Route of infectious bronchitis virus vaccination determines the type and magnitude of immune responses in table egg laying hens

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Abstract
Chicken immune responses to infectious bronchitis virus (IBV) vaccination can depend on route of administration, vaccine strain and bird age. Typically for layer chickens, IBV vaccinations are administered by spray in the hatchery at day-old and boosted at intervals with live vaccines via drinking water (DW). Knowledge of live attenuated IBV vaccine virus kinetics and the immune response in egg-laying hens is exceptionally limited. Here, we demonstrated dissemination of vaccine viruses and differences in hen innate, mucosal, cellular and humoral immune responses following vaccination with Massachusetts or 793B strains, administered by DW or oculonasal (ON) routes. Detection of IBV in the Mass-vaccinated groups was greater during early time-points, however, 793B was detected more frequently at later timepoints. Viral RNA loads in the Harderian gland and turbinate tissues were significantly higher for ON-Mass compared to all other vaccinated groups. Lachrymal fluid IgY levels were significantly greater than the control at 14 days post-vaccination (dpv) for both vaccine serotypes, and IgA mRNA levels were significantly greater in ON-vaccinated groups compared to DW-vaccinated groups, demonstrating robust mucosal immune responses. Cell mediated immune gene transcripts (CD8-α and CD8-β) were up-regulated in turbinate and trachea tissues. For both vaccines, dissemination and vaccine virus clearance was slower when given by DW compared to the ON route. For ON administration, both vaccines induced comparable levels of mucosal immunity. The Mass vaccine induced cellular immunity to similar levels regardless of vaccination method. When given either by ON or DW, 793B vaccination induced significantly higher levels of humoral immunity.

Keywords: Avian coronavirus, infectious bronchitis virus, vaccination, layer chicken, gene transcription, innate, mucosal, cellular and humoral immunity

Introduction
Commercial laying hens infected with infectious bronchitis virus (IBV) can exhibit both respiratory [1] and reproductive system [2, 3] complications. Oviduct infection can lead to reduced laying production and lower egg quality [4–7], resulting in economic losses for poultry farmers. In the United Kingdom (UK), infectious bronchitis (IB) affected approximately 22.5 million chickens at an estimated total cost of £12.6 million from egg production losses between 2005 and 2019 [8, 9]. Egg quality problems include reduced shell thickness, a mottled and discoloured shell, and watery albumin [10]. Prevention of infectious bronchitis (IB) in egg-laying chickens is mainly through the use of inactivated and live vaccines during the rearing period, and the vaccine strains administered are dependent on the prevalent endemic strains [11].

Live attenuated IBV vaccines are used to induce local immunity, which is important as a part of an effective
host mucosal immune response [12–14]. Live vaccines are administered to day-old chicks and then at intervals, to achieve early and continuous mucosal protection. At 4–6 weeks before the onset of lay, each bird also receives an inactivated vaccine to boost humoral antibody levels [15–17]. The gold standard approach for IBV vaccination is oculonasal (ON) [18, 19], however, this route is not practical for vaccine administration to a large number of birds. For this reason, in the field, live vaccines are typically administered via drinking water [20, 21], coarse spray [22] or aerosols at day-old or within the first week of age [19]. Though publications are available on the assessment of live vaccines in young birds, to date, there has not been any report on the impact of administering monovalent vaccines in older layer hens.

The Massachusetts (Mass) serotype vaccine was the first to be produced, and strains belonging to Mass (such as H120 and Ma5) are used throughout the world for laying hens [23–25]. Another serotype that is widely distributed in Europe, Asia, and other parts of the world is 793B [26]. Strains identified as the 793B serotype, including 793B, CR88 and 1/96, emerged in the 1990s and have been associated with welfare and economic problems in flocks [27, 28].

The innate, mucosal, cellular and immune responses are vital for protection against virulent IBVs. In terms of early immunity following IBV vaccination, previous work has outlined transcriptional expression of toll-like receptors (TLRs), melanoma differentiation-associated protein 5 (MDA5), type I interferons, and pro-inflammatory cytokines [14, 29].

While previous studies have investigated specific-pathogen-free (SPF) or broiler chickens, to date, there have been no publications investigating host immune responses in the respiratory or head-associated lymphoid tissues (HALT) of egg-laying hens following IBV vaccination. We report vaccine virus distribution in a number of tissues (HALT) of egg-laying hens following IBV vaccination by titrating in tracheal organ cultures (TOCs) to determine the mean tissue culture infectious dose (TCID50); Mass (10^4.75 TCID50/mL) and 793B (10^4.45 TCID50/mL) [30].

**Materials and methods**

**Ethical statement**

All experimental procedures were performed according to the UK legislation governing experimental animals under the project license P8E4FC2C9. Experimental procedures were approved by the University of Liverpool’s ethical review process.

**Layer chickens**

Seventy-two 41-week-old light brown Lohman layer chickens were obtained from a commercial UK farm and kept up to three weeks under strict biosecurity measures in the University of Liverpool. When these birds were in the rearing farm, the flock received live IBV Massachusetts vaccination at day-old, followed by a live 793B vaccine at day-14. Subsequently, an inactivated IBV M41-D274 was given at 15 weeks of age. No IB or other vaccines were applied during the laying period. For this study, birds were transported to University of Liverpool experimental house one week prior to vaccination to allow for acclimatisation to conditions. Hens were reared on deep litter with nest boxes, and bird-friendly toys, antibiotic-free water and feed were provided ad libitum.

