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Detection of Avian Metapneumovirus Subtypes A and B in Moroccan Broiler

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Farms

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Running title: aMPV A & B in Morocco

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Abstract

Background: Avian Metapneumovirus (aMPV) has been proven to be a widespread infectious respiratory pathogen affecting turkeys and chickens, with co-predominance of the subtypes A and B.

Objectives: No official reports exist in Morocco about the subtypes of aMPV circulating. Hence, using quantitative Reverse-Transcriptase Polymerase Chain Reaction (qRT-PCR) subtypes-specific A and B, we aimed at detecting and identifying the potential subtype (s) circulating.

Methods: We conducted a longitudinal study on three broiler flocks strictly not vaccinated against aMPV and located in two different geographical regions, and two flocks that expressed typical swollen head syndrome (SHS) and sampled once. Furthermore, we sampled dead birds of one flock confirmed seropositive from a previous study. 118 swabs pooled in 24 samples were subjected to the ribonucleic acid (RNA) extraction and amplified using a triplex RT-PCR for specific detection of aMPV subtypes A and B.

40 Additionally, serum samples were taken at slaughtering age to cross-check the molecular results. A total of 84 sera were analyzed with a commercial indirect Enzyme-Linked Immunosorbent Assay (ELISA) kit to detect and titer antibodies against the two subtypes.

Results: Avian Metapneumovirus was detected by qRT-PCR in all the flocks. 87.50% of the samples were positive for aMPV-B, and 16.67% for aMPV-A and aMPV-B simultaneously. All

45 the flocks showed seropositivity, confirming the molecular findings.

Conclusion: The present investigation is the first molecular study in Morocco to elucidate the circulation of aMPV-A and aMPV-B in broiler farms in Morocco with a dominance of aMPV-B and the possibility of co-presence of both subtypes.

Keywords: avian Metapneumovirus, broilers, Morocco, RT-PCR, subtypes.

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Introduction

Avian Metapneumovirus (aMPV) is a significant virus that causes respiratory and reproductive distress in chickens and turkeys. This leads to impaired performance and increased mortality, particularly when accompanied by secondary infections. While the virus has traditionally been downplayed in broilers, recent field investigations have highlighted the direct implication of aMPV in respiratory problems (Al-Hasan et *al.*, 2022; Nguyen et *al.*, 2021; Franzo et *al.*, 2017;

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Tucciarone et *al.*, 2018).

Previously, aMPV was classified into four subtypes (aMPV-A to D) based on variations in the glycoprotein (G), responsible for surface attachment, and the antigenic differences between strains (Cook and Cavanagh, 2002). However, the discovery of two divergent viruses in a monk parakeet and a gull has raised the possibility of new subtype candidates (Canuti et *al.*, 2019; Retallack et *al.*, 2019).

Although subtype B is more prevalent than subtype A (Mernizi et *al.*, 2023), both aMPV-A and aMPV-B are widespread except in Australia and Canada (Suarez et *al.*, 2019). In the United States, only subtype C has been reported in turkeys so far, without evidence of spread in broilers per se (Cha et *al.*, 2013). aMPV-C has also been confirmed in Asia (Kwon et *al.*, 2010) and recently in wild birds in Italy, but it belongs to a different lineage known as the Eurasian sub-lineage (Graziosi et *al.*, 2022; Tucciarone et *al.*, 2022; Toquín et *al.*, 2006). On the other hand, subtype D has only been detected in turkeys in France (Bäyon-Auboyer et *al.*, 2000).

- 70 Diagnosing aMPV infection based solely on clinical signs is irrelevant due to the nonpathognomonic symptoms, especially in broilers. Isolating the virus is time-consuming, laborintensive, and requires a viable virus. Therefore, routine detection of aMPV can be achieved through serology, molecular tests, or a combination of both, depending on the timing of the sample (Lemaitre et *al.*, 2018). Serological tests such as ELISA detect antibodies produced after
- ⁷⁵ infection (Rautenschlein et *al.*, 2011), but the results are delayed by at least two weeks for seroconversion (Guionie et *al.*, 2007). Molecular techniques such as PCR, which detect viral genetic material, are preferred during the infectious phase and provide sensitive, specific, and rapid results (Franzo et *al.*, 2014). They can also differentiate between subtypes using specific gene-sequence-based real-time qRT-PCR (Cook & Cavanagh, 2002; Guionie et *al.*, 2007).
- In Morocco, official subtyping data is not available, as research is limited to national serological evidence and the identification of several risk factors associated with seropositivity (Mernizi et *al.*, 2023). Therefore, the present study aims to highlight the circulation of aMPV-A and aMPV-B in broiler flocks using a quantitative triplex RT-PCR targeting the G gene, which allows for differentiation between these two subtypes. These findings will be supported by confirmation through blood sera tested with ELISA.

