *E. coli* (q) through:

\[
s = 1 - (1 - q)^k
\]
The distribution $s$ across the whole dataset using Method 1 for different values of $q$ is shown in Figure S4. While the value of $q$ did change the sensitivity associated with each observation, this change had only a negligible effect on the coefficient estimates (Figure S1A) and AUC (Fig S1B) over the range 0.01 – 1.00. The reported values used a $q$ of 0.01. The R-code for this link function is provided in Supplementary (Figure S5).

**Model for predicting presence of *E. coli* resistant to non-cephalosporins**

For each resistance phenotype – amoxicillin, ciprofloxacin, streptomycin, and tetracycline - a random intercept model with farm as the random effect was fitted using the R package BRMS on the full dataset (https://cran.r-project.org/web/packages/brms/index.html). All variables (see above) were used as predictors of resistant *E. coli* in a sample but were split into two groups. The first group (the ‘main’ variables) comprised farm-level ABU and average monthly temperature at the time of sample collection as these were the main variables hypothesised to influence resistance. For all models, total ABU, streptomycin usage, tetracycline usage, amoxicillin usage, fluoroquinolone usage and cefalexin usage were included as predictors. For each model, the usage of additional antibacterial drugs was tested if they were hypothesised to be important in selecting for resistance to the relevant model: for the amoxicillin model, first-generation cephalosporins, penicillins, potentiated amoxicillin; for the ciprofloxacin model, novobiocin, third- and fourth-generation cephalosporins; for the streptomycin and tetracycline models, no additional ABU variables were tested. The second group of variables (the ‘regularised’ variables) contained the remaining farm management variables.
For the first group (the ‘main’ variables), uninformative priors (normal distribution with a mean 0, standard deviation 5) were used; for the second group (the ‘regularised’ variables), a regularising prior (horseshoe prior with a single degree of freedom) was used. The mean intercept was also given a diffuse prior (normal distribution with mean 0, standard deviation 5) while the standard deviation of the random effects was given a half-Student-T distribution with three degrees of freedom and a scale factor of 10 (the default in BRMS). Four independent Markov chains were sampled with 1,000 warmup iterations and 10,000 sampling iterations. The target acceptance criterion was increased from the default 0.8 to 0.98 to decrease the chance of sampling divergencies in all models. Measurement error based on *E. coli* density was accounted for as described above. All reported results used a q of 0.01.

In addition, the measurement error of temperature was considered because only an average monthly temperature was available. Data where daily temperature was recorded for a (different) twelve-month period were used to calculate an estimated average monthly standard deviation over the twelve months (2.89). The ‘me’ function in the package ‘BRMS’ (https://rdrr.io/cran/brms/man/me.html) was used, whereby the monthly temperature variable was assumed to vary with a standard deviation of 2.89 (0.61 when scaled as the temperature was scaled to mean 0 and standard deviation 1 before entering the model; the actual standard deviation of the temperature was 4.7).

To further test the robustness of the model outputs, all models were re-run using sceptical priors, whereby the prior was set to the opposite of what would be expected. So, for all main variables, if there was an association, a positive association would be expected given prior knowledge. To test the models with sceptical priors, a prior for the main variables was given as a normal distribution with
mean -0.5 and a narrow standard deviation 2 (i.e. assuming a negative correlation with low variation; the opposite of expected).

**Bayesian model checking**

Convergence is assumed to be good with a Gelman-Rubin statistic (Rhat) of 1.00 for all variables across all models. Figure S6 shows trace plots for the associated variables, also providing evidence of good convergence. There were no divergence issues reported for any of the models. Table S4 shows full results from all models, including odds ratios, 95% credible intervals and effective sample sizes. Figure S7 shows the posterior distributions of the associated variables. Re-running the models with sceptical priors did not alter the model conclusions (Table S5).

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**Transparency declarations**

D.C.B. was president of the British Cattle Veterinary Association 2018-19.

Otherwise, the authors declare no competing interests. Farming and veterinary businesses who contributed data and permitted access for sample collection were not involved in the design of this study or in data analysis and were not involved in drafting the manuscript for publication.

**Author Contributions**

Conceived the Study: D.C.B., K.K.R., M.B.A.


Initial Drafting of Manuscript: H.S., K.K.R., M.B.A.