**Attenuated live IBV vaccines**

Two commercial live IBV vaccines, Massachusetts (Mass) serotype (H120, Boehringer Ingelheim Animal Health Limited, UK) and 793B serotype (4/91, MSD Animal Health, UK) were prepared according to manufacturer’s instructions. One vial of each vaccine was thoroughly mixed in 100 mL of chilled distilled water and kept on crushed ice before use. Each hen received 100 μL of the vaccine or sterile distilled water (SDW) via the oculonasal (ON) route. For the drinking water (DW) group, to ensure uptake of full vaccine dosage, the vaccine was administered orally. The dosage was maintained as administered in commercial farms. Vaccine strains were titrated in tracheal organ cultures (TOCs) to determine an absence of IBV by RT-PCR. Upon arrival, seventy two 41-week-old layer chickens were divided into six groups (n = 12 per group); (A1) DW-unvaccinated control, (A2) ON-unvaccinated control, (B1) DW-Mass, (B2) ON-Mass, (C1) DW-793B and (C2) ON-793B. Prior to vaccination, egg production and clinical signs were monitored for one week. At 42-weeks-old, OP swabs were collected from 10 birds in each group to confirm absence of IBV, avian metapneumovirus (aMPV), Mycoplasma synoviae and Mycoplasma gallisepticum by PCR.

Experimental design

Oropharyngeal (OP) swabs were collected one week before the hens arrived at the University (40 weeks of age) to confirm an absence of IBV by RT-PCR. Upon arrival, seventy two 41-week-old layer chickens were divided into six groups (n = 12 per group); (A1) DW-unvaccinated control, (A2) ON-unvaccinated control, (B1) DW-Mass, (B2) ON-Mass, (C1) DW-793B and (C2) ON-793B. Prior to vaccination, egg production and clinical signs were monitored for one week. At 42-weeks-old, OP swabs were collected from 10 birds in each group to confirm absence of IBV, avian metapneumovirus (aMPV), Mycoplasma synoviae and Mycoplasma gallisepticum by PCR. Groups A1 and A2 were sham-inoculated with 0.1 mL of vaccine-free SDW. Groups B1, B2, C1 and C2 were vaccinated with 0.1 mL of live Mass (10^3.75 TCID50/mL) or 793B (10^4.45 TCID50/mL). Following vaccination, OP and cloacal (CL) swabs were collected from five chickens at 1, 3, 5, 7, and 14 dpi for virus detection and quantification by quantitative real-time RT-PCR (qRT-PCR). At 7 and 14 dpi, lachrymal fluid and blood were collected from five birds in each group to assay for anti-IBV antibodies by indirect ELISA. Three birds from each group
were humanely euthanized at 1, 3, 5, and 14 dpv. The Harderian gland, turbinate, choanal cleft, trachea, caecal tonsil and kidneys were collected and stored at −20 °C in RNALater™ (Qiagen, Crawley, UK) for quantification of viral load and/or host gene expression analysis by qRT-PCR.

**Humoral immune responses by indirect ELISA**

Sera were analysed using a commercial IBV ELISA kit (IDEXX, Westbrook, Maine, USA) to determine anti-IBV antibodies according to the manufacturer’s guidelines. Antibody titres were determined by converting the sample/positive ratio according to a formula provided by the manufacturer, with a positive ELISA titre cut-off determined as 396.

**Mucosal immune responses by indirect ELISA**

Lachrymal fluid was assayed for IBV-specific IgA and IgY using an indirect monoclonal ELISA [31–33]. Each well of a flat bottom 96-well microplate (STARLAB®, UK) was coated with 100 µL of purified 2.5 µg/mL IBV M41 antigen in 50 mM sodium carbonate/bicarbonate buffer (pH 9.6). Plates were incubated for 1 h at 37 °C, and then overnight at 4 °C. Wells were blocked with 200 µL phosphate buffer saline (PBS) containing 3% non-fat skimmed milk powder. Lachrymal fluid samples were tested in triplicate at a single dilution of 1:10 in PBS containing 0.05% tween-20 (PBST) (Sigma Aldrich®, Dorset, UK). Mouse monoclonal antibodies against either chicken IgA or IgY (BIO-RAD®, Hertfordshire, UK) were added at a dilution of 1:1000 (50 µL) as the secondary antibody, and incubated for an hour at 37 °C. This was followed by goat anti-mouse IgG horse-radish peroxidase-conjugate (BIO-RAD®) at a dilution of 1:10 000 (50 µL), and 1 hour incubation at 37 °C. Tetramethylbenzidine (TMB) (Sigma Aldrich®) substrate was added to each well (50 µL) and incubated in the dark for 15 min to allow for colour development. The reaction was stopped by adding 50 µL of sodium hydrochloric acid (0.5 M HCL), and plates were analysed at 450 nm. Corrected optical density (COD) values were calculated by deducting the OD values of non-antigen coated (blank) wells for each sample [31, 34].

**Quantification of viral load from swabs and tissues**

Viral RNA was extracted from the swab and tissue samples, using the QIAamp viral RNA mini kit and the RNeasy Mini kit (Qiagen, UK) respectively, according to manufacturer’s instructions. Quantification of viral RNA was carried out by qRT-PCR, using an IBV 3’ untranslated region (UTR) gene-specific primer and probe as previously described [35]. Obtained Ct values were converted to log relative equivalent units (REU) of viral RNA by a standard curve generated from using five ten-fold dilutions of RNA extracted from M41 virus-positive allantoic fluid [36, 37].

**Measurement of host gene transcription**

Extracted RNA was tested by qRT-PCR for expression of pro-inflammatory cytokine IL-6, innate immune pattern recognition receptors (TLR3 and MDA5), interferon beta (IFN-β) [36–38], mucosal immune responses (IgA and IgY) and cellular immune responses (CD8-α and CD8-β) [14, 39, 40]. Each cDNA sample was tested in triplicate using LightCycler 480 SYBR Green I Master mix and gene specific primers (Table 1). For IL-6, TLR3, MDA5 and IFN-β, data was normalized using a relative standard curve method to 18S ribosomal RNA expression [41] and data presented as the log2 fold difference in gene expression of vaccinated against control samples. For IgA, IgY, CD8-α and CD8-β, data was normalized against 18S, and the fold change was calculated using the double delta Ct (ΔΔCt) method. Significant up-regulation or down-regulation was reported when compared with the control group, unless otherwise stated.