Materials and Methods

Sampling protocol

The investigation was designed as a longitudinal study focusing on broilers that were strictly not vaccinated against aMPV. The study involved three flocks, namely 2-N38, 2-N39, and 2-TS416

90 (designated with internal codification), and located in two areas in Morocco. From 3 to 5 weeks of age, each flock was swabbed every 3 or 4 days, specifically from the trachea. On each occasion, ten birds were randomly chosen and sampled.

At the request of the responsible veterinarians, tracheal and turbinate swabs were collected from two neighboring flocks located in another geographic area. These flocks were over five weeks of

95 age and had reported cases of SHS (Swollen Head Syndrome). A one-time sampling of ten randomly selected birds per flock was conducted seven days after the appearance of clinical signs.

Furthermore, molecular analysis was conducted to verify the aMPV serological positive results obtained from a previous study. Swabs were collected post-mortem from the preserved trachea and inner side of the skin of eight bird skull heads that exhibited typical swelling. These birds belonged to a flock that had already been tested and confirmed positive for aMPV using serology.

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To complement the detection of aMPV, a serology test was also performed for all the flocks. Twenty sera were collected at the age of slaughter for each flock in the longitudinal study, and twelve sera were sampled simultaneously with the swabs for the one-off sampling flocks. Although both types of samples were obtained from the same birds as mentioned earlier, the collected tissues and serum were not paired or individually identified.

Fresh blood samples were collected from the brachial wing veins of the birds by puncturing the alar veins. The samples were then stored in sterile tubes and transported to the Avian Pathology

110 Unit of the Hassan II Agronomic and Veterinary Institute in Rabat. Serum extraction and preservation in Eppendorf tubes at -20°C were carried out for subsequent analysis using a commercial indirect ELISA kit capable of detecting and titrating antibodies against both subtypes A and B, CIVTEST AVI TRT®, HIPRA S.A. (Amer, Spain).

The mean titers, validity tests, and coefficients of variation were automatically calculated by flock and sample series using the HIPRASOFT® 5.0 software from HIPRA S.A. (Amer, Spain).

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Table 1 provides an overview of the sampling protocol for the different flocks included in the study.

Table 1. Sampling protocol for the longitudinal study and the one-off sampling flock

| Flock | Age (days) | Swabbed organ(s) | Swabs (n) | Sera (n) |
|-------|------------|------------------|-----------|----------|
| 2-N38 | 23 | Trachea | 10 | |
| | 26 | | 10 | |

| 2 | 8 | 10 | |
|-------------|-------------------------------|----|--------------|
| | 0 | 10 | |
| | | | C . |
| 3 | 5 | | 20 |
| 2 | 3 | 10 | \mathbf{O} |
| 2 2 2 | 6 Trachea | 10 | |
| 2-1039 | 8 | 10 | |
| 3 | 5 | | 20 |
| 2 | 6 | 10 | |
| 2-TS416 3 | 1 Trachea | 10 | |
| 3 | 5 | 10 | 20 |
| 2-MK36 3 | 7 Trachea and nasal turbinate | 10 | 12 |
| 2-MK37 4 | 2 Trachea and nasal turbinate | 10 | 12 |
| 1 TS50(*) 2 | 6 Trachea: subcutis (heads) | 08 | |

(*) The samples of this flock, belong to a sero-epidemiological study performed in 2021

Samples preparation and ribonucleic acid extraction

Following identification and date referencing, swabs were immediately placed in an icebox without any transport media and delivered to the Avian Pathology Unit of the Hassan II Agronomic and Veterinary Institute in Rabat. The samples were then stored at -20°C until the extraction of ribonucleic acid (RNA) was performed.

The skull heads were also stored at the Avian Pathology Unit of the Hassan II Agronomic and Veterinary Institute in Rabat, maintained at a temperature of -20°C. The skin was preserved to increase the likelihood of detecting the genetic material of the aMPV.