Corrected and Approved Manuscript: All Authors
### Table 1. Prevalence of *E. coli* carrying *bla*<sub>CTX-M</sub> at farm and sample levels

<table>
<thead>
<tr>
<th>Sample type</th>
<th>Farm level</th>
<th>Sample level</th>
</tr>
</thead>
<tbody>
<tr>
<td>Overall</td>
<td>Total sample size</td>
<td>53</td>
</tr>
<tr>
<td></td>
<td>Total (%) positive for CTX-M-carrying <em>E. coli</em></td>
<td>42 (79.2%)</td>
</tr>
<tr>
<td>Adult</td>
<td>Total sample size</td>
<td>52</td>
</tr>
<tr>
<td></td>
<td>Total (%) positive for CTX-M-carrying <em>E. coli</em></td>
<td>25 (48.1%)</td>
</tr>
<tr>
<td>Dry Cow</td>
<td>Total sample size</td>
<td>46</td>
</tr>
<tr>
<td></td>
<td>Total (%) positive for CTX-M-carrying <em>E. coli</em></td>
<td>7 (15.2%)</td>
</tr>
<tr>
<td>Calf</td>
<td>Total sample size</td>
<td>51</td>
</tr>
<tr>
<td></td>
<td>Total (%) positive for CTX-M-carrying <em>E. coli</em></td>
<td>33 (64.7%)</td>
</tr>
<tr>
<td>Heifer</td>
<td>Total sample size</td>
<td>41</td>
</tr>
<tr>
<td></td>
<td>Total (%) positive for CTX-M-carrying <em>E. coli</em></td>
<td>18 (44%)</td>
</tr>
<tr>
<td>Pastureland</td>
<td>Total sample size</td>
<td>47</td>
</tr>
<tr>
<td></td>
<td>Total (%) positive for CTX-M-carrying <em>E. coli</em></td>
<td>8 (17%)</td>
</tr>
<tr>
<td>Pastureland that is publicly accessible (Footpath)</td>
<td>Total sample size</td>
<td>41</td>
</tr>
<tr>
<td></td>
<td>Total (%) positive for CTX-M-carrying <em>E. coli</em></td>
<td>8 (20.0%)</td>
</tr>
</tbody>
</table>
Table 2. Fixed effects from the multilevel, multivariable logistic regression model predicting \( \text{bla}_{\text{CTX-M}} \) \( E. \text{coli} \) positivity performed on Calf samples

<table>
<thead>
<tr>
<th>Risk factor</th>
<th>Odds ratio [95% confidence interval]</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Use of cefquinome dry cow therapy in the last six months</td>
<td>4.18 [2.11, 8.25]</td>
<td>0.0003</td>
</tr>
<tr>
<td>Daily water trough cleaning</td>
<td>0.44 [0.29, 0.69]</td>
<td>0.0002</td>
</tr>
<tr>
<td>Average monthly temperature</td>
<td>1.57 [1.20, 2.06]</td>
<td>0.0008</td>
</tr>
<tr>
<td>Use of framycetin dry cow therapy in the last six months</td>
<td>1.91 [1.01, 3.61]</td>
<td>0.04</td>
</tr>
</tbody>
</table>
Table 3. Fixed effects from the multilevel, multivariable logistic regression model predicting $bla_{CTX-M}$ *E. coli* positivity performed on the full dataset

<table>
<thead>
<tr>
<th>Risk factor</th>
<th>Odds ratio [95% confidence interval]</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample taken from the environment of pre-weaned heifers</td>
<td>4.52 [3.25, 6.27]</td>
<td>&lt;0.0000001</td>
</tr>
<tr>
<td>Average monthly temperature</td>
<td>1.61 [1.36, 1.90]</td>
<td>0.0000001</td>
</tr>
<tr>
<td>Sample taken from pastureland</td>
<td>0.32 [0.17, 0.61]</td>
<td>0.0004</td>
</tr>
<tr>
<td>Feeding of maize silage</td>
<td>3.28 [1.50, 7.18]</td>
<td>0.002</td>
</tr>
</tbody>
</table>
Table 4. Farm and sample-level prevalence of resistance to non-cephalosporins