**Statistical analysis**

Data were confirmed to be normally distributed and analysed using GraphPad™ Prism version 6.00. Significant differences between groups were analysed using univariate ANOVA, along with the homogeneity of variance test, to confirm statistical differences within the data set, followed by post hoc Tukey’s testing to compare between each group. When groups had p < 0.05 for the homogeneity of variance test, Tamhane’s T2 was applied post hoc instead of Tukey’s. For IgA, IgY, CD8-α and CD8-β gene transcript fold change comparisons, vaccinated groups were compared to the control group using the t test. Differences between groups were considered significant at p < 0.05 unless otherwise stated.

**Results**

**Humoral immunity: IBV-specific ELISA**

At 7 dpv, a significantly higher titre was observed in the DW-793B group (8065 ± 792) in comparison with the control (4970 ± 759) (Figure 1). For DW vaccinated birds at 14 dpv, there was a significantly higher antibody titre in the 793B group (8076 ± 759) compared with the Mass-vaccinated (5465 ± 762) and control (4062 ± 218) groups. Significantly higher antibody titres were observed at 14 dpv in ON vaccinated hens [Mass (8306 ± 636) or 793B (7999 ± 461)] compared to the DW-Mass (5465 ± 762) and control (4062 ± 218) groups (Figure 1). No significant differences were noted with the same groups between both sampling days.
Mucosal immunity: IBV-specific lachrymal IgA and IgY titres

Significantly higher IBV-specific IgA levels compared to the control (0.256 ± 0.05) were only observed in the DW-Mass group (0.744 ± 0.19) at 14 dpv (Figure 2A), or when compared to the DW-Mass group at 7 dpv (0.334 ± 0.09).

In 793B-vaccinated hens (DW and ON) and ON-Mass vaccinated birds, IgY titres were significantly higher at 14 dpv compared to the control group. For DW-vaccinated groups, the average IgY titre in the 793B group (2.07 ± 0.07) was significantly higher in comparison to the Mass group (1.55 ± 0.1) at 7 dpv (Figure 2B).

Mucosal immunity: total mRNA expression of IgA in the turbinate and trachea

**Turbinate**
In both ON-vaccinated groups, there was significant up-regulation in the mRNA expression of IgA at all-time points compared with DW-vaccinated chickens, with the exception of DW-Mass at 1 dpv. In the ON-793B group,
there was significant up-regulation of IgA mRNA expression at 1 dpv compared to the ON-Mass vaccinated group (Figure 3A). All DW-vaccinated groups were significantly down-regulated compared with the control, whereas only the ON-Mass group was down-regulated at 1 dpv, with no other differences noted.

**Trachea**

There was significant up-regulation of IgA mRNA expression in the trachea of both ON-vaccinated groups at 3 and 5 dpv, and only at 1 dpv in the ON-793B vaccinated chickens, compared to DW-vaccinated hens. In the ON-793B group, there was significant up-regulation of IgA mRNA expression at 1 and 3 dpv compared to ON-Mass vaccinated group. At 5 dpv, expression of IgA mRNA was significantly higher in the ON-Mass group compared to the ON-793B vaccinated birds (Figure 3B). All DW groups were significantly down-regulated compared to the control, whereas the ON-793B group was significantly up-regulated at 3 and 5 dpv.

**Mucosal immunity: total mRNA expression of IgY in the turbinate and trachea**

**Turbinate**

IgY mRNA expression at 3 dpv was significantly higher in the ON-793B-vaccinated group in comparison to both Mass-vaccinated groups. Moreover, expression of IgY mRNA was significantly greater at 5 dpv in 793B-DW birds compared with the Mass-DW group (Figure 3C). At 1 and 5 dpv, the ON-Mass and DW-Mass groups were significantly down-regulated compared with the control respectively, whereas both 793B groups were significantly up-regulated at 3 dpv.

**Trachea**

At 3 dpv, IgY transcripts in the DW-Mass group were significantly lower than the ON-793B group. No other statistical differences were noted (Figure 3D). At 3 and 5 dpv, the DW-Mass group was significantly down-regulated compared to the control.

**Viral RNA quantification: swabs and tissues**

**Individual OP swabs**

Viral RNA loads in OP swabs in the control group were below the detection limit. For vaccinated groups, RNA load peaked at 3 dpv for DW-793B birds (3.25 log REU), while the viral RNA load in both ON vaccinated groups peaked at 7 dpv (Mass = 4.34 log REU; 793B = 3.07 log REU) (Figure 4). A significant increase was identified from 5 to 7 dpv in the ON-Mass group, with a significant decrease seen in the DW-793B group from 3 to 14dpv.

**Individual CL swabs**

Viral RNA loads in all CL swabs were below the detection limits in the control group. RNA levels peaked at 3 dpv in the Mass-DW group (1.911 log REU) and 5 dpv in the 793B-DW group (2.499 log REU). There was a significant increase in viral load from 1 to 3 dpv in the Mass-DW group, and a significant reduction from 5 to 14 dpv in the Mass-ON group (Figure 5).
**Harderian gland (HG)**

Viral load was significantly higher at 1 and 3 dpv in the ON-Mass group compared to other groups (Figure 6A). Both DW-vaccinated groups had a significantly higher viral load compared to ON-vaccinated groups at 5 dpv (Figure 6A). In the DW-Mass vaccinated group, viral copies were significantly higher at 5 dpv compared to other sampling days. Both 793B vaccinated groups and the ON-Mass group were negative at 14 dpv.

**Turbinate**

Viral RNA load was significantly higher for all groups at 3 dpv in comparison to the other time points, with the exception of ON-Mass. There were significantly higher viral loads in the ON-Mass-vaccinated birds at 1 dpv, compared to all other groups, and in both DW groups compared to the ON-vaccinated group at 5 dpv (Figure 6B).
Choanal cleft
Viral load was significantly higher at 3 dpv than all the other sampling days in the Mass-vaccinated groups and the ON-793B group. Significantly higher levels were noted at 5 dpv in both DW-vaccinated groups compared to ON-793B birds (Figure 6C).

Trachea
Viral copy numbers were significantly higher at 3 dpv for all groups, compared to other time points, with the exception of DW-Mass, which was significantly higher at 5 dpv. In addition, the ON-Mass birds were significantly lower at 1 and 5 dpv compared to other groups (Figure 6D).