For each flock, every five swabs (or four swabs in the case of post-mortem samples) were pooled together, resulting in a total of 24 pools.

The RNA extraction from the dry pools was carried out using the PureLinkTM Viral RNA/DNA Mini Kit® from Thermo Fisher Scientific (Waltham, Massachusetts, USA), following the manufacturer's instructions.

It is important to note that RNA ideally should be stored at -80°C. However, due to the unavailability of equipment capable of reaching such low temperatures, the RNA samples were only stored at -20°C.

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Reverse-Transcription & amplification

The RNA amplification was processed using a single-step triplex real-time Reverse Transcriptase PCR (RTRT-PCR) targeting the G gene. In this technique, the RNA sequences of

140 aMPV subtype A and B, along with an endogenous control, were transcribed in reverse and amplified in a single tube using specific primer pairs in the polymerase chain reaction.

To detect the amplified RNA of aMPV-A, aMPV-B, and the control's endogenous target gene (beta-actin), TaqMan probes labeled with fluorescent dyes (Fam, Cy5, and HEX) were utilized. The thermocycler measured the emitted fluorescence during the amplification process.

- 145 The endogenous control was based on the detection of beta-actin, a "housekeeping" protein present in the host cells from which the samples originated. The target beta-actin gene (endogenous RNA) was co-amplified (HEX channel) in each reaction. This control allowed for the assessment of sampling adequacy, sample storage and shipping, sample preparation, and the execution of the real-time RT-PCR itself.
- 150 A positive control was included to ensure the specificity and efficiency of reagents, the RT-PCR reaction, and the thermocycler. On the other hand, a negative control was used to exclude any contamination.

The Kylt® aMPV A&B kit from AniCon Labor GmbH (Emstek, Germany) was employed for the amplification, following the manufacturer's instructions. The data obtained from the

155amplification was automatically processed using the ThermoCycler AriaMx® Real-Time PCRSystem software from Agilent Technologies (Santa Clara, California, USA).

Test evaluation

The analysis of the test began by conducting a validity check for the entire real-time RT-PCR series. This check ensured that the negative control samples for the FAM, HEX, and Cy5

160 channels were negative, while the positive control samples for these channels were positive with values greater than 15 and equal to or less than 35.

Additionally, internal control was used to validate each sample reaction and its real test result. The Cq value of the internal control channel (HEX) was compared to determine the validity of the sample reaction.

165 Finally, the specific status of aMPV subtypes A and B for each sample was analyzed using the FAM and Cy5 channels, respectively.



Table 2 shows the detection of aMPV in the different flocks and the number of positive results 170 per pool.

| | | | samplings | 5 | | |
|---------|--------|-----------|--------------------------|-------|-------|--|
| Flock | Age | Number of | Positive pools | | | |
| | (days) | pools | aMPV-A aMPV-B aMPV-A & B | | | |
| | | | | | | |
| | 23 | 02 | 00/02 | 02/02 | 00/02 | |
| 2-N38 | 26 | 02 | 00/02 | 02/02 | 00/02 | |
| | 28 | 02 | 00/02 | 02/02 | 00/02 | |
| | 23 | 02 | 00/02 | 02/02 | 00/02 | |
| 2-N39 | 26 | 02 | 00/02 | 02/02 | 00/02 | |
| | 28 | 02 | 00/02 | 02/02 | 00/02 | |
| | 26 | 02 | 00/02 | 00/02 | 00/02 | |
| 2-TS416 | 31 | 02 | 00/02 | 02/02 | 00/02 | |

 Table 2. Detection of aMPV-A and aMPV-B by age from the longitudinal study and one-off samplings

| | 35 | 02 | 00/02 | 02/02 | 00/02 | |
|--------|----|----|-------|-------|-------|-----------------|
| 2-MK36 | 37 | 02 | 02/02 | 02/02 | 02/02 | $\overline{\ }$ |
| 2-MK37 | 42 | 02 | 02/02 | 02/02 | 02/02 | \mathbf{O} |
| 1-TS59 | 36 | 02 | 00/02 | 01/02 | 00/02 | |
| Total | | 24 | 04/24 | 21/24 | 04/24 | |

The subtype B of aMPV was detected in all six flocks included in the study. It was found in a total of 21 out of 24 pools, corresponding to 87.50% of the samples, and indicating a high

prevalence.

