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#### Antibiotic-Resistance Pathogenic and Genes of Pasteurella multocida Isolated from Goats in the Mekong Delta, Vietnam

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### ABSTRACT

Pasteurella multocida (P. multocida) is one of the predominant pathogens that mostly cause respiratory diseases in domestic animals, such as goats. To determine P. multocida serotypes and the prevalence of pathogenic and antibiotic-resistance genes the PCR method was used. A total of 143 isolated P. multocida strains were collected from 289 healthy hybrid Boer-Saanen goats' nasal samples in the Mekong Delta, Vietnam, from March to June 2023. A total of 143 P. multocida strains, serotype B accounted for the highest proportion (51.05%), followed by serotype A (14.69%), and the lowest was serotype E (0.70%) while (39.86%) of strains could not be determined serotypes. Among the six virulence genes surveyed, the sodA gene (56.64%) had the highest presence, while the ompH gene (4.20%) had the lowest presence. Pathogenic genes were present mainly in serotypes A and B; tbpA was frequently detected in serotype A (66.67%), and sodA was commonly detected in serotype B (56.16%). There were 14 virulence gene combinations in 59/109 (54.13%) serotyped P. multocida strains, and the pattern of sodA + toxA + *tbpA* was prevalent at the highest rate (12.84%). Moreover, among the eight investigated antibiotic resistance genes, the sull gene had the highest presence rate (74.13%), compared to the tetA gene with the lowest presence rate (13.29%). Gene sullI was mainly detected on strains belonging to serotypes A (80.95%), B (83.56%), and F (77.78%). A total of (77.98%) of serotyped P. multocida strains indicated multi-harbor from two to six antibioticresistance genes, and the most common pattern was aadB + sulli (10.09%). The prevalence of five pathogenic P. multocida serotypes harboring diverse antibiotic-resistance genes isolated from nasal samples could be a critical issue in treating and preventing the respiratory diseases caused by *P. multocida* in goats in the Mekong Delta.

Keywords: Antibiotic resistance, Goat, Pasterella multocida, Pathogenicity, Mekong Delta

### **INTRODUCTION**

Respiratory disease caused by Pasteurella multocida is among the most common infections in ruminant animals. Goats and other small ruminants are at a moderate risk of contracting this pathogen due to exposure to physical stress or uncomfortable environmental conditions (Mohamed and Abdelsalam, 2008). P. multocida is more frequently associated with the outbreak of acute pneumonia and death of goats in all age groups than Mannheimia haemolytica is in previous reports (Falade, 2002). Respiratory diseases in goats cause economic losses arising from mortality and morbidity. The respiratory disease mortality rate caused by P. multocida is 10% or more (Smith and Sherman, 2009). Girma et al. (2023) reported 27,563 and 37,522 cases of sheep and goats with pneumonia, in Southern Ethiopia from 2016 to 2021. There were a few reports on P. multocida isolated from small ruminants in Vietnam. In a previous report, P. multocida was detected in healthy goats raised on medium-scale farms in Can Tho City, the Mekong Delta at 49.48% (Nguyen et al., 2024). These observations implied the significance of pneumonic pasteurellosis in small ruminants.

Pasteurella multocida is classified into five serotypes, A, B, D, E, and F, according to capsular antigen, and 16 serotypes according to lipopolysaccharide antigen. The capsular antigen is considered an essential form of virulence of P. multocida, which allows P. multocida to avoid innate host defense systems (Boyce et al., 2000). Each serotype has different circulation in different animals. Serotypes A and D are two serotypes that commonly appear in cases of pneumonia and pasteurellosis in goats (Rawat et al., 2009; Tabatabaei and Abdolahi, 2023). Besides, additional factors essential for the proliferation and maturation of P. multocida have been discovered. These encompass P. multocida toxin (PMT), fimbriae, adhesins, the capacity to metabolize sialic acid, outer membrane proteins, and hyaluronidases (Harper et al., 2006). Previous studies reported that the prevalence of some virulence-associated genes, including colonization

factors (*ptfA*, *fimA*, *hsf2*), iron acquisition factors (*exbB*, *exbD*, *tonB*, *Fur*), superoxide dismutase (*sodA*, *sodC*), and outer membrane proteins (*ompA*, *ompH*, *oma87*, *plpB*), were frequently detected in the pig origin *P. multocida* isolates (May et al., 2001; Peng et al., 2016; 2018). Mombeni et al. (2021) surveyed virulence genes on *P. multocida* strains isolated from goats in Iran and showed that the *sodA* gene had the highest presence rate (100.00%), followed by *toxA*, *nanH*, and *ompH*, with the same ratio of 61.90%.

