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CURSO DE MESTRADO



**CARACTERIZAÇÃO DE *Escherichia coli* PRODUTORAS
DE BETA-LACTAMASES DE ESPECTRO ESTENDIDO
ISOLADAS DE BEZERROS**

YASMIN GARCIA MARANGONI

Campo Grande – MS
2023

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Characterization of Escherichia coli producers of extended spectrum beta-lactamases isolated from calves.

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Dissertação apresentada ao Programa de Pós-Graduação em Ciências Veterinárias da Universidade Federal de Mato Grosso do Sul, como requisito parcial para a obtenção do título de Mestre em Ciências Veterinárias.

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CERTIFICADO

Certificamos que a proposta intitulada "Caracterização epidemiológica, patológica e molecular de diarreias em bezerros nos biomas Cerrado e Pantanal sul-matogrossense", registrada com o nº 1.134/2020, sob a responsabilidade de **CARLOS ALBERTO DO NASCIMENTO RAMOS** - que envolve a utilização de animais pertencentes ao filo Chordata, subfilo Vertebrata, para fins de pesquisa científica – encontra-se de acordo com os preceitos da Lei nº 11.794, de 8 de outubro de 2008, do Decreto nº 6.899, de 15 de julho de 2009, e com as normas editadas pelo Conselho Nacional de Controle de Experimentação Animal (CONCEA), e foi aprovada pela COMISSÃO DE ÉTICA NO USO DE ANIMAIS/CEUA DA UNIVERSIDADE FEDERAL DE MATO GROSSO DO SUL/UFMS, na 5ª reunião ordinária do dia 30/07/2020.

FINALIDADE	() Ensino (x) Pesquisa Científica
Vigência da autorização	01/09/2020 a 31/07/2023
Espécie/Linhagem/Raça	Bos taurus/ Bos indicus/ Nelore e/ou cruzamentos
Nº de animais	400
Peso/idade	Não se aplica/ Até 90 dias
Sexo	Fêmeas e machos
Origem	Pantanal e Cerrado

Fábio José Carvalho Faria

Coordenador da CEUA/UFMS

Campo Grande, 11 de agosto de 2020.



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RESUMO

A resistência antimicrobiana não é um fenômeno recente, mas é um problema crítico de saúde pública mundial. A produção de beta-lactamases de espectro estendido (ESBL), pelo grupo das Enterobacteriales, é o mecanismo de resistência bacteriana adquirida cada vez mais citado nas últimas décadas, presentes na interface humano-animal-ambiente. Este estudo teve como objetivo verificar em amostras fecais de bezerros, a presença de cepas de *Escherichia coli* diarreio gênicas produtoras de ESBL, identificando os genes de resistência *bla*_{CTX-M-2}, *bla*_{CTX-M-8}, *bla*_{CTX-M-15}, *bla*_{SHV} e *bla*_{TEM} pelo método de Reação em Cadeia da Polimerase (PCR) e fenotipicamente pelo método de triagem de disco de aproximação. Também se objetivou a padronização de um método de identificação rápido, para isolados de *E. coli* multirresistentes, usando o método de Espectroscopia de Infravermelho com Transformada de Fourier (FT-IR). Duzentos e oito isolados de *E. coli* diarreio gênicas foram submetidos à PCR, sendo a presença de pelo menos um dos genes citados verificada em 35,6% do total (n=74). A distribuição dos genes nas amostras positivas ocorreu da seguinte forma: 3,84% positivas para *bla*_{CTX-M-2}, 5,28% para *bla*_{SHV} e 29,3% para o gene *bla*_{TEM}. Os genes *bla*_{CTX-M-15} e *bla*_{CTX-M-8} não foram observados no estudo. Quanto à suscetibilidade aos antimicrobianos, as amostras apresentaram altas taxas de resistência ao grupo dos beta-lactâmicos, bem como para tetraciclina (80%), no qual 50,9% (106/208) foram fenotipicamente multirresistentes, mostrando que animais portadores de cepas bacterianas com a enzima ESBL, podem ser um reservatório constante de disseminação. Adicionalmente, 40 isolados de *E. coli* diarreio gênicas sabidamente multirresistentes foram selecionados para serem mensurados no FT-IR, com espectros colhidos na faixa de 4000 a 700 cm⁻¹, 3000 a 2800 cm⁻¹ e 1800 a 800 cm⁻¹. O método de FT-IR para o aprendizado de máquina foi capaz de identificar os isolados de *E. coli* diarreio gênicas multirresistentes com 70% de precisão no teste de validação, demonstrando que o FT-IR poderia ser uma ferramenta de triagem para estudos bacteriológicos e diagnóstico.

Palavras-chave: beta-lactâmicos, ESBL, FT-IR, pecuária e resistência à antibióticos.

MARANGONI, Y.G. Characterization of *Escherichia coli* producers of extended spectrum beta-lactamases isolated from calves. 2023. Master's degree – Postgraduate Program in Veterinary Science. Faculdade de Medicina Veterinária e Zootecnia, Universidade Federal de Mato Grosso do Sul, Campo Grande, MS, 2023.

ABSTRACT

Antimicrobial resistance is not a recent phenomenon, but it is a critical problem of world public health. The production of extended spectrum beta-lactamase (ESBL) by the Enterobacteriales group is the increasingly cited bacterial resistance mechanism in recent decades, present in the human-animal-environment interface. This study aimed to verify in fecal calf samples, the presence of ESBL-producing *Escherichia coli* strains, identifying *bla*_{CTX-M-2}, *bla*_{CTX-M-8}, *bla*_{CTX-M-15}, *bla*_{SHV} and *bla*_{TEM} by the Polymerase Chain Reaction method (PCR) and phenotypically by the approach disc screening method. It also was an aim of a rapid identification method for multiresistant *E. coli* isolates, using the Fourier Transform Infrared Spectroscopy (FTIR) method. Diarrheogenic *E. coli* isolates, and eight isolates were subjected to PCR, the presence of at least one of the genes mentioned in 35.6% of the total (n = 74). A distributed genes from positive samples occurred as follows: 3.84% were positive for *bla*_{CTX-M-2}, 5.28% for *bla*_{SHV} gene and 29.3% for the *bla*_{TEM} gene. The *bla*_{CTX-M-15} and *bla*_{CTX-M-8} genes were not observed in the study. Regarding susceptibility to antimicrobials, the samples showed high resistance rates to the beta-lactam group, as well as for tetracycline (80%), in which 50.9% (106/208) were phenotypically multi-resistant, showing that strains carriers with strains bacterial with ESBL enzyme can be a constant dissemination reservoir. Additionally, 40 isolates of *E. coli* diarrheogenic multiresistant were selected to be measured in the FTIR, with spectra harvested from 4000 to 700 cm⁻¹, 3000 to 2800 cm⁻¹, and 1800 to 800 cm⁻¹. The FTIR method for machine learning was able to identify the isolates of *E. coli* diarrheogenic multi-resistant with 70% accuracy in validation tests, demonstrating that the FTIR could be a screening tool for bacteriological and diagnostic studies.

Keywords: beta-lactams, ESBL, FTIR, livestock and antibiotic resistance.

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CAPÍTULO 1

1. INTRODUÇÃO GERAL

A identificação de bactérias multirresistentes e seus genes de virulência na pecuária tem grande importância em saúde pública mundial, assim como a identificação de possíveis genes de resistência à antimicrobianos (AGRs) (HAN et al., 2017).

O rebanho comercial brasileiro cresce com taxas altas, chegando a 218,2 milhões de cabeças de gado. A região Centro-Oeste, respondeu por mais de 75 milhões dessas cabeças. Em termos municipais, Corumbá, Mato Grosso do Sul, segue em segunda posição com o maior rebanho nacional (IBGE, 2020).

A enterite em bezerros, provocada pela bactéria *Escherichia coli* tem sido relatada desde o início do século XX como um dos principais agentes envolvidos nas diarreias de origem infecciosa. Considerando que o estado de Mato Grosso do Sul é uma importante região de pecuária de corte, este agente etiológico e suas cepas diarreiogênicas passaram a ser estudadas e identificadas no estado (SMITH; ORCUTT, 1925; TUTIJA et al., 2022).

A Organização Mundial da Saúde, OMS (2017) lista agentes bacterianos e suas respectivas resistências antimicrobianas na medicina humana, como um alerta à população mundial a crescente resistência global aos medicamentos antimicrobianos, entretanto, dentro da medicina veterinária já podemos observar a cosmopolitização destes AGRs listados pela OMS, como a produção de beta-lactamase de espectro estendido (ESBL) pela bactéria *E. coli* na interface humano-animal-ambiente (MELO et al., 2018; PALMEIRA et al., 2020).

Atualmente, por não existirem programas de vigilância epidemiológica de abrangência nacional ativos referentes à resistência bacteriana e a seus mecanismos, torna-se difícil estimar a proporção de cepas multirresistentes e aquelas produtoras de ESBL na bovinocultura (SANTAJIT; INDRAWATTANA, 2016; CHONG et al., 2018).

Com o surgimento de ferramentas da biologia molecular, podemos realizar análises detalhadas das sequências de ácidos nucleicos para identificação de AGRs, porém sua abrangência, seu poder discriminatório e simplicidade são variáveis e, para muitas análises, o alto custo e o tempo dificultam sua implementação na rotina laboratorial (SABAT et al., 2013).

35 Hoje é altamente desejável a busca pela implementação de novos métodos
36 para detectar microrganismos e AGRs, com resultados rápidos, reduzido custo
37 com consumíveis e fácil portabilidade (BENGTSSON-PALME et al., 2017).

38 Os estudos de espectroscopia óptica, como a Espectroscopia no
39 Infravermelho com Transformada de Fourier (FT-IR), representam uma
40 ferramenta poderosa para atingir esses objetivos, bem como em estudo futuros,
41 nos processos de vigilância epidemiológica nacional referentes à resistência
42 bacteriana e a seus mecanismos, usando abordagens baseadas em análises
43 vibracionais em conjunto com a bioinformática (NOVAIS et al., 2019).

44 Diante deste contexto relatado, este trabalho objetivou estudar a presença de
45 cepas de *E. coli* diarreiogênicas produtoras de ESBL no rebanho do estado de
46 Mato Grosso do Sul, Brasil, em conjunto com o aperfeiçoamento do diagnóstico
47 de resistência antimicrobiana por técnicas de fotônica.

48

49 **1.1 OBJETIVO GERAL**

50 O presente estudo teve por objetivo investigar fenotipicamente e
51 genotipicamente a presença de linhagens de *E. coli* produtoras de ESBL das
52 famílias CTX-M (*bla*_{CTX-M-2}, *bla*_{CTX-M-8} e *bla*_{CTX-M-15}), SHV (*bla*_{SHV}) e TEM (*bla*_{TEM})
53 na microbiota intestinal de bezerros, provenientes de fazendas das quatro
54 macrorregiões do estado de Mato Grosso do Sul, Brasil, bem como realizar a
55 padronização para o método de FT-IR em isolados de *E. coli* multirresistentes.

56

57 **1.2 OBJETIVOS ESPECÍFICOS**

- 58 • Avaliar *in vitro* o perfil de suscetibilidade aos antimicrobianos em isolados
59 de *E. coli* diarreiogênicas pelo método de disco difusão;
- 60 • Identificar fenotipicamente a presença de estirpes produtoras de ESBL;
- 61 • Identificar genotipicamente por meio da técnica de Reação em Cadeia da
62 Polimerase (PCR), os genes codificadores de ESBL: *bla*_{CTX-M-2}, *bla*_{CTX-M-8},
63 *bla*_{CTX-M-15}, *bla*_{SHV} e *bla*_{TEM};
- 64 • Avaliar o desempenho da técnica de Espectroscopia de Infravermelho por
65 Transformada de Fourier (FT-IR) para identificar cepas multirresistentes
66 de *E. coli*.

67

68 2 REVISÃO DE LITERATURA

69

70 2.1. Resistência antimicrobiana

71 A introdução de antibióticos no uso clínico foi indiscutivelmente o maior
72 avanço médico do século 20 (KATZ; BALTZ, 2016).

73 Em 1904, Paul Ehrlich e Alexander Fleming iniciaram a moderna “era dos
74 antibióticos”. Ehrlich iniciou um programa de rastreamento sistemático para
75 encontrar um medicamento contra a sífilis, baseadas em arsênico sintético
76 salvarsan, porém os primeiros relatos de resistência ao salvarsan foram em 1930
77 (EHRlich, 1913).

78 Salvarsan foi substituído pela sulfonamida Prontosil (OTTEN, 1986),
79 considerado o primeiro antimicrobiano de amplo espectro com uso clínico,
80 entretanto foi amplamente substituído pela descoberta da penicilina por
81 Alexander Fleming em 1928 (FLEMING, 1929). A partir de 1945, a penicilina foi
82 purificada por um grupo de pesquisadores da Universidade de Oxford, e após
83 esse período, houve relatos crescentes de resistência a este medicamento
84 (ABRAHAM et al., 1941).

85 O início da “era de ouro” da descoberta de antibióticos, ocorreu entre as
86 décadas de 1940 e 1960. A maioria desses compostos ainda está em uso clínico,
87 mas sua eficácia foi prejudicada pelo aumento da resistência microbiana. Na
88 verdade, a descoberta rápida e relativamente fácil de várias classes de
89 antibióticos durante um período relativamente curto, levou ao uso excessivo
90 desses medicamentos (KATZ; BALTZ, 2016).

91 Existem diferentes mecanismos de resistência antimicrobiana: a resistência
92 intrínseca e a resistência adquirida. A resistência intrínseca ocorre de forma
93 natural, como parte de um processo de evolução bacteriana. Alguns exemplos
94 são membros da ordem Enterobacteriales, na qual a bactéria *E. coli* está
95 inserida, resistentes as classes dos macrolídeos, glicopeptídeos, bem como ao
96 ácido fusídico, linezolidina e a penicilina G, considerados antimicrobianos de
97 grande uso na rotina clínica (WHO, 2012; EUCAST, 2020; CLSI, 2021).

98 Com frequência as bactérias utilizam mais de uma estratégia para evitar a
99 ação dos antimicrobianos. Na resistência adquirida, a bactéria altera seu material
100 genético de duas formas: pela mutação no seu material genético; ou introdução
101 de um material genético estranho, como genes de resistência, que podem ser

102 transferidos entre gêneros ou espécies diferentes de bactérias, principalmente
 103 devido à pressão seletiva exercida pelo uso indiscriminado de antimicrobianos
 104 (LIVERMORE et al., 2001; WHO, 2012; EUCAST, 2020).

105 O relatório O'Neill (2016) previu que, sem ação urgente, dez milhões de
 106 pessoas, por ano morrerão de infecções causadas por bactérias multirresistentes
 107 à antimicrobianos até 2050.

108 Como um alerta à população mundial a crescente resistência aos
 109 medicamentos antimicrobianos, a Organização Mundial da Saúde (OMS), lista
 110 os agentes bacterianos e suas respectivas resistências antimicrobianas (Tabela
 111 1), utilizando critérios para a seleção das bactérias, como: a) quão letal as
 112 infecções que esses microrganismos causam são; b) se o seu tratamento requer
 113 longas estadias hospitalares; d) como eles se espalham facilmente entre os
 114 animais, dos animais aos seres humanos, e de pessoa para pessoa; e) se os
 115 microrganismos podem ser prevenidos; f) quantas opções de tratamento ainda
 116 existem para esses microrganismos (WHO, 2017).

117

118 Tabela 1: Agentes bacterianos e suas respectivas resistências antimicrobianas na medicina
 119 humana, segundo a Organização Mundial da Saúde (2017).

Prioridade crítica	Prioridade alta	Prioridade média
<i>Acinetobacter baumannii</i> resistente à carbapenêmicos	<i>Enterococcus faecium</i> resistente à vancomicina	<i>Streptococcus pneumoniae</i> não penicilina suscetível
<i>Pseudomonas aeruginosa</i> resistente à carbapenêmicos	<i>Helicobacter pylori</i> resistentes à claritromicina	<i>Haemophilus influenzae</i> resistentes à ampicilina
<i>Enterobacteriaceae</i> resistentes à 3ª geração de cefalosporinas e carbapenêmicos.	<i>Salmonella</i> sp resistentes à fluoroquinolonas	<i>Shigella</i> sp resistentes à fluoroquinolona
	<i>Staphylococcus aureus</i> resistente à vancomicina e metilina	
	<i>Campylobacter</i> sp resistentes à fluoroquinolona	
	<i>Neisseria gonorrhoeae</i> resistentes à 3ª geração de cefalosporinas e fluoroquinolona	

120 Fonte: WHO (2017).

121

122 Dentro da medicina veterinária já podemos observar a cosmopolização de *A.*
 123 *baumannii* (POMBA et al., 2015; EWERS et al., 2017), *P. aeruginosa*
 124 (FERNANDES et al., 2018; ELSHAFIEE et al., 2019), *Staphylococcus aureus*

125 (OLIVEIRA et al., 2016); *Salmonellas* species (COLOBATIU et al., 2015; PRIBUL
126 et al., 2017), *Campylobacter* species (FRASAO et al., 2015), bem como algumas
127 bactérias membros do grupo Enterobacteriales, como a *E. coli* (MELO et al.,
128 2018; PALMEIRA et al., 2020) na interface humano-animal-ambiente.

129 A bactéria *E. coli* é considerada comensal uma vez que, apenas uma
130 pequena parte das cepas apresentam patogenicidade responsável por
131 enfermidades. A diferenciação de cepas patogênicas (diarreio gênicas) de cepas
132 não patogênicas, é baseada na produção de fatores de virulência, sendo cinco
133 os principais patótipos de *E. coli* diarreio gênica para bezerros: *E. coli*
134 enterotoxigênica (ETEC), *E. coli* enteropatogênica (EPEC), *E. coli*
135 enterohemorrágica (EHEC), *E. coli* produtora de toxina Shiga (STEC), *E. coli*
136 necrotoxigênica (NTEC). Cada categoria possui capacidade de causar síndrome
137 clínica, com características epidemiológicas e patológicas específicas
138 (BLANCHARD, 2012; COURA et al., 2015).

139 Quando há disseminação de cepas de *E. coli* diarreio gênicas no rebanho
140 bovino, podem surgir cepas bacterianas capazes de expressarem enzimas que
141 conferem resistência a múltiplos antimicrobianos, em resposta ao amplo uso
142 destas moléculas para tratar os casos de diarreia (BLANCHARD, 2012; COURA
143 et al., 2015; POIREL et al., 2018).

144 A classe de antimicrobianos beta-lactâmicos constitui a primeira via de
145 escolha para o tratamento de diarreia, devido a sua boa eficácia terapêutica e
146 baixa toxicidade. Representantes dessa classe são o grupo das penicilinas
147 (amoxicilina, associada ou não à clavulanato de potássio e ampicilina) e
148 cefalosporinas (1ª e 3ª geração), ambos com a presença do anel beta-lactâmico
149 em sua estrutura molecular (CONSTABLE, 2004; SPINOSA et al., 2014).

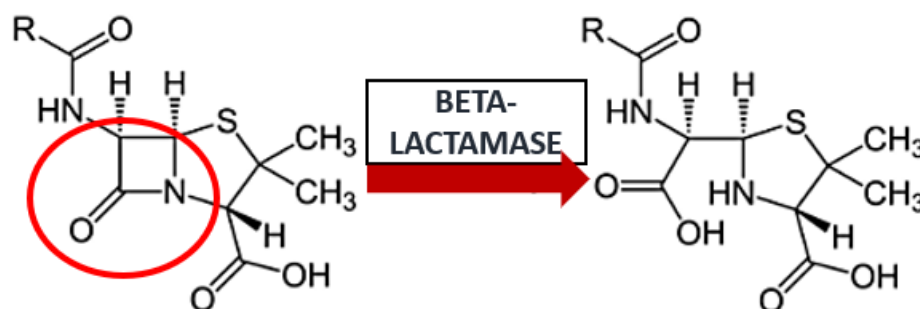
150 O mecanismo principal de ação dos beta-lactâmicos é a inibição da última
151 etapa de síntese de peptídeoglicano para a formação da parede celular
152 bacteriana, logo estes antimicrobianos somente são capazes de atuar em
153 bactérias durante a fase de crescimento logarítmica, quando, então, há
154 necessidade da síntese da parede celular (SPINOSA et al., 2014; ANDRADE;
155 DARINI, 2016).

156 O mecanismo de resistência mais importante para os beta-lactâmicos é a
157 produção de betalactamases pelas bactérias. As betalactamases produzidas por
158 diferentes bactérias possuem propriedades físicas, químicas e funcionais

159 variadas, sendo algumas betalactamases específicas para penicilinas
160 (penicilinases), outras para cefalosporinas (cefaloporinases) e outras que podem
161 atuar em ambos os grupos de antimicrobianos, como as beta-lactamases de
162 espectro estendido (ESBL) (SPINOSA et. al, 2014).

163 Diversos trabalhos têm apontado para o crescimento mundial das ESBL em
164 bactérias Gram-negativas, como membros do grupo Enterobacteriales (ELLER
165 et al., 2014; ROCHA et al., 2016; GUNDRAN et al., 2019; SUKMAWINATA et al.,
166 2020; WIDODO et al., 2020). Estas enzimas quando produzidas pelas bactérias
167 Gram-negativas são secretadas no espaço periplasmático (SPINOSA et. al,
168 2014) podendo hidrolisar o anel beta-lactâmico irreversivelmente da ligação
169 amina do anel beta-lactâmico (Figura 1), impedindo a ação dos beta-lactâmicos
170 na síntese da parede celular bacteriana (BUSH; JACOBY, 2010), hidrolisando
171 todas as penicilinas, cefalosporinas, incluindo as de terceira e quarta geração, e
172 o aztreonam (COQUE et al., 2008).

173



174

175 Figura 1: Esquema bioquímico demonstrando a ação da enzima beta-lactamase na ligação amina
176 do anel beta-lactâmico. Arquivo pessoal.

177

178 A presença de ESBL tem um significado clínico importante, pois são
179 codificadas por plasmídeo, e estes frequentemente carregam genes com co-
180 resistência a outras classes de antimicrobianos, como, por exemplo,
181 tetraciclina, aminoglicosídeos e fluoroquinolonas. Portanto, as opções de
182 antimicrobianos no tratamento de organismos produtores de ESBL são
183 extremamente limitadas (PATERSON; BONOMO, 2005; ADEYANKINNU et al.,
184 2014).

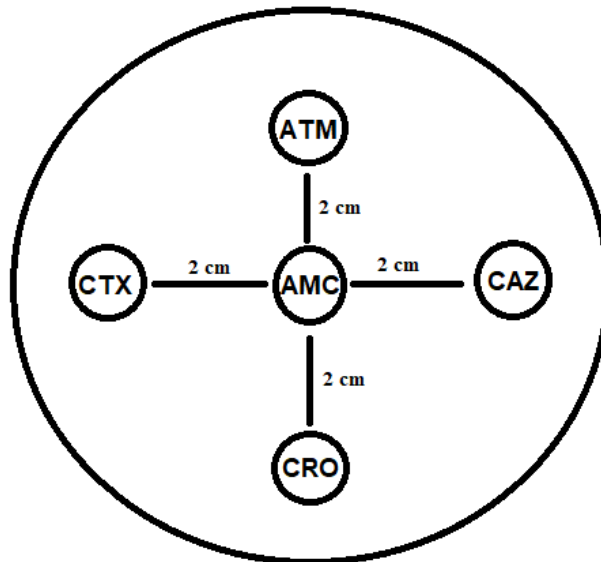
185 Dentre as variantes enzimáticas descritas, as enzimas da família SHV e TEM
186 foram inicialmente reconhecidas em diferentes espécies bacterianas Gram-
187 negativas, descritas em humanos a partir da década de 1960 (CANTÓN et al.,

188 2012), porém as enzimas da família CTX-M têm despertado atenção por sua alta
189 incidência e grande capacidade de propagação (PITOUT et al., 2005; WIDODO
190 et al., 2020).

191 As fezes de animais de produção, que possuem cepas com ARGs, como a
192 enzima ESBL, podem ser reservatórios constantes de disseminação aos
193 humanos (CARATTOLI, 2008; SCHMIDT et al., 2013), ao meio ambiente e a
194 outros animais (BOONYASIRI et al., 2014; TEKINER; OZPINAR, 2016), levando
195 à atrasos na terapia apropriada, aumento da morbidade, mortalidade, e
196 conseqüentemente, em perdas econômicas aos produtores (SANTAJIT;
197 INDRAWATTANA, 2016; CHONG et al., 2018).

198 Padrões de resistência entre bactérias têm sido estudados utilizando o cultivo
199 a partir de métodos fenotípicos, como o teste de triagem de disco de
200 aproximação (JARLIER et al., 1988) (Figura 2), bem como métodos moleculares,
201 como a Reação em Cadeia da Polimerase (PCR), localizando os genes *bla*_{CTX-M},
202 *bla*_{SHV} e *bla*_{TEM} (PATERSON; BONOMO, 2005).

203



204

205 Figura 2: Ilustração da posição ideal das cefalosporinas, como cefotaxima (CTX), ceftazidima
206 (CAZ) e ceftriaxona (CRO), o monobactâmico aztronam (ATM), e o inibidor de beta-lactamase
207 amoxicilina + ácido clavulônico (AMC) no teste de triagem de disco de aproximação, padronizado
208 pelo CLSI (2021). Arquivo pessoal.

209

210 Entretanto, hoje é altamente desejável a busca pela implementação de novos
211 métodos de identificação bacteriana, dos quais não utilizem uma série de testes

212 para a identificação definitiva do organismo. Uma abordagem diferente para a
213 identificação rápida de microrganismos é baseada em técnicas espectroscópicas
214 (MAQUELIN et al., 2002).

215

216 **2.2 Espectroscopia óptica**

217 Do ponto de vista histórico, entre 1665 e 1666, Isaac Newton, um físico inglês,
218 passou a introduzir a espectroscopia ao observar a luz do sol. A luz ao passar
219 de meio de propagação para outro, como um prisma de vidro, poderia ser
220 decomposta em diversas cores (NEWTON, 1972).

221 No ano de 1802, o interesse pela espectroscopia aumentou
222 consideravelmente, quando um grupo passou a observar que substâncias
223 emitiam cores diferentes quando colocadas sob a chama, com o objetivo de
224 identificar os elementos químicos presentes. A partir de 1859, os químicos
225 Robert Wilhelm Bunsen e Gustav Kirchhoff tiveram um papel fundamental na
226 construção do primeiro espectroscópio, descobrindo que cada elemento químico
227 produz um padrão único de linhas espectrais, distribuída ao longo dos
228 comprimentos de onda (BUNSEN; KIRCHHOFF, 1862).

229 Esses e muitos estudos posteriores abriram caminhos para a utilização dessa
230 metodologia para a análise de substâncias, como a descoberta do cloreto de
231 céσιο e o cloreto de rubídio, bem como para a identificação de bactérias em
232 diferentes níveis taxonômicos, desde gêneros, espécies a sorotipo/sorogrupo
233 (BUNSEN; KIRCHHOFF, 1862; HELM et al. 1991; MAQUELIN et al., 2002).

234 Durante a década de 1990, houve novos desenvolvimentos de
235 equipamentos, procedimentos experimentais e análise de dados em software, o
236 que impulsionaram a novas publicações sobre a técnica de espectroscopia no
237 infravermelho com transformada de Fourier (FT-IR) para identificação bacteriana
238 (HELM et al. 1991).

239

240 **2.2.1 Espectroscopia no Infravermelho com Transformada de Fourier** 241 **(FT-IR).**

242 Técnicas espectroscópicas destrutivas, como a Espectrometria de Massa
243 com Ionização por Dessorção a Laser Assistida por Matriz (MALDI-TOF MS) tem
244 mostrado resultados promissores para a identificação de grupos bacterianos
245 Gram-negativos e Gram-positivos, porém inconsistências entre os protocolos

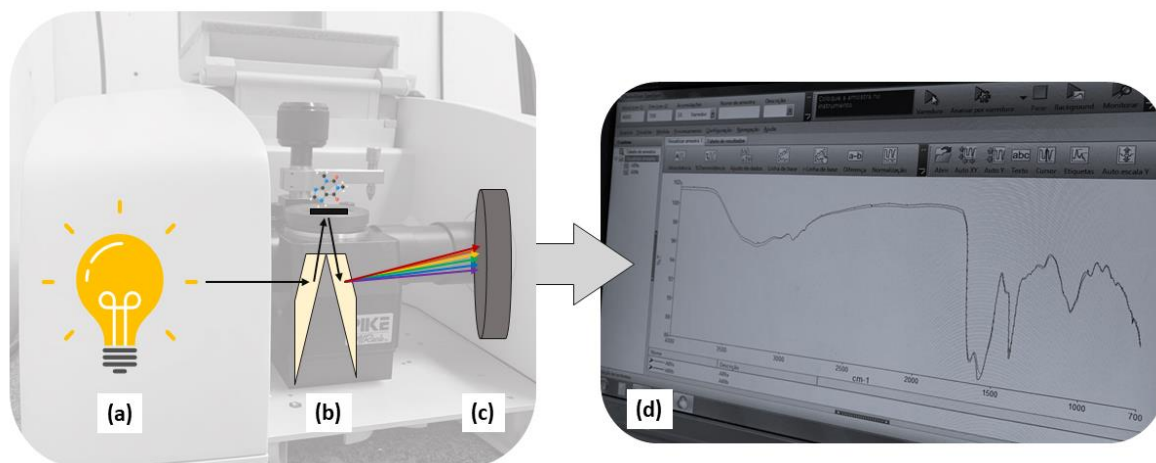
246 experimentais ou de análise de dados adotados podem dificultar o alcance de
247 reprodutibilidade, padronização e, principalmente, a portabilidade para
248 laboratórios de rotina de microbiologia clínica (SAUGET et al., 2017).

249 Considerando o bom desempenho em diferenciação de cepas, alto
250 rendimento, custo-benefício, mínimo tempo de trabalho, e fácil portabilidade,
251 métodos não destrutivos - àqueles que deixam a amostra intacta durante a
252 análise - podem ser desenvolvidos com base em espectroscopias vibracionais,
253 como o FT-IR (MAQUELIN et al., 2002; NOVAIS et al., 2019).

254 Na região do infravermelho (100 a 10000 cm^{-1}), ocorre a absorção da
255 radiação infravermelha por um determinado tipo de amostra como, por exemplo,
256 células bacterianas, causando excitação e vibração dos seus diferentes
257 compostos químicos, com o objetivo de determinar a composição molecular
258 (ácidos nucleicos, proteínas, lipídios e carboidratos) destas células bacterianas
259 como um todo, semelhantes a impressões digitais ou espectro que podem ser
260 utilizados para identificar a identidade do microrganismo (HELM et al. 1991;
261 SETTLE, 1997; STUART, 2004).

262 O método de FT-IR, utiliza um equipamento chamado de espectrômetro
263 (Figura 3). No espectrômetro, a fonte de luz emite um feixe de luz com energia e
264 comprimento de onda bem definido. Este feixe de luz ao passar pela amostra
265 analisada pode interagir com o meio e ser transmitido para um fotodetector.
266 Logo, um espectro característico da absorção de radiação eletromagnética da
267 amostra é obtido, e por meio de cálculos matemáticos, da transformada de
268 Fourier, a distância do comprimento óptico pode ser convertida para o valor da
269 frequência de radiação e vice-versa, garantindo a otimização das funções da
270 espectroscopia no infravermelho, permitindo maior sensibilidade e velocidade de
271 análise (STUART, 2004).

272



273

274 Figura 3: Ilustração básica da funcionalidade do espectrômetro, demonstrando a (a) fonte de luz,
 275 (b) interação da luz com a amostra analisada, (c) transmissão para um fotodetector, (d) registro
 276 do espectro da absorção de radiação eletromagnética, ou seja, "impressão digital. Arquivo
 277 pessoal.
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O potencial do método de FT-IR para diferenciar e identificar grupos de cepas dentro de espécies Gram-positivas, como *Staphylococcus aureus* e Gram-negativas, principalmente em *E. coli*, sendo explorado inicialmente por Helm et al. (1991).

No estudo de Beutin et al. (2007), foi possível demonstrar que o FT-IR pode revelar diferenças estruturais de superfície entre dois grupos de cepas de *E. coli* O123. Para Abu-Aqil et al. (2022) e Abu-Aqil et al. (2023), foi utilizado o FT-IR e o aprendizado de máquina para identificar *E. coli* diretamente da urina de pacientes hospitalizados, bem como determinar o perfil de suscetibilidade aos antibióticos e a presença de isolados produtores de beta-lactamases de espectro estendido.

O estudo de Helm et al. (1991) determinou cinco janelas espectrais correspondentes à absorção expressa em números de onda (cm^{-1}): janela 1, $3000\text{-}2800\text{cm}^{-1}$ dominada por vibrações presentes em ácidos graxos (lipídios); janela 2, $1800\text{-}1500\text{cm}^{-1}$, dominada por vibrações das bandas amida I e amida II (proteínas e peptídeos); janela 3, $1500\text{-}1200\text{cm}^{-1}$, com informações de proteínas, ácidos graxos e compostos de fosfato (região mista); janela 4, $1200\text{-}900\text{cm}^{-1}$, bandas de carboidratos presentes na parede celular (polissacarídeos); janela 5, $900\text{-}700\text{cm}^{-1}$, mostrando alguns padrões espectrais específicos, ainda não atribuídos a componentes celulares ou grupos funcionais (região de impressão digital).

300 Esses padrões específicos de impressão digital tornam este método uma
301 alternativa para identificação rápida de bactérias, tipagem e a triagem no nível
302 de subespécies, sendo as janelas 3 e 4, o ponto chave para a identificação
303 bacteriana (MEYERS, 2006; SHI et al., 2020).

304 Apesar da grande aplicabilidade de FT-IR em microbiologia nos últimos anos,
305 esta técnica pode ser considerada pouco convencional e seu potencial nas
306 rotinas de microbiologia pode ser subestimado, pela aceitação e adaptação
307 desta metodologia pela comunidade microbiológica. Entretanto, o método de FT-
308 IR poderia agregar valor à caixa de ferramentas da identificação bacteriana e,
309 em estudos futuros, nos processos de vigilância epidemiológica nacional
310 referentes à resistência bacteriana e a seus mecanismos (NOVAIS et al., 2019).

311

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CAPÍTULO 2

4. (ARTIGO I)

[Artigo formatado de acordo com as normas da Revista Arquivo Brasileiro de Medicina Veterinária e Zootecnia]

IDENTIFICAÇÃO FENOTÍPICA E GENOTÍPICA DE *Escherichia coli* PRODUTORAS DE BETA-LACTAMASES DE ESPECTRO ESTENDIDO (ESBL) ISOLADAS DE FEZES DE BEZERROS

Phenotypic and genotypic characterization of Escherichia coli producing extended spectrum beta-lactamases (ESBL) isolated from calf feces.

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RESUMO

A produção de beta-lactamases por Enterobacteriales é o mecanismo de resistência mais comum aos antibióticos beta-lactâmicos. As beta-lactamases de espectro estendido (ESBL), são consideradas uma urgência clínica e epidemiológica mundial. Este estudo teve como objetivo verificar a presença de cepas de *Escherichia coli* diarreiogênicas, isoladas de bezerros, produtoras de ESBL. Amostras de fezes de 208 bezerros, de duas faixas etárias distintas, foram obtidas e cultivadas em meio de enriquecimento para isolamento de *E. coli*. A análise fenotípica foi realizada pelo método de triagem de disco de aproximação empregando discos de amoxicilina + ácido clavulônico (AMC), ceftriaxona (CRO), cefotaxime (CTX), ceftazidima (CAZ) e aztreonam (ATM). Foi realizada a Reação em Cadeia da Polimerase (PCR), para identificação dos genes de resistência *bla*_{CTX-M-2}, *bla*_{CTX-M-8}, *bla*_{CTX-M-15}, *bla*_{SHV} e *bla*_{TEM}. A presença de pelo menos um dos genes citados foi verificada em 35,6% do total (n=74). A distribuição dos genes nas amostras positivas ocorreu da seguinte forma: 3,84% (n=8) para *bla*_{CTX-M-2}, 5,28% (n=11) para *bla*_{SHV} e 29,3% (n=61) para o gene *bla*_{TEM}. Os genes *bla*_{CTX-M-15} e *bla*_{CTX-M-8}

606 não foram observados no estudo. Quanto à suscetibilidade aos antibióticos, as amostras
607 apresentaram altas taxas de resistência ao grupo dos beta-lactâmicos, bem como para
608 tetraciclina (80%). Foi possível verificar que 50,9% (106/208) apresentaram perfil de
609 multirresistência. Este estudo demonstra que bezerros podem ser reservatórios de cepas
610 de *E. coli* produtoras de ESBL, constituindo um risco à saúde humana e animal, pelo risco
611 de disseminação desses isolados no ambiente.

612 **Palavras-chave:** beta-lactâmicos, ESBL, pecuária e resistência à antibióticos.

613

614

ABSTRACT

615

616 Beta-lactamase production by Enterobacteriales is the most common resistance
617 mechanism for beta-lactam antibiotics. Extended spectrum beta-lactamase (ESBL) are
618 considered a world clinical and epidemiological urgency. This study aimed to verify the
619 presence of diarrheagenic *Escherichia coli* strains, isolated from calves, ESBL producers.
620 Fecal samples from 208 calves, from two distinct age groups, were obtained and
621 cultivated in enrichment for *E. coli* isolation. Phenotypic analysis was performed by the
622 approximation disc screening method employing amoxicillin discs + clavulonic acid
623 (AMC), ceftriaxone (CRO), cefotaxime (CTX), ceftazidima (CAZ) and aztreonam
624 (ATM). Polymerase Chain Reaction (PCR) was performed to identify the *bla*_{CTX-M-2},
625 *bla*_{CTX-M-8}, *bla*_{CTX-M-15}, *bla*_{SHV} and *bla*_{TEM} resistance genes. The presence of at least one of
626 the genes cited was verified in 35.6% of the total (n = 74). The distribution of genes in
627 the positive samples occurred as follows: 3.84% (n = 8) to *bla*_{CTX-M-2}, 5.28% (n = 11) for
628 *bla*_{SHV} and 29.3% (n = 61) for *bla*_{TEM} gene. The *bla*_{CTX-M-15} and *bla*_{CTX-M-8} genes were not
629 observed in the study. Regarding susceptibility to antibiotics, the samples showed high
630 resistance rates to the beta-lactam group, as well as for tetracycline (80%). It was possible
631 to verify that 50.9% (106/208) had multipurpose profile. This study demonstrates that
632 calves can be *E. coli* strains reservoirs of ESBL producing, constituting a risk to human
633 and animal health, due to the risk of dissemination of these isolates in the environment.

634 **Keywords:** beta-lactams, ESBL, livestock and antibiotic resistance.

635

636

INTRODUCTION

637

638 The identification of multiresistant bacteria and their virulence genes in livestock
639 has great importance in public health worldwide, as well as the identification of possible

640 antibiotic resistance genes, resulting from exacerbated therapeutic conduct and selective
641 pressure (Han *et al.*, 2017).

642 Diarrhea in calves is one of the main causes of economic losses in Brazilian and
643 worldwide livestock. Since the beginning of the 20th century, the worldwide scientific
644 community has pointed to the bacterium *Escherichia coli* as one of the main agents
645 involved in diarrhea of infectious origin, therefore, this bacterial genus and its
646 diarrheagenic strains have been studied and identified (Smith; Orcutt, 1925; Coura *et al.*,
647 2015; Tutija *et al.*, 2020).

648 The differentiation between *E. coli* pathogenic strains from non-pathogenic strains
649 is based on the production of virulence factors, as well as on the identification of the
650 mechanisms related to disease. In response to the widespread use of antimicrobials in the
651 treatment of enteritis cases in calves, spread of virulence factors between herds may
652 occur. This event may be relevant for the emergence and dissemination of bacteria
653 capable of expressing enzymes that confer resistance to multiple antibiotics (Coura *et al.*,
654 2015; Poirel *et al.*, 2018).

655 Several studies have pointed to the worldwide growth of this resistance by
656 members of Enterobacteriales, such as the emergence of ESBL, which can irreversibly
657 hydrolyze the beta-lactam ring of the amine bond, preventing its action on transpeptidases
658 (Bush and Jacoby, 2010). They can hydrolyze all penicillins, cephalosporins including
659 third and fourth generation and aztreonam (Coque *et al.*, 2008).

660 Feces of production animals, which have strains with antimicrobial resistance
661 genes, such as the ESBL enzyme, can be constant reservoirs of dissemination to humans
662 (Carattoli, 2008; Schmidt *et al.*, 2013), the environment and other animals (Boonyasiri *et*
663 *al.*, 2014; Tekiner; Ozpinar, 2016; Melo *et al.*, 2018; Brisola *et al.*, 2019), leading to
664 delays in appropriate therapy, increased morbidity, mortality, and consequently,
665 economic losses to producers and industry (Santajit; Indrawattana, 2016; Chong *et al.*,
666 2018).

667 Thus, the aim of this study was to evaluate the presence of diarrheagenic *E. coli*
668 strains producing ESBL, from an important beef cattle producing region in Brazil.

669

670

MATERIAL AND METHODS

671

672 Two hundred and eight isolates (n = 208) of diarrheagenic *E. coli* were used in
673 this study. These isolates were previously characterized as diarrheagenic *E. coli* by Tutija
674 *et al.* (2020).

675 These isolates came from stool and rectal swabs, from beef calves, with ages
676 varying from one to 60 days. The study was conducted at the Veterinary Bacteriology
677 Laboratory at the Universidade Federal de Mato Grosso do Sul (UFMS), from March
678 2021 to December 2022.

679 In vitro susceptibility to antimicrobials was verified, according to the antibiogram
680 method standardized by the CSLI (Clinical Laboratory Standard Institute) (2021), which
681 is based on the method originally described by Bauer *et al.* (1966). The most used
682 antibiotics in the field reported by the veterinarians themselves were used, such as:
683 amoxicillin + clavulanic acid (AMC), amoxicillin (AMO), cephalixin (CFE),
684 enrofloxacin (ENO), florfenicol (FLF), gentamicin (GEN), norfloxacin (NOR), penicillin
685 G (PEN), sulfamethaxazol+trimethoprim (SUT) and tetracycline (TET).

686 As for the phenotypic analysis of ESBL-encoding genes, the approximation disk
687 screening technique according to Jarlier *et al.* (1988), was performed using amoxicillin +
688 clavulanic acid (AMC), ceftriaxone (CRO), cefotaxime (CTX), ceftazidime (CAZ) and
689 aztreonam (ATM) discs. A sample of *E. coli* ATCC® 25922 was used as a control,
690 verifying the interaction of the AMC antibiotic with the others.

691 Subsequently, molecular analysis of the diarrheagenic *E. coli* isolates was
692 performed to identify the genes encoding the ESBL enzyme, such as *bla*_{CTX-M} (including
693 *bla*_{CTX-M-2}, *bla*_{CTX-M-8} and *bla*_{CTX-M-15}), *bla*_{SHV} and *bla*_{TEM}, using Reaction Polymerase in
694 Chain (PCR).

695 The isolates were suspended in BHI broth (brain and heart infusion), incubated in
696 a culture oven for 24 hours. After this time, the centrifugation was performed and the
697 extraction of the sediment DNA started, according to the protocol described by Adwan
698 (2014). For the analysis of DNA purity and its quantification, the NanoDrop™ OneC
699 (Thermo Fischer Scientific) was used.

700 For the amplification of target genes (Tab. 1), the reactions were performed with
701 a final volume of 25µL, containing 2.5µL of 10X buffer (20 Mm Tris-HCl, pH 8.3, 50
702 mM KCl), 1.5 mM MgCl₂, 0.2 mM dNTP, 1.25 U of *Taq* DNA polymerase (5 U/µL), 10
703 pmol of each primer (100 ng/µL) and 2µL of DNA (approximately 50 ng).

704 Amplification conditions for the CTX-M-like enzyme included: 1) initial
705 denaturation at 94°C for 5 minutes; 2) denaturation at 94°C for 1 minute; 3) annealing at

706 a specific temperature for each primer, for 1 minute; 4) extension at 72°C for 1 minute;
 707 5) repetitions of steps 2, 3 and 4, 29 times; ending with step 6, the final extension at 72°C,
 708 for 5 minutes.

709 As for the SHV-type enzyme, they included: 1) initial denaturation at 94°C for 5
 710 minutes; 2) denaturation at 94°C for 30 seconds; 3) annealing at a specific temperature
 711 for each initiator, for 30 seconds; 4) extension at 72°C for 1 minute; 5) repetitions of steps
 712 2, 3 and 4, 32 times; with final extension at 72°C for 10 minutes.

713 For the TEM-type enzyme, they included: 1) initial denaturation at 94°C for 5
 714 minutes; 2) denaturation at 95°C for 30 seconds; 3) annealing at a specific temperature
 715 for each primer, for 1 minute; 4) extension at 72°C for 2 minutes; 5) repetitions of steps
 716 2, 3 and 4, 32 times; with final extension at 72°C for 5 minutes.

717

718 Table 1. Initiators to identify the genes encoding ESBLs.

Genes	Sequence 5'-3'	T (°C) *	bp**	References
<i>bla_{CTX-M-2}</i>	F5'-GCGACCTGGTTAACTACAATC-3' R5'-CGGTAGTATTGCCCTTAAGCC-3'	55	351	Tollentino <i>et al.</i> , 2011
<i>bla_{CTX-M-8}</i>	F5'-CTGGAGAAAAGCAGCGGGGG-3' R5'-ACCCAGGATGTGGGTAGCCC-3'	55	320	Bonnet <i>et al.</i> , 2000
<i>bla_{CTX-M-15}</i>	F5'-CACACGTGGAATTTAGGGACT-3' R5'-GCCGTCTAAGGCGATAAACA-3'	56	550	Muzaheed <i>et al.</i> , 2008
<i>bla_{SHV}</i>	F5'-AGGATTGACTGCCTTTTTG-3' R5'-ATTTGCTGATTTGCTCG3'	54	392	Colom <i>et al.</i> , 2003
<i>bla_{TEM}</i>	F5'-TTGGGTGCACGAGTGGGTTA-3' R5'-TAATTGTTGCCGGAAGCTA-3'	54	465	Gangoue-Pieboji <i>et al.</i> , 2005

719 *T (°C): annealing temperature; ** bp: base pairs.

720

721 Subsequently, the amplified products were analyzed by electrophoresis in agarose
 722 gel 1%, together with *E. coli* ATCC® 25922. Molecular weight marker (Ladder 100bp,
 723 Ludwig Biotec, Brazil), and Gel Red nucleic acid stain (Biotium, USA) were used. The
 724 amplified products were visualized under ultraviolet light, with the aid of a transilluminator
 725 (Transiluminator UV for gel - LTB HE, Locus, Brazil).

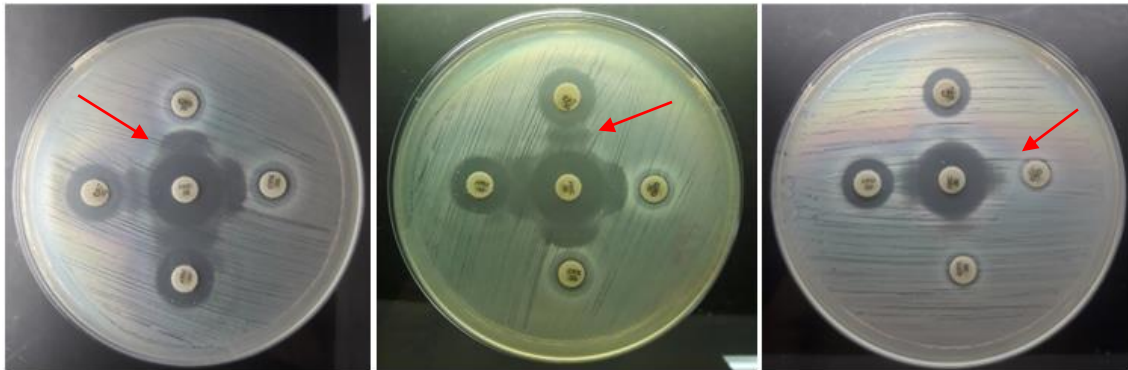
726

727

RESULTS

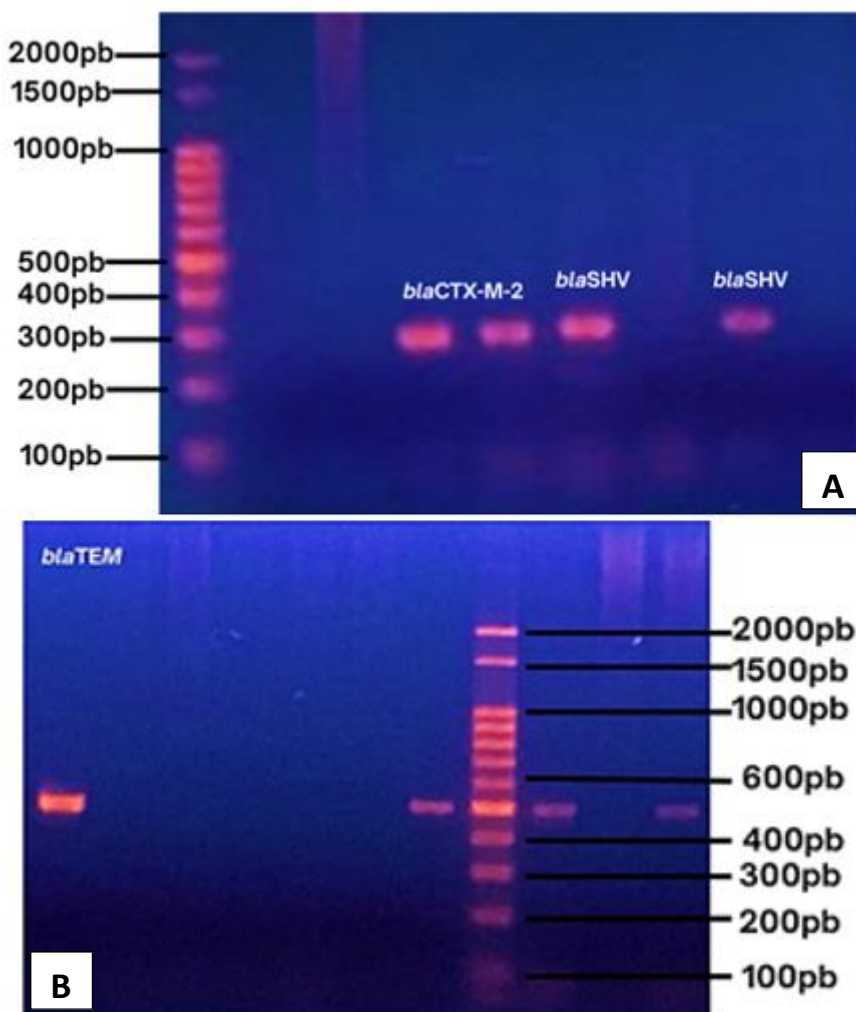
728

729 In the phenotypic analysis, an increase in the diameter of the inhibition zone
730 towards the AMC disk or the appearance of the phantom zone was observed in 31.25%
731 (65/208) samples, as shown in the figure 1.
732



733
734 Figure 1: Positive approximation disk screening technique demonstrating the phantom zone or absence of
735 inhibition halo (red arrow) to verify samples possibly producing the ESBL enzyme.
736

737 Genotypic analysis by the PCR method for the search for ESBL coding enzymes
738 showed positivity in 3.84% (8/208) of the samples for the *bla*_{CTX-M-2} enzymes, 5.28%
739 (11/208) for the gene *bla*_{SHV} and 29.3% (61/208) for the *bla*_{TEM} gene (Figure 2). The
740 *bla*_{CTX-M-15} and *bla*_{CTX-M-8} genes were not observed in the study.
741



742

743 Figure 2: Electrophoresis of positive samples on a 1% agarose gel, using a molecular marker of 100 base
 744 pairs, visualizing the amplification of the *bla*_{CTX-M-2} and *bla*_{SHV} (A), as well as the *bla*_{TEM} gene (B).

745

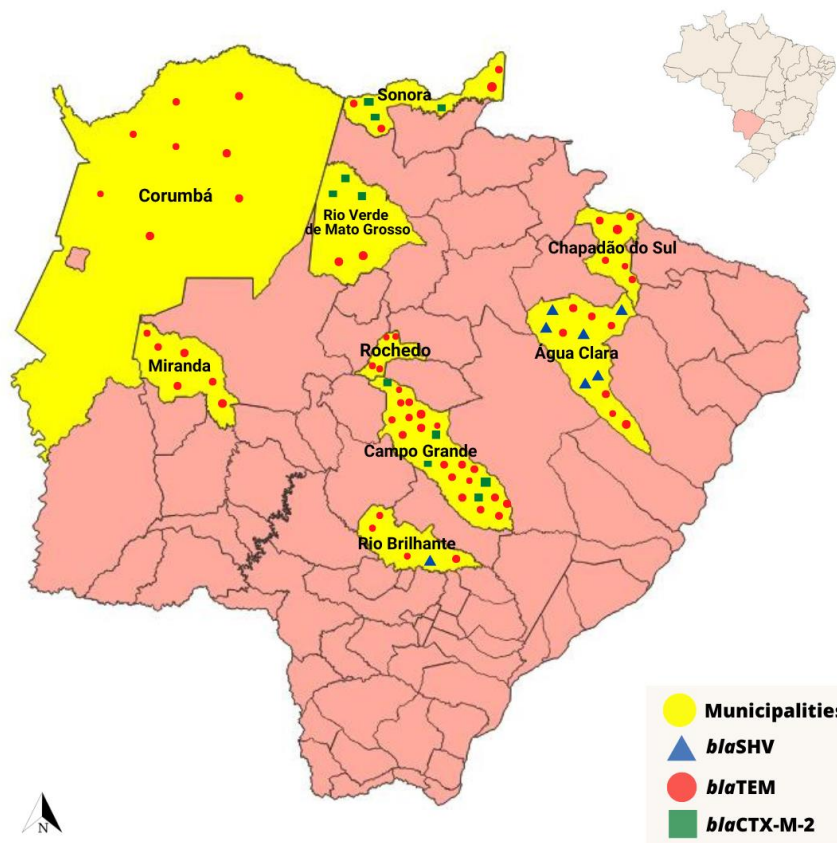
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Through genetic identification, it was possible to observe that in the four macro regions of the state of Mato Grosso do Sul, at least one ESBL coding gene was located (Figure 3).



750

751 Figure 3: Map of the state of Mato Grosso do Sul, Brazil, showing the positive municipalities (yellow) for
 752 the genotypic identification of the ESBL-encoding genes: *bla*_{CTX-M-2} (triangle), *bla*_{SHV} (square) and *bla*_{TEM}
 753 (circle).

754

755 In this study, it was possible to observe that 65.7% of the positive samples for
 756 ESBL coding genes are from calves between 1 and 30 days of age and 34.2% between 31
 757 and 60 days.

758 In 35.6% (74/208) of the samples positive for the enzymatic variants CTX-M,
 759 SHV and TEM, the frequency of in vitro antimicrobial resistance of the class of beta-
 760 lactams tested can be observed (Tab. 2A) and, additionally, the correlation of the
 761 frequency of in vitro antimicrobial resistance of antimicrobials of the aminoglycosides,
 762 fluoroquinolones, tetracyclines, amphenicols and sulfonamides classes, with the
 763 localization of CTX-M, SHV and TEM family genes (Tab. 2B).

764

765 Table 2. Correlation of the frequency of antimicrobial resistance to beta-lactams (A) by the disk diffusion
 766 method and antimicrobials from the classes of aminoglycosides, fluoroquinolones, tetracyclines,
 767 amphenicols and sulfonamides (B), with the location of the *bla*_{CTX-M-2} genes, *bla*_{SHV} and *bla*_{TEM} from
 768 diarrheagenic *E. coli* isolates by the PCR method.

Antibiotics	Number of resistant samples (%)		
	<i>bla</i> _{CTX-M-2}	<i>bla</i> _{SHV}	<i>bla</i> _{TEM}
Penicilin G	8 (100%)	11 (100%)	61 (100%)
Amoxicillin	7 (87.5%)	4 (36.36%)	43 (70.49%)

Cephalexin	7 (87.5%)	6 (54.54%)	23 (37.70%)
Amoxicillin + Clavulanic Acid	3 (37.5%)	0	16 (26.22%)
B			
Tetracycline	7 (87.5%)	7 (63.63%)	55 (90.16%)
Sulfamethoxazole+Trimethoprim	4 (50%)	4 (36.36%)	35 (57.37%)
Enrofloxacin	4 (50%)	6 (54.54%)	22 (36.06%)
Norfloxacin	3 (37.5%)	5 (45.45%)	17 (27.86%)
Gentamicin	3 (37.5%)	5 (45.45%)	14 (22.95%)
Florfenicol	2 (25%)	0	7 (11.47%)

769

770 After performing the antibiogram, the frequency of in vitro antimicrobial
 771 resistance of the 208 samples was observed (Tab. 3), in which the highest resistance rate
 772 was for penicillin G (100%), followed by tetracycline (80%), amoxicilin (61%),
 773 sulfametaxazole+trimethoprim (45.7%) and cephalixin (45%) and with less frequency of
 774 antimicrobial resistance, florfenicol (12%).

775

776 Table 3. Frequency of in vitro antimicrobial resistance in 208 diarrheagenic *Escherichia coli* isolates from
 777 stool and rectal swab samples.

Antibiotics	Number of resistant samples (%)
Penicilin	208 (100%)
Tetracycline	166 (80%)
Amoxicillin	126 (61%)
Cephalexin	94 (45%)
Sulfamethoxazole+Trimethoprim	95 (45.7%)
Enrofloxacin	66 (32%)
Norfloxacin	50 (24%)
Amoxicillin + Clavulanic Acid	43 (20.6%)
Gentamicin	39 (19%)
Florfenicol	25 (12%)

778

779 One hundred and six (50.9%) samples showed multidrug-resistance to three or
 780 more classes of antibiotics, in which no antibiotic was 100% effective, of these, 48.11%
 781 were positive for the phenotypic disk approximation test, and 42.45% molecularly
 782 positive for ESBL enzymes in relation to the presence of multidrug-resistance.

783 Twenty-eight (13.5%) samples of diarrheagenic *E. coli* were positive for the disk-
 784 proximation screening test and for the presence of one of the genes encoding ESBL.
 785 Forty-five samples (21.6%) did not express the phantom zone or lack of inhibition zone
 786 and were molecularly positive. However, 32 (15.4%) samples tested positive for the
 787 screening test and were molecularly negative.

788

789

DISCUSSION

790

791 Production animals, such as cattle, are considered one of the main reservoirs of *E.*
792 *coli* producing ESBL (Chong *et al.*, 2018).

793 According to the age of the animals, Brunton *et al.* (2014), demonstrated in their
794 study that resistance genes may be less prevalent in relation to the growth of the animals,
795 due to the increase in the physical area in which they remain, hindering the transmission
796 of resistance genes between them. In this case, the animals in the study were not weaned
797 and were in the same physical area, according to the selected farm. The limitation of the
798 age range in the present study, up to 60-days-old, may have influenced the non-alteration
799 of the indices. When extending this age range to more than 60 days, it is possible that the
800 variation mentioned by Brunton *et al.* (2014) could be noticed.

801 The situation of ESBL incidence in the world has undergone several changes, and
802 in the new epidemiological scenario, the prevalence of enzymes of the CTX-M family
803 stands out, both in hospital isolates, the community, and animals. This worldwide spread
804 was described as “the CTX-M pandemic” (Cantón *et al.*, 2012).

805 In the present study, enzymes of the CTX-M family were highlighted in 8 (3.84%)
806 of the 208 fecal samples from calves. The study by Sukmawinata *et al.* (2020) confirmed
807 the presence of this family in the feces of thoroughbred horses in Japan, as well as the
808 study by Gundran *et al.* (2019) which obtained 89.86% gene positivity in isolates from
809 broiler chickens in the Philippines, confirming the cosmopolitanization of the production
810 of ESBL type CTX-M.

811 In the molecular analysis, the prevalence of genes encoding ESBL was higher for
812 the *bla*_{TEM} gene (29.3%), followed by *bla*_{SHV} (5.28%) and, lastly, by the *bla*_{CTX-M-2} gene
813 (3.84%). The *bla*_{CTX-M-15} and *bla*_{CTX-M-8} genes were not observed in this study.

814 In Brazilian cattle, the *bla*_{CTX-M-15} and *bla*_{CTX-M-2} genes were found in the Southeast
815 region, mainly in the states of São Paulo and Rio de Janeiro (Rocha *et al.*, 2016), as well
816 as in the Northeast region (Palmeira *et al.*, 2020). Sukmawinata *et al.* (2020) reported a
817 high prevalence of the *bla*_{CTX-M-2} gene, however, among the few studies that investigated
818 the prevalence of ESBL in calves, Eller *et al.* (2014) reported a high prevalence of *bla*_{CTX-}
819 _{M8}, unlike our study.

820 The studies reported in human and veterinary medicine by Cergole-Novella *et al.*
821 (2010), Rocha *et al.* (2016), Widodo *et al.* (2020), and Ejaz *et al.* (2021) demonstrated
822 that the prevalence of the *bla*_{CTX-M}, *bla*_{TEM} and *bla*_{SHV} gene is increasing among ESBL-
823 producing bacteria, that corroborates our study. They are common in humans (Aworh *et*
824 *al.*, 2022), especially in nosocomial infections; in animals - dogs, cats, horses, chickens,

825 pigs, cattle, wild animals (Ewers *et al.*, 2010; Melo *et al.*, 2018; Brisola *et al.*, 2019) - and
826 in the environment (Cantón *et al.*, 2012).

827 For Aworh *et al.* (2022), healthcare professionals should be aware that people who
828 have close contact with calves and their feces are a high-risk group for spreading *E. coli*
829 with ESBL, when compared to the general population. In their study, in addition to
830 isolating *E. coli* with ESBL in 45.4% of calves' feces from an abattoir, they observed the
831 presence of this enzyme in workers (41.2%) and in the work environment (13.4%).

832 The WHO (2017) lists bacterial agents and their respective antimicrobial
833 resistance in human medicine, but within veterinary medicine we can already observe the
834 cosmopolitanization of *Acinetobacter baumannii* (Pomba *et al.*, 2015; Ewers *et al.*, 2017),
835 *Pseudomonas aeruginosa* (Fernandes *et al.*, 2018; Elshafiee *et al.*, 2019), as well as
836 ESBL-producing *E. coli* (Melo *et al.*, 2018; Brisola *et al.*, 2019; Palmeira *et al.*, 2020) in
837 the critical priority group, and in the high priority group, *Staphylococcus aureus* (Oliveira
838 *et al.*, 2016), *Salmonella* species (Colobatiu *et al.*, 2015; Pribul *et al.*, 2017) and
839 *Campylobacter* species (Frasao *et al.*, 2015), at the human-animal-environment interface.

840 The production of ESBL by *E. coli* bacteria is a major public health problem, as
841 infections by these bacteria can result in failure of beta-lactam therapy (Chong *et al.*,
842 2018). Another reason is the phenomenon of co-resistance to other antibiotics, as they are
843 encoded by plasmids. Profiles of resistance genes for aminoglycoside, macrolides,
844 tetracyclines and fluoroquinolones antibiotics were observed by Carey *et al.* (2022) for
845 the *bla_{CTX-M}* gene, which may facilitate co-selection processes, limiting current
846 therapeutic options (Paterson; Bonomo, 2005; Cantón *et al.*, 2012; Adeyankinnu *et al.*,
847 2014).

848 According to a report by the OIE (2021), tetracyclines were the most used
849 antibiotics in food-producing animals, followed by macrolides and penicillins.

850 Regarding antimicrobial resistance to the beta-lactam class and the location of the
851 genes that encode ESBL, we could observe that the highest index of resistance was for
852 penicillin G (100%), similar results to those of Shahrani *et al.* (2014), who evaluated the
853 antimicrobial resistance profile correlated with the identification of the STEC virulence
854 gene in samples of diarrhea from calves. However, it is known that members of the
855 Enterobacterales group, such as *E. coli*, are intrinsically resistant to penicillin G, that is,
856 this drug was being administered incorrectly, possibly due to the lack of knowledge of
857 the bacteria's intrinsic resistance.

858 Brower *et al.* (2017), Rahmahani *et al.* (2020), and Jarrige *et al.* (2020), observed
859 a profile of significant resistance to tetracycline in poultry farms (47%), domestic
860 chickens (48%), from cloacal swab, and in milk dairy calves (68%), results that
861 corroborate the findings of this study, presenting antimicrobial resistance in 80%
862 (166/208) for tetracycline. In addition, records from the farms tested showed that, at some
863 point during management, tetracycline was administered to the calves in the study.

864 Bacteria showed the lowest frequency of resistance to florfenicol, with only 12%
865 (25/208) of inhibition failure in the tested samples, as well as the study by Nespolo *et al.*
866 (2014), who analyzed 52 bovine isolates destined for slaughter, obtained a resistance
867 index of 1.92% to florfenicol.

868 *E. coli* is also prone to developing broad drug resistance, as it has a great capacity
869 to accumulate resistance genes, mainly by horizontal gene transfer, which leads to
870 infections that are difficult to treat (Poirel *et al.*, 2018). In the present study it was
871 observed three samples with two tested genes, *bla_{SHV}* and *bla_{TEM}*, and three samples with
872 *bla_{CTX-M-2}* and *bla_{TEM}*, as well as their multidrug resistance to six classes of antibiotics,
873 such as beta-lactams, fluoroquinolones, tetracycline, aminoglycoside, sulfonamides and
874 amphenocol.

875 The multidrug-resistance profile, characterized by its resistance to three or more
876 classes of antimicrobials (Magiorakos *et al.*, 2011), was found in 50.9% (106/208) of the
877 isolates, results like Cergole-Novella *et al.* (2010) and Brisola *et al.* (2019), in which
878 multidrug resistance was observed in eight of the 12 samples (66.7%) and 50 of the 135
879 samples (37.4%), respectively.

880 Al-Tamimi *et al.* (2019) proposes to use the phenotypic synergy test to detect the
881 synergistic ability of clavulanic acid with cephalosporins and monobactam in ESBL-
882 positive isolates. However, in our study there were divergences between positive samples
883 in the phenotypic test (15.4%) and negative in the genotypic analysis, which could explain
884 this phenomenon is the use of specific primers for the CTX-M, SHV and TEM families
885 in this study, and the literature mentions other genes that can encode ESBL enzymes such
886 as *bla_{CMY}* (Brisola *et al.*, 2019) and *bla_{OXA}* (Palmeira *et al.*, 2020).

887 In the phenotypic synergism test, the ideal distance for high synergistic activity is
888 20 to 25 mm between cephalosporins/monobactam and amoxicillin and clavulanic acid
889 (Jarlier *et al.*, 1988; Bengtsson-Palme *et al.*, 2017). In the present study, 45 (21.6%)
890 samples did not express the phantom zone or any zone of inhibition and were molecularly

891 positive, possibly greater distances could decrease the sensitivity of the test, hiding most
892 of the resistance factors during the laboratory routine (Al-Tamimi *et al.*, 2019).

893 Many production animals, carrying these strains of ESBL, can be constant
894 reservoirs of dissemination for humans, such as, for example, the contamination of
895 products of animal origin (Tekiner; Ozpinar, 2016).

896 Currently, there are no national programs for monitoring bacterial resistance and
897 its mechanisms in livestock, which makes it difficult to estimate the proportion of ESBL-
898 producing strains in the federations. In turn, human and animal infections by ESBL-
899 producing bacteria is associated with increased mortality, morbidity, high cost of
900 hospitalization, delay in adequate therapy and, consequently, economic losses for
901 producers (Santajit; Indrawattana, 2016; Chong *et al.*, 2018).

902

903

CONCLUSION

904

905 This study presents the first report on the identification of the *bla*_{CTX-M-2}, *bla*_{SHV}
906 and *bla*_{TEM} genes in stool samples from calves from Mato Grosso do Sul, Brazil, alerting
907 veterinarians about the use and knowledge of antibiotics in livestock, indicating the need
908 to establish strategies for surveillance and tracking of antibiotic resistance in the country.

909

910

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911

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1143 5. (ARTIGO II)

1144

1145 [Artigo formatado de acordo com as normas da revista *Journal of Biophotonics*]

1146

1147 **IDENTIFICATION OF MULTI-RESISTANT DIARRHEAGENIC *Escherichia*** 1148 ***coli* BY USING FTIR SPECTROSCOPY AND MACHINE LEARNING: A** 1149 **FEASIBLE STRATEGY TO IMPROVE THE GROUP CLASSIFICATION**

1150

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1159

1160 **ABSTRACT**

1161

1162 The identification of multidrug-resistant strains from *E. coli* species responsible for
1163 diarrhea in calves still faces many laboratory limitations and is necessary for properly
1164 monitoring the microorganism spread and control. Then, there is a need for the
1165 development of a screening tool for bacterial strain identification in microbiology
1166 laboratories, which must show easy implementation, fast response, and accurate results.
1167 The use of FTIR spectroscopy to identify microorganisms has been successfully
1168 demonstrated in the literature, including many bacterial strains, here we explored the
1169 FTIR potential for multidrug-resistant *E. coli* identification. First, we applied principal
1170 component analysis to observe the group formation tendency, the first results showed no
1171 clustering tendency with a messy sample score distribution, then we improved these
1172 results by properly selecting the main principal components. Finally, by using machine
1173 learning algorithms, a predicting model showed 70% overall accuracy in the validation
1174 test, which can be considered a great result for a screening tool. The present results

1175 demonstrate the viability of the method for microorganism identification, which can
1176 significantly contribute to its control.

1177 **Keywords:** multi-resistant bacteria, diarrheagenic, *Escherichia coli*, FTIR, machine
1178 learning.

1179

1180

INTRODUCTION

1181

1182 The introduction of antibiotics into clinical use was the greatest medical advance of
1183 the 20th century, and the rapid discovery of several classes of antibiotics in a relatively
1184 short period has led to the overuse of these drugs, which help promotes increasing
1185 antimicrobial resistance and antibiotics effectiveness loss [1]. The O'Neill report [2]
1186 predicted that without urgent action, ten million people a year will die from drug-resistant
1187 infections by 2050. As a warning to the world's population about the growing global
1188 resistance to antimicrobials, the WHO, World Health Organization [3] lists bacterial
1189 agents and their respective resistance to antimicrobials, divided into critical, high, and
1190 medium priority.

1191 Important strategies have been adopted in this scenario for world public health
1192 maintenance, between them the identification of multidrug-resistant bacteria in herds of
1193 animals [4], since we have observed the migration of this microorganism to a human-
1194 animal-environment interface, such as *Acinetobacter baumannii* [5, 6], *Pseudomonas*
1195 *aeruginosa* [7, 8], *Staphylococcus aureus* [9]; *Salmonellas species* [10, 11],
1196 *Campylobacter species* [12], and *Escherichia coli* [13, 14, 15].

1197 Besides, diarrhea in calves is one of the main causes of economic losses in Brazilian
1198 and worldwide livestock. Since the beginning of the 20th century, the worldwide
1199 scientific community points to the bacteria *E. coli* as one of the main agents involved in
1200 diarrhea of infectious origin; therefore, this bacterial genus and its pathogenic strains
1201 began to be studied and identified [16, 17, 18, 19].

1202 The *E. coli* bacteria is considered a commensal of the intestinal flora, only a small
1203 part of the strains has pathogenicity responsible for diseases and the differentiation of
1204 pathogenic (diarrheagenic) strains from non-pathogenic strains is based on the production
1205 of virulence factors [17]. Once *E. coli* bacteria with virulence factors spread in the cattle
1206 herd, serious concerns arise due to its capability of expressing enzymes that confer
1207 resistance to multiple antimicrobials [20, 17, 21]. Animal feces, which have multiresistant

1208 strains, serve as constant reservoirs for bacteria dissemination to humans [22, 23], the
1209 environment, and other animals [24, 25] which hinders the adequate therapy, increased
1210 morbidity, mortality, and cause economic losses to producers and industry [26, 27].

1211 Phenotypic and molecular methods have been used in bacterial culture to analyze the
1212 nucleic acid sequences and identify microorganisms with multidrug resistance
1213 characteristics, but their variable discriminatory capacity, high cost, and time-consuming
1214 experimental routine are difficult the implementation as a standard laboratory procedure
1215 [28, 29].

1216 In an attempt to overcome such limitations, several studies have been developed by
1217 using Fourier Transform Infrared Spectroscopy (FTIR) as a screening test [30, 31], which
1218 can obtain information about the sample chemical compounds through their vibrational
1219 molecular modes, enabling the identification of sample molecular composition (nucleic
1220 acids, proteins, lipids, and carbohydrates) of these bacterial cells [32, 33, 34].

1221 The potential use of FTIR spectroscopy to discriminate, classify and identify
1222 microorganisms has been successfully demonstrated in the literature, including many
1223 bacterial strains from Gram-positive, and Gram-negative species [32, 35, 36, 37, 38, 39,
1224 40, 41]. The specific spectral patterns observed for each molecular group can be revealed
1225 with multivariate analysis and/or machine learning algorithms aid, which enable FTIR
1226 spectroscopy as an alternative for rapid bacterial identification at the subspecies level [42,
1227 43]. Despite its great applicability for microorganism identification in recent years, this
1228 technique's potential in microbiology routines is still underestimated, due to the difficult
1229 acceptance and comprehension of the methodology by the microbiological community
1230 [31].

1231 Therefore, here we explore the use of FTIR spectroscopy associated with machine
1232 learning algorithms for data analysis to identify and classify multidrug-resistant strains
1233 from *E. coli* species responsible for diarrhea in calves, which may add value to the
1234 microbiology laboratory toolbox and society [26, 27, 31]. Our study is focused on the
1235 development of a simple laboratory routine to sample preparation and data acquisition,
1236 then our data analysis aims to improve the method's accuracy for future implementation
1237 as a screening tool in bacterial strain identification in microbiology laboratories.

1238

1239

METHODS

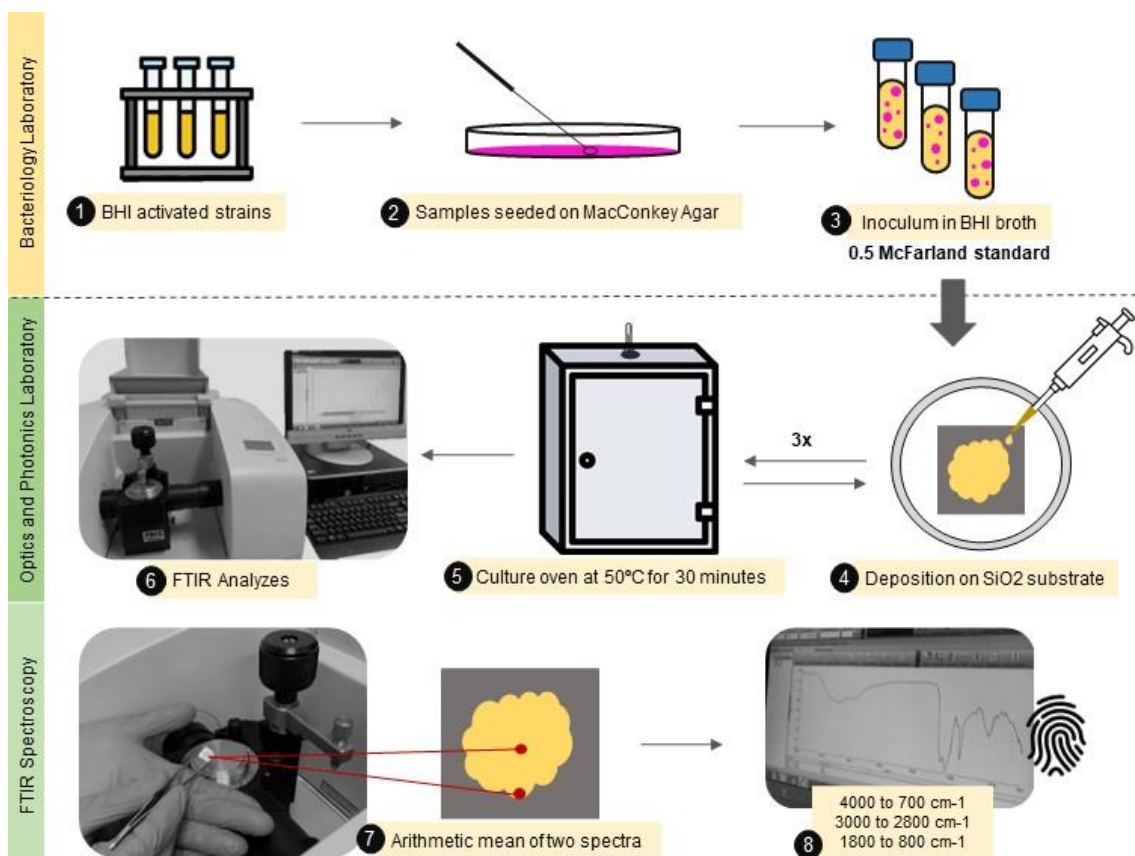
1240

1241 *Sample preparation and FTIR spectra acquisition*

1242 A total of 80 *Escherichia coli* isolates were analyzed. The sample pool was
 1243 divided into 40 multidrug-resistant (MR) *E. coli* isolates and 40 replicates of *E. coli*
 1244 ATCC® 25922. First, they were screened for analysis and placed in Brain Heart Infusion
 1245 (BHI) broth for activation, subsequently, seeded on MacConkey Agar to verify purity and
 1246 isolate selected colonies. Then, the colonies of MR *E. coli* and ATCC *E. coli* were
 1247 suspended in 1 mL of BHI broth, and a resulting turbid inoculum was adjusted to the 0.5
 1248 McFarland scale (10^8 CFU/mL) before the measurements.

1249 A small aliquot (30 μ L) of the bacterial isolate was carefully deposited onto a flat
 1250 silicon substrate (SiO_2) by casting, followed by drying at 50°C per 30 minutes – this
 1251 procedure was repeated three times until the obtention of a thick film. Each sample was
 1252 produced in duplicate, and the samples were analyzed in two different spots from the
 1253 center to the border, to avoid inhomogeneity in the sample composition due to the drying
 1254 process [44]. Finally, the mid-infrared spectra were obtained by using an attenuated total
 1255 reflectance accessory (ATR) at Fourier Transform Infrared Spectrophotometer (Spectrum
 1256 100, Perkin Elmer). The spectra were collected from 4000 to 700 cm^{-1} , with 4 cm^{-1}
 1257 resolution and 10 scans. This entire experimental roadmap is illustrated in Figure 1.

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1260 Figure 1: Experimental scheme describing the main steps from the sample (*E. coli*)
1261 preparation to FTIR spectra acquisition.
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1263 ***Data analysis and sample classification***

1264 The data analysis was performed in Python (version 3.9.12), using the Scikit-learn
1265 package (version 1.1.2) [45]. First, the FTIR spectra from two different spots of duplicate
1266 samples were averaged and then subjected to Standard Normal Variate (SNV) pre-
1267 processing method, which removes the variation from the baseline and rescales the
1268 spectral intensity, to prevent interference in the data analysis due to random experimental
1269 variations [46].

1270 The average FTIR-SNV spectra for MR *E. coli* and ATCC *E. coli* group, in the
1271 4000 to 800 cm^{-1} range, were submitted to principal component analysis (PCA) [47]. PCA
1272 is an unsupervised method that will project our pre-processed data set into a new
1273 dimension (PCs – principal components) which aims to maximize the data variance, this
1274 dimensionally reduced data set retains most of the information from the original variables
1275 and shows how each data sample is distributed in this new dimension (score plot),
1276 allowing cluster the similar samples and distinguish groups. Each PC represents a
1277 percentage of the data variance and, allows us to analyze the main spectral range that most
1278 contributes to the data variance percentage through the loading plot. PCA is an important
1279 step to visualize the group classification tendency. Here we analyzed three different
1280 ranges: (i) 4000 to 800 cm^{-1} ; (ii) 3000 to 2800 cm^{-1} , and (iii) 1800 to 800 cm^{-1} , to use only
1281 those vibrational modes that improve the group clustering and classification and eliminate
1282 highly correlated data [48].

1283 The sample classification is performed by prediction models built by machine
1284 learning (ML) algorithms using PC output data from 70% sample set. Before sample
1285 classification tests, we must determine the ideal number of PCs used by ML algorithms
1286 to avoid overfitting and underfitting [49, 50]. Here we used the Feature Selection
1287 Recursive Feature Elimination (RFE), which selects the main PCs that most contribute to
1288 achieving high accuracy and remove other PCs with the weakest contribution to correct
1289 sample classification in ML tests [51]. The use or removal of a determined PC was made
1290 based on the accuracy achieved by using Linear Discriminant Analysis (LDA) to classify
1291 the samples in a Leave One Out Cross-validation (LOOCV) test.

1292 In a brief description, Discriminant Analysis (DA) classifies the sample based on
1293 the distance between the sample data and the contour built by using a linear (L) or

1294 quadratic (Q) function to separate the classes (group) [52]. In LOOCV, one sample is
1295 taken from the data set, and the others are used to build the prediction model (training).
1296 Then, the prediction model accuracy is tested by using the sample data withdrawal from
1297 the data set. The procedure is repeated until all sample data have been tested [53].

1298 After determining the ideal number of PCs, and which PCs most contribute to
1299 sample classification in each spectral range analyzed, a LOOCV test was performed – by
1300 using the respective RFE-PCs data for each range – based on three different methods: (i)
1301 DA (described above); (ii) k-Nearest Neighbor (KNN), which uses the Euclidean distance
1302 between k closest neighbors to classify the sample [54]; and (iii) Support Vector Machine
1303 (SVM), which organizes each sample class through the optimization of a hyperplane –
1304 the hyperplane can be linear or nonlinear, being optimized to reach high performance –
1305 between the classes [55]. Finally, we were able to determine the best spectral range, ML
1306 algorithm, and PCs to build a predicting model, whose ability for generalization was
1307 tested in an external validation test by using 30% of the sample set.

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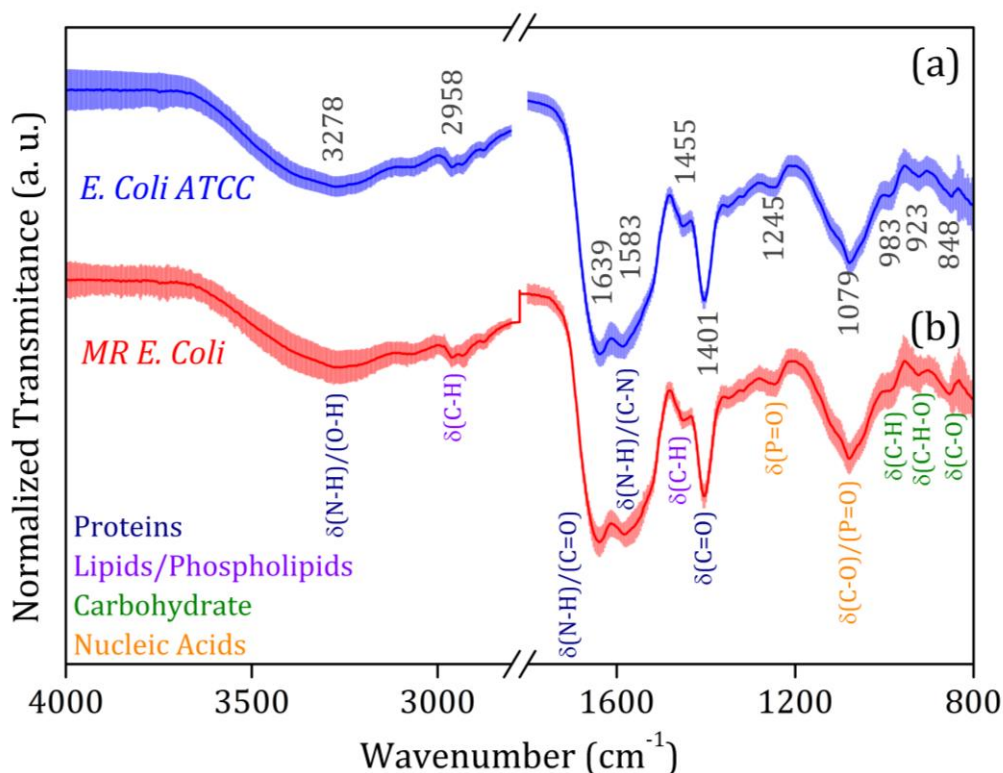
1309

RESULTS AND DISCUSSION

1310

1311 The average FTIR-SNV spectra for *E. coli* ATCC, and MR *E. coli* samples, Figure
1312 2, exhibit great similarity between them, with a small standard deviation into the data set.
1313 It suggests that direct identification of the isolate is not possible, and the data set shows
1314 great coherence. There is a remarkable presence of the three most pronounced bands in
1315 the spectra, the first around 1600 cm^{-1} is the result of overlapped bands 1639 and 1583
1316 cm^{-1} , assigned to Amides I and II from proteins, which is relatively wide, suggesting the
1317 presence of a third band, mainly due to the small shoulder observed around 1500 cm^{-1} .
1318 The vibrational bands assigned to Amides I (N-H and C=O) and II (N-H and C-N) have
1319 shown great contributions to biological sample classification in the literature [56, 57].
1320 The second band around 1400 cm^{-1} can be assigned to C-H and C=O vibrational modes
1321 from lipids and proteins, while the third band around 1080 cm^{-1} is usually assigned to
1322 nucleic acid and phospholipids. Also, weak bands assigned to C-H, and C-H-O vibrational
1323 modes from carbohydrate below 1000 cm^{-1} was identified in the spectra [58, 59].

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1326 **Figure 2:** Average FTIR-SNV spectra for (a) *E. coli* ATCC (blue line), and (b) multidrug-resistant
 1327 (MR) *E. coli* (red line) isolates. The main vibrational assignments are indicated in detail and their
 1328 related molecular groups (proteins, lipids, carbohydrates, and fatty acids) are identified by colors.
 1329 The symbol δ indicates the deformation vibrations.

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Figure 3 shows the principal component analysis results for the FTIR-SNV spectral data from *E. coli* ATCC, and MR *E. coli* sample groups. On the left side, we can observe the score plot, and on the right side, the loading plot for (i) 4000 to 800 cm^{-1} ; (ii) 3000 to 2800 cm^{-1} , and (iii) 1800 to 800 cm^{-1} range. For all cases analyzed two main characteristics stand out, the score plot exhibits a single cluster for both groups, which hinders the future group classification, and the loading shows few regions with a small contribution to the data variance, suggesting a small contribution of the bands for data variance in PC1 and PC2. We found PC1 and PC2 responsible for 79.2%, 96.8%, and 74.9% of data variance at 4000 to 800 cm^{-1} , 3000 to 2800 cm^{-1} , and 1800 to 800 cm^{-1} range, respectively, besides this great contribution for data variance previous studies have shown that the first PCs can be ignored, and high order PCs can be used to improve the group classification for ML algorithms [60].

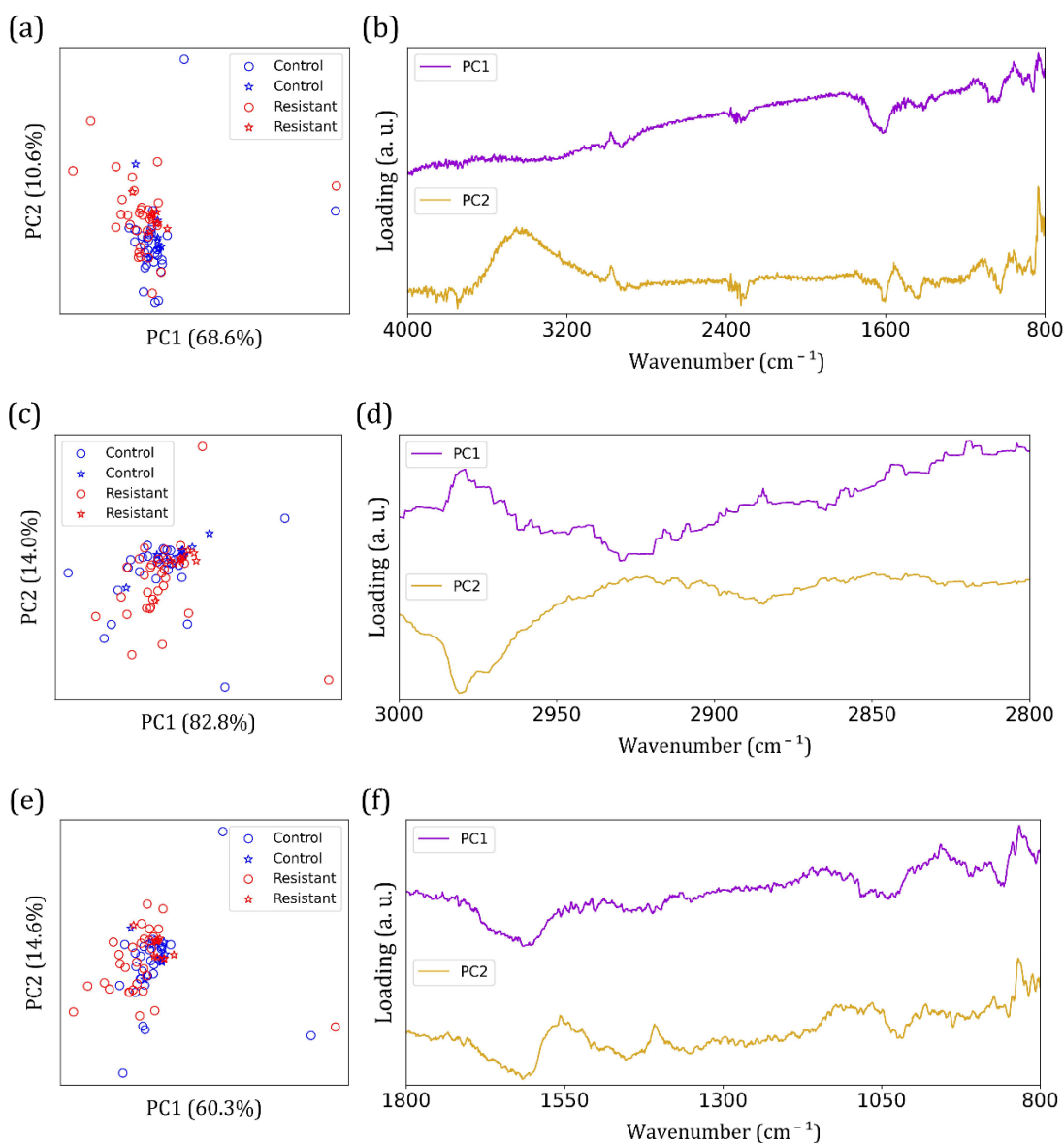


Figure 3: Principal Component Analysis for *E. coli* ATCC (blue circles/stars), and multidrug-resistant (MR) *E. coli* (red circles/stars) isolates average FTIR-SNV spectra. The score plot and respective loading for the three different ranges were analyzed: (a-b) 4000 to 800 cm⁻¹; (c-d) 3000 to 2800 cm⁻¹; and (e-f) 1800 to 800 cm⁻¹. The circles represent 70% of the sample data set used to build the predicting model, and the stars represent 30% of the sample data set used for external validation.

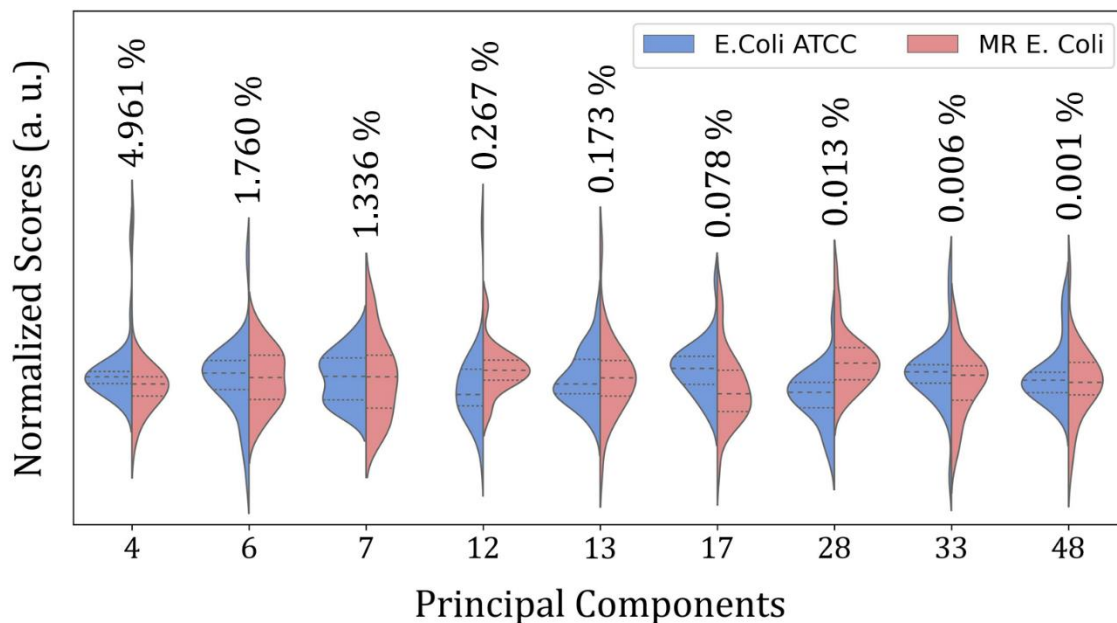
Usually, the proper choice of spectral range helps to improve group classification and clustering formation [61], since we use only spectral information that most contributes to clustering from those with highly correlated data, which hinders cluster formation. But here we couldn't succeed with this strategy probably because of the high similarities between the groups involved, and the highly correlated data present. The presence of antibiotic resistance genes in the MR *E. coli* group was mainly identified as resulting of the ESBL enzyme – previous study [62] - which is responsible for amine

1361 beta-lactam ring hydrolysis, resulting in antibiotics inactivation on the bacterial cell walls.
1362 Since it represents a break in the C-N bond (Amide group) and the appearance of N-H
1363 and O-H bonding, almost no difference in the IR spectra will be observed. Because, the
1364 C-N and N-H vibrational modes are superimposed around 1580 cm^{-1} , and O-H vibrational
1365 mode, around 3278 cm^{-1} , is too wide to provide enough information for group
1366 differentiation.

1367 Then, an alternative to improve group clustering and separation is the use of
1368 Feature Selection Recursive Feature Elimination (RFE) to find the best PCs (data
1369 projection) that most contribute to data separation. The RFE can select PCs that most
1370 contribute to achieving high accuracy and remove those with a small contribution. Here,
1371 the main PCs were determined by the overall accuracy achieved in a Leave One Out
1372 Cross-validation (LOOCV) test by using Linear Discriminant Analysis (LDA) [60]. The
1373 main PCs found for our data were: PC2, PC3, PC7, PC8, and PC19 for the 4000 to 800
1374 cm^{-1} range, responsible for 20.26% of data variance; PC5, PC13, PC21, PC36, and PC46
1375 for the 3000 to 2800 cm^{-1} range, responsible for 0.49% of data variance; and PC4, PC6,
1376 PC7, PC12, PC13, PC17, PC28, PC33, and PC48 for the 1800 to 800 cm^{-1} range
1377 responsible for 8.58% of data variance.

1378 Figure 4 shows the violin plot for the 9 most relevant PCs, in 1800 to 800 cm^{-1}
1379 range, with a clear monomodal distribution for the score data project over its respective
1380 PC for both sample groups, except for PC7, which exhibits a wide distribution for MR *E.*
1381 *coli* group, and a double peak maximum for *E. coli* ATCC, similar behavior observed for
1382 MR *E. coli* group in PC6. But the more important characteristic to be observed is the
1383 median value (dashed thick line around the peak center), which assumes a better distinct
1384 position projection over the axis for these PCs compared to the others and improves the
1385 clustering and group classification of our data in the LOOCV test with LDA.

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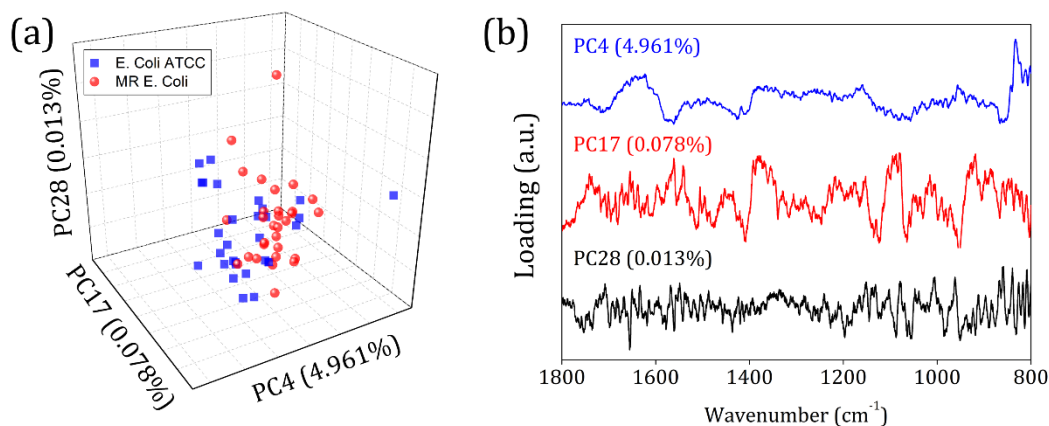
1387

1388 **Figure 4:** Violin plot for nine main PCs normalized scores selected by RFE in 1800 to 800 cm^{-1}
 1389 range from *E. coli* ATCC® 25922 (blue), and multidrug-resistant (MR) *E. coli* (red) isolates. The
 1390 thick dashed line represents the data distribution median. The data variance percentage of each
 1391 PC is described above each plot.
 1392

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1393 Figure 5 exemplifies the importance of these RFE-PCs. The score plot and loading
 1394 for PC4, PC17, and PC28 demonstrate the improvement of data clustering and group
 1395 separation. They were responsible for only 5.05% of the data variance, similar results
 1396 have been reported in the literature [60]. Besides that, the loading plot for each PC, Figure
 1397 5 (right column), shows a contribution for data variance in the 1600 to 1000 cm^{-1} range
 1398 in accordance with the main bands identified in the FTIR-SNV spectra (Figure 2) assigned
 1399 to protein and phospholipids molecules. In this case, with great similarity among the
 1400 samples, then between the spectra – high data correlation – the first PCs can be
 1401 responsible for a great percentage of data variance, with a small contribution to group
 1402 separation. However, low data variance PCs can still be useful for group classification
 1403 and better influence the algorithm performance, as demonstrated by Figure 5 results.
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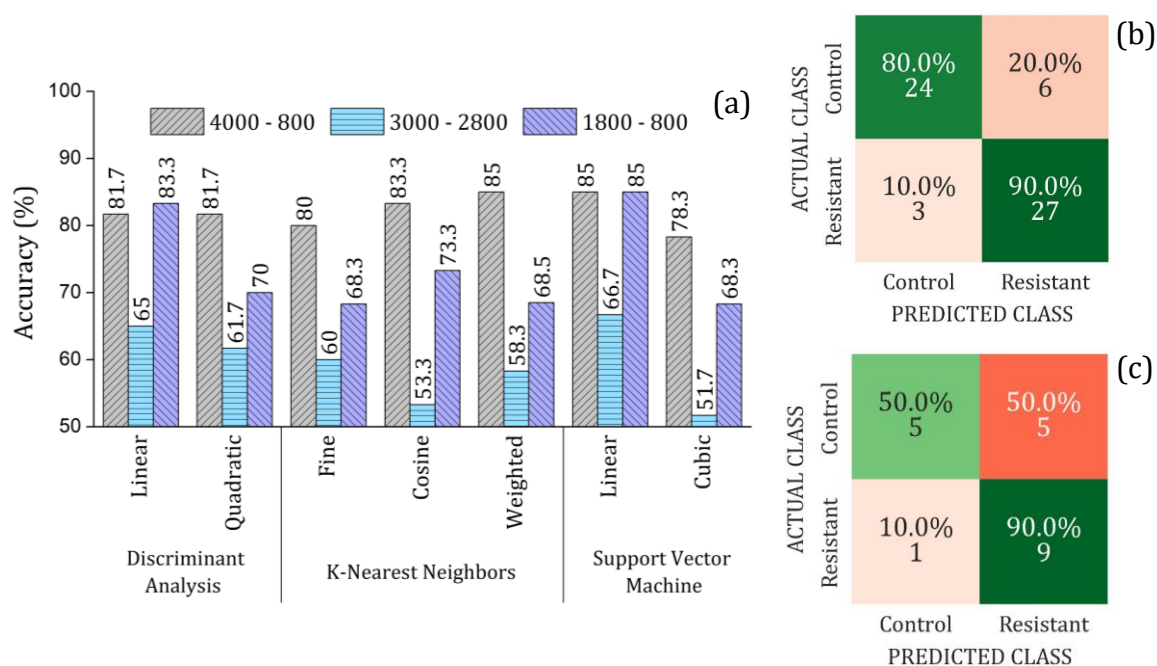
1405

1406 **Figure 5:** (a) score plot clustering improvement due to PCs selection by RFE., and (b) loading
 1407 for PC4, PC17, and PC28 from *E. coli* ATCC® 25922 (blue square), and (b) multidrug-resistant
 1408 (MR) *E. coli* (red circles) isolates.

1409

1410 The RFE-PCs were submitted to machine learning algorithms (DA, SVM, and
 1411 KNN) to build a prediction model for sample classification, the overall accuracy achieved
 1412 for each model by using different functions for classification in the LOOCV test is
 1413 summarized in Figure 6 (a). The maximum overall accuracy achieved by the predicting
 1414 models was 85% for Linear SVM by using 9 PCs – responsible for 8.59% of data variance
 1415 – within the 1800 to 800 cm^{-1} range, and 5 PCs in the 4000 to 800 cm^{-1} range. The
 1416 Weighted KNN also achieved an overall accuracy of 85% in the 4000 to 800 cm^{-1} range
 1417 with 5 PCs, which was responsible for 20.26% of data variance.

1418



1419 **Figure 6:** (a) Overall accuracy obtained in the LOOCV tests for Discriminant analysis (DA), K-
 1420 nearest neighbor (KNN), and Support vector machine (SVM) algorithms. Different ranges were

1421 analyzed with respective RFE-PCs: (i) 4000 to 800 cm^{-1} (gray bar color with right inclined line
1422 pattern) using 5 PCs; (ii) 3000 to 2800 cm^{-1} (light blue color with horizontal line pattern) by using
1423 5 PCs; (iii) 1800 to 800 cm^{-1} (purple color with left inclined line pattern) by using 9 PCs. The
1424 highest overall accuracy was 85% for Linear SVM at (4000 to 800 cm^{-1}) and (1800 to 800 cm^{-1})
1425 ranges, and Weighted KNN at (4000 to 800 cm^{-1}) range. Confusion matrix for the performance of
1426 Linear SVM algorithm in 1800 to 800 cm^{-1} range by using 9 PCs (b) LOOCV test with 70% data
1427 set, and (c) external validation test with 30% data set. The input PCA data from average FTIR-
1428 SNV spectra of *E. coli* ATCC® 25922, and multidrug-resistant (MR) *E. coli* isolates.
1429

1430 Besides both models achieving the same overall accuracy of 85% in the 4000 to
1431 800 cm^{-1} range, the W-KNN and L-SVM showed a much lower generalizability of the
1432 model in the external validation test, when compared to L-SVM in the 1800 to 800 cm^{-1}
1433 range. The external validation overall accuracy for L-SVM was 55%, and only 50% for
1434 W-KNN (see Supp. Mat.). However, the L-SVM with 85% of overall accuracy in the
1435 1800 to 800 cm^{-1} range, achieved 90% and 80% of sensitivity and specificity,
1436 respectively, Figure 6(b). Then, in the external validation test, Figure 6(c), the overall
1437 accuracy decreased to 70%, and achieved 90% and 50% of sensitivity and specificity,
1438 respectively. As can be observed in Figure 6(c), the prediction model failed to identify *E.*
1439 *coli* ATCC samples, which is less problematic for a trial method considering the need for
1440 large-scale tests to a fast decision in livestock management.

1441 The current study presented an easy and suitable methodology for the
1442 identification of MR-*E. coli* bacteria by FTIR spectroscopy and machine learning
1443 algorithms but must keep in mind that the methodology and results can be still improved,
1444 so the possibility for new studies is open. Here, we discussed the FTIR limitations to
1445 evidenced spectral changes due role played by ESBL enzyme to create MR bacteria, so
1446 new photonic techniques must be explored for better data acquisition. In our study, we
1447 explore a promising route to build a prediction model with good performance in the
1448 external validation test, but we can still search for the best hyperparameters for the
1449 algorithm or even explore new pre-processing data methods and algorithms to build the
1450 prediction model and improve its generalization capacity in the external validation test.

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CONCLUSIONS

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1454 The FTIR spectra, in the 1800 to 800 cm^{-1} range, of multi-resistant diarrhegenic
1455 *Escherichia coli*, isolates obtained from calves' feces showed great potential for
1456 microorganism identification. The usual principal component analysis was not able to
1457 provide a promising clustering formation for future sample classification. Then, we

1458 applied the Feature Selection Recursive Feature Elimination (RFE) algorithm, from
 1459 which the most important PCs were chosen before being used in machine learning
 1460 algorithms. The score plot of PC4 x PC17 x PC28 clearly demonstrated the improvement
 1461 of clustering tendency and group separation, here 9 PCs were selected by RFE and used
 1462 to build a prediction model by using Linear SVM. The prediction model showed an 85%
 1463 overall accuracy in the LOOCV test, and achieved 70% overall accuracy, with 90%
 1464 sensitivity and 50% specificity in the external validation test.

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1467

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