ELISA

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The proportion of pools testing positive for subtype A was 4 out of 24, corresponding to 16.67% of the samples. The subtype A of aMPV was always detected simultaneously with subtype B.

180 The following Figure illustrates the results obtained from the indirect ELISA test to detect and titer antibodies against aMPV subtypes A and B. The sera originated from the five flocks sampled, namely: 2-N38, 2-N39, 2-MK37, 2-MK38, and 2-TS416.



Figure. Detection and titration of antibodies against aMPV-A and aMPV-B by indirect 185 ELISA in flocks from the longitudinal study (2-N38, 2-N39, and 2-7S416) and one-off sampling (2-MK37 and 2-MK38) using the CIVTEST AVI TRT®, with kit's cut-off = 196, and Geometric Mean Titers (GMT) are represented by a cross mark "x"

Based on the cut-off value of the CIVTEST AVI TRT® kit, 54 out of the total 84 sera samples tested positive, indicating a seropositivity rate of 64.28%.

190 It is worth noting that all the five flocks included in the study had geometric mean titers (GMT) above the cut-off value of 196. This means that the antibody levels in all the flocks were higher than the threshold considered for seropositivity. Therefore, all the flocks demonstrated a seropositive status for aMPV, indicating previous exposure to the virus.

Discussion

- 195 Although aMPV infection has been known as the causative agent of the swollen head syndrome (SHS) in broilers in Morocco for over 30 years, its role in respiratory problems as a primary agent has not been clearly established. The present field investigation provided the first evidence of the circulation of aMPV in Moroccan broiler farms using RT-PCR and ELISA, confirming the presence of the virus and identifying its most important and prevalent subtypes.
- The study revealed that subtype B of aMPV was detected in all the flocks studied, indicating its widespread presence. In contrast, subtype A was only identified in two neighboring farms where clinical cases of SHS had been reported. These findings suggest that subtype B is the predominant circulating subtype, which is consistent with its high prevalence reported in North Africa and the Mediterranean Basin (Lachheb et *al.*, 2022; Sid et *al.*, 2015; Franzo et *al.*, 2017;
 2020; Tucciarone et *al.*, 2017). Subtype B has been recognized for its high spreading capacity in the region.

Our findings are consistent with previous reports that highlighted the dominance of aMPV subtype B over subtype A (Banet-Noach et *al.*, 2005; Dos Santos et *al.*, 2012) or non-B subtyped aMPV in general (Darebaghi et *al.*, 2021). This further supports that subtype B is more widespread compared to subtype A.

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The longitudinal study conducted in our investigation demonstrated the relevance of the results obtained. We were able to detect aMPV in flocks 2-N38 and 2-N39 starting from three weeks of age and onwards, even in the absence of respiratory symptoms. However, in the case of flock 2-TS416, respiratory symptoms were observed at five weeks of age, while the one-time sampled

215 flocks, 2-MK36 and 2-MK37, exhibited clinical symptoms of swollen head syndrome. It is worth noting that the sampling protocol for flock 2-TS416 was slightly delayed due to logistical issues and the unavailability of responsible individuals.

In field conditions, determining the exact onset time of infection and thus identifying the optimal sampling time can be challenging. In Morocco, all chicks usually come from breeders that have been immunized against aMPV using live and killed vaccines. As a result, it is expected that flocks wouldn't be exposed to field virus challenge earlier than three weeks of age, as the levels of maternal antibodies decline progressively and disappear only after two weeks from hatching (Rubbenstroth and Rautenschlein, 2009). Furthermore, it is known that the highest quantity of virus is shed by infected birds from three to

five days post-infection (Catelli et *al.*, 1998). Considering these factors, the repetition of sample collection in our study was conducted within a short period, which was less than five days. This approach aimed to increase the likelihood of detecting the genetic material of aMPV during the active shedding phase.

Nevertheless, identifying aMPV at such young age could be linked to a general decreased immunocompetence predisposition, resulting from the intense genetic selection for rapid growth and high production rates (Nikbakht Brujeni, 2022a; Nikbakht Brujeni et *al.*, 2022b).

