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Avian coronavirus infectious bronchitis attenuated live vaccines undergo selection of subpopulations and mutations following vaccination

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KEYWORDS

Coronavirus; Avian infectious bronchitis virus; Vaccine virus adaptation; Selection; Mutation; Molecular typing; Spike gene sequence; Serotype; Quasispecies

Summary In this study, we were interested in determining if high titered egg adapted modified live infectious bronchitis virus (IBV) vaccines contain spike gene related quasispecies that undergo selection in chickens, following vaccination. We sequenced the spike glycoprotein of 12 IBV vaccines (5 different serotypes from 3 different manufacturers) directly from the vaccine vial, then compared that sequence with reisolated viruses from vaccinated and contact-exposed birds over time. We found differences in the S1 sequence within the same vaccine serotype from different manufacturers, differences in S1 sequence between different vaccine serials from the same manufacturer, and intra-vaccine differences or quasispecies. Comparing the sequence data of the reisolated viruses with the original vaccine virus, we were able to identify in vivo selection of viral subpopulations as well as mutations. To our knowledge, this is the first report showing selection of a more fit virus subpopulation as well as mutations associated with replication of modified live IBV vaccine viruses in chickens. This information is important for our understanding of how attenuated virus vaccines, including potential vaccines against the SARS-CoV, can ensure long-term survival of the virus and can lead to changes in pathogenesis and emergence of new viral pathogens. This information is also valuable for the development of safer modified live coronavirus vaccines. Published by Elsevier Ltd.

Introduction

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Coronaviruses are an important group of enveloped single stranded RNA viruses that cause highly contagious respiratory and enteric diseases in a variety of animals and in humans. Coronaviruses belong to the order *Nidovirales*, family *Coronaviridae*, and are separated into 3 groups based on genetic and antigenic characteristics [1]. Avian infectious bronchitis virus (IBV) belongs to the group 3 coronaviruses along with the other avian coronaviruses, whereas coronaviruses in groups 1 and 2 produce disease in many species of mammals and in humans. Confusion around the origin of severe acute respiratory syndrome coronavirus (SARS-CoV) has lead to a proposed group 4; however, genetic studies suggest that SARS-CoV diverged from the group 2 viruses, thus a group 2b classification was suggested [2].

Avian infectious bronchitis virus has an enormous economic impact on commercial poultry causing highly contagious disease in chickens of all ages. Since IBV was first described by Schalk and Hawn in the 1930s, reviewed in ref. [3], many serotypes have been identified worldwide. There are currently 4 reported structural proteins associated with the IBV virion; nucleocapsid, membrane, small membrane and spike. The spike glycoprotein on the outside of the virus contains epitopes associated with serotype differences and binding of neutralizing antibodies, and it plays a role in attachment and entry into the host cell.

Coronaviruses including IBV have been shown to exist as a mixture of genetic mutants within an isolate [4,5]. These so called quasispecies are generated through genetic mutations and recombination events that occur when the viral RNA-dependent RNA-polymerase, which lacks proofreading capabilities, replicates the viral genome [6]. For SARS-CoV, non-synonymous and synonymous substitution rates have been estimated to be as high as 3.3×10^{-3} and 4.67×10^{-3} per site per year, respectively [7]. Evolutionary and mutation rates for IBV in the face of vaccination were determined to be 2.5 and 1.5% per year, respectively, in the hypervariable region of the spike glycoprotein; however, in the absence of vaccines, IBV lineages appear to evolve at a much slower rate (0.3%) [8]. The group 3 coronaviruses, which exist only in avian species were found to have a relatively recent divergence date (1925) and exponential population growth [2]. In general, constant population size is associated with natural reservoirs of the virus, which remain largely asymptomatic in the host, whereas a change from constant to exponential population growth indicates emergence of a relatively new virus associated with interspecies transmission and disease [2].

