STUDY THE EFFECT OF MYCOPLASMA CONTAMINATION OF EGGS USED IN VIRUS TITRATION AND EFFICACY OF SOME LIVE ATTENUATED POULTRY VIRAL VACCINES

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ABSTRACT

Objective: The study of Mycoplasma gallisepticum (MG) infection is needed, not only to understand the disease process but also to understand the interference with the evaluation of some live viral poultry vaccines. This study aims to investigate the titration and potency of some live attenuated poultry viral vaccines; Newcastle disease, infectious bronchitis, infectious bursal disease, and Reo in both specific pathogen-free (SPF) embryonated chicken eggs (ECEs) and chickens.

Methods: Titration of live attenuated viral poultry vaccines in ECEs was carried out by dividing the inoculated eggs into four groups; the pre-, simultaneously-, post-, and non-MG contaminated. MG effect on the potency test was carried out using seventeen groups of SPF chickens (25 chicken/group) placed into separate isolators. Each live attenuated viral poultry vaccine was inoculated into 4 groups.

Results: The highest titer of these vaccines that appeared in MG pre-contaminated ECEs were 10^11, 10^7, 10^9, and 10^3, respectively. The lowest vaccine titers that appeared in non-MG contaminated ECEs were 10^7, 10^9, 10^4, and 10^6, respectively. Although the potency of these previous vaccines indicated that the highest antibodies titer that appeared in MG pre-infected vaccinated chickens were 7.5 log_10 enzyme-linked immunosorbent assay unit (EU), and 42 EU, respectively; the lowest antibodies titer that appeared in non-MG infected vaccinated chickens were 6.5 log_10 EU, 17 EU, and 10 EU, respectively.

Conclusion: The present study findings underline the importance of using Mycoplasma-free eggs or chicken for the production of virus vaccines.

Keywords: Mycoplasma gallisepticum, Newcastle disease virus, Infectious bronchitis virus, Infectious bursal disease virus, Reo virus, Chicken, Specific pathogen-free eggs.

INTRODUCTION

One of the most important mycoplasmas isolated from domestic avian species is Mycoplasma gallisepticum (MG) [1]. MG causes chronic respiratory disease of chickens and infectious sinusitis in turkeys, resulting in economic losses. It is transmitted through the egg and is well-known for its interactions with other infectious agents [1].

Mycoplasmas, particularly species of the genera Mycoplasma, are frequent contaminants of vaccine substrates, that is, continuous cell lines, and less frequently, animal-derived tissues and primary cell cultures [2]. Mycoplasma may affect the cell-mediated immune system by inducing either suppression or stimulation of B and T lymphocytes and inducing cytokines [3].

Newcastle disease (ND), infectious bronchitis (IB), infectious bursal disease (IBD), and Reovirus strains can be categorized as velogenic (highly virulent) and mesogenic (intermediate virulence) [4]. These diseases were reported as the most important viral diseases of poultry in the world including developing countries [5-7]. ND virus strains are spread rapidly between bird flocks so vaccination is necessary and is routinely performed using live-virus vaccines against ND virus (lentogenic strain LaSota), which is commonly produced using embryonated chicken eggs (ECEs) [4]. So that, mixed infections involving Mycoplasma, viruses, and bacteria are well-recognized in chickens. Synergism has been demonstrated between MG and the viruses of ND and IB [8]. Both MG and ND virus are able to successfully propagate in ECEs. The chicken embryo could represent a model system for the analysis of consecutive Mycoplasma synoviae (MS) -lentogenic-ND virus co-infection that may occur after ND virus infection (e.g., vaccination) of MS-infected host. Co-infections with MS and different avian viruses, concerning clinical signs, have been described by Kleven [1] and Landman and Feberwee [9].

Avian IB infection may precipitate latent Mycoplasma infection. Many studies [11,12] emphasize the importance of using Mycoplasma-free eggs for the production of virus vaccines. There was at least one hundred-fold increase in the multiplication of MG in eggs simultaneously or previously infected with avian IB virus (IBV) made by inoculation of eggs in the allantoic sacs of 10-day embryonated hen's eggs [11].