An interesting finding in our study was the simultaneous co-circulation of aMPV-A and aMPV-B in the two sampled flocks, namely 2-MK36 and 2-MK37. Similar observations have been reported in Israel (Banet-Noach et *al.*, 2005) and Brazil (Chacón et *al.*, 2011). However, in our

235 case, these two neighboring flocks belonged to an area known exclusively for broiler production, making it initially unlikely to find more than one subtype of aMPV, especially considering the absence of commercial turkeys or layers in the vicinity.

Additionally, it was surprising to detect the genetic material of aMPV one week after the recovery from SHS, assuming the virus should be rapidly cleared and bacterial complications would limit its detection. However, it has been demonstrated that convalescent flocks can

experience reinfection by aMPV due to the continuous circulation of the virus within the flock or across the surrounding area (Al-Hasan et *al.*, 2022).

In our study, the absence of an extended investigation involving other pathogens such as Infectious Bronchitis Virus (IBV) or *Mycoplasma gallisepticum*, known to be prevalent in respiratory complexes (Muofaq Khalaf and Jawad Ali, 2023; Motamed and Bashashati, 2022; Hajijafari Anaraki et *al.*, 2021), assumed that the flocks investigated were aMPV mono-infected only. This plausible attribution may have contributed to the primordial lack of noticeable signs or complications during the visit, or paradoxically, the importance of aMPV as a major respiratory pathogen (Tucciarone et *al.*, 2018), in the case of the flock 2-TS416 that showed respiratory distress. That same observation potentially indicates a state of recovery within the flocks, despite contradicting the reported occurrence of SHS in flocks 2-MK37 and 2-MK38, which typically involve bacterial secondary infections.

The serological analysis conducted in our study was highly relevant as it demonstrated positive antibody titers in flocks that were not vaccinated against aMPV and subsequently, presumably exposed to field virus. Remarkably, all the flocks exhibited antibody levels higher than those typically observed in naïve birds or following vaccination. While titers are generally expected to be higher and more homogenous after infection, the possibility of the presence of immunosuppressive agents such as the Gumboro disease virus may have lowered the levels of anti-aMPV antibodies (Sharifi et *al.*, 2022). This is particularly noteworthy because flocks 2-MK37 and 2-MK38 were reported to have shown symptoms consistent with clinical Gumboro

cases, despite the absence of overt signs or conclusive evidence during our visit.

Therefore, the positive serological results obtained in our study support and validate our molecular findings, considering the high national seroprevalence of aMPV in Morocco (Mernizi et *al.*, 2022). These findings further emphasize the importance of considering both molecular and

serological approaches to comprehensively assess the presence and impact of aMPV in broilers.

Conclusion

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The present study provides valuable insights into the prevalence and co-circulation of aMPV subtypes in Moroccan broiler farms, highlighting the predominance of subtype B. The concurrent presence of subtypes A and B within the same flock is an interesting finding and emphasizes the need for further molecular characterization of the circulating subtypes in the country. Currently, there is limited information about the epidemiological situation of aMPV in Morocco compared to other regions of the world, including neighboring countries, and the origins of the field strains remain unclear.

To address these knowledge gaps, future research should focus on extending the molecular investigation and incorporating strain sequencing. By doing so, a more comprehensive understanding of the characteristics of aMPV isolates in Morocco can be gained, enabling the development of targeted strategies for controlling and preventing aMPV infections in broiler farms. This ongoing work will contribute to updating our understanding of aMPV epidemiology and inform the implementation of effective management measures in the poultry industry.

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References

- Al-Hasan, B.A., Alhatami, A.O., Abdulwahab, H.M., Bustani, G.S., Hameed, M.A., Jawad, A.H. (2022). First report of Avian metapneumovirus type B in Iraqi broiler flocks with swollen
- head syndrome. Veterinary World, 15, 16–21. doi:10.14202/vetworld.2022.16-21. PMID: 35369601.
 - Banet-Noach, C., Simanov, L., Perk, S. (2005). Characterization of Israeli avian metapneumovirus strains in turkeys and chickens. Avian Pathology, 34(3), 220–226. doi:10.1080/03079450500112625. PMID: 16191705.
- Bäyon-Auboyer, M.H., Arnauld, C. Toquin, D., Eterradossi, N. (2000). Nucleotide sequences of the F, L and G protein genes of two non-A/non-B avian pneumoviruses (APV) reveal a novel APV subgroup. The Journal of General Virology, 81:2723–2733. doi:10.1099/0022-1317-81-11-2723. PMID: 11038385.
- Canuti, M., Kroyer, A.N.K., Ojkic, D., Whitney, H.G., Robertson, G.J., Lang, A.S. (2019).
 Discovery and characterization of novel rna viruses in aquatic North American wild birds.
 Viruses, 11(9), 1–14. <u>doi:10.3390/v11090768. PMID: 31438486.</u>