Caecal tonsil
The viral load was significantly higher in the ON-Mass group at 14 dpv compared to all other groups (Figure 6E). No further significant differences were noted.
Kidney
For 793B, both DW and ON groups were IBV-positive at 1 dpv, whereas only the DW-793B group was positive at 3, 5 and 14 dpv. The Mass groups were only IBV positive from 3 dpv, with the Mass-ON group negative at 14 dpv. In addition, the Mass-DW group was significantly higher than the 793B-DW group at 14 dpv (Figure 6F).

Differential mRNA expression: TLR3 and MDA5
**Harderian gland (HG)**
There was significant up-regulation in the mRNA expression of TLR3 in Mass-vaccinated birds via both routes of vaccination at 3 dpv, and at 5 dpv in DW-793B and ON-Mass birds compared to other groups (Figure 7). Expression of MDA5 was significantly up-regulated at 3 and 5 dpv in ON-Mass birds compared to other groups, with the exception of the DW-Mass group at 3 dpv.
Turbinate
Significant down-regulation was noted in TLR3 mRNA expression at 1 and 3 dpv in DW-793B vaccinated chickens, compared to all other groups. This was followed by significant down-regulation in all groups at 5 dpv. There was significant down-regulation of MDA5 transcription at 1 dpv in the DW-793B and Mass-vaccinated birds.

Trachea
There was significant up-regulation of TLR3 mRNA expression at 5 dpv in both Mass-vaccinated and DW-793B groups. Measurement of MDA5 transcription showed significant up-regulation at 5 dpv in Mass-vaccinated chickens via both immunisation routes (Figure 7).

Differential mRNA expression: IFN-β
Harderian gland (HG)
There was significant up-regulation of IFN-β at all-time points in ON-Mass-vaccinated birds, and at 1 and 3 dpv in DW-Mass-vaccinated birds. Significant up-regulation of IFN-β mRNA expression was observed in the DW-793B vaccinated group at 1 dpv, and at 3 dpv in the ON-vaccinated groups (Figure 7).

Turbinate
There was significant down-regulation of IFN-β mRNA expression in both Mass vaccinated groups and in the DW-793B group at 1 dpv, followed by significant down-regulation in all groups at 5 dpv. Significant up-regulation of IFN-β mRNA was observed in the DW-Mass group at 3 dpv.

Trachea
Expression of IFN-β mRNA was significantly up-regulated in both DW vaccinated groups at 3 and 5 dpv and in all vaccinated groups at 5 dpv.

Differential mRNA expression: IL-6
Harderian gland (HG)
There was significant up-regulation of IL-6 mRNA expression at 3 dpv in the 793B-vaccinated groups compared to other groups, with the exception of DW-Mass (Figure 7).

Turbinate
At 3 dpv, IL-6 was significantly up-regulated in the DW-Mass and both 793B-vaccinated groups. At 5 dpv, there was significant down-regulation of IL-6 mRNA expression in the ON-Mass group.

Trachea
Expression of IL-6 mRNA was significantly up-regulated in ON-Mass (5 dpv) and DW-793B (3 dpv) vaccinated chickens (Figure 7).

Cell-mediated immune responses: mRNA expression of CD8-α and CD8-β
Turbinate
Expression of CD8-α and CD8-β mRNA was significantly down-regulated in all vaccinated groups on the majority of sampling days. In contrast, CD8-α mRNA expression was significantly up-regulated at 3 dpv in the DW-Mass group compared to other vaccinated groups (Figure 8). Both 793B-vaccinated groups were significantly down-regulated at 3 dpv compared with the control, whereas the DW-Mass group was down-regulated at 1 and 5 dpv. For CD8-β, mRNA expression was significantly greater in the ON-793B group compared to all other groups at 1 dpv. All groups were significantly down-regulated compared to the control, with the exception of ON-793B at 1 and 3 dpv.

Trachea
Significant up-regulation of CD8-α was noted at 3 dpv in the ON-Mass group compared to both 793B-vaccinated groups (Figure 8). At 5 dpv, the ON-793B group was significantly down-regulated compared to all other vaccinated groups. All groups at 5 dpv, with the exception of ON-793B, were significantly up-regulated compared to the control. Levels of CD8-β transcripts in the DW-Mass, DW-793B and ON-793B group was significantly down-regulated at 5 dpv compared to the ON-Mass group. The ON-793B group was significantly down-regulated compared to the control at 1 and 3 dpv, whereas the DW-793B group was down-regulated at 3 and 5 dpv.

Discussion
Live attenuated IBV vaccines have been widely used in young commercial broiler [45] and rearing layer/breeder chickens [46], and are normally not administered in hens in lay. Despite this, in recent years, the “off-licence” use of live IBV vaccines in egg-laying birds has increased in many countries. In the literature, we found no information on viral loads of infectious bronchitis vaccine viruses or immune responses in head-associated lymphoid and respiratory tissues of commercial egg-laying hens. To address this, we investigated the viral load, innate, mucosal, and cellular immune responses following vaccination via two different routes—oculonasal and drinking water. We compared responses following vaccination via the oral route to those vaccinated via the oculonasal route, as individual dosing of birds through the respiratory route often gave optimal immune responses and protection against a number of IBV challenge viruses [18]. Globally, an
Figure 7 Expression profile of host genes in Harderian gland, turbinate, choanal cleft and trachea tissue following IBV Mass or IBV 793B vaccination of layer chickens. Data is shown as the log₂ fold change when compared to the non-vaccinated group. Different letters indicate significant differences (p < 0.05) in fold change between vaccine strains and vaccination route on the same sample day.
increasing number of producers are administering live IBV vaccines via drinking water [47], as it is convenient for farm workers to prepare and administer the vaccine and as it is less stressful to the birds compared to the intranasal or intraocular routes due to reduced animal handling. Drinking water administration is also presumed to cause less irritation to the respiratory mucosa, and normally has little to no impact on egg production [48].

To diagnose IBV, swab samples (OP and/or CL) and tissues are often examined for the virus, either by RT-PCR, qRT-PCR or virus isolation. In this study, following ON vaccine administration, the viral load of individual OP, CL and tissue samples were examined at intervals by qRT-PCR. From OP swabs, the Mass vaccine was detected for a shorter period (up to 7 dpv) compared to 793B, which persisted up to 14 dpv. In contrast, for the CL swabs, Mass vaccine was detected earlier, but infrequently (1, 5 and 14 dpv), compared to 793B which was only detected at 7–14 dpv. The earlier detection of Mass vaccine, followed by later detection of 793B vaccine viruses, has been reported previously.

Figure 8 Relative mRNA expression of A CD8-α in turbinate tissue, B CD8-α in trachea tissue, C CD8-β in turbinate tissue and D CD8-β in trachea tissue at 1, 3, and 5 dpv. Data are expressed as log₂ mean fold change compared to the control group ± SEM. Expression is calculated based on double delta Ct (ΔΔCt) values. Different letters indicate significant differences (p < 0.05) within time points, determined using one-way ANOVA.
in broiler chickens [49]. Following ON administration, viral loads for both strains were quantified in HG, turbinate, choanal cleft, trachea, caecal tonsil and kidney tissues, however, levels differed depending on tissue type. Significantly higher viral loads of Mass vaccine were often found in the HG, turbinate, caecal tonsil and kidneys. Interestingly, in the trachea, a higher 793B viral load was found at 1 and 5 dpv, with neither strain detected at 14 dpv. It appears that in most tissues, the Mass vaccine was able to establish itself more efficiently than 793B when administered by the ON route. Following DW application, IBV detection and Mass and 793B vaccine viral loads in OP swabs were similar to ON administration. In contrast, for CL swabs, Mass and 793B vaccines viruses were persistently detected at higher levels compared to ON application. This may indicate potential virus replication in the gastrointestinal and/or renal tissues. Interestingly, following DW application, no significant differences in viral loads between each vaccine were seen in the HG, turbinate, choanal cleft or caecal tonsil. For the trachea, the 793B vaccine had a significantly higher viral load at 3 dpv, whereas the Mass vaccine showed significantly higher levels at 5 and 14 dpv. For kidney tissues, both vaccines persisted up to 14 dpv, however, Mass was significantly higher than 793B at 3, 5 and 14 dpv. Overall, it appears that with DW application, the Mass vaccine disseminated slowly to tissues, and persisted at higher viral loads in the trachea and kidneys, when compared to 793B.

All birds in this study received live IB Mass vaccine at day-old, an inactivated IB 793B vaccine at 14-days-old, and a M41 + D274 vaccine at 15–18 weeks of age [15–17]. No other live or killed IBV vaccines were administered after the birds were transferred into an “all-in all-out” free-range layer farm. Following ON vaccination of either strain, significantly higher levels of humoral antibodies were found at 14 dpv in both vaccinated groups (Mass—8306.4 and 793B—7998.8) compared to the control (4062). In contrast to ON vaccination, when the vaccines were given by DW, antibody levels in the 793B group (8075) were significantly higher than the Mass group (5464.6) at 14 dpv. In egg-laying hens, it has been shown that serum anti-IBV antibody levels have a significant influence in providing protection against a drop in egg production and quality [19, 46].

For mucosal immunity, immunoglobulin A (IgA) antibodies play an important role in conferring protection against IBV, and the levels of IBV-specific IgA in lachrymal fluid have been reported before [53–56]. In this study, mucosal immunity was assessed by quantifying IgA and IgY antibodies in lachrymal fluid. No significant increase in IBV-specific IgA was found in any of the ON-vaccinated groups. In contrast, lachrymal anti-IBV IgA levels in the Mass vaccine group given via DW had a significant increase at 14 dpv. Previous studies have reported increased levels of IgA in lachrymal fluid following ocular and/or DW vaccination of day-old SPF or broiler chicks [14, 57, 58]. High levels of IBV-specific IgA have also been associated with a degree of protection against virulent IBV [14]. For IgY (the equivalent of IgG in mammals), detection in lachrymal fluid could have been due to transudation [59] and/or local secretion by B-cells [60]. In the current study, it appears that ocuonosal inoculation of either vaccine resulted in a significant increase in lachrymal IgY by 14 dpv. This is potentially due to greater IBV replication and inflammation in the upper respiratory and lymphoid tissues, including the HG [61]. These results are similar to those found by Gallego et al. who demonstrated that soluble antigen administration in seven-week-old chickens via the ocular route is an efficient application for producing a local immune response in the HG [62]. The significantly higher levels of lachrymal IgY at 14 dpv also corresponded with the overall increase of humoral IBV antibody titres [13, 14, 51], reflecting increased transudation of IgY into the lacrymal fluid. This appears to be particularly true for the 793B vaccine, as we only detected significant IgY levels in the DW-793B group, and not for the DW-Mass group. The local memory immune response has been shown to be dominated by IgY [13]. Therefore, higher levels of IgY in egg-laying hens would be beneficial in minimising IBV infection, subsequently avoiding or reducing egg production losses, and sustaining a good overall flock health.

Although indirect ELISAs (as described above) are preferred for measuring mucosal immunity, it is not always possible for several reasons. This includes the requirement of IgA/IgY monoclonal antibodies, IBV antigen purification and standardization, the small quantity of lachrymal fluid which is collectable from each bird, the laborious process of tracheal wash collection, and the inclusion of upper respiratory tissues (e.g., turbinate) where no fluid could be collected. For these reasons, we also attempted to measure the regulation of IgA and IgY mRNA transcription in tissues of turbinate and trachea. This allowed for quantification of mucosal immunity parameters in tissue samples. For changes in IgA or IgY mRNA transcription, comparisons between the ON-vaccinated groups showed up or down regulation depending on the vaccine strain and sampling timepoint. For turbinate tissues of Mass-vaccinated hens, IgA and IgY mRNA was down-regulated at all sampling points, whereas those vaccinated with 793B showed a significant up-regulation at 3 (IgA and IgY) and 5 (IgA) dpv. In contrast, for the trachea, up-regulation of IgA mRNA, but not IgY, was found for both Mass or 793B vaccinated hens. Interestingly,
up-regulation of IgA mRNA in the Mass group was significantly higher than the 793B group at 5 dpv. This may suggest consistent and higher replication of both Mass and 793B in the trachea, whereas the turbinate appears to support better replication of 793B. This was consistent with viral load data in turbinate and trachea tissue as described above. In contrast to ON application, IgA mRNA expression in the trachea and turbinate following DW administration showed down-regulation at all sampling points. This could have been due to slow dissemination of the vaccines and/or lower viral load in the HG and respiratory tissues. The early up-regulation of IgY mRNA at 3 dpv in the turbinate and trachea appears to be associated with the peak of 793B viral load. Also, an increase of IL-6 was noted at the same sampling point, indicating increased inflammation.