When mildly virulent virus strains infect birds, which have already been infected with Mycoplasma, interactions between the host, Mycoplasma cells, and the virus may result in effects that differ from those which are expected after single pathogen infection [13]. Thus, severe inflammatory reactions and synergistic pathogen interactions can occur after vaccinations with live virus vaccines [13].

The current study aimed to clarify the effect of MG contamination in specific pathogen-free (SPF) eggs or infected chickens on virus titration and efficacy of some live attenuated viral poultry vaccines.
METHODS

SPF eggs
Fertile SPF eggs were obtained from the national project for production of SPF eggs (Nile SPF eggs), Koom Oshiem, Fayoum, Egypt. Furthermore, they were hatched in the Central Laboratory for Evaluation of Veterinary Biologics (CLEVB), Cairo, Egypt.

Chickens
Four hundred forty-five of 14 days SPF chickens were obtained from the national project for production of SPF eggs (Nile SPF eggs), Koom Oshiem, Fayoum, Egypt.

MG strain and its growth media
MG strain was obtained from Animal Health Institute in Dokki, Mycoplasma department. This strain was propagated according to Naylor et al. [14]. The number of colony forming units (CFU) was determined by the standard procedures of Rodwell and Whitcomb [15] to obtain 10^6 CFU of MG strain to be inoculated in different groups of ECEs and chickens.

MG antigen and antiserum
MG-colored antigens and chicken anti-MG sera were used for serum plate agglutination test (SPA). They were kindly obtained from Mycoplasma Department, Animal Health Research Institute, Dokki - Giza, Egypt.

Live attenuated viral poultry vaccines
Four types of live attenuated viral poultry vaccines (ND, IB, IBD, and Reo virus vaccines) were obtained from CLEVB. The vaccines were kept at refrigerator (4-8°C) till use.

Virus strains
Newcastle disease virus (NDV) strain genotype 7 accession No.KM288609, standard local viral strains of IB, IBD and Reo were obtained from Viral Strain Bank of CLEVB that used for challenge tests.

Experimental design
Titration of ND, IB, IBD, and reo live attenuated viral poultry vaccines in ECEs
The titration of ND vaccine
Ten-fold serial dilutions (10^−1-10^−12) of the vaccine were prepared in sterile saline with antibiotic. Of each dilution, 0.2 ml was inoculated in 20 eggs via allantoic cavity per egg (in 9- to 11-day-old embryonated SPF chicken eggs according to Senne [16]). The inoculated eggs of each dilution were divided into four groups (pre-, simultaneously-, post-, and non-MG contaminated) as follow:

Group A (pre-MG contaminated): Five eggs inoculated with 0.2 ml of 5×10^6 CFU of MG strain first at 7-day-old and then at 9-day-old inoculated with NDV via allantoic cavity.

Group B (simultaneously-MG contaminated): Five eggs inoculated simultaneously by 0.2 ml of 5×10^6 CFU of MG strain with the vaccine.

Group C (post-MG contaminated): Five eggs inoculated with 0.2 ml of each dilution of NDV first at 9-day-old and then 0.2 ml of 5×10^6 CFU of MG strain via allantoic cavity.

Group D (non-MG contaminated): Five eggs per each dilution inoculated with 0.2 ml of NDV at 9-day-old via allantoic cavity.

The inoculated eggs were incubated at 37°C with 60% humidity and candled daily for 6 days later. Dead embryos (within 24 hrs) were discarded and considered non-specific. Then, any dead embryos after these 24 hrs were removed and kept in refrigerator at 4°C and recorded daily for calculation of the titer of the vaccine at the end of the sixth day and all remaining eggs were put in refrigerator. Haemagglutination test (HI) was applied on the allantoic fluid of the inoculated eggs, and the positives were counted, and the titer was calculated [17].

The titration of IB vaccine
All previous steps of the previous experiment were applied with the exception that the vaccine of IB dilutions was from 10^−1 to 10^−6. In addition, the titer of vaccine was calculated according to criteria of embryo [17].

The titration of IBD and Reo vaccine
All previous steps used in the titration of IB vaccine were applied.