Catelli, E., Cook, J.K.A., Chesher, J., Orbell, S.J., Woods, M.A., Baxendale, W., Huggins, M.B. (1998). The use of virus isolation, histopathology and immunoperoxidase techniques to

study the dissemination of a chicken isolate of avian pneumovirus in chickens. Avian

- 310 Pathology, 27(6), 632–640. <u>doi:10.1080/03079459808419395</u>. <u>PMID: 18484053</u>.
 - Cha, R.M., Yu, Q., Zsak, L. (2013). The pathogenicity of avian metapneumovirus subtype C wild bird isolates in domestic turkeys. Virology Journal, 10(1), 1–8. doi:10.1186/1743-422X-10-38. PMID: 23363433.
 - Chacón, J.L., Mizuma, A.M., Vejarano, A.M.P., Toquín, D., Eterradossi, N., Patnayak, D.P.,
- 315 Goyal, S.M., Ferreira, A.J.P. (2011). Avian Metapneumovirus Subtypes Circulating in Brazilian Vaccinated and Nonvaccinated Chicken and Turkey Farms. Avian Diseases, Mar;55(1):82–89. doi:10.1637/9501-081310-Reg.1. PMID: 21500641.
 - Cook, J.K.A., Cavanagh, D. (2002). Detection and differentiation of avian pneumoviruses (metapneumoviruses). Avian Pathology, 31(2), 117–132. doi:10.1080/03079450120118603.
- 320 PMID: 12396356.
 - Darebaghi, A., Emadi Chashmi, H., Kafshdozan, K., Hosseini, H.A. (2021). Molecular Detection of Avian Metapneumovirus in Semnan Broiler Farms. Iranian Journal of Veterinary Medicine, 15(1), 27-34. doi:10.22059/ijvm.2020.302556.1005087.
- Dos Santos, M.B., Martini, M.C., Ferreira, H.L., Da Silva, L.H.A., Fellipe, P.A., Spilki, F.R.,
 Arns, C.W. (2012). Brazilian avian metapneumovirus subtypes A and B: Experimental infection of broilers and evaluation of vaccine efficacy. Pesquisa Veterinária Brasileira, 32(12), 1257–1262. doi:10.1590/S0100-736X2012001200008.

- Franzo, G., Legnardi, M., Mescolini, G., Tucciarone, C.M., Lupini, C., Quaglia, G., Catelli, E., Cecchinato, M. (2020). Avian Metapneumovirus subtype B around Europe: a phylodynamic
- reconstruction. Veterinary Research, 51(1), 1–10. doi:10.1186/s13567-020-00817-6. PMID:
 32641149.
 - Franzo, G., Tucciarone, C.M., Enache, M., Bejan, V. Ramon, G., Koutoulis, K.C., Cecchinato, M. (2017). First Report of Avian Metapneumovirus Subtype B Field Strain in a Romanian Broiler Flock during an Outbreak of Respiratory Disease. Avian Diseases, 61(2), 250–254. doi:10.1637/11557-121216-ResNote.1. PMID: 28665722.

- Franzo, G., Drigo, M., Lupini, C., Catelli, E., Laconi, A., Listorti, V., Bonci, M., Naylor, C.J., Martini, M., Cecchinato, M. (2014). A sensitive, reproducible, and economic real-time reverse transcription PCR detecting avian metapneumovirus subtypes A and B. Avian Diseases, 58(2), 216–222. doi:10.1637/10676-092413-Reg.1. PMID: 25055624.
- Graziosi, G., Lupini, C., Catelli, E. (2022). Disentangling the role of wild birds in avian metapneumovirus (aMPV) epidemiology: A systematic review and meta-analysis. Transboundary and Emerging Diseases, 69(6), 3285–3299. doi:10.1111/tbed.14680. PMID: 35960706.
- Guionie, O., Toquín, D., Sellal, E., Bouley, S., Zwingelstein, F., Allée, C., Bougeard, S.
 Lemière, S., Eterradossi, N. (2007). Laboratory evaluation of a quantitative real-time reverse transcription PCR assay for the detection and identification of the four subgroups of avian

metapneumovirus. Journal of Virological Methods, 139(2), 150–158. doi:10.1016/j.jviromet.2006.09.022. PMID: 17126416.