An actively changing genetic population allows rapid adaptation to the host through selection of the most fit viral subpopulation. Selection ensures long-term survival of the virus at the cell, organism, and host population level, and can lead to changes in pathogenesis and emergence of new viral pathogens [9]. For economic reasons, vaccine manufacturers attempt to produce high titered (>1 \times 10⁸ embryo infectious dose₅₀ (EID₅₀)/ml) IBV vaccines so they can be packaged in 5000 or even 10,000 dose vials. The average IBV vaccine dose can vary but is generally approximately 1×10^4 EID₅₀. Since the number of intra-isolate genome subpopulations are reported to increase dramatically when the virus titer increases [9], we were interested in determining if quasispecies of high titered modified live IBV vaccines contain specific spike gene mutations that undergo selection in chickens, following vaccination. In addition, we wanted to determine if molecular changes were solely the result of selection or if mutations also occur following vaccination.

To study the dynamics of IBV vaccine genetic adaptation to the host, we sequenced the spike glycoprotein of 12 IBV vaccines (5 different serotypes from 3 different manufacturers) directly from the vaccine vial. Then, we vaccinated chickens with those viruses and examined the spike glycoprotein sequence of the vaccines reisolated from vaccinated and from contact-exposed birds over time. Comparing the sequence data of the reisolated viruses with the original vaccine virus, we were able to access *in vivo* selection of subpopulations and genetic mutations.

Materials and methods

Vaccines

Twelve IBV commercial vaccines were obtained from three different manufacturers designated A, B and C herein, and had a titer of at least 1×10^8 EID₅₀/ml. The vaccines used in this study from manufacturer A were Ark-DPI (Ark/A), GA98 (GA/A) and Mass (Mass/A). Vaccines from manufacturer B were Ark-DPI (Ark/B), Mass (Mass/B) and a mixture of Mass and Conn (Conn/B). The vaccines from manufacturer C were Ark-DPI (Ark/C), DE072 (Del/C), GA98 (GA/C), Mass (Mass/C), a mixture of Mass and Conn (Conn/C), and MassD (MassD/C). The lyopholyzed vaccines were resuspended in 10 ml of diethylpyrocarbonate (DEPC) treated water. The DE072 vaccine, stored in liquid nitrogen, was thawed and diluted using 1 ml of DEPC treated water.

Birds and housing

Specific-pathogen-free (SPF) white leghorn chicks (Merial, Gainesville, GA) were housed in positive-pressure Horsfal isolation units. Feed and water were provided *ad libitum*, and the birds were examined twice daily.

Experimental design

To guard against potential cross-contamination, the 12 vaccines were assigned to 4 different groups ensuring that there were no vaccines of the same serotype in each group. Two different experiments were conducted with vaccine groups 1 and 2 in the first experiment and vaccine groups 3 and 4 in the second experiment. Vaccines used in experiment 1 were for group 1 Del/C, GA/C, Mass/A, and for group 2 Ark/C, GA/A, and Mass/C. Vaccines used in experiment 2 were for group 3 Ark/B, Conn/B, Mass/B, and for group 4 Ark/A, Conn/C, and MassD/C. Each vaccine group was tested in a separate filtered air positive pressure room containing the isolators. One-week old chicks were randomly divided into groups of 10 birds for each vaccine type. Five birds in each group were inoculated by eye-drop with $30 \,\mu l$ of vaccine (equal to one dose, according to the manufacturers recommendation), the other 5 birds served as contact-exposed birds. A negative control group of 5 birds was included for each group of vaccines tested.

Tracheal swabs from all birds were collected at 3, 6 and 9 days post-vaccination (dpv), and at necropsy. Birds in experiment 1 were necropsied at 14 dpv while birds in experiment 2 were necropsied at 13 dpv. Swabs from 5 vaccinated birds and 5 contact-exposed birds for each of the vaccine types were pooled separately in 1 ml PBS (pH 7.4). The swabs were

stored at $-80\,^\circ\text{C}$ until they were analyzed for IBV RNA by RT-PCR.