MG effect on the potency test for the evaluation of live attenuated viral poultry vaccines
Seventeen groups of SPF chickens (25 chicken/group) were placed into separate隔离ors. Each live attenuated viral poultry vaccine was inoculated into 4 groups as follows: Group 1 inoculated with MG at 14 days old SPF chickens 0.25 ml intranasally (i.n.) and 0.25 ml s.c. route of 10^6 CFU of prepared MG. The inoculated eggs were incubated at 37°C in CO_2 incubator for 48-72 hrs or until the color of the broth changed. After that loopful of the culture broth was plated on PPLO agar plate [19], incubate the plate at 37°C in CO_2 incubator for 5-7 days. The plate was examined daily under stereo-zone microscope for identification of fried egg colonies specific to Mycoplasma.

MG re-isolation and identification
Re-isolation
The re-isolation of MG was done by taken egg yolk samples from the contaminated ECEs (pre-, simultaneously- and post-contamination). Swabs were taken from the eyes, nostrils, and pharyngeal of infected chickens (pre-, simultaneously- and post-infection). The egg yolk samples and different swabs were dipped into pleuropneumonia-like organisms (PPLO) broth medium (Difco Laboratories) to enhance isolation of Mycoplasma [18]. The cultured broth was incubated at 37°C in CO_2 incubator for 48-72 hrs or until the color of the broth changed. After that loopful of the culture broth was plated on PPLO agar plate [19], incubate the plate at 37°C in CO_2 incubator for 5-7 days. The plate was examined daily under stereo-zone microscope for identification of fried egg colonies specific to Mycoplasma.

MG identification
Egg yolk and serum samples [20] were collected randomly from contaminated ECEs and infected chickens in pre, simultaneously and post groups, then egg yolk and serum samples were prepared for SPA test to ensure the presence of MG.

Egg yolk and blood samples collection and preparation
The egg yolk and serum preparation were done according to Office International Des Epizooties (OIE) [21].

Haemagglutination inhibition (HI) test
It was applied for calculating the titer of ND virus in the collected allantoic fluid of all inoculated groups of ECEs according to OIE [21]. All serum samples from Groups 1-4 were collected for estimation of antibody titers against ND virus using HI test [22]. The test was performed according to the procedure of OIE (2002). In brief, two-fold serial dilution of 25 µl serum was made with phosphate-buffered saline (PBS) in V-bottomed microtiter plates (Nunc) up to 10^6 well. 25 µl of 4 hemagglutinating (HA) units of virus or antigen was added up to 11^6 well. The plates were kept at room temperature for more than 30 minutes to facilitate antigen-antibody reaction. Then, 50 µl of 1% (v/v) chicken red blood cells (RBCs) suspension was added to each well. The 11^6 well contains antigen and RBCs as the positive control, and the 12^6 well contains only RBCs as the negative control. After gentle mixing, the RBCs were allowed to settle at room temperature for 40 minutes, and agglutination was assessed by tilting the plates. The samples showing peculiar central button shaped settling of RBCs were
recorded as positive and maximum dilution of each sample causing HI was considered the end point, which was used to estimate the HI titer. The HI titer of each serum sample was expressed as reciprocal of the serum dilution.

**Enzyme-linked immunosorbent assay (ELISA)**

It was conducted as described by Jordan et al. [23]. IBV AB ELISA kit was used for estimation of antibodies titers against IBV in serum samples collected from Groups 5 to 8.

AB ELISA kit calculated the antibodies titer against IBD virus (IBDV) in serum samples collected from Groups 9 to 12. While ReoV AB ELISA kit was used for estimation of antibodies titers against Reovirus in serum samples collected from Groups 13 to 16.

**Challenge test**

At 28 days post-vaccination, the vaccinated and control groups were challenged then kept under observation for 10-14 days post-challenge.