Hajijafari Anaraki, M., Sheikhi, N., Haghbin Nazarpak, H., Nikbakht Brujeni, G.H. (2022). Real

- Time Detection of Different Variant Strains of Infectious Bronchitis Virus in Trachea, Lung and Kidney of Infected Broiler Chickens. Iranian Journal of Veterinary Medicine, 16(1), 57-63. doi:10.22059/IJVM.2021.321594.1005165.
 - Kwon, J.S., Lee, H.J., Jeong, S.H., Park, J.Y., Hong, Y.H., Lee, Y.J., Youn, H.S., Lee, D.W., Do,S.H., Park, S.Y., Choi, I.S., Lee, J.B., Song, C.S. (2010). Isolation and characterization of
- avian metapneumovirus from chickens in Korea. Journal of Veterinary Science, 11(1), 59–
 66. doi:10.4142/jvs.2010.11.1.59. PMID: 20195066.
 - Lachheb, J., Bouslama, Z., Nsiri, J., Badr, C., Al Gallas, N., Souissi, S., Khazri, I., Larbi, I., Kaboudi, K., Ghram, A. (2022). Phylogenetic and phylodynamic analyses of subtype-B metapneumovirus from chickens in Tunisia. Poult. Sci., 102(1), 102253.
- doi:10.1016/j.psj.2022.102253 PMID: 36455491.
 - Lemaitre, E., Allée, C., Vabret, A., Eterradossi, N., Brown, P.A. (2018). Single reaction, real time RT-PCR detection of all known avian and human metapneumoviruses. Journal of Virological Methods, 251, 61–68. doi:10.1016/j.jviromet.2017.10.010. PMID: 29030071.

Luis, J., Mizuma, M., Vejarano, M.P., Toquín, D., Eterradossi, N., Patnayak, D.P., Goyal, SM.,
Ferreira, A.J.P. (2011). Avian Metapneumovirus Subtypes Circulating in Brazilian

Vaccinated and Nonvaccinated Chicken and Turkey Farms. Avian Diseases, Mar;55(1), 82–9. doi:10.1637/9501-081310-Reg.1. PMID: 21500641.

Mernizi, A., Fathi, H., Criado, J.L., Bouslikhane, Ghram, A., M., Mouahid, M., Nassik, S. (2023). Avian Metapneumovirus Review: A Focus on Broilers. Poultry Science Journal, 11(1), 1–17. doi:10.22069/psj.2022.20165.1814.

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- Mernizi, A, Bouziane, S. Fathi, H., Criado, J.L., Bouslikhane, M., Ghram, A., Catelli, E., Mouahid, M., Nassik, S. (2023). First seroepidemiological and risk factor survey of avian metapneumovirus circulation in Moroccan broiler farms. Veterinarski Glasnik, 77(1), 51-68, doi:10.2298/VETGL220307009M.
- Motamed, N., Bashashati, M. (2022). Comparative Genetic Features of S1 Gene from Six
 Massachusetts and Variant Avian 2 Infectious Bronchitis Viruses Isolated in Iran. Archives of Razi Institute, Oct 31;77(5):1611-1619. doi: 10.22092/ARI.2022.356836.1924. PMID: 37123127.
- Muofaq Khalaf, S., Jawad Ali, A. (2023). Immune Response and Histological Changes in
 Broilers Chickens Vaccinated with Mycoplasma gallisepticum Vaccines. Archives of Razi
 Institute, Mar/Ap 31;78(2):769-775. doi:10.22092/ARI.2022.359729.2464.

Nikbakht Brujeni, B., Houshmand, P., Esmailnejad, A., Abasabadi, F. (2022). Major Histocompatibility Complex as Marker-assisted Selection for Breeding Immunocompetent Animal. Iranian Journal of Veterinary Medicine, 16(3), 211-227. doi:10.22059/IJVM.2022.339598.1005251.

- Nikbakht Brujeni, N. (2022). Novel Insights into Infection and Immunity. Iranian Journal of Veterinary Medicine, 16(2), 99-100. doi: 10.22059/IJVM.2022.337927.1005234.
- Nguyen, V.G., Chung, H.C., Do, H.Q., Nguyen, T.T., Cao, T.B.P., Truong, H.T., Mai, T.N., Le, T.T., Nguyen, T.H., Le, T.L., Huynh, M.L. (2021). Serological and molecular characterization of avian metapneumovirus in chickens in Northern Vietnam. Veterinary Sciences, 8(10). doi:10.3390/vetsci8100206. PMID: 34679036.

390

- Rautenschlein, S., Aung, Y.H., Haase, C. (2011). Local and systemic immune responses following infection of broiler-type chickens with avian Metapneumovirus subtypes A and B. Veterinary Immunology and Immunopathology, 140(1–2), 10–22. doi:10.1016/j.vetimm.2010.11.003. PMID: 21183227.
- Retallack, H., Clubb, S., DeRisi, J.L. (2019). Genome Sequence of a Divergent Avian Metapneumovirus from a Monk Parakeet (*Myiopsitta monachus*). Microbiology Resource Announcements, 8(16), 99–101. doi:10.1128/mra.00284-19. PMID: 31000555.
- Rubbenstroth, D., Rautenschlein, S. (2009). Investigations on the protective role of passively
 transferred antibodies against avian metapneumovirus infection in turkeys. Avian Pathology, 38:427–436. doi:10.1080/03079450903349204. PMID: 19937531.

Sharifi, A., Allymehr, M., Talebi, A. (2022). Concurrent Occurrence of Infectious Bursal Disease and Multicausal Respiratory Infections Caused by Newcastle Disease and Avian Metapneumovirus in Broilers. Archives of Razi Institute, Jun 30;77(3):1007-1016. doi:10.22092/ARI.2021.354272.1631. PMID: 36618283.

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- Sid, H., Benachour, K., Rautenschlein, S. (2015). Co-infection with Multiple Respiratory Pathogens Contributes to Increased Mortality Rates in Algerian Poultry Flocks. Avian Diseases, 59(3), 440–446. doi:10.1637/11063-031615-Case.1. PMID: 26478165.
- Suarez, D.L., Miller, P.J., Koch, G., Mundt, E., Rautenschlein, S. (2019). Newcastle disease,
- other avian paramyxoviruses, and avian metapneumovirus infections. In Diseases of Poultry. Swayne, D.E., Boulianne, M., Logue, C.M., McDougald, L.R., Nair, V., Suarez, D.L., Wit, S., Grimes, T., Johnson, D., Kromm, M., Prajitno, T.Y., Rubinoff, I., Zavala, G. (eds.). (14th Edition). Wiley, New Jersey, pp. 111–166. doi:10.1002/9781119371199.ch3.
 - Toquín, D., Guionie, O., Jestin, V., Zwingelstein, F., Allee, C., Eterradossi, N. (2006). European
- and American subgroup C isolates of avian metapneumovirus belong to different genetic lineages. Virus Genes, Feb;32(1):97–103. doi: 10.1007/s11262-005-5850-3. PMID: 16525740.

Tucciarone, C.M., Franzo, G. Legnardi, M., Pasotto, D., Lupini, C., Catelli, E., Quaglia, G., Graziosi, G., Dal Molin, E., Gobbo, F., Cecchinato, M. (2022). Molecular Survey on A, B, C

- 420 and New Avian Metapneumovirus (aMPV) Subtypes in Wild Birds of Northern-Central Italy. Veterinary Sciences, 9(7). doi:10.3390/vetsci9070373. PMID: 35878390.
 - Tucciarone, C.M., Franzo, G., Lupini, C., Alejo, C.T., Listorti, V., Mescolini, G., Brandão, P.E.,
 Martini, M., Catelli, E., Cecchinato, M. (2018). Avian Metapneumovirus circulation in
 Italian broiler farms. Poultry Science, 97(2), 503–509. doi:10.3382/ps/pex350. PMID:
 29253264.
 - Tucciarone, C.M., Andreopoulou, M., Franzo, G., Prentza, G., Chaligiannis, I., Cecchinato, M. (2017). First Identification and Molecular Characterization of Avian Metapneumovirus Subtype B from Chickens in Greece. Avian Diseases, 61(3), 409–413. doi:10.1637/11631-032017-CaseR. PMID: 28957007.

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