Early immune responses following virulent IBV infection or vaccination in young SPF and broiler chicks have been published before [36, 63, 64]. In this study, for the first time, certain parameters of the early immune response in Mass or 793B vaccinated egg-laying hens are reported. Findings showed that mRNA transcription of TLR3 was significantly up-regulated in the HG (3–5 dpv) and trachea (5 dpv) following either ON or DW inoculation. Similarly, MDA5 was significantly increased in the HG and trachea. Up-regulated transcription of both genes has been previously demonstrated in the respiratory tissues of SPF and broiler chicks [61, 65]. Significantly higher transcripts in Mass-vaccinated hens in contrast to the 793B groups, reflects a potentially slower 793B dissemination and replication, and recognition by the host [66]. In contrast, for DW application, differences in magnitude or pattern were not consistently noted in the HG, turbinate or trachea between Mass and 793B groups. This could have potentially been associated with a lower viral load in these tissues when either vaccine was given by DW in comparison to ON.

Although cell-mediated immunity (CMI) is an essential part of the immune response to IBV, there is no information available on the CMI responses following live IB vaccination in egg-laying hens. In this study, following ON application, the Mass vaccine produced a stronger induction of CMI compared to the 793B vaccine, as demonstrated by significantly raised CD8-α transcripts in the trachea at 3–5 dpv. This is likely due to the Mass vaccine virus’s ability to disseminate and replicate more efficiently in respiratory tissues. It has been reported that CD8+ cells expressing the CD8-α chain are involved in elimination of IBV-infected cells during early infection [57, 67, 68]. Significant up-regulation (compared to the control) of CD8-α mRNA in the trachea of the Mass ON group at 3–5 dpv, compared to only 5 dpv for 793B, again reflects the ability of the Mass vaccine to replicate and stimulate local cellular immune responses. Based on this study, it appears that by 5 dpv, the host cellular response to Mass vaccination, either given by DW or ON, were comparable.

In conclusion, this study represents the first investigation to evaluate the viral load of Mass or 793B attenuated live vaccines, and host immune responses to these vaccines, in egg-laying hens. This study has demonstrated that the pattern of IBV replication differs according to tissue type, vaccine strain, and inoculation route. For both vaccines, the dissemination and clearance of the vaccine viruses were slower when given by DW compared to the ON route. When given by the ON route, both vaccines were able to induce comparable levels of mucosal immunity. The Mass IBV vaccine induces cellular immunity to similar levels regardless of vaccination method. When given either by ON or DW, the 793B vaccine induced significantly higher levels of humoral immunity. These findings can be used for improved IB vaccination strategies in egg-laying hens to prevent drop in egg production and/or quality.

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Authors’ contributions
MAR conducted sampling collection, laboratory analysis and manuscript preparation. CB aided with laboratory assays, data analysis and manuscript preparation, and KG conceived and designed the experiment, supervised the laboratory and data analysis, and finalised the manuscript. All authors have read and approved the final manuscript.

Declarations
Competing interests
The authors declare that they have no competing interests.