Chicken groups vaccinated by ND vaccine were challenged intramuscularly with a local virulent strain of ND virus containing at least 10<sup>0.1</sup> embryo lethal doses (ELD<sub>50</sub>/bird [24]). The highest titer of IB antibodies by ELISA test on collected serum at the 28 days post inoculation was appeared in Mycoplasma pre-infected group (36 ELISA unit [EU]) while the lowest titers were recorded in NDV inoculated alone group (6.5 log). The protection percentage against NDV revealed that the Mycoplasma pre-infected, ND challenged group was more protected group (100%) than simultaneously and post Mycoplasma infected group (95%) and the non-infected ND challenged group recorded lowest protection percent (90%). Fig. 4 illustrated the postmortem lesions showed the presence of petechial hemorrhage all over mucus membranes in challenged unvaccinated group that are characteristic to NDV. Fig. 5 represents SPF chickens showed nervous manifestations after challenged unvaccinated group with NDV.

The results in Table 3 recorded that the highest titer of IBD antibodies by ELISA test on collected serum at the 28 days post inoculation was in Mycoplasma pre-infected group (42 EU) while the lowest titers were recorded in IBDV inoculated without MG-infected group (17 EU). The protection percentage against IBDV revealed that all MG-infected IBDV challenged groups showed higher protection percentage (100%) than the MG non-infected IBDV challenged group that recorded (90%). The highest titer of Reo antibodies by ELISA test on collected serum at the 28 days post inoculation was in Mycoplasma pre-infected group (14 EU) while the lowest titers were recorded in Reo inoculated without MG-infected group (10 EU). The highest protection percentage against Reovirus was in Reo challenged MG (Pre- and simultaneously infected group (100%) while the MG non-infected Reo challenged group was the lowest protected group (90%).

**RESULTS**

**Mycoplasma re-isolation from ECEs and chicken swabs**

All samples taken from egg yolk and swabs, from both contaminated eggs and infected chickens of pre-, simultaneous- and post-MG groups, were gave fried egg colonies, by examination under stereo-zone microscope. While samples obtained from non-MG contaminated and infected groups were not gave any colonies.

**Mycoplasma identification by SPA test**

All collected samples of egg yolk and serum samples from both contaminated eggs and infected chickens of pre-, simultaneous- and post-MG groups were gave clear agglutination by SPA test while there was any agglutination in samples obtained from non-MG contaminated and infected groups.

**Titration of live attenuated vaccines in SPF-ECEs among MG-infected groups**

Table 1 indicated that the highest titer of ND virus by HA test, applied on allantoic fluid of ECEs, was appeared in Mycoplasma pre-contaminated group (10<sup>3</sup>). The lowest titers were recorded in Mycoplasma non-infected group (10<sup>0</sup>). The highest titer of IBV was appeared in Mycoplasma pre-contaminated group (10<sup>7.5</sup>), whereas the lowest titers were recorded in IBV MG non-infected group (10<sup>0</sup>).

The highest titer of IBDV was appeared in Mycoplasma pre-contaminated group (10<sup>6</sup>), whereas the lowest titers were recorded in IBDV inoculated without MG contamination group (10<sup>0</sup>). Finally, the highest titer of Reo virus was appeared in Mycoplasma pre-contaminated group (10<sup>7.5</sup>), whereas the lowest titers were recorded in Reo virus inoculated without MG contamination group (10<sup>0</sup>).

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bursa of fabricius with yellowish peribursal oedema in unvaccinated chicken challenged with IBDV. Fig. 8c shows hemorrhages on thigh muscles in unvaccinated chicken challenged with IBDV. Fig. 9 of SPF chicken shows severe diarrhea in unvaccinated group challenged with IBDV. In Fig. 10, the red rings showed malabsorption syndrome (characteristics to Reo disease) in proventriculus of unvaccinated chicken challenged with Reo virus. Fig. 11 illustrates the SPF chicken showed limpness associated with tenosynovitis in challenged unvaccinated group with Reo virus.