Passage in embryonating eggs

The last positive tracheal swab by RT-PCR from each vaccinated and contact-exposed group of birds was passaged a maximum of ten times in 9–11 day old specific pathogen free (SPF) embryonating eggs. Three eggs per group were inoculated with 0.1 ml of PBS from the tracheal swabs into the chorioallantoic sac (CAS) as described [10]. The eggs were incubated at 37 °C and allantoic fluid was harvested from all eggs with live embryos at 48 h post-inoculation, pooled and kept at -80 °C. Subsequent passage was done by inoculating 0.2 ml of allantoic fluid from the previous passage into the CAS of 9–11 day old embryonating eggs and the allantoic fluid was harvested at 48 h post-inoculation as above.

Viral RNA extraction

Viral RNA was extracted from allantoic fluid, rehydrated or diluted vaccine, and tracheal swabs using the High Pure RNA Isolation Kit (Roche Diagnostics, Indianapolis, IN) according to manufacturer's protocol.

RT-PCR amplification and cloning of the S1 gene

Reverse transcriptase-polymerase chain reaction was carried out using the Titan One Tube RT-PCR kit (Roche Diagnostics, Indianapolis, IN) according to manufacturer's protocol with primers NEWS10LIG05' [11] and Degenerate3' [12]. Primers specific for the Conn serotype [13] were used to screen for the presence of that virus in swabs collected from birds vaccinated with vaccine containing both the Mass and Conn types and in swabs from the corresponding contact birds. The PCR products were agarose gel purified using the QIAquick gel extraction kit (Qiagen, Valencia, CA) according to the manufacturer's protocol. To aid in sequence analysis, purified PCR products from Conn/B and Conn/C, which were mixtures of Mass and Conn vaccines, and Ark/B were cloned into the TOPOXL vector using TOPOXL cloning kit (Invitrogen, Carlsbad, CA) according to manufacturer's protocol. The IBV real-time RT-PCR analysis on tracheal swabs was performed as previously described [4,14].

Sequencing and sequence analysis of the S1 gene

Viral S1 gene sequence was determined for two different production lots for each IBV vaccine examined. The S1 gene of reisolated viruses from the first and last RT-PCR positive tracheal swabs only was also sequenced for both vaccinated and contact-exposed birds. The S1 gene of reisolated virus from Ark/A vaccinated birds was only positive by real time RT-PCR. The pooled tracheal swabs collected from the 13 dpv vaccinated birds were sequenced. In addition, each embryo passaged virus was sequenced to determine if genetic changes in the reisolated vaccine viruses would revert to the original vaccine sequence. Purified RT-PCR products or at least 4 cloned RT-PCR products for each virus were sequenced using the BigDye[®] Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems, Foster City, CA) according to manufacturer's protocol. Sequencing reactions were purified using Centri-Sep Columns (Princeton Separations, Inc, Adelphia, NJ) or with the Performa DTR Ultra Dye Terminator removal system (Edge BioSystems, Gaithersburg, MD). Sequence reactions were examined at the Molecular Genetics Instrumentation Facility (University of Georgia, Athens, GA).

The S1 sequences from each virus were compiled using the EditSeq and MegAlign program (DNASTAR, Inc. Madison, WI) and open reading frames were predicted using ORF finder at National Center for Biotechnology Information (NCBI, http://www.ncbi.nlm.nih.gov/gorf/gorf.html). The BlastX program (http://www.ncbi.nlm.nih.gov/BLAST/) was used to search GenBank (National Center for Biotechnology Information, http://www.ncbi.nlm.nih.gov/) for homologous IBV S1 sequences.

Results

S1 sequence analysis of vaccine viruses

All nucleotide sequences reported herein have been submitted to NCBI, GenBank (http://www.ncbi.nlm.nih.gov/) and the accession numbers are listed in Table 1. To determine if mutations or selection of a more fit virus subpopulation was responsible for the changes observed when IBV adapts to an *in vivo* environment, we first needed to establish the exact sequence of the vaccine viruses directly from the manufacturer. The S1 gene deduced amino acid sequence for different production lots of the Ark/A and Ark/C vaccine viruses were the same, whereas Ark/B vaccine production lots had 1 amino acid difference (Pro524Ser) in the S1 gene. In addition, sequence differentes were observed between Arkansas vaccines from the different manufacturers (Table 2).