<table>
<thead>
<tr>
<th>Antibacterial drug</th>
<th>Farm level resistance</th>
<th>Sample level resistance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amoxicillin</td>
<td>53/53 (100%)</td>
<td>2754/4145 (66%)</td>
</tr>
<tr>
<td>Ciprofloxacin</td>
<td>49/53 (92%)</td>
<td>263/4145 (6%)</td>
</tr>
<tr>
<td>Streptomycin</td>
<td>53/53 (100%)</td>
<td>1475/4145 (36%)</td>
</tr>
<tr>
<td>Tetracycline</td>
<td>53/53 (100%)</td>
<td>2874/4145 (69%)</td>
</tr>
</tbody>
</table>
Table 5. Fixed effects from a Bayesian model predicting resistant *E. coli* positivity performed on the full dataset

<table>
<thead>
<tr>
<th>Risk factor</th>
<th>Odds ratio [95% credible interval]</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Amoxicillin resistance</strong></td>
<td></td>
</tr>
<tr>
<td>Average monthly temperature</td>
<td>1.91 [1.57, 2.35]</td>
</tr>
<tr>
<td>Sample taken from the environment of pre-weaned heifers</td>
<td>1.99 [1.29, 2.98]</td>
</tr>
<tr>
<td>Sample taken from pastureland</td>
<td>0.27 [0.20, 0.37]</td>
</tr>
<tr>
<td><strong>Ciprofloxacin resistance</strong></td>
<td></td>
</tr>
<tr>
<td>Average monthly temperature</td>
<td>2.14 [1.63, 2.87]</td>
</tr>
<tr>
<td>Sample taken from the environment of pre-weaned heifers</td>
<td>4.13 [2.79, 6.46]</td>
</tr>
<tr>
<td><strong>Streptomycin resistance</strong></td>
<td></td>
</tr>
<tr>
<td>Average monthly temperature</td>
<td>1.53 [1.32, 1.77]</td>
</tr>
<tr>
<td>Sample taken from the environment of pre-weaned heifers</td>
<td>1.95 [1.46, 2.51]</td>
</tr>
<tr>
<td><strong>Tetracycline resistance</strong></td>
<td></td>
</tr>
<tr>
<td>Average monthly temperature</td>
<td>1.98 [1.55, 2.55]</td>
</tr>
<tr>
<td>Sample taken from the environment of pre-weaned heifers</td>
<td>3.39 [2.00, 5.82]</td>
</tr>
<tr>
<td>Sample taken from pastureland</td>
<td>0.24 [0.15, 0.35]</td>
</tr>
<tr>
<td>Sample taken during the calving season</td>
<td>2.02 [1.10, 3.29]</td>
</tr>
<tr>
<td>Average monthly rainfall</td>
<td>1.26 [1.08, 1.47]</td>
</tr>
<tr>
<td>Total farm streptomycin use in 2017 measured in mg/PCU</td>
<td>0.76 [0.60, 0.97]</td>
</tr>
<tr>
<td>Total farm ABU in 2017 measured in mg/PCU</td>
<td>1.47 [1.01, 2.13]</td>
</tr>
</tbody>
</table>
References


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35. Carvalho CM, Polson NG, Scott JG. 2010. The horseshoe estimator for sparse signals, Biometrika 97:465-480

**Figure Legends**

**Figure 1.** Average monthly temperature vs. presence or absence of *E. coli* positive for *bla*\textsubscript{CTX-M} in samples from (A) pre-weaned calves and (B) all faecally contaminated dairy farm environments. Each sample is represented by a dot. A multilevel, multivariable logistic regression model revealed a positive association with increased temperature in both cases (p=<0.0001).

**Figure 2.** Percentage of farms with (A) *E. coli* positive for *bla*\textsubscript{CTX-M} in samples (B) ciprofloxacin-resistant *E. coli*. Data are presented by month (bars) and overlayed by a graph of average monthly temperature (dots) representing a year during the middle period of this study. Samples from calves have been excluded.
Figure 1

A

CTX-M Present

CTX-M Absent

Average Monthly Temperature (Celsius)

B

CTX-M Present

CTX-M Absent

Average Monthly Temperature (Celsius)
Figure 2

A

% Farms Positive for CTX-M

MONTH

B

% Farms Positive for CIP Resistance

MONTH