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References
1. Gelb J Jr, Wolf J, Moran C (1991) Variant serotypes of infectious bronchitis virus isolated from commercial layer and broiler chickens. Avian Dis 35:82–87
2. Cavanagh D (2007) Coronavirus avian infectious bronchitis virus. Vet Res 38:281–297
3. Cavanagh D, Gelb J (2008) Infectious bronchitis. In: Swayne D (ed) Diseases of poultry. Blackwell Publishing, Ames
4. Box P, Beresford A, Roberts B (1980) Protection of laying hens against infectious bronchitis with inactivated emulsion vaccines. Vet Rec 106:264–268
5. Cook JK (1971) Recovery of infectious bronchitis virus from eggs and chicks produced by experimentally inoculated hens. J Comp Pathol 81:203–211
6. Broadfoot DI, Pomeroy BS, Smith W Jr (1954) Effect of infectious bronchitis on egg production. J Am Vet Med Assoc 124:128–130
7. Van Eck J (1983) Effects of experimental infection of fowl with EDS'76 virus, infectious bronchitis virus and/or fowl adenovirus on laying performance. Vet Q 5:11–25
8. APHA (2019) GB avian quarterly report. Disease surveillance and emerging threats. https://assets.publishing.service.gov.uk/government/uploads/attachment_data/file/806086/pub-survey-a0119.pdf. Accessed 11 June 2020
9. DEFRA (2005) Defra report on the Economic Assessment of Livestock Diseases in the United Kingdom (UK)-Z0102. http://sciencedefra.gov.uk/Default.aspx?Menu=Menu&Module=More&Location=Home&ProjectID=9781&search=y&publisher=1&searchText=Z0102&sort=5&trino=project. Accessed 11 June 2020
10. Sevoian M, Levine P (1957) Effects of infectious bronchitis on the reproductive tracts, egg production, and egg quality of laying chickens. Avian Dis 1:136–164
11. De Wit J, Cook JK, Van der Heijden HM (2011) Infectious bronchitis virus variants: a review of the history, current situation and control measures. Avian Pathol 40:223–235
12. Chhabra B, Forrester A, Lemiere S, Awad F, Chantrey J, Ganapathy K (2015) Mucosal, cellular, and humoral immune responses induced by different live infectious bronchitis virus vaccine regimes and protection conferred against infectious bronchitis virus Q1 strain. Clin Vaccine Immunol 22:1050–1059
13. Guo X, Rosa AJ, Chen DG, Wang X (2008) Molecular mechanisms of primary and secondary mucosal immunity using avian infectious bronchitis virus as a model system. Vet Immunol Immunopathol 121:332–343
14. Okino CH, Alessi AC, Montassier MD, F, Rosa AJ, Wang X, Montassier HJ (2013) Humoral and cell-mediated immune responses to different doses of attenuated vaccine against avian infectious bronchitis virus. Vet Immunol Immunopathol 161:259–267
15. Bande F, Arshad SS, Hair Bejo M, Moenei H, Omar AR (2015) Progress and challenges toward the development of vaccines against avian infectious bronchitis virus. J Immunol Res 2015:424860
16. Box P, Ellis K (1985) Infectious bronchitis in laying hens: interference with response to emulsion vaccine by attenuated live vaccine. Avian Pathol 14:9–22
17. Finney P, Box P, Holmes H (1990) Studies with a bivalent infectious bronchitis killed virus vaccine. Avian Pathol 19:435–450
18. De Wit J (2013) Underestimation of the difficulties of vaccination against viral respiratory diseases by mass application methods. Proceedings of the XVIII International Congress of the World Veterinary Poultry Association, Nantes, pp 19–23
19. Jackwood MM, de Wit S (2013) Infectious bronchitis. In: Swayne D (ed) Diseases of poultry, 13th edn. Blackwell, Wiley
20. Gough R, Alexander D (1979) Comparison of duration of immunity in chickens infected with a live infectious bronchitis vaccine by three different routes. Res Vet Sci 26:329–332
21. Ratanaesathkul C, Cumming R (1983) Immune response of chickens to various routes of administration of Australian infectious bronchitis vaccine. Aust Vet J 60:214–216
22. Matthijs MGR, van Eck JHH, De Wit J, Bouma A, Stegeman J (2005) Effect of IBV-H120 vaccination in broilers on colibacillosis susceptibility after infection with a virulent Massachusetts-type IBV strain. Avian Dis 49:540–545
23. Blijenla G, Cook JK, Gelb J, Jack WJ (2004) Development and use of the H strain of avian infectious bronchitis virus from the Netherlands as a vaccine: a review. Avian Pathol 33:550–557
24. Jackwood MW (2012) Review of infectious bronchitis virus around the world. Avian Dis 56:634–641
25. Jordan B (2017) Vaccination against infectious bronchitis virus: a continuous challenge. Vet Microbiol 206:137–143
26. Worthington KJ, Currie R, Jones RC (2008) A reverse transcriptase-polymerase chain reaction survey of infectious bronchitis virus genotypes in Western Europe from 2002 to 2006. Avian Pathol 37:247–257
27. Gough R, Randal C, Dagless M, Alexander D, Cox W, Pearson D (1992) A ‘new’ strain of infectious bronchitis virus infecting domestic fowl in Great Britain. Vet Rec 130:493–494
28. Parsons D, Ellis M, Cavanagh D, Cook J (1999) Characterisation of an infectious bronchitis virus isolated from vaccinated broiler breeder flocks. Vet Rec 131:408–411
29. Śmiałek M, Welenc J, Koncicki A (2016) Systemic and local immune mechanisms stimulated in the course of chicken infectious bronchitis. Med Weter 72:358–363
30. Reed LJ, Muench H (1938) A simple method of estimating fifty per cent endpoints. Am J Hyg 27:493–497
31. Ganapathy K, Cargill PW, Jones RC (2005) A comparison of methods of inducing lachrymation and tear collection in chickens for detection of virus-specific immunglobulins after infection with infectious bronchitis virus. Avian Pathol 34:248–251
32. Mockett AP, Cook JK (1986) The detection of specific IgM to infectious bronchitis virus in chicken serum using an ELISA. Avian Pathol 15:437–446
33. Raj GD, Jones RC (1996) Local antibody production in the oviduct and gut of hens infected with a variant strain of infectious bronchitis virus. Vet Immunol Immunopathol 53:147–161
34. Fournier-Carauana J, Poineer B, Haond G, Jallet C, Fuchs T, Tordo N, Perrin P (2003) Inactivated rabies vaccine control and release use of an ELISA method. Biologicals 31:9–16
35. Jones RS, Ellis R, Cox W, Errington J, Fuller C, Irvine R, Wakeley P (2011) Development and validation of RT-PCR tests for the detection and S1 genotyping of infectious bronchitis virus and other closely related gammacoronaviruses within clinical samples. Transbound Emerg Dis 58:411–420
36. Chhabra B, Ball C, Chantrey J, Ganapathy K (2018) Differential innate immune responses induced by classical and variant infectious bronchitis viruses in specific pathogen free chicks. Dev Comp Immunol 87:16–23
37. Løndt BZ, Brookes SM, Kelly MD, Nash BJ, Brown IH (2013) Failure to infect pigs co-housed with ducks or chickens infected experimentally with A/ turkey/Turkey/1/2005 (H5N1) highly pathogenic avian influenza virus. Vet Micro 162:944–948
38. Chhabra R, Kuchipudi SV, Chantrey J, Ganapathy K (2016) Pathogenicity and tissue tropism of infectious bronchitis virus is associated with elevated apoptosis and innate immune responses. Vetrol 488:232–241
39. Livak KJ, Schmittgen TD (2001) Analysis of relative gene expression data using real-time quantitative PCR and the 2−ΔΔCT method. Methods 25:402–408
40. Okino CH, Mores MA, Trevischom IM, Coldebellom A, Montassier HJ, Brentano LL (2017) Early immune responses and development of pathogenesis of avian infectious bronchitis viruses with different virulence profiles. PLoS One 12:e0172275
41. Kuchipudi SV, Tellabati M, Nelli RK, Perez BB, Sebastian S, Dunham SP, Chang KC, White GA, Slomka MJ, Brookes SM, Brown IH (2012) 18S rRNA is a reliable normalisation gene for real time PCR based on influenza virus infected cells. Virol J 9:230
42. Ndegeya EN, Joner SK, Taro H, Van Ginkel FW, Van Santen VL (2012) The proportion of specific viral subpopulations in attenuated Arkansas Delmarva poultry industry infectious bronchitis viruses influence vaccine outcome. Avian Dis 56:642–653
43. Zheng W, Ikazi J, Furusawa S, Yoshimura Y (2001) A sensitive non-radioactive in situ hybridization method for the detection of chicken IgG gamma-chain mRNAs: a technique suitable for detecting of variety of mRNAs in tissue sections. Biol Proc Online 3:1–7
44. Kuchipudi SV, Tellabati M, Sebastian S, Løndt BZ, Jansen C, Verveke L, Brookes SM, Brown IH, Dunham SP, Chang KC (2014) Highly pathogenic avian influenza virus infection in chickens but not ducks is associated with elevated host immune and pro-inflammatory responses. Vet Res 45:118
45. Jackwood MW, Clark R, Cheng S, Jordan BJ (2020) Protection following simultaneous vaccination with three or four different attenuated live vaccine types against infectious bronchitis virus. Avian Path 49:335–341
46. De Wit J, Malo A, Cook JK (2019) Induction of IBV strain-specific neutralizing antibodies and broad spectrum protection in layer pullets primed with IBV Massachusetts (Mass) and 793B vaccines prior to injection of inactivated vaccine containing Mass antigen. Avian Pathol 48:135–147
47. Tizard IR (2020) Vaccination against coronaviruses in domestic animals. Vaccine 38:133–5130
48. Hewson K, Robertson T, Steer P, Devlin J, Noormohammadi A, Ignjatovic J (2014) Assessment of the potential relationship between egg quality and infectious bronchitis virus infection in Australian layer flocks. Aust Vet J 92:132–138
49. Ball C, Awad F, Hutton S, Forrester A, Baylis M, Ganapathy K (2017) Infectious bronchitis vaccine virus detection and part-S1 genetic variation following single or dual inoculation in broiler chicks. Avian Pathol 46:309–318
50. Davelaar F, Kousenhooven B (1977) Influence of maternal antibodies on vaccination of chicks of different ages against infectious bronchitis. Avian Pathol 6:41–50
51. Raggi LG, Lee GG (1965) Lack of correlation between infectivity, serologic response and challenge results in immunization with an avian infectious bronchitis vaccine. J Immunol 94:538–543
52. Sulaiman A, Roberts J (2011) Infectious bronchitis vaccination protocols for laying hens. Media Peternakan 34:159–159
53. Cook K, Otsuki K, da Silva MN, Ellis M, Huggins M (1992) The secretory antibody response of inbred lines of chicken to avian infectious bronchitis virus infection. Avian Pathol 21:681–692
54. Orr-Burks N, Gulley SL, Gallardo RA, Toro H, Van Ginkel FW (2014) Immunoglobulin A as an early humoral responder after mucosal avian coronavirus vaccination. Avian Dis 58:279–286
55. Toro H, Fernandez I (1994) Avian infectious bronchitis: specific lachrymal IgA level and resistance against challenge. Zentralbl Veterinarmed B 41:467–472
56. van Ginkel FW, van Santen VL, Gulley SL, Toro H (2008) Infectious bronchitis virus in the chicken Harderian gland and lachrymal fluid: viral load, infectivity, immune cell responses, and effects of viral immunodeficiency. Avian Dis 52:608–617
57. Smialek M, Tykalowski B, Dziewulska D, Stenzel T, Koncicki A (2017) Immunological aspects of the efficiency of protectotype vaccination strategy against chicken infectious bronchitis. BMC Vet Res 13:44
58. Toro H, Espinoza C, Ponce V, Rojas V, Morales M, Kaleta E (1997) Infectious bronchitis: effect of viral doses and routes on specific lacrimal and serum antibody responses in chickens. Avian Dis 41:379–387
59. Toro H, Lavaud P, Vallejos P, Ferreira A (1993) Transfer of IgG from serum to lachrymal fluid in chickens. Avian Dis 37:60–66
60. Smialek M, Tykalowski B, Stenzel T, Koncicki A (2011) Local immunity of the respiratory mucosal system in chickens and turkeys. Pol J Vet Sci 14:291–297
61. Deist MS, Lamont SJ (2018) What makes the Harderian gland transcriptome different from other chicken immune tissues? A gene expression comparative analysis. Front Physiol 9:492
62. Gallego M, Del Cacho E, Arnal C, Bascuas J (1992) Local immune response in the chicken Harderian gland to antigen given by different ocular routes. Res Vet Sci 52:38–43
63. Chhabra R, Chantrey J, Ganapathy K (2015) Immune responses to virulent and vaccine strains of infectious bronchitis viruses in chickens. Viral Immunol 28:478–488
64. Manswr B, Ball C, Forrester A, Chantrey J, Ganapathy K (2020) Immunopathogenesis of infectious bronchitis virus Q1 in specific pathogen free chicks. Microb Pathog 149:104535
65. Manswr B, Ball C, Forrester A, Chantrey J, Ganapathy K (2021) Host immune response to infectious bronchitis virus Q1 in two commercial broiler chicken lines. Res Vet Sci 136:587–594
66. Lin S-Y, Li Y-T, Chen Y-T, Chen T-C, Hu C-M, Chen H-W (2016) Identification of an infectious bronchitis coronavirus strain exhibiting a classical genotype but altered antigenicity, pathogenicity, and innate immunity profile. Sci Rep 6:37725
67. Kameka AM, Haddadi S, Kim DS, Cork SC, Abdul-Careem MF (2014) Induction of innate immune response following infectious bronchitis corona virus infection in the respiratory tract of chickens. Virol 450–451:114–121
68. Okino CH, Santos Icd, Fernando FS, Alessi AC, Wang XQ, Montassier HJ (2014) Inflammatory and cell-mediated immune responses in the respiratory tract of chickens to infection with avian infectious bronchitis virus. Viral Immunol 27:383–391

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