On the other hand, antibiotic resistance is becoming a global concern as more and more multidrug-resistant bacteria appear. The cause is the overuse of antibiotics when treating animal diseases or using antibiotics as growth stimulants in animals (Martin et al., 2015). Kandimalla et al. (2022) revealed that *P. multocida* strains isolated from sheep and goats were susceptible to ceftriaxone, cefoperazone/sulbactum, ceftiofur, cloxacillin, ciprofloxacin, enrofloxacin, levofloxacin and tetracycline (100.00%) but were resistant to erythromycin (41.67%), gentamycin (66.67%). Nguyen et al. (2023) reported that *P. multocida* isolated from sheep in Central Vietnam was resistant to tetracycline (51.22%), ampicillin (53.66%), and erythromycin (65.85%). The frequent impact of antibiotics leads to mutations and the formation of genetic drug-resistance factors in bacteria. Surveying the presence of genes encoding drug-resistance factors in *P. multocida* gives a more general view of antibiotic resistance issues.

Diseases of domestic animals are still a massive issue in the Mekong Delta, Vietnam. There have been a few reports on the prevalence of pathogenic and antibiotic-resistance genes of *P. multocida* isolated from small ruminants; however, there were no reports in goats. This causes challenges in controlling and preventing diseases in goats. Therefore, this study aims to clarify the characteristic prevalence of *P. multocida* serotypes originating from goats and their pathogenicity and antibiotic resistance genes in the Mekong Delta, Vietnam.

# MATERIALS AND METHODS

## **Ethical approval**

The procedure for collecting fluid swab samples on goats was performed according to NAHMS (National Animal Health Monitoring System) guidelines (USDA, 2022), and *P. multocida* strains were isolated according to Vietnamese National Standard TCVN 8400-14:2011. In the author's previous study, samples were collected according to the guidelines outlined in the Helsinki Declaration and the animal welfare and safety procedures of Can Tho University, Vietnam.

# Identification of Pasteurella multocida serotypes

A total of 143 *P. multocida* strains were previously isolated from 289 nasal swabs of healthy hybrid Boer-Saanen meat and dairy goats at all ages in small-scale farms in the Mekong Delta, Vietnam, in 2023. *Pasteurella* spp. were isolated from nasal fluid samples in goats and isolated on blood agar with 5% of sheep blood according to Vietnamese National Standard TCVN 8400-14:2011. Then, *P. multocida* strains were identified using the PCR method to detect gene *Pm1231*. Those identified strains were kept in the Veterinary Food Hygiene Laboratory, Faculty of Veterinary Medicine, College of Agriculture, Can Tho University to conduct this study.

*Pasteurella multocida* strains were subcultured on trypticase soy agar (TSA, Merck. Germany) at 37°C for 24 h to extract DNA. The DNA of *P. multocida* strains was extracted using the heat-shock method and stored at -20°C for the following experiments (Ahmed and Dablool, 2017).

*Pasteurella multocida* serotypes were determined by performing PCR reactions with primers of genes encoding for each serotype, including *hyaD-hyaC* (serotype A), *bcbD* (serotype B), *dcbF* (serotype D), *ecbJ* (serotype E), and *fcbD* (serotype F). The PCR conditions and primer sequences followed the description of Townsend et al. (2001).

The PCR mixture for one reaction included Mastermix 2X (BIO25042, Bioline, Meridian Bioscience, USA, 12.5  $\mu$ l), forward primer (0.5  $\mu$ l), reverse primer (0.5  $\mu$ L), distilled water (9.5  $\mu$ L), and DNA template (2.0  $\mu$ L).

### **Determination of pathogenic genes**

This study determined six pathogenic genes encoded for capsular and lipopolysaccharide antigens, including *sodA*, *toxA*, *tbpA*, *ptfA*, *pfhA*, and *ompH*. The PCR conditions and primer sequences followed the description of Doughty et al. (2000) and Ewers et al. (2006). Among genes, the annealing temperature was 55°C for *sodA*, *toxA*, *tbpA*, *ptfA*, and *ompH*, while it was 58°C for *pfhA*.

The *P. multocida* strains, previously isolated from cattle in the Mekong Delta, were used as a control. The MyTaq Mix 2X (BIO25042, Bioline, Meridian Bioscience, USA) was used in those experiments as described in the above method.

#### **Determination of antibiotic-resistance genes**

The PCR assay was used to detect eight antibiotic-resistance genes representative of beta-lactam (*blaROB-1*, *blaOXA*), aminoglycoside (*aadB*, *strA*), tetracycline (*tetA*, *tetB*), sulfonamide (*sulII*), and macrolide (*mph*). The PCR conditions and primer sequences followed the description of Randall et al. (2004), Saenz et al. (2004), Carattoli et al. (2005), Momtaz et al. (2012), Klima et al. (2014), and Abo-Almagd et al. (2023). The *P. multocida* strains, previously isolated from cattle in the Mekong Delta, were used as a positive control. The PCR procedure was conducted as described in the above experiment of detection of *P. multocida* serotypes.

## Statistical analysis

The difference in the prevalence of pathogenic and antibiotic-resistance genes in *P. multocida* isolated from goats was statistically analyzed at a significance rate of 95% using the Pearson Chi-square test in the Minitab 17.0 software.

### RESULTS

Of 143 *P. multocida* strains, serotype B was the most predominant serotype (51.05%), followed by serotype A (14.69%), serotype F (6.29%), serotype D (3.50%), and serotype E (0.70%, p < 0.05). There were 39.86% of *P. multocida* strains, which could not determine serotypes in this study (Table 1).

Of the six pathogenic genes examined (Table 2), *sodA* was present at the highest rate (56.64%), followed by *toxA* (45.45%), *tbpA* (30.77%), *ptfA* (10.49%), *pfhA* (4,90%), and *ompH* (4.20%, p < 0.05). Most pathogenic genes were found in *P. multocida* strains belonging to serotypes A and B.

Of 109 serotyped *P. multocida* strains, gene *tbpA* was frequently detected in serotype A (66.67%), and *sodA* was commonly in serotype B (56.16%); however, only one strain belonging to serotype E harbored gene *sodA* (Table 3).

There were 59/109 (54.13%) serotyped *P. multocida* strains that harbored a combination of two to three pathogenic genes (Table 4). Among gene combinations, the *sodA* + *toxA* + *tbpA* pattern was the most common (12.84%), followed by *sodA* + *toxA* (11.93%).

Of the eight antibiotic-resistance genes examined (Table 5), gene *sulII* was detected at the highest rate (74.13%), followed by *aadB* (42.66%), *strA* (33.57%), *mph* (28.67%), *blaROB-1* (21.68%), *bloOXA* (17.48%), *tetB* (16.08%), and *tetA* (13.29%).

Moreover, *sulII* and *aadB* genes were also recorded at the highest rate in serotyped *P. multocida* strains belonging to all serotypes, followed by *mph* gene (Table 6). Besides, 85/109 (77.98%) serotyped *P. multocida* strains harbored combinations of two to six antibiotic-resistance genes (Table 7). Among gene patterns, the pattern of aadB + sulII was the most common (10.09%).

Serotype	Encoded gene	No. of positive strains	Percentage
А	hyaD-hyaC	21	14.69
В	bcbD	73	51.05
D	dcbF	5	3.50
E	ecbJ	1	0.70
F	fcbD	9	6.29
Untyped		57	39.86

Table 1. Distribution of Pasteurella multocida serotypes isolated from hybrid Boer-Saanen goats in the Mekong Delta,
Vietnam from March to June 2023 (n=143)

Untyped: P. multocida strains were not determined in the serotypes (A, B, D, E, and F) using the specific primers in this study. No: Number

**Table 2.** Prevalence of pathogenic genes in *Pasteurella multocida* strains isolated from hybrid Boer-Saanen goats in the Mekong Delta, Vietnam from March to June 2023 (n=143)

Pathogenic gene	No. of positive strains	Percentage
sodA	81	56.64
toxA	65	45.45
tbpA	44	30.77
ptfA	15	10.49
pfhA	7	4.90
ompH	6	4.20

No: Number

	No. of positive strains (%)	Serotype A	Serotype	Serotype D	Serotype E	Serotype F	Total
Gene		( <b>n</b> = 21)	<b>B</b> $(n = 73)$	( <b>n</b> = 5)	( <b>n</b> = 1)	( <b>n</b> = 9)	(n = 109)
sodA		10 (47.62)	41 (56.16)	2 (40.00)	1 (100.00)	5 (55.56)	59 (54.13)
toxA		9 (42.86)	36 (49.32)	2 (40.00)	0 (0.00)	4 (44.44)	51 (46.79)
tbpA		14 (66.67)	29 (39.73)	2 (40.00)	0 (0.00)	4 (44.44)	49 (44.95)
ptfA		1 (4.76)	9 (12.33)	1 (20.00)	0 (0.00)	1 (11.11)	12 (11.01)
pfhA		0 (0.00)	3 (4.11)	0 (0.00)	0 (0.00)	0 (0.00)	3 (2.75)
ompH		1 (4.76)	4 (5.48)	0 (0.00)	0 (0.00)	0 (0.00)	5 (4.59)

**Table 3.** Distribution of pathogenic genes in serotyped *Pasteurella multocida* strains isolated from hybrid Boer-Saanen goats in the Mekong Delta, Vietnam from March to June 2023

No: Number

**Table 4.** Pathogenic gene patterns of serotyped *Pasteurella multocida* strains from hybrid Boer-Saanen goats in the Mekong Delta, Vietnam from March to June 2023 (n=109)

No. of genes	Pattern	No. of strains	Percentage
	sodA + toxA	13	11.93
	sodA + tbpA	5	4.59
	sodA + ptfA	4	3.67
	sodA + pfhA	1	0.92
2	toxA + pfhA	1	0.92
	toxA + tbpA	10	9.17
	toxA + ompH	1	0.92
	ptfA + tbpA	1	0.92
	ompH + tbpA	1	0.92
	sodA + toxA + tbpA	14	12.84
	sodA + toxA + ptfA	1	0.92
3	sodA + ptfA + tbpA	3	2.75
	sodA + toxA + ompH	3	2.75
	toxA + ptfA + tbpA	1	0.92
Total		59	54.13

No: Number

**Table 5.** Prevalence of antibiotic-resistance genes in *Pasteurella multocida* strains isolated from hybrid Boer-Saanen goats in the Mekong Delta, Vietnam from March to June 2023 (n=143).

Antibiotic group	Gene	No. of positive strains	Percentage
Beta-lactam	blaROB-1	31	21.68
Deta-factalli	blaOXA	25	17.48
Aminoglycoside	aadB	61	42.66
Ammogrycoside	strA	48	33.57
Totrogyaling	tetA	19	13.29
Tetracycline	tetB	23	16.08
Sulfonamide	sulII	106	74.13
Macrolide	mph	41	28.67

No: Number

**Table 6.** Distribution of antibiotic-resistance genes in serotyped *Pasteurella multocida* strains from hybrid Boer-Saanengoats in the Mekong Delta, Vietnam from March to June 2023

Gene	No. of positive strains (%)	Serotype A (n = 21)	Serotype B (n = 73)	Serotype D (n = 5)	Serotype E (n = 1)	Serotype F (n = 9)	Total (n = 109)
blaROB-1		5 (23.81)	16 (21.92)	1 (20.00)	0 (0.00)	0 (0.00)	22 (20.18)
blaOXA		2 (9.52)	13 (17.81)	1 (20.00)	1 (100.00)	1 (11.11)	18 (16.51)
aadB		12 (57.14)	30 (41.10)	1 (20.00)	1 (100.00)	6 (66.67)	50 (45.87)
strA		8 (38.10)	27 (36.99)	2 (40.00)	1 (100.00)	1 (11.11)	39 (35.78)
tetA		3 (14.29)	11 (15.07)	0 (0.00)	0 (0.00)	1 (11.11)	15 (13.76)
tetB		6 (28.57)	14 (19.18)	2 (40.00)	0 (0.00)	0 (0.00)	22 (20.18)
sulII		17 (80.95)	61 (83.56)	1 (20.00)	0 (0.00)	7 (77.78)	86 (78.90)
mph		9 (42.86)	26 (35.62)	2 (40.00)	0 (0.00)	5 (55.56)	42 (38.53)

No: Number

No. of genes	Gene patterns	No. of strains	Percentage
	blaROB-1 + sulII	1	0.92
	blaROB-1 + strA	1	0.92
	aadB+ sulII	11	10.09
	aadB+ mph	3	2.75
2	strA + sulII	2	1.83
	strA + tetB	1	0.92
	strA + mph	1	0.92
	tetA + sulII	2	1.83
	sulII + mph	1	0.92
	blaROB-1 + tetA + sulII	1	0.92
	blaROB-1 + strA + sulII	3	2.75
	blaROB-1 + strA + tetB	1	0.92
	blaROB-1 + sulII + tetB	1	0.92
	blaOXA + aadB+ sulII	4	3.67
	blaOXA + strA + sulII	1	0.92
	blaOXA + aadB + mph	3	2.75
	blaOXA + sulII + tetB	2	1.83
3	blaOXA + aadB + strA	1	0.92
	aadB+ tetA + sulII	1	0.92
	aadB+ sulII + mph	8	7.34
	strA + tetB + mph	1	0.92
	strA + sulII + mph	1	0.92
	strA + tetA + sulII	1	0.92
	strA + sulII + tetB	3	2.75
	sulII + tetB + mph	2	1.83
	blaROB-1 + strA + tetB + mph	1	0.92
	blaROB-1 + blaOXA + sulII + mph	1	0.92
	blaROB-1 + aadB+ strA + sulII	2	1.83
	blaROB-1 + strA + tetA + sullI	3	2.75
	blaROB-1 + blaOXA + tetA + sulII	1	0.92
4	blaOXA + aadB + sullI + mph	1	0.92
•	aadB+ strA + sulII + mph	4	3.67
	aadB+ sulII + tetB + mph	2	1.83
	aadB+ strA + sulII + tetB	1	0.92
	aadB+ tetA + sulII + mph	2	1.83
	strA + tetA + sulII + mph	1	0.92
	blaROB-1 + blaOXA + aadB + sulII + mph	1	0.92
	blaROB-1 + strA + sulII + tetB + mph	3	2.75
5	blaROB-1 + blaOXA + strA + tetA + sullI	1	0.92
-	aadB+ strA + tetA + sullI + tetB	1	0.92
	aadB + strA + sullI + tetB + mph	2	1.83
6	blaROB-1 + aadB+ strA + sulII + tetB + mph	- 1	0.92
Total	····· ··· ····· ······················	85	77.98

**Table 7.** Multiple antibiotic-resistance gene patterns of serotyped *Pasteurella multocida* strains (on the strains harbored from two antibiotic-resistance genes) from hybrid Boer-Saanen goats in the Mekong Delta, Vietnam from March to June 2023 (n=109).

No: Number

# DISCUSSION

In this study, *P. multocida* serotypes A and B were more prevalent than the remaining serotypes. Shayegh et al. (2009) and Mombeni et al. (2021) indicated that *P. multocida* strains isolated from goats in Iran belonged mainly to two serotypes, A and D. Serotypes B and E are two serogroups reported to commonly cause hemorrhagic infections in ruminant carriers. Serogroup B was often found in the nasopharyngeal fluid of livestock in Southeast Asia, while serogroup E was more common in Africa (Markey et al., 2013). Aski and Tabatabaei (2016) recorded the prevalence of three serotypes, A, B, and D, in *P. multocida* strains isolated from healthy and clinically infected goats. *P. multocida* serotype B, specifically serotype B:2, was a common serotype detected in cases of infected cattle. However, 39.86% of *P. multocida* strains were not determined serotypes in this study. The reason could be due to the specific primers or the characteristic structure of capsular antigens in those *P. multocida* strains. Further study should be done to clarify and confirm the prevalence of diverse serotypes of *P. multocida* strains isolated from goats using other primers or serotyping methods.

Furthermore, three genes, *sodA*, *toxA*, and *tbpA*, were more commonly detected in *P. multocida* strains. The *sodA* gene was frequently detected in *P. multocida* isolated from poultry, pigs, and rabbits in previous studies (Furian et al., 2015; Li et al., 2018; Mahrous et al., 2022). Therefore, it is difficult to determine whether *sodA* is a characteristic gene in *P. multocida* strains isolated from goats or was related to serotypes A and B. According to Rimac et al. (2017), the *tbpA* gene is closely related to *P. multocida* strains isolated from ruminants with pneumonia and sepsis. Katsuda et al. (2013) also indicated a close relationship between serotype A strains and the *tbpA* gene. Research by Nguyen et al. (2023) showed that the *tbpA* gene was detected in serotypes A, B, and D strains in clinical pneumonic pasteurellosis sheep in central Vietnam at 48.78%, 7.32%, and 21.95%, respectively. The above results suggested that there might be a relationship between serotypes A and B and the *tbpA* gene, especially in diseased animals. On the other hand, Pullinger et al. (2003) reported that gene *toxA* can be transferred horizontally, and the *toxA* gene was determined to be concerning serotypes A and D strains (Furian et al., 2015). Cid et al. (2019) reported that the *toxA* gene encodes the *P. multocida* toxin (PMT), which is a dermonecrotic protein in the virulence factor of capsular type D.

*Pasturela multocida* causes progressive atrophic rhinitis in pigs and is significantly found in ovine pneumonia isolates in Spain. Besides, the detection of the *toxA* gene could serve as a reliable indicator of the toxigenic fitness of *P*. *multocida* (Cid et al., 2019). Thus, the high presence of the *toxA* gene in *P. multocida* strains isolated from goats showed that those *P. multocida* strains harboring PMT toxin could cause pasteurellosis in goats in the Mekong Delta, Vietnam.

Of serotyped strains, three genotypes with high prevalence rates included sodA + toxA + tbpA, sodA + toxA, and toxA + tbpA, with rates of 12.84%, 11.93%, and 9.17% respectively. The toxA was present in most of the common patterns. Pullinger et al. (2003) showed that the toxA gene was encoded in the genome of a latent bacteriophage to allow the toxA gene to circulate easily among *P. multocida* strains of many different serotypes. Bernal et al. (2023) recorded that all *P. multocida* strains isolated from cows with respiratory disease carried more than two virulence genes. The appearance of serotypes carrying diverse virulent genes makes it difficult to prevent and control diseases in goats in the Mekong Delta.

Among antibiotic-resistance genes, gene *sulII* had the highest presence rate (74.13%). Gene *sulII* was not generally considered part of a separate genetic element; it was found on large conjugative plasmids and was associated with resistance to other antibiotics (Bean et al., 2009; Wu et al., 2010). Antibiotic-resistant genes could be silent resistance genes that are not frequently exhibited or exhibited at low levels, even when exposed to antibiotics. They are nonessential residues and do not play an essential role in the bacterial life cycle (Stasiak et al., 2021). However, antibiotic-resistance genes *blaROB-1*, *blaOXA*, *tet A*, and *tetB* had a relatively low presence rate in *P. multocida* strains in this study. In another report, Babetsa et al. (2012) indicated that *P. multocida* strains isolated from bovine, ovine, caprine, and swine pneumonic lungs were resistant to tetracycline in Greece. Among those tetracycline-resistance *P. multocida* strains, 72.22% of strains carried the *tetH* gene, and 22.22% of strains carried the *tetA* and *tetM* were not found. The current study results showed a difference in the prevalence of antibiotic-resistance genes in *P. multocida* isolated from goats in the Mekong Delta. Thus, further research should be conducted to clarify the diverse prevalences of antibiotic-resistance genes in *P. multocida* isolated from goats in this region.

Of serotyped *P. multocida* strains, the *sull1* had the highest presence rate in serotypes A, B, and F. The gene *blaROB-1* and *tetB* genes were not found in serotype F strains. Most antibiotic-resistance genes are present in the bacterial genome through random gene transfer using mobile genetic elements (Bennett, 2008). Besides, genes *tetA* and *tetB* are two representatives of genetic factors for tetracycline resistance through the formation of ABC (ATP-binding cassette superfamily) transporters (Reynolds et al., 2016). Rendueles et al. (2018) revealed that capsular antigens were related to antibiotic resistance genes, especially genes related to antibiotic efflux pumps. Previous studies mainly focused on the relationship between serotypes and virulent genes; however, a few studies have shown the relationship between

serotypes and antibiotic-resistance genes in *P. multocida* (Harper et al., 2006; Katsuda et al., 2013; Cid et al., 2019; Nguyen et al., 2023). Thus, the current study results seemed to be the first report on the prevalence of antibiotic-resistance genes in *P. multocida* serotypes isolated from healthy goats in the Mekong Delta, Vietnam.

Moreover, 77.98% *P. multocida* strains isolated from goats in the Mekong Delta showed several antibiotic-resistance gene patterns. Among 42 antibiotic-resistance gene combinations of *P. multocida* strains, the most common patterns across serotypes were aadB + sulII (10.09%) and aadB + sulII + mph (7.34%) in this study. The aadB gene is usually associated with type 1 integron or plasmids, and the *sulII* gene is mainly located on small non-conjugative plasmids or large multiresistant plasmids that can be horizontally transferred (Antunes et al., 2005; Naderi et al., 2023). The common prevalence of *aadB* and *sulII* genes in several antibiotic-resistance gene patterns in serotypes showed that *aadB* and *sulII* might be a typical resistance gene cluster in *P. multocida* isolated from goats in the Mekong Delta, Vietnam. In addition, the diversity of antibiotic-resistance gene patterns in those *P. multocida* strains exhibited the high ability of multidrug resistance of those strains. It could cause difficulty in treating and preventing the respiratory diseases caused by *P. multocida* in goats in this region.

# CONCLUSION

This study showed that five pathogenic *P. multocida* serotypes were detected in healthy hybrid Boer-Saanen goats in the Mekong Delta, Vietnam. The results indicated that *P. multocida* serotypes A and B were the dominant serotypes in goats in the Mekong Delta. Moreover, those *P. multocida* strains harbored diverse pathogenic genes and antibiotic-resistance genes with serval gene patterns. The pathogenic gene (*sodA*) and the antibiotic-resistance gene (*aadB*) were the most detected in all *P. multocida* serotypes in this study. In addition, the pathogenic gene pattern of *sodA* + *toxA* + *tbpA* and the antibiotic-resistance gene pattern of *aadB* + *sulII* were frequently prevalent in those *P. multocida* strains. It revealed that *P. multocida* strains isolated from goats were potential pathogens causing severe diseases in goats in the Mekong Delta. Therefore, the control of pathogenic and antibiotic-resistant *P. multocida* is essential to prevent and treat pasteurellosis in goats.

## DECLARATION

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### **Conflicts of Interests**

The authors declare that we do not have any conflicts of interest.

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#### Authors' contributions

Thuan K. Nguyen, Thuong T. Nguyen, and Trung T. Truong conceptualized, designed, and supervised the research. Thuan K. Nguyen and Thuong T. Nguyen critically reviewed the study. Thuan K. Nguyen, Thuong T. Nguyen, Chi T.H. Nguyen, Vy L.P. Nguyen, and Trung T. Truong collected samples and conducted experiments. Chi T.H. Nguyen and Vy L.P. Nguyen analyzed and interpreted the data generated. All authors revised and approved the submitted manuscript.

## **Competing interests**

The authors declare that they have no conflict of interest.

## Availability of data and materials

The authors of this article confirm that all data supporting the findings of this research are available upon reasonable request.

# **Ethical considerations**

The authors considered farmers' ethical concerns and consent before conducting the study. This article was written originally without any copy from data of published articles and books.

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