There were 26 amino acid differences in the S1 sequences between the two production lots for the Conn/B vaccine (Table 3). For the Conn/C vaccine, we observed 5 amino acid differences between the different production lots. In addition, one of the Conn/C vaccine production lots had a deletion of amino acid Asn at position 58.

There were no differences observed between the S1 sequences obtained from the two vaccine production lots for the Del/C, GA/A, GA/C, Mass/A, Mass/B, and Mass/C. There were 13 amino acid sequence differences between the two production lots of the MassD/C vaccine virus (Table 4). MassD/C vial 1 was closest to Mass/A with only 1 amino acid sequence difference between them, whereas MassD/C vial 2 was closest to Mass/C and Mass/B with 6 and 7 amino acid sequence differences, respectively. In addition, there were differences in the amino acid sequences between the Mass vaccines from the different manufacturers. We observed 14 and 11 amino acid sequence differences between Mass vaccines from manufacturer A and the Mass vaccines from manufacturers B and C, respectively. We also found 2 amino acid sequence differences between Mass vaccines from manufacturers B and C. Finally, there were 28 amino acid sequence differences and one deletion between the GA vaccines from manufacturers A and C (Table 5).

IBV vaccine virus	S1 sequence source	NCBI accession number
Ark/A	First vaccine lot Second vaccine lot 13 dpv ^b vaccinated birds	EU283045 EU283046 EU283047
Ark/B	First vaccine lot Second vaccine lot 3 dpv vaccinated 9 dpv vaccinated 6 dpv contact birds 13 dpv contact birds	EU283048 EU283049 EU283050 EU283051 EU283052 EU283053
Ark/C	First vaccine lot Second vaccine lot 6 dpv vaccinated birds	EU283054 EU283055 EU283056
Conn/B	First vaccine lot Second vaccine lot 6 dpv contact birds 13 dpv contact birds	EU283057 EU283058 EU283059 EU283060
Conn/C	First vaccine lot Second vaccine lot	EU283061 EU283062
Del/C	First vaccine lot Second vaccine lot Reisolated vaccine virus ^c	EU283063 EU283064 EU283065
GA/A	First vaccine lot Second vaccine lot 3 dpv vaccinated birds 9 dpv vaccinated birds	EU283066 EU283067 EU283068 EU283069
GA/C	First vaccine lot Second vaccine lot Reisolated vaccine virus	EU283070 EU283071 EU283072
Mass/A	First vaccine lot Second vaccine lot Reisolated vaccine virus	EU283073 EU283074 EU283075
Mass/B	First vaccine lot Second vaccine lot 3 dpv vaccinated 9 dpv vaccinated 6 dpv contact birds 13 dpv contact birds	EU283076 EU283077 EU283078 EU283079 EU283080 EU283081
Mass/C	First vaccine lot Second vaccine lot Reisolated vaccine virus	EU283082 EU283083 EU283084
MassD/C	First vaccine lot Second vaccine lot 3 dpv vaccinated birds 6 dpv contact birds	EU283085 EU283086 EU283087 EU283088

Table 1 NCBI^a accession numbers for IBV S1 gene sequences obtained directly from virus in vaccine vials, and reisolated vaccine virus from vaccinated and contact-exposed birds

^a NCBI = National Center for Biotechnology Information (http://www.ncbi.nlm.nih.gov/).
^b dpv = day post-vaccination.

^c Reisolated vaccine virus indicates that all viruses reisolated from vaccinated and contact-exposed birds had the same sequence.

vaccines from different manufacturers						
Residue position	Ark/A	Ark/B	Ark/C			
43	Tyr	His	Tyr			
171	Tyr	His	Tyr			
213	Ser	Ala	Ser			
249	Phe	Ser	Phe			
323	Arg	Arg	Tyr			
410	Lys	Glu	Lys			
524	Ser	Pro (Ser) ^a	Ser			

Table 2Pairwise comparisons of amino acids between Arkvaccines from different manufacturers

 $^{\rm a}$ Amino acid in brackets indicates the residue observed in vaccine vial 2.

Virus detection from vaccinated and contact-exposed birds

Tracheal swabs collected from vaccinated and contactexposed birds were tested for IBV by RT-PCR and real-time RT-PCR and the results are presented in Table 6. Tracheal swabs collected from all the negative control birds were negative for IBV by RT-PCR and real-time RT-PCR. Based on sequence data from reisolated viruses, there was no crosscontamination detected between the vaccines. The tracheal swabs from Ark/B, Ark/C, Mass/B, Mass/C, and MassD/C vaccinated birds were positive for IBV at all sampling times. The tracheal swabs from Del/C, GA/C, GA/A, and Mass/A vaccinated birds were positive on days 3, 6, and 9 dpv but not at necropsy (14 dpv Expt. 1 or 13 dpv Expt. 2). Tracheal swabs for all of the contact-exposed birds in groups Ark/B, Conn/B, Mass/A, and Mass/B were positive for IBV at all sampling times. Swabs from Del/C, GA/A, Mass/C, and MassD/C contact-exposed birds were positive for IBV by RT-PCR or real time RT-PCR except at 3 dpv. Tracheal swabs from Ark/C contact-exposed birds were negative at 3, 6, and 9 dpv but positive at necropsy (14 dpv Expt. 1 or 13 dpv Expt. 2). Using RT-PCR analysis, tracheal swabs collected from Ark/A vaccinated and contact-exposed birds were negative for IBV. However, real-time RT-PCR analysis, which is more sensitive than RT-PCR, showed that the swabs from Ark/A vaccinated birds but not contact-exposed birds were positive for virus.

Table 3 Pairwise comparisons of amino acids between Conn vaccines from different manufacturers

Residue position	Conn/B vial 1	Conn/B vial 2	Conn/C vial 1	Conn/C vial 2
10	Ala	Thr	Thr	Thr
58	Asn	Asn	Asn	Deleted
74	Ser	Asn	Asn	Asn
81	Thr	Ser	Thr	Thr
88	Asp	Asp	Gly	Asp
114	His	His	Tyr	His
161	Leu	Phe	Phe	Phe
188	Lys	Asn	Asn	Asn
214	Asp	Asn	Asp	Asp
243	lle	Thr	Thr	Thr
260	Val	lle	lle	lle
264	Phe	Leu	Leu	Leu
265	Thr	Lys	Lys	Lys
268	Tyr	Asn	Asn	Asn
270	Ser	Thr	Thr	Thr
281	Pro	Leu	Leu	Leu
309	Ser	Gly	Gly	Gly
330	Leu	Pro	Pro	Pro
331	Glu	Lys	Glu	Glu
343	Ser	Leu	Ser	Ser
373	Glu	Gly	Gly	Gly
376	Leu	Ser	Ser	Ser
381	Val	Val	lle	Val
383	Ser	Leu	Leu	Leu
387	Asp	Lys	Lys	Lys
388	His	Ser	Ser	Ser
389	Asn	Asp	Asp	Asp
402	Gly	Asp	Asp	Asp
406	lle	lle	Thr	Thr
430	Asp	Asp	Gly	Asp
460	Leu	Met	Met	Met
478	Tyr	Tyr	Cys	Tyr
481	Asn	Thr	Thr	Thr

Table 4	Pairwise co	mparisons of	amino	acids	between	Mass	vaccines	from	different	: manufacturers	

Residue position	Mass/A	Mass/B	Mass/C	MassD/C vial 1	MassD/C vial 2
19	Ala	Val	Val	Ala	Ala
38	Asp	Asn	Asn	Asp	Asn
64	Gly	Glu	Gly	Gly	Gly
69	lle	Thr	Thr	lle	Thr
73	Gly	Asp	Gly	Gly	Gly
75	Val	Val	Val	Val	Ala
76	Val	Val	Val	Val	Ala
101	Tyr	His	His	Tyr	His
109	Val	Val	Val	Val	Glu
130	Ser	Phe	Phe	Ser	Phe
179	Ala	Glu	Ala	Glu	Glu
205	Arg	Lys	Lys	Arg	Lys
246	Thr	lle	lle	Thr	lle
271	Asn	Tyr	Tyr	Asn	Tyr
273	Thr	Ser	Ser	Thr	Arