¹ Updates in managing *Salmonella* Dublin

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7 Abstract

Salmonella Dublin, a host-adapted *Salmonella* serotype in cattle, has become substantially more
 prevalent in dairy and calf-rearing facilities in the US and Canada since 2012. S. Dublin bacteria

- 10 isolated from US and Canadian farms commonly exhibit multidrug-resistant characteristics. This
- multidrug resistance substantially complicates the treatment and control of salmonellosis due to
 S. Dublin infection. Because it is a zoonotic disease, S Dublin infection in cattle also presents a
- 13 potential risk to human health. In cattle, S. Dublin infection results in high morbidity and
- 14 mortality rates in young calves and decreases the performance of mature animals. Clinical signs
- 15 usually include pneumonia, respiratory distress, and hyperthermia. Diagnosis is based on
- 16 bacterial identification via culture or PCR assay, or serological testing. Treatment involves
- 17 correcting dehydration and electrolyte imbalances and decreasing inflammation; the use of
- 18 antimicrobials is controversial. Prevention and control are via enhanced biosecurity practices.
- 19
- 20 Keywords:salmonellosis, calf health, zoonosis, latent carrier
- 21

22 Introduction

- 23 Salmonella enterica subspecies enterica serovar Dublin (S. Dublin) is a Gram-negative
- 24 bacterium commonly affecting dairy cattle. *Salmonella* Dublin is host-adapted to cattle, where it
- 25 can cause severe disease and compromise the welfare of young and mature bovine, and the
- 26 economic return of the producer [1-4]. Moreover, S. Dublin is a zoonosis that can cause severe
- disease in humans[5 6]. Some countries like Denmark initiated a surveillance and control
- program since 2002, and as a result, the prevalence of *S*. Dublin was reduced from 25 to 7%
- from 2002 to 2015 [7]. In countries without a control program, however, the prevalence of
- 30 infections is high[8]. Also, S. Dublin has been the most frequently identified serotype among
- bovine *Salmonella* isolates from clinical samples submitted to veterinary diagnostic laboratories
 in the U.S. and U.K. [9-12].
 - 33
- 34 In the U.S., S. Dublin has become one of cattle's most important multi-drug resistant (MDR)
- bacteria [5 13]. The MDR has complicated the treatment of clinically sick animals and has
- 36 become a threat to human medicine [14]. In addition, S. Dublin may be difficult to control and
- 37 eradicate from positive herds, as infection may persist in latent carriers and intermittently be
- 38 shed to the environment [2].
- 39

40 Importance of Salmonella Dublin

- 41 Prevalence in dairy farms
- 42 Salmonella Dublin is present worldwide, but estimates of the proportion of S. Dublin-infected
- 43 herds vary greatly by country (Table 1). Some European countries have established a S. Dublin
- 44 control and eradication that includes routine testing of all farms [15-17]. Although no country is

- 45 free from salmonellosis, 9 E.U. countries report only sporadic cases. Some countries, namely
- 46 Finland, Norway, and Sweden, have additional restrictions for cattle trade in place [18].
- 47 Conversely, more limited information regarding the prevalence of *S*. Dublin is available in
- 48 countries without control programs. However, S. Dublin has been identified as one of the most
- 49 common isolates of *Salmonella* spp. in dairy farms in the U.S., Germany, and the U.K. [9-12 19].
- 50
- 51 In 2014, the USDA's National Animal Health Monitoring System (NAHMS) conducted a cross-
- 52 sectional study including 234 farms nationwide. *Salmonella* Dublin was present in 0.7%, 6.7%,
- and 1.8% of the operations, milk samples, and milk filters, respectively [20]. Additionally, the
- 54 University of Minnesota Veterinary Diagnostic Laboratory (VDL) determined that S. Dublin was
- 55 the most prevalent serotype isolated from bovine samples between 2005 and 2014, representing
- 56 31.8% of all isolates examined from 880 dairy farms from the upper Midwest [9]. Likewise, *S.*
- 57 Dublin was the most prevalent serotype in bovine samples in the University of Wisconsin VDL,
- accounting for 23% of all isolates from 2006 to 2015 [10]. Similarly, S. Dublin has been the most
- 59 common *Salmonella* serovar isolated from bovine samples at the Michigan State University VDL (0, 1) between 2018 and 2022 measurements in 10.2007 of all begins $S_{1} = -\frac{1}{2}$
- 60 between 2018 and 2022, representing 10-20% of all bovine *Salmonella* isolations (Table 2).In
- 61 Germany and Italy, however, S. Typhimurium was the most frequently isolated serovar in cattle
- 62 samples collected as part of official outbreak investigations, followed by serovar Dublin
- 63 accounting for 30-40% of samples [19 21].
- 64

65 Human health hazard

- 66 Salmonella Dublin is a zoonotic bacterium that can cause rare but severe illness in humans, and
- 67 it is characterized by acute gastroenteritis and bacteremia [5]. The case fatality for S. Dublin has
- 68 been reported as the highest compared to other *Salmonella enterica* serotypes and has been
- 69 described as 6 times greater than *Salmonella* Typhimurium [22]. The consumption of raw milk
- 70 or raw dairy products has been associated with outbreaks of human salmonellosis caused by
- 71 serovar Dublin [23-26]. However, farm personnel, veterinarians, and any person in direct contact
- 72 with cattle are at risk of infection by accidentally ingesting animal feces or fluids [27].
- 73
- 74 The US Foodborne Disease Active Surveillance Network determined an increase in the incidence
- of human *S*. Dublin by 7.6 times from 1968 to 2013 [5]. The same study determined an increase
- in hospitalization from 68 to 78% and an increase in mortality from 2.7 to 4.2% when comparing
- 1996-2004 with 2005-2013 [5]. As discussed in the following section, S. Dublin has been
- characterized as a multidrug-resistant bacterium to common antibiotics used to treat bacterial
- 79 infections in humans and animals. Therefore, S. Dublin is a pathogen that can affect human
- 80 health severely and compromise medical treatment. Therefore, it is fundamental to prevent and
- 81 reduce the risk of infection from cattle to farm workers, animal caretakers, and from animal-
- 82 derived food to humans.
- 83
- 84 Antimicrobial resistance
- 85 The prevalence of MDR *S*. Dublin is associated with geographical location. While *S*. Dublin is
- 86 considered one of the most common MDR serotypes in the US [13], MDR is not common in the
- 87 European S. Dublin isolates [28]. However, S. Dublin MDR can reduce the success of treatments,
- delay recovery, and increase mortality and costs in humans and cattle [14].
- 89

90 In North America, *S.* Dublin has a 43% higher MDR prevalence than other *Salmonella* isolates

91 [5]. The National Antimicrobial Resistance Monitoring System (NARMS) reported that among

92 S. Dublin isolates, 84% were resistant to 5 or more classes of antimicrobial drugs, and 57% were

- resistant to 7 or more [5]. Furthermore, a 29 to 79% increase was observed in the proportion of isolates resistant to one or more antimicrobial classes when comparing 1996-2004 with 2005-
- isolates resistant to one or more antimicrobial classes when comparing 1996-2004 with 2005-2013 [5].
- 96

97 US isolates of S. Dublin are generally susceptible to gentamicin, amikacin, cefoxitin, cephalothin, 98 enrofloxacin, meropenem, and azithromycin [6 13]. Even though this pathogen is susceptible to 99 enrofloxacin, this drug is only allowed to treat bovine respiratory disease pathogens (specifically 100 Mannheimiahaemolytica, Pasteurella multocida, Histophilussomni and Mycoplasma bovis) in 101 non-lactating cows and dairy replacements younger than 20 months. Hence, enrofloxacin is not 102 labeled as a treatment for S. Dublin infections, and the extra-label use of this drug is prohibited 103 for food animals in the U.S. Although most producers and veterinarians would treat respiratory 104 disease without a pathogen isolation diagnosis, current U.S. regulations imply that enrofloxacin 105 cannot be used when S. Dublin is suspected or confirmed. This complicates the proper treatment 106 of sick calves and potentially might increase the use of drugs that S. Dublin has a reduced 107 susceptibility to. The antimicrobial susceptibility pattern of S. Dublin isolated has largely 108 remained unchanged in recent years (Table 2), with S. Dublin being generally susceptible to only 109 four antimicrobials. Among those four, only Trimethoprim/Sulfamethoxazole has been labeled

