Coexistence of Virulence and Antibiotic Resistance Genes in Pasteurella multocida Isolated from Diseased Rabbits

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Abstract

Pasteurella multocida (P. multocida) is considered a predominant pathogenic bacterial agent causing respiratory manifestations (snuffles) in rabbits with considerable economic losses and unfavorable prognosis in Egypt. A few recent P. multocida strains exhibit resistance to the most commonly used antibiotics in the veterinary field. Therefore, the present study was carried out to investigate the prevalence of both virulence and antibiotic resistance genes among P. multocida isolated from diseased rabbits in Sharkia Governorate, Egypt. Only 10 out of 100 tested rabbits' nasal swabs were finally confirmed positive for P. multocida of serogroup A by polymerase chain reaction (PCR). Antibiotic susceptibility testing of the recovered isolates revealed that they were all multidrug resistant (MDR) with a predominance of resistance to amoxicillin, ampicillin, amoxicillin/clavulanic acid, neomycin and tetracycline (100% each), followed by kanamycin and streptomycin (90% each). All the recovered isolates were further subjected to PCR screening of some common virulence and antibiotic resistance genes of interest. With the exception of toxA gene, the other virulence associated genes (ptfA, Omp87 and nanB) were found among all the examined isolates. Totally, all MDR P. multocida isolates contained at least one antibiotic resistance gene with aphA1 being the most prevalent (100%), followed by ermX gene (40%). Antibiotic resistance genotyping demonstrated the presence of multiple antibiotic resistance genes among majority of the isolates (40%) with only one isolate harboring 4 genes encoding identical resistance phenotypes. Evidently, all MDR P. multocida isolates possessed at least 3 virulence genes accompanied by the attendance of antibiotic resistance genes. These findings evidenced that rabbits are potential sources of pathogenic P. multocida strains harboring virulence and antibiotic resistance genes. Therefore, it is evident that there is an urgent need for the judicious use of antibiotics in rabbits’ treatment systems to successfully mitigate the propagation of drug resistance across P. multocida species.

Keywords: P. multocida, Rabbits, Multi-drug resistant, Antibiotic Resistance Genes, Virulence Factors.

Introduction

Rabbit pasteurellosis is a highly contagious disease of domestic rabbits caused by Pasteurella multocida (P. multocida) [1]. The disease is characterized by various clinical manifestations including rhinitis (snuffles) with purulent nasal discharge, abscesses, otitis media, pneumonia, pyometra, orchitis and septicemia [2,3]. The stress and immunodeficiency of hosts play a major role in the development of pasteurellosis by auto-infections [4].

P. multocida is a Gram negative commensal pathogen in a wide host range including mammals, birds and humans [5]. Serologically, it is classified into five capsular serogroups (A, B, D, E and F) and 16 lipopolysaccharide (LPS) somatic serotypes (1-16) [6]. Pasteurellosis in rabbits is mainly caused by the capsular type A and to a lesser extent type D P. multocida strains [7]. Capsular type F strains have been also recorded in rabbits [8].

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The polysaccharide capsule and LPS are of the major important virulence determinants contributed in the pathogenesis of *P. multocida* [9]. However, many other putative virulence factors are related to pathogenicity including adherence and colonization factors, fimbriae, iron regulating and acquisition proteins, exotoxins and extracellular enzymes [10].

Using the antimicrobial therapy for controlling rabbit pasteurellosis is failing to bring an end to many infections due to the emergence of multi-drug resistant (MDR) strains with a great potential for transfer of this resistance from animals to human. As a result, the antibiotic resistance phenomenon among pathogenic bacteria has become a growing problem in veterinary and human medicininal fields [11]. This leads to increased treatment costs, prolonged illness and sometimes death with adverse effects on the economy of Egypt [12]. Therefore, pre-testing of antibiotics susceptibility is essential to select the effective drugs to be used by the veterinarians in the field [13].

Data on the antibiotic resistance and virulence properties of individual *P. multocida* strains would help to implement appropriate preventive strategies. Assessing the distribution of resistance genes represents a more detailed and potentially beneficial additional tool for better understanding antimicrobial resistance, particularly in Egypt, where existing data is quite limited. Additionally, the description of co-occurrence between virulence and resistance in *P. multocida* from rabbits has not been reported in Egypt. These knowledge gaps encourage the need for researches in these serious issues.

Therefore, the present investigation aimed to assess the occurrence of *P. multocida* in rabbit farms from different localities at Sharkia Governorate as well as their possession of four important virulence-associated genes (*ptfA, Omp87*, *nanB* and *toxA*) putatively involved in the pathogenesis of *P. multocida*. Moreover, we documented the phenotypic and genotypic characteristics of antibiotic resistance among the recovered *P. multocida* isolates to clarify the relationship between the phenotypic and genotypic resistance profiles. Finally, we attempted to associate the co-presence of virulence and antibiotic resistance which is becoming more profitable for the pathogenic bacteria.

**Materials and Methods**

**Clinical data and specimens collection**

The study was approved by the Committee of Animal Welfare and Research Ethics, Faculty of Veterinary Medicine, Zagazig University. It was performed on a total of 100 clinically diseased rabbits originated from five rabbitries in different localities at Sharkia Governorate. They were admitted to the clinic of Avian and Rabbit Medicine Department, Faculty of Veterinary Medicine, Zagazig University with respiratory manifestations (oculonasal discharge, sneezing and coughing) during the period from March 2017 to August 2018. Owing to the major role of stress in the development of pasteurellosis in rabbits, relevant information was obtained from the owners of the diseased rabbits before bacteriological examination. The five investigated rabbitries revealed the following stress factors; improper rabbit hut (a small or a glass cage), poor quality of feeding as carrots, overcrowding, improper or excessive handling, transportation, poor ventilation, extreme temperature, pregnancy, parturition, heavy lactation and bad management (excessive ammonia level). The external nares of the diseased rabbits were swabbed using sterile cotton-tipped swabs.

**Isolation and identification of Pasteurella multocida**

*P. multocida* was isolated and identified according to the conventional protocols stated previously [14]. Briefly, each swab was inoculated into brain heart infusion (BHI) broth (Oxoid, Hampshire, UK) and incubated aerobically at 37°C for 24 h. Subsequently, a loopfull from the enriched broth was streaked on each of blood agar media supplemented with 5% fresh sheep blood and MacConkey’s agar plates (Oxoid, Hampshire, UK). After overnight incubation at 37°C under aerobic conditions, all the isolates that did not grow on MacConkey’s agar and showed typical colonies of *Pasteurella* spp. on blood agar were further confirmed using a series of
biochemical tests including oxidase, catalase, indole and urease production according to the standard laboratory procedures [15]. Presumptive isolates were confirmed as *P. multocida* by PCR amplification of the species specific *kmi1* gene fragment [16]. The capsular types of the confirmed *P. multocida* isolates were then determined using the multiplex PCR assay [17].

**Antibiotic susceptibility testing**

All the recovered *P. multocida* isolates were examined for their antibiotic susceptibility patterns against 10 antibiotic agents of four different groups by the Kirby–Bauer disk diffusion method [18]. Overnight-grown cultures were spread onto Mueller Hinton agar plates to form a bacterial lawn; thereafter, antibiotic discs (Oxoid, Hampshire, UK) were evenly placed and then the plates were incubated at 37 °C for 24 h.

The tested antibiotic agents were commonly used in poultry industry including ampicillin (AM, 10 µg), amoxicillin (AX, 25 µg), amoxicillin/clavulanic acid (AMC, 20/10 µg), neomycin (N, 30 µg), gentamicin (CN, 10 µg), kanamycin (K, 30 µg), streptomycin (S, 10 µg), erythromycin (E, 15 µg), tetracycline (TE, 30 µg) and doxycycline (DO, 30 µg). The inhibition zones' diameters were measured and scored as sensitive (S), intermediate (I) and resistant (R) according to the European Committee on Antimicrobial Susceptibility Testing (EUCAST) clinical breakpoint table version 7.1 2017 [19]. Isolates showing resistance to at least 3 different antibiotic groups were categorized as MDR.

**Molecular detection of virulence and resistance associated genes**

The presence of four virulence genes encoding adhesins (*ptfA*), protectins (*Omp87*), sialidases (*nanB*) and dermonecrototoxin (*toxA*) were examined in all recovered *P. multocida* isolates by individual PCR protocols utilizing specific oligonucleotide primers [20,21].

For detecting the corresponding antibiotic resistance genes in all phenotypic resistant isolates, target genes conferring resistance to β-lactams (*blaROB-1*), aminoglycosides (*aphA1*), macrolide (*ermX*) and tetracyclines (*tetH*) were screened by PCR assay. The primer sequences and cycling conditions were as described previously [22]. All runs were repeated 3 times in parallel with the relevant positive and negative controls. The details of the oligonucleotide primers used to detect virulence and resistance genes are depicted in Table 1.
Table 1: Primers used for the detection of *P. multocida* serogroups and their virulence and resistance-associated genes

<table>
<thead>
<tr>
<th>Gene function</th>
<th>Target gene</th>
<th>Primer sequence (5′-3′)</th>
<th>Amplicon size (bp)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>P. multocida</em> species specific</td>
<td>KMT1</td>
<td>KMT1T7-ATCCGCTATTTACCCAGTG&lt;br&gt;KMT1SP6-ATCCGCTATTTACCCAGTG</td>
<td>460</td>
<td>[16]</td>
</tr>
<tr>
<td>A</td>
<td>hyaD-hyaC</td>
<td>F- TGCCAAAATCGCAATCGTAC&lt;br&gt;R- TTGCCATCATTTGCAG</td>
<td>1,044</td>
<td>[17]</td>
</tr>
<tr>
<td>B</td>
<td>bcbD</td>
<td>F- CATTTATCCAAGCTCCAC&lt;br&gt;R- GCCGGAGATTTCAATCC</td>
<td>760</td>
<td>[17]</td>
</tr>
<tr>
<td><em>P. multocida</em> serogroup</td>
<td>D</td>
<td>dcBF-TTACAAAAGGAAGACTAGGAGGCC&lt;br&gt;R- CATCTACCCACTCAACACATCAG</td>
<td>657</td>
<td>[17]</td>
</tr>
<tr>
<td>E</td>
<td>ecbJ</td>
<td>F- TCCGCAAGAAAATTATTGACTC&lt;br&gt;R- GCTTGCTGTGATTTGCT</td>
<td>511</td>
<td>[17]</td>
</tr>
<tr>
<td>F</td>
<td>fcbD</td>
<td>F- AATCGGAAGACCGAGAAATCAG&lt;br&gt;R- TCCGCGCTCAATTACTCTG</td>
<td>851</td>
<td>[17]</td>
</tr>
<tr>
<td>Adhesins</td>
<td>ptfA</td>
<td>F- TGTGGAATCTTACAGC&lt;br&gt;R- TCATGAAATTTAATCTGTC</td>
<td>488</td>
<td>[20]</td>
</tr>
<tr>
<td>Protectins</td>
<td>Omp87</td>
<td>F- AGGTGAAAGGTTATG&lt;br&gt;R- TACCTAACTCAACCTAC</td>
<td>200</td>
<td>[21]</td>
</tr>
<tr>
<td>Sialidases</td>
<td>nanB</td>
<td>F- CATTCACCAACACCTCT&lt;br&gt;R- GGACACTGTGACCTGCAC</td>
<td>555</td>
<td>[20]</td>
</tr>
<tr>
<td>Toxins</td>
<td>toxA</td>
<td>F- CCATGAGAAGC&lt;br&gt;R- GAATGCCACACCTCTCTAG</td>
<td>864</td>
<td>[20]</td>
</tr>
<tr>
<td>β-lactams</td>
<td>bla&lt;sub&gt;ROB-1&lt;/sub&gt;</td>
<td>F- AATACCCCTRTGAC&lt;br&gt;R- TCCTATGCTGAC</td>
<td>685</td>
<td>[22]</td>
</tr>
<tr>
<td>Aminoglycosides</td>
<td>aphA</td>
<td>F- TTATGCCCTTTCCGACCT&lt;br&gt;R- GAGAAACTCAACGGGACGAG</td>
<td>489</td>
<td>[22]</td>
</tr>
<tr>
<td>Macrolides</td>
<td>ermX</td>
<td>F- GAGATCGRACGAG&lt;br&gt;R- GTGTCGCCACATGCCGTCA</td>
<td>488</td>
<td>[22]</td>
</tr>
<tr>
<td>Tetracyclines</td>
<td>tetH</td>
<td>F- ATACTGCTGTAC&lt;br&gt;R- TCCCAATAAGCAGCT</td>
<td>1076</td>
<td>[22]</td>
</tr>
</tbody>
</table>

**Results**

**Frequency and identification of *P. multocida* among diseased rabbits**

Ten out of 100 rabbits (10%) examined in this study were positive for *P. multocida*. On blood agar medium, the colonies were smooth, glistening, iridescent, dew drop, convex and non-haemolytic; meanwhile, there was no growth on MacConkey’s agar plates. Microscopic examination of the colonies revealed that they were Gram-negative, non-motile, non-spore forming and coccobacilli shaped bacteria. The 10 isolates produced positive reactions with indole, oxidase and catalase tests, while they were negative for urease utilization test. The above examined criteria identified those isolates as *Pasteurella* spp. All the isolates were confirmed as *P. multocida* based on the PCR amplifications of kmt1 species specific gene fragments. All the isolates were of serogroup A as was identified by multiplex PCR for the detection of their capsular types.

**Antibiotic sensitivity test**

The distribution of antibiotic resistance among the analyzed isolates is presented in Figure 1A. All the isolates exhibited full resistance to ampicillin, amoxicillin, amoxicillin/clavulanic acid, neomycin and tetracycline (100%), followed by kanamycin and streptomycin (90%, each) and erythromycin and doxycycline (80% each). Resistance to at least 7 antibiotic agents was detected in *P. multocida* isolates with a remarkable MDR pattern being observed across all the tested isolates. Interestingly, four isolates (40%) were resistant to all tested antibiotics (Table 2).
Table 2. Phenotypic antibiotic resistance and presence of virulence and antibiotic resistance genes among ten *P. multocida* isolates of naturally infected rabbits

<table>
<thead>
<tr>
<th>Isolate code No.</th>
<th>Phenotypic antibiotic resistance profile</th>
<th>Antibiotic resistance gene(s)</th>
<th>Virulence genes</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>AM, AX, AMC, CN, N, S, K, E, TE</td>
<td>aphA1</td>
<td>ptfA, Omp87, nanB</td>
</tr>
<tr>
<td>2</td>
<td>AM, AX, AMC, CN, N, S, K, E, TE</td>
<td>aphA1</td>
<td>ptfA, Omp87, nanB</td>
</tr>
<tr>
<td>3</td>
<td>AM, AX, AMC, CN, N, S, K, E, TE, DO</td>
<td>aphA1,ermX</td>
<td>ptfA, Omp87, nanB</td>
</tr>
<tr>
<td>4</td>
<td>AM, AX, AMC, CN, N, S, K, E, TE, DO</td>
<td>aphA1, ermX, blaROB-1, tetH</td>
<td>ptfA, Omp87, nanB</td>
</tr>
<tr>
<td>5</td>
<td>AM, AX, AMC, CN, N, S, K, E, TE, DO</td>
<td>aphA1</td>
<td>ptfA, Omp87, nanB, toxA</td>
</tr>
<tr>
<td>6</td>
<td>AM, AX, AMC, N, S, E, TE, DO</td>
<td>aphA1</td>
<td>ptfA, Omp87, nanB</td>
</tr>
<tr>
<td>7</td>
<td>AM, AX, AMC, N, S, K, E, TE, DO</td>
<td>aphA1</td>
<td>ptfA, Omp87, nanB, toxA</td>
</tr>
<tr>
<td>8</td>
<td>AM, AX, AMC, CN, N, S, K, TE, DO</td>
<td>aphA1, ermX, blaROB-1</td>
<td>ptfA, Omp87, nanB</td>
</tr>
<tr>
<td>9</td>
<td>AM, AX, AMC, CN, N, S, K, E, TE, DO</td>
<td>aphA1, ermX</td>
<td>ptfA, Omp87, nanB, toxA</td>
</tr>
<tr>
<td>10</td>
<td>AM, AX, AMC, N, K, TE, DO</td>
<td>aphA1</td>
<td>ptfA, Omp87, nanB, toxA</td>
</tr>
</tbody>
</table>


Figure 1: Frequency of antibiotic resistance (A) and virulence and antibiotic resistance genes (B) among *P. multocida* isolates from diseased rabbits. AM: ampicillin, AX: amoxicillin, AMC: amoxicillin/clavulanic acid, CN: gentamicin, N: neomycin, K: kanamycin, S: streptomycin, E: erythromycin, TE: tetracycline, Do: doxycycline.
Virulence and resistance gene profiles of *P. multocida* isolates

The results confirmed that, with the exception of *toxA* gene, the other virulence associated genes were found among all examined field isolates (Figure 1B). According to the virulence genes analysis, four screened isolates harbored all examined genes (*ptfA*, *Omp87*, *nanB* and *toxA*) (Table 2); meanwhile, the remaining 6 isolates possessed only 3 of these virulence associated genes (*ptfA*, *Omp87* and *nanB*).

The PCR results indicated that all 10 *P. multocida* isolates contained antibiotic resistance genes regardless of their susceptibility patterns using disc diffusion assay. The resistance genes commonly detected were those encoding resistance to aminoglycosides (*aphA1*, 100%), followed by macrolides (*ermX*, 40%). The other genes conferring resistance to β-lactams and tetracyclines were present to lesser extents including *blaROB-1* (20%) and *tetH* (10%), respectively (Figure 1B). Except *aphA1* gene which was detected in all the 10 aminoglycosides resistant isolates, the PCR detections of antibiotic resistance genes were partially consistent with the phenotypic antibiotic susceptibility criteria because majority of the genes were not detected in isolates those had the potential to confer the relevant phenotypic resistance. Notably, 4 MDR isolates (40%) carried multiple genes with only one isolate harboring 4 genes encoding identical resistance phenotypes (Table 2). All MDR *P. multocida* isolates possessed at least 3 virulence genes accompanied by the existence of antibiotic resistance genes (Table 2). The expected PCR products for the examined virulence and resistance genes and their lengths are shown in Figure 2A-H.

![Figure 2: Agarose gel electrophoresis showing typical amplification products for virulence genes of *P. multocida* isolated from rabbits; *ptfA* (A), *Omp87* (B), *nanB* (C) and *toxA* (D) and antibiotic resistance genes; *blaROB-1* (E), *aphA1* (F), *ermX* (G) and *tetH* (H). Lane L: DNA molecular size marker (100-bp), lanes 1-10: *P. multocida* isolates from diseased rabbits, lane Pos.: positive control, lane Neg.: negative control. PCR amplification products and their sizes in base pairs (bp) are indicated.](image-url)
Discussion

Rabbit pasteurellosis continues to be a great threat facing rabbit industry in Egypt with a possibility of fatal consequences. In the current study, we demonstrated the occurrence of P. multocida in clinically diseased rabbits which originated from different localities at Sharkia Governorate, Egypt. Additionally, we demonstrated the distribution of MDR P. multocida isolates as well as their virulence and resistance genes’ profiles. We then monitored the co-occurrence of both virulence and resistance genes among the analyzed isolates. Ten P. multocida isolates were isolated from 100 nasal swabs collected from diseased rabbits with respiratory manifestations (10%). This result was nearly similar to those obtained by previous reports in Kafr El Sheikh (9.4%) [23] and Brazil (7.4-10%) [12]. On the other hand, the isolation percentage of P. multocida in the current study was low when compared with other previous reports in different Governorates in Egypt; Sharkia (47.5%) [24] and Kaliobeya (27%) [25]. The variation in the frequencies of P. multocida may be attributed to differences in geographical distributions, environmental conditions, stress factors, health and immune status of each examined rabbit and genetic resistance.

The recovered isolates were subjected to phenotypic identification depending on standard microbiological techniques as those from previous scientific literatures [26,27]. All the recovered isolates were confirmed as P. multocida by PCR amplifications of 460-bp products of kmt1 gene confirming that PCR is very important for rapid and specific characterization of P. multocida as was obtained previously [16]. Capsular typing using the multiplex PCR technique revealed that all P. multocida isolates (100%) were of capsular type A yielding the expected amplicons of 1044 bp. This finding is in agreement with other researches, where P. multocida capsular type A were most commonly isolated from rabbits [28,29].

The antimicrobial susceptibility data from the current study indicated higher resistance rates to ampicillin, amoxicillin, neomycin and tetracycline (100%) reported in previous studies conducted in India [30], where all P. multocida isolates of rabbits suffering from pasteurellosis were resistant to neomycin, penicillin and ampicillin and in Egypt [31], where all P. multocida isolated from diseased rabbits were resistant to tetracycline (100% each). These findings are probably considered as a consequence of the extensive use of these antibiotics in rabbit farms. On the other hand, the data on the antibiotic susceptibility of rabbit P. multocida isolates available in another literature conducted in Italy showed a relatively low frequency of resistance against ampicillin and tetracycline [32]. Different geographic locations with their particular isolated strains might have caused these variations.

Concerning the resistance profiles in our study, MDR phenomenon was observed among all the recovered isolates. This observation comes in parallel with a recent report conducted in Egypt [31], where all the examined rabbit P. multocida strains demonstrated remarkably MDR patterns. The presence of multiresistant P. multocida isolates recovered from rabbits affected by pasteurellosis has already been reported in Italy [32]. In another study in Brazil, 47.8% (22/46) of rabbit P. multocida strains were resistant to at least one of the tested drugs [12]. The antibiotic resistance is a growing problem probably attributed to the extensive and indiscriminate usage of antibiotics for prevention and treatment of pasteurellosis in the field.

In this study, P. multocida isolates were screened for the existence of four critical virulence associated genes (ptfA, Omp87, nanB and toxA) involved in bacterial pathogenesis. The results confirmed that, with the exception of toxA gene, the other virulence associated genes were found among all the isolates. Other researches worldwide confirmed our findings and reported the occurrence of selected virulence genes encoding dermonecrotxin production (toxA), fimbriae adhesions (ptfA), extracellular enzymes (nanB) and outer membrane protein synthesis (Omp87) in P. multocida isolated from rabbits [12,33].
The high prevalence rate of ptfA gene detected in this study (100%) was expected due to it is assumed to be a key factor in fixing the bacteria on the epithelial cells' surfaces. This result agrees with the finding of other researches [12]. Moreover, nanB gene was found in all the examined isolates as was previously evidenced [33]. This could be attributed to the fact that sialidases (nanB) play an important role in colonization to epithelial surfaces of the host and they also promote the virulence of bacteria by unmasking the key receptors and reducing mucin effectiveness. With respect to toxA gene, only four isolates harbored this gene. This may be attributed to the fact that the gene which encodes dermonecrototoxin is initially detected in serotype D isolates and it was found to be associated with atrophic rhinitis in pigs [34]. Later on, it was detected in serotype A strains from pigs and other hosts [35]. Moreover, even in toxigenic isolates of pigs, the presence of this gene evidenced to be low and it is usually missing after few subcultures.

Monitoring the antibiotic resistance trends among bacterial pathogens isolated from humans and animals is necessary to inform the public policy regarding the appropriate use of the antibiotics in both human and veterinary medicine [36]. To our knowledge, there were no published data regarding the prevalence of antibiotic resistance genes among _P. multocida_ rabbit isolates. The current study appeared also to be the first to report the relationship between the phenotypic and genotypic resistance profiles of the recovered isolates.

The _aphA1, ermX, blaROB-1_ and _tetH_ genes are among the aminoglycoside, macrolides, β-lactams and tetracyclines resistance determinants, respectively. They have been commonly observed in some bacterial pathogens, but data about these genes prevalence in _P. multocida_ of rabbits are limited, particularly in Egypt. The current study demonstrated a high prevalence of positive _aphA1_ _P. multocida_ isolates (100%). This gene was also detected in all 23 _P. multocida_ isolates from bovine respiratory infections in a previous study conducted in China [37]. Another recent Canadian study has also detected this gene in most _Mannheimia haemolytica_ isolated from nasal swabs of cattle [22]. In the current study, genes encoding resistance to β-lactams (_blaROB-1_) and tetracycline (_tetH_) showed frequencies (20 and 10%, respectively) lower than those reported in studies carried out in Spain [38], where all 13 _P. multocida_ isolates from diseased pigs carried _blaROB-1_ and in the United States and Canada [39], where 25 out of 31 _pasteurella_ isolates from cattle, turkey and pigs (80.64%) were found to contain _tet(H)_ gene.

Notably, most of the screened genes were not observed in the phenotypic resistant isolates except _aphA1_ gene which was detected in all of the 10 aminoglycosides resistant isolates. These results denote that there may be other antibiotic resistance genes encoding resistance that were not examined in the current study.

Finally, this study demonstrated the co-occurrence of virulence and antibiotic resistance genes among the analyzed isolates. The co-occurrence of these concerning trends confirmed that acquisition of antibiotic resistance has been accompanied by increased virulence. These findings were discussed in several reports [40,41], where the increased bacterial resistance is accompanied by an increased in their virulence. Fundamentally, the bacterial virulence is directly correlated with the evolution of antibiotic resistance from the theoretical point of view that just when the host manifests the disease clinical signs. This means that when the virulent bacterial pathogens are present, antibiotic treatment is administrated; however, in the absence of bacterial infection, exposure to antibiotics is much lower, so the possibility of emerging antibiotic resistance due to the lack of antibiotic pressure is becoming also lower.

**Conclusion**

Based on the previous findings, the current investigation demonstrated the high incidence of MDR and virulent _P. multocida_ isolates among the clinically diseased rabbits. Our observations highlight the role of rabbits as potential sources of _P. multocida_ with a wide range of virulence determinants and antibiotic resistance genes with subsequent major negative public health implications. These results should be used to emphasize the
responsibility of the veterinarians and relevant authorities to develop prudent use guidelines to minimize the emergence and spread of MDR bacterial strains originating from rabbits in Egypt.

Conflict of Interest
None of the authors have any conflict of interest to declare.

References


مصابحة جينات ضراوة ومقاومة المضادات الحيوية في الباستريلا مالتوسيدا المعزولة من الأرانب المريضة

تعتبر الباستريلا مالتوسيدا من العوامل البكتيرية السائدة للعرايز التنفسية في الأرانب مما يؤدي إلى خسائر اقتصادية كبيرة وسوء تشبه العرض في مصر. ومع ظهور القليل من عوارض الباستريلا مالتوسيدا المقاومة للعديد من المضادات الحيوية الشائعة الاستخدام في المجال البيطري حديثاً، ذلك أجريت الدراسة الحالية لمعرفة مدى انتشار كلا من جينات ضراوة والمقاومة للمضادات الحيوية في عوارض الباستريلا مالتوسيدا المعزولة من الأرانب المريضة بمحافظة الشرقية. تم تأكد فقط من 100 مسحة أنتي بحثية مختبرة من الأرانب على النهاية الباستريلا مالتوسيدا من النوع أ بواسطة تفاعل أنزيم البلمرة المتباعد.

وقد كشف اختيار جينات المضادات الحيوية للعوارض أن كلها كانت مقاومة للعديد من المضادات الحيوية مع شيوع المقاومة للأموكسيسيلين، الأمبيسيلين، الأموكسيسيلين / كلافوتيك أميد، النيپاميسين والتراسكيلين (100% لكل منها)، بليم الكاناميدين والستريتوسين (100% لكل منها). كما تم الكشف عن بعض جينات الضراوة والمقاومة للمضادات الحيوية ذات الأهمية في جميع العوارض عن طريق تفاعل أنزيم البلمرة المتباعد. واستثناء جين toxA، فقد تم العثور على باقي الجينات الأخرى ذات الضراوة (nanB و Omp87 و ptfA) في جميع العوارض المحفوصة. وبشكل عام، فقد احتوت جميع العوارض مقدار المقاومة للمضادات الحيوية على جين واحد من جينات مقاومة المضادات الحيوية على الأقل مع انتشار جين aphA1 (100%). أظهرت الدراسات الجينية لمقاومة المضادات الحيوية وجود جينات مقاومة متعددة للمضادات الحيوية في غالبية العوارض (94%)، مع وجود عزلة واحدة فقط بها 4 جينات مقاومة ترمز إلى نفس أنماط المقاومة الظاهرة. وبصورته واضحة فقد حُدد جميع عوارض الباستريلا مالتوسيدا مرتدة المقاومة للمضادات الحيوية على 3 جينات ضراوة على الأقل مصحوبة بوجود جينات مقاومة للمضادات الحيوية. أثبتت هذه النتائج أن الأرانب هي مصدر محتملة لعوارض الباستريلا مالتوسيدا المسببة للأمراض المحتوية على كلا من جينات الضراوة والمقاومة للمضادات الحيوية. ولذلك، من الواضح أن هناك حاجة لاستخدام الحكيم للمضادات الحيوية في أسلحة علاج الأرانب من أجل تخفيض بنجاح من انتشار المقاومة المتعددة للأدوية عبر أنواع الباستريلا مالتوسيدا.