**DISCUSSION**

The current study investigated the effect of MG contamination to fertile SPF eggs and chickens on the titration and potency of some live attenuated poultry viral vaccines (ND, IB, IBD and Reo virus) was carried out. ECEs are among the most useful and available forms of living animal tissue for the isolation and identification of viruses, for titrating viruses, and for quantity cultivation in the production of viral vaccines [25]. Therefore, inoculated ECEs with MG in three ways, pre-, simultaneously- and post- of the inoculated vaccines were chosen, in the current study. It is clear that MG inoculations enhance the replication of viruses and increase their titers in ECEs. These results are supported with Bolha et al. [26] who reported that when mildly virulent virus strains infect birds

![Fig. 2: (a) Normal embryo liver, (b) Yellowish-white discoloration of embryo liver indicated the characteristic lesion of infectious bursal disease virus](image)

![Fig. 3: (a) Normal embryo, (b) Strawberry like embryos due to Reo virus inoculation](image)

![Fig. 4: Postmortem lesions showed the presence of petechial haemorrhage all over mucous membranes of the proventriculus in challenged unvaccinated group which are characteristic to Newcastle disease virus](image)

![Fig. 5: Specific pathogen-free chickens showed nervous manifestations after challenged unvaccinated group with Newcastle disease virus](image)

![Fig. 6: Arrows refer to bronchioles containing caseous plugs in infectious bronchitis virus challenged unvaccinated chicken which characteristics to infectious bronchitis disease](image)

![Fig. 7: Specific pathogen-free chicken show gasping of air in unvaccinated group challenged with infectious bronchitis virus](image)
that have already been infected with Mycoplasma, interactions between the host, Mycoplasma cells, and virus may result in effects that differ from those which are expected after single pathogen infection.

The present study depended on the fact that Mycoplasma can be transmitted vertically to eggs. MG seemed to play a role in increasing the titer of inoculated viruses (ND, IB, IBD, and Reo) in eggs that might be due to changing the pH of the egg content by the metabolite products of Mycoplasma obtained from its multiplication inside the egg [12]. MG can multiply on the chorioallantoic or allantoic sac despite its inoculation intra-yolk [27]. The two routes of egg inoculation (intra yolk and intra allantoic) for inoculation of ECEs were used in the current study.

Mixed infections involving Mycoplasma, viruses, and bacteria are well recognized in chickens. Synergism has been demonstrated between MG and the viruses of ND and IB and Escherichia coli [8].

The current study results showed the highest titer of ND virus by HA test, applied on allantoic fluid of ECEs, was appeared in Mycoplasma pre-contaminated group while the lowest titers were recorded in ND inoculated without MG contamination group. Co-infections were more severe than single infections [28]. Both MG and ND virus can successfully propagate in ECEs. The chicken embryo could represent a model system for the analysis of consecutive MS-lentogenic ND virus co-infection that may occur after the virus infection (e.g., vaccination) as concluded by Bolha et al. [26].

### Table 1: Titration of live attenuated vaccines in SPF ECEs

<table>
<thead>
<tr>
<th>Infected groups</th>
<th>Vaccine</th>
<th>ND</th>
<th>IB</th>
<th>IBD</th>
<th>Reo</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pre-MG</td>
<td>$10^{11}$</td>
<td>$10^{7.5}$</td>
<td>$10^{7.0}$</td>
<td>$10^{7.5}$</td>
<td></td>
</tr>
<tr>
<td>Simultaneous-MG</td>
<td>$10^{10}$</td>
<td>$10^{7.0}$</td>
<td>$10^{7.2}$</td>
<td>$10^{7.0}$</td>
<td></td>
</tr>
<tr>
<td>Post-MG</td>
<td>$10^{10}$</td>
<td>$10^{4.0}$</td>
<td>$10^{7.0}$</td>
<td>$10^{6.0}$</td>
<td></td>
</tr>
<tr>
<td>Non-MG</td>
<td>$10^{6.0}$</td>
<td>$10^{4.0}$</td>
<td>$10^{7.2}$</td>
<td>$10^{6.0}$</td>
<td></td>
</tr>
</tbody>
</table>

SPF: Specific pathogen-free, ECEs: Embryonated chicken eggs, MG: Mycoplasma gallisepticum, IBD: Infectious bursal disease, IB: Infectious bronchitis

### Table 2: Antibody titers tested by HI test and protection percent among the vaccinated groups

<table>
<thead>
<tr>
<th>Groups</th>
<th>14th PV</th>
<th>21st PV</th>
<th>28th PV</th>
<th>Protection percent</th>
<th>t-test</th>
<th>Sig. (two-tailed)</th>
<th>SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1</td>
<td>$5.4 \text{ log}_{10}$</td>
<td>$6.5 \text{ log}_{10}$</td>
<td>$7.5 \text{ log}_{10}$</td>
<td>100</td>
<td>0.035</td>
<td>7.0000</td>
<td></td>
</tr>
<tr>
<td>G2</td>
<td>$4.9 \text{ log}_{10}$</td>
<td>$6.2 \text{ log}_{10}$</td>
<td>$7.2 \text{ log}_{10}$</td>
<td>95</td>
<td>0.012</td>
<td>1.17898</td>
<td></td>
</tr>
<tr>
<td>G3</td>
<td>$4.5 \text{ log}_{10}$</td>
<td>$5.8 \text{ log}_{10}$</td>
<td>$6.8 \text{ log}_{10}$</td>
<td>95</td>
<td>0.013</td>
<td>1.153256</td>
<td></td>
</tr>
<tr>
<td>G4</td>
<td>$4.2 \text{ log}_{10}$</td>
<td>$5.5 \text{ log}_{10}$</td>
<td>$6.5 \text{ log}_{10}$</td>
<td>90</td>
<td>0.015</td>
<td>1.153256</td>
<td></td>
</tr>
</tbody>
</table>

PV: Post-vaccination, G1: The infected with MG then vaccinated with Newcastle disease virus vaccine (NDVV), G2: The vaccinated with NDVV and simultaneously infected with MG, G3: The vaccinated with then infected with MG, G4: The vaccinated only with NDVV, Sig.: Significance (0.01-0.05), SD: Standard deviation, HI: Haemagglutination test, MG: Mycoplasma gallisepticum

### Table 3: Antibody titers tested by ELISA and protection percent among the vaccinated groups

<table>
<thead>
<tr>
<th>Groups</th>
<th>14th PV</th>
<th>21st PV</th>
<th>28th PV</th>
<th>Protection percent*</th>
<th>t-test</th>
<th>Sig. (two-tailed)</th>
<th>SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>G5</td>
<td>25 EU</td>
<td>35 EU</td>
<td>36 EU</td>
<td>100</td>
<td>0.012</td>
<td>6.082763</td>
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<tr>
<td>G6</td>
<td>20 EU</td>
<td>15 EU</td>
<td>22 EU</td>
<td>95</td>
<td>0.046</td>
<td>6.027714</td>
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</tr>
<tr>
<td>G7</td>
<td>8 EU</td>
<td>11 EU</td>
<td>14 EU</td>
<td>95</td>
<td>0.024</td>
<td>3.000000</td>
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</tr>
<tr>
<td>G8</td>
<td>7 EU</td>
<td>10 EU</td>
<td>12 EU</td>
<td>90</td>
<td>0.022</td>
<td>2.516611</td>
<td></td>
</tr>
<tr>
<td>G9</td>
<td>18 EU</td>
<td>30 EU</td>
<td>42 EU</td>
<td>100</td>
<td>0.049</td>
<td>12.000000</td>
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<tr>
<td>G10</td>
<td>10 EU</td>
<td>19 EU</td>
<td>25 EU</td>
<td>100</td>
<td>0.054</td>
<td>7.549834</td>
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<tr>
<td>G11</td>
<td>9 EU</td>
<td>16 EU</td>
<td>20 EU</td>
<td>100</td>
<td>0.043</td>
<td>5.567764</td>
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<tr>
<td>G12</td>
<td>7 EU</td>
<td>12 EU</td>
<td>17EU</td>
<td>90</td>
<td>0.053</td>
<td>4.041452</td>
<td></td>
</tr>
<tr>
<td>G13</td>
<td>6 EU</td>
<td>11 EU</td>
<td>14 EU</td>
<td>100</td>
<td>0.047</td>
<td>3.055050</td>
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<tr>
<td>G14</td>
<td>6 EU</td>
<td>10 EU</td>
<td>12 EU</td>
<td>100</td>
<td>0.034</td>
<td>3.055050</td>
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</tr>
<tr>
<td>G15</td>
<td>6 EU</td>
<td>10 EU</td>
<td>11 EU</td>
<td>95</td>
<td>0.028</td>
<td>3.055050</td>
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<tr>
<td>G16</td>
<td>4 EU</td>
<td>8 EU</td>
<td>10 EU</td>
<td>90</td>
<td>0.053</td>
<td>3.055050</td>
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</tr>
</tbody>
</table>