- 110 for treating Salmonella infections.
- 111

112 Pathogenesis and clinical signs of Salmonella Dublin infection in cattle

113 Salmonella Dublin infection in cattle can cause respiratory disease and septicemia. The disease is

114 transmitted by two major routes: oral and vertical (Figure 1). In the oral route, susceptible cattle 115 ingest the bacteria through contact with materials contaminated by feces or other bodily fluids

(e.g., milk, saliva, nasal secretions) from infected animals. In the vertical route, infected pregnant

117 cows transmit the disease to their offspring in utero. This can result in abortion in the last

118 trimester of gestation or the birth of congenitally infected calves. Aerosolized transmission is

- also possible, especially among calves housed in tight, confined spaces.
- 120

121 Once an animal is infected, *S*. Dublin colonizes the digestive tract and moves to the mesenteric 122 lymph nodes. From there, it can disseminate and cause systemic disease. Adaptation of *S*. Dublin 123 to cattle is attributed to the selective survival of strains capable of evading the host's immune

- response. In these instances, the inflammatory response to infection in the intestine is ineffective
- 125 in preventing systemic dissemination of infection. Because of the more invasive capacity of *S*.
- 126 Dublin, clinical signs of infection with this serotype are more severe than they are with

127 salmonellosis from other, less pathogenic, bovine-adapted *Salmonella* serotypes, such as Cerro.

128

129 The clinical signs of *S*. Dublin infection depend on the affected patient's age and the pathogen's

- 130 endemicity in the herd. Although S. Dublin infection can affect cattle of all ages, it is most
- 131 common in calves aged 2–12 weeks. In naive herds, the pathogen is rapidly transmitted, and an
- 132 outbreak ensues. Although most Salmonella infections present as GI disease, S. Dublin infection

133 is often a respiratory illness. Typical clinical signs of *S*. Dublin infection in calves include:

- hyperthermia (fever)
- obtundation (listlessness)

- 136 anorexia ٠
- 137 pneumonia •
- 138 respiratory distress (e.g., elevated respiratory rate, coughing) •
- 139 • dehydration
- septicemia 140 •

141 Arthritis (swollen joints) and meningoencephalitis can also occur in calves after bloodborne

142 transmission of the bacteria. Bloody diarrhea is also possible but not very common. A peracute 143 presentation may occur in calves, and sudden death in 1-2 days may result from endotoxic

144 shock. Calves 6–8 weeks old that survive acute infection can develop chronic infection

145 characterized by poor growth rate, ill thrift, lameness due to arthritis, and loose stool. Morbidity,

146 mortality, and case fatality rates for S. Dublin infection outbreaks in dairy calves are 10.5-

- 147 34.8%, 2.3–18.2%, and 26.4%, respectively [3 29].
- 148

152

149 In adult cattle, typical clinical signs of S. Dublin infection include:

- 150 slight fever • 151
 - mild diarrhea •
 - sudden decrease in milk production

153 Less typically, S. Dublin infection in adults can cause bloody diarrhea and, in rare instances,

154 death. Pregnant cattle may abort as a result of bacteremia. S. Dublin infection in adult cattle can

155 generate persistent infections without clinical signs. These latent carriers can periodically shed

156 the pathogen in feces or fluids during times of stress or when immunocompromised, contributing

157 to disease transmission in affected herds. 158

Diagnosis 159

Bacterial identification 160

Bacteriological culture has been useful for isolating and identifying S. Dublin to trace infections 161

162 and active shedders [2 30]. Bacteriological culture can be performed utilizing a variety of

163 samples, including feces and fluids from live animals, organs from necropsies, aborted fetuses, or

164 environmental samples. This method aims to isolate live bacteria [2]. Thus, the procedure

- 165 involves pre-enrichment and selective enrichment to allow bacterial growth, followed by plating
- 166 and confirmation [2]. This method has been described as more relevant in acute infections and 167 clinically ill animals, as the correct isolation will depend on the number of bacteria in the sample
- 168 [2 30 31]. For that reason, the sensitivity of this assay has been described as low [32], and it has
- 169 a limitation that latent carriers might be undetected due to the intermittent fecal shedding of S.
- 170 Dublin. Bacteriological culture using samples from dung pits, drinking water, milk filters, and
- 171 feces of clinically ill animals was associated with a sensitivity of 45, 5, 7, and 38% for detecting
- 172 S. Dublin, respectively [33].
- 173

174 In post-mortem examination of clinically ill animals, the collection of tissues from the lungs,

- 175 spleen, liver, intestine loops, gallbladder, intestinal content, and lymph nodes increases the
- 176 probability of bacteria isolation [3 34]. A potentially more sensitive and faster method for the
- 177 detection of genetic material of Salmonella is the polymerase chain reaction test (PCR) or real-
- 178 time PCR [35]. Persson, et al. [36] described an S. Dublin-specific real-time PCR. The procedure
- 179 for this method requires a pre-enrichment of the sample from lysates or extracted DNA [35]. To
- 180 increase sensitivity, a DNA extraction is recommended [35]. However, the specificity of the
- 181 assay in comparison to the numerous other Salmonella serotypes is yet to be determined

182

- 183 Serology
- 184 The detection of immunoglobulins against *S*. Dublin is performed through an Enzyme-linked
- 185 immunosorbent assay (ELISA). This method has a lower cost than bacteriological culture, and it
- 186 can be used as a monitoring strategy in the herd to identify latent carriers during control and
- 187 eradication programs[37 38]. *Salmonella* Dublin is part of the D-serogroup of *Salmonella* and
- has the antigenic factors O1, O9, and O12; therefore, cross-reaction between serovars sharing O antigens may occur [39]. The ELISA is based on detecting immunoglobulins directed to the LPS
- antigens may occur [39]. The ELISA is based on detecting immunoglobulins directed to the LPS
 O- antigen from serum, milk, and bulk tank milk (BTM) samples [40 41]. The kit is
- O- antigen from serum, milk, and bulk tank milk (BTM) samples [40 41]. The kit is
 commercially available in several diagnostic laboratories across the USto monitor*Salmonella*
- 192 infections in cattle herds. The results provided in this ELISA are semi-quantitative for antibody
- 193 concentration as they are expressed in ODC% (optical density coefficient). The interpretation of
- the result is based on an estimated cut-off point to determine positive animals depending on the
- sample. The ODC% cut-off for serum, milk from an individual, or BTM is 35 ODC%. A positive
- 196 correlation exists between the ODC% and antibody concentration in a sample. In BTM, the
- 197 greater the ODC%, the higher the spread of infection in the herd [42]. Sequential samples should
- 198 be obtained from individual animals using milk or serum samples to identify latent carriers of *S*.
- 199 Dublin due to their intermittent and low-intensity shedding. The limitations of this assay include
- 200 that the sensitivity and specificity are age-dependent, as it performs better as a diagnostic test in
- animals older than 100 days [32]. Additionally, milk samples have the limitation that only
- 202 lactating cows can be tested [2 33].
- 203
- 204 Necropsy
- There are no pathognomonic lesions in internal organs for infections with *S*. Dublin. However, while considering the age of the animal and the clinical signs, a necropsy may be helpful to guide
- 207 diagnosis or for sample collection. In calves with clinical presentation, the gross pathologic
- 208 findings in the lungs include pulmonary congestion, suppurative pneumonia, and chronic
- bronchopneumonia, depending on the severity of the clinical case [13 34]. The intestinal lesions
- 210 may include diffuse catarrhal hemorrhagic enteritis, ileitis, and mesenteric lymphadenitis [3 34].
- The intestinal content is watery, malodorous, and may contain mucous, blood, or fibrin clots [3
- 34]. Moreover, the liver is enlarged with rounded edges, hemorrhagic areas on the capsular
 surface, and gelatinous gallbladder edema[3]. In some cases, swollen joints may be a finding
- 213 surfa 214 [13].
- 214 215

216 Treatment

- 217 There is no targeted treatment for S. Dublin infection beyond the general recommendations for
- any *S. enterica* infection, which are to correct dehydration and electrolyte imbalances and to
- decrease inflammation. Calves with systemic infection should be administered NSAIDs (e.g.,
- flunixin meglumine, 1 mg/lb (2.2 mg/kg), IV, every 24 hours; or meloxicam, 0.23 mg/lb (0.5
- 221 mg/kg), IV or SC, every 24 hours for up to 5 days) to manage inflammation.
- 222
- 223 The administration of antimicrobials for treating *S*. Dublin infection is controversial for several
- reasons. First, appropriate antimicrobial selection is challenging because most *S*. Dublin strains
- are multidrug-resistant. US strains are frequently not susceptible to antimicrobials labeled for use
- 226 in calves with septicemia. Thus, in most cases, treatment with antimicrobials would require

- 227 extra-label administration of these drugs and determination of withholding periods for meat
- 228 under the direction of a licensed veterinarian.
- 229
- 230 Second, using an antimicrobial deemed potentially effective, based on the susceptibility of *S*.
- 231 Dublin to the drug, is usually not permitted to treat S. Dublin infection. For example, US isolates
- of *S*. Dublin are usually susceptible to enrofloxacin; however, the use of enrofloxacin to treat *S*.
- Dublin infection is extra-label drug use, which is prohibited for fluoroquinolones in food-
- 234 producing animals in the US.
- 235
- 236 Finally, there is a risk of enhancing pathogen resistance to antimicrobials with continuous
- administration, and cattle treated with antimicrobials are more likely to become latent carriers ofS. Dublin that contribute to further transmission of infection.
- 238

240 Prevention and control strategies

- 241 Prevention and control goals for S Dublin infection in cattle are to 1) minimize pathogen
- 242 exposure and 2) maximize pathogen resistance. Sanitation and biosecurity are critically
- 243 important for achieving these goals.
- 244
- 245 Farm management practices
- The following farm management practices can help minimize transmission of S Dublin infectionamong cattle [8]:
- 248 249
 - providing clean, dry calving pens and avoiding large group-calving areas
- removing calves from contact with their dams' feces as soon as possible after birth
- placing calves in a clean environment, where they have no contact with other calves or adult cattle
- maintaining strict control of colostrum management
- feeding pasteurized, rather than raw, milk to calves
- identifying and isolating newly sick cattle immediately, and ensuring that farm personnel
 handle sick cattle separately
- sanitizing and disinfecting all equipment used between animals
- ensuring that personnel wash hands, boots, and any common equipment used between groups of animals
- 260

261 Sanitation

- 262 Research has demonstrated that practices associated with the cleaning and disinfection of the
- environment are key elements in the prevention and control of *S*. Dublin [38 43 44]. Thus, when
- 264 cattle become infected with *S*. Dublin, it is essential to thoroughly clean and disinfect the
- 265 environment. All organic material (e.g., bedding, contaminated feed, feces) must be removed,
- and all surfaces must be completely washed down with water plus a detergent cleaner to remove
- any organic residues. A disinfectant should then be applied to ensure proper contact time.
- 268
- 269 Disinfectants used to combat Salmonella spp. include halogens (e.g., dilute chlorine bleach),
- 270 phenols, quaternary ammonium compounds, and oxidizing agents (e.g., potassium

- 271 peroxymonosulfate). Pressure washers should be avoided because they can transmit aerosolized
- 272 bacteria to both calves and personnel operating the washers.
- 273
- 274 Biosecurity
- 275 The purchase of cattle, particularly from multiple sources, is a major risk factor for introducing
- 276 S. Dublin into a herd [38 45-47]. Given the intermittent shedding of carriers, quarantine
- screening using fecal testing has a low sensitivity. Clinically ill cattle should be isolated from the
- herd and not returned too quickly to the main herd after clinical signs abate.
- 279
- 280 Because the bacterium can also be transmitted via inanimate objects (e.g., boots, clothes, and
- equipment), strict biosecurity practices should be implemented for visitors to the farm. *S.* Dublin can infect rodents; therefore, rodent control and protection of feed stores are important
- 282 can infect fodents, therefore, fodent control and protection of feed store283 biosecurity measures.
- 284
- 285 Vaccination
- 286 Commercial and autologous vaccines have been used to control S. Dublin in herds. However,
- 287 published studies have not evaluated autologous vaccines for their efficacy in preventing and
- reducing the clinical signs or the shedding of *S*. Dublin in dairy animals. A commercially
- available modified-live vaccine (EnterVene-D, Boehringer Ingelheim) is recommended for
- animals older than two weeks with a booster after 12 to 16 days. The benefits of an attenuated-
- 291 live S. Dublin vaccine are associated with a robust response at the mucosal level due to its action
- on lymphoid tissue in the gut and a robust cell-mediated immune response due to intracellular
- 293 proliferation [48 49]. Recent research also suggests that siderophore receptor vaccines might be
- immunogenic in newborn calves [50].
- 295
- The age for the first dose can be too late as calves may get infected with *S*. Dublin at birth or in the first hours of life. Moreover, limited research addresses the dam vaccination as an approach for producing antibodies that can be delivered to the newborn calf through colostrum [51]. The evidence suggests that specific antibodies for *S*. Dublin are in a higher concentration in the colostrum of cows vaccinated 30 days before dry-off than in non-vaccinated cows [51]. However, it remains unknown if those antibodies have a protective effect on the newborn calf. A recent study also explored the effect of vaccinating *S*. Dublin latent carriers with the commercial
- 303 attenuated-live vaccine on vertical transmission. In this study, latent carriers vaccinated at dry-off
- 304 with a live culture *Salmonella* Dublin commercial vaccine were 5 times less likely to give birth 305 to a seropositive calf [52].
- 306
- 307 Alternative routes of vaccine administration have also been explored. Research evaluating
- 308 intranasal and oral vaccination of 4-day-old calves suggests these are safe routes [53 54]. Using
- 309 these extra-label routes of administration reduced the disease severity as calves administered the
- 310 vaccine had a reduced mortality rate compared to unvaccinated calves [54]. However, the
- 311 incidence of pneumonia, abnormal fecal scores, and the fecal shedding of *S*. Dublin were not
- 312 reduced [53 54]. Furthermore, no differences were observed in the average daily gain or antibody
- 313 concentration at 10 weeks and 10 months of life compared to control calves [54]. Importantly,
- earlier studies noted that oral vaccination required a larger dose to induce a measurable immune
- response and was not protective against challenge [55]. Thus, existing evidence does not support
- the use of this alternative routes of administration.

- 317
- 318 Additionally, few studies assessed the cross-protection between *Salmonella enterica* with
- 319 modified-live vaccines. Mohler, et al. [34] found that calves younger than 2 weeks of life orally
- 320 vaccinated with modified-live S. Typhimurium had less severe clinical signs, improved appetite,
- and reduced fecal shedding when challenged with *S*. Dublin compared to control calves.
- However, calves in that study were challenged with a dose of *S*. Dublin to induce disease and
- 323 minimize mortality, and respiratory clinical signs were not assessed. Similar results were found
- using an attenuated-live S. Typhimurium on diarrhea and shedding of S. Newport and S. Cerro
- [48]. Moreover, there is a study assessing the vaccination of the dry cow with an S. Newport
 bacterin to provide cross-protection in an S. Typhimurium challenge in calves fed colostrum
- 327 from vaccinated dams. Despite higher serological titers, no difference in mortality, clinical signs,
- hematology, and fecal cultures were observed in calves fed colostrum from vaccinated cows and
- the control group [56]. Based on this research, the cross-protection between *Salmonella* spp. and
- 330 potential protection against S. Dublin in dairy herds is still in development.
- 331

332 Conclusions

- 333 *S.* Dublin severely affects cattle and human health. Recent reports indicate its prevalence has
- increased in several countries in the last several years, making it an emergent pathogen.
- 335 Information on pathogenicity, antimicrobial resistance, risk factors, and preventive management
- 336 practices is available. However, more research is still needed on the effectiveness of strategies
- that could be implemented in dairy facilities to prevent and control *S*. Dublin.
- 338

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- 344

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