PV: Post-vaccination, G5-8: Vaccinated with IB VV, G9-12: Vaccinated with IBD VV, G13-16: Vaccinated with Reo VV, G5-16: They were vaccinated with virus vaccine as mentioned before in Groups 1-4; *: 6th days post challenge, EU: ELISA unit, Sig.: Significance (0.01-0.05), SD: Standard deviation, ELIZA: Enzyme-linked immunosorbent assay

**Fig. 8:** (a) Kidneys of unvaccinated chicken affected by a severe urate diathesis due to challenge with infectious bursal disease virus (IBDV), (b) enlarged bursa of fabricius with yellowish peribursal oedema in unvaccinated chicken challenged with IBDV, (c) hemorrhages on thigh muscles in unvaccinated chicken challenged with infectious bursal disease virus.
Severity of clinical respiratory symptoms associated with MS, which exacerbated by co-infection with various respiratory viruses, such as ND and IB were reported Landman and Feberwee [9]. This is compatible with the present results, which revealed the highest titer of ND antibodies by HI test at the 28 days post-inoculation in Mycoplasma pre-infected group while the lowest titers were recorded in NDV inoculated alone.

The effect of MG on potency of IBV by was investigated by three ways (pre-, simultaneously- and post-vaccination). Inoculation the IB vaccine

this following a part of the model of Landman and Feberwee [9], as clinical symptoms due to IBV were expected to peak. Hopkins and Yoder [29] suggested that chicken-passage mild IB vaccine virus markedly increased the incidence of airsacculitis compared with non-passage vaccine virus so that chickens vaccinated with IB vaccine have higher AB titer and more protected against the IB disease alone. The effect of the previous infection of 8-week-old chicken with IBDV on their susceptibility to MS and Mycoplasma gallinaceum (MGN) then the antibody titers against MS and IBDV were detected by using HI and immune diffusion test [30]. Concomitant infection of MGn acts synergistically with MS and that previous exposure to IBDV increases the susceptibility of the synovial tissue to MS infection.

It was recorded that, birds infected with IBDV, that were later inoculated with MS (day 14), NDV (days 14 and 28), experienced an increased incidence and greater severity of airsacculitis than did chickens which were not exposed to IBDV [31]. Sarueng et al. [32] stated that severity of the respiratory disease resulting from infection by IBV is increased when there is co-infection with other infectious agents such as E. coli, MG and/or MS occurs. The present study recorded the protection percentage against IBDV revealing that all MG-infected IBD challenged groups showed higher protection percentage than the MG non-infected IBD challenged group. IBV has been shown to interfere with the serological response to MS [31]. Reproduced synovitis in chickens was in groups challenged with both IBDV and MS, but not with MS alone [30].

Many previous studies [33-35] helped support the overall importance of the present results. Reo viruses have been shown to interact with MS in the production of synovitis. This virus neutralizing antibodies were detected in group infected with Reo virus together with the Mycoplasma, at 3 weeks and persisted until the end of the experiment at 15 weeks.

Fig. 9: Specific pathogen-free chicken showed severe diarrhea in unvaccinated group challenged with infectious bursal disease virus

Fig. 10: The red rings showed malabsorption syndrome (characteristics to Reo disease) in proventriculus of unvaccinated chicken challenged with Reo virus

Fig. 11: Specific pathogen-free chicken show limpness associated with tenosynovitis in challenged unvaccinated group with Reo virus
CONCLUSION
The aim of this study was trying to explain the adverse effect of Mycoplasma on the titration and potency of viral vaccines, and the results revealed that MG infection could lead to false evaluation of these vaccines. The interaction between Mycoplasmas and viruses has been considered to be important not only in natural infection but also in vaccination process failure.

Compliance with ethics requirements
Chickens care as well as experimental protocols were in compliance with guidelines of ethical standards released by CLEBV on animal care and use. All efforts were made to minimize the numbers of animals and their suffering in this study following the guidelines on laboratory animal care and use.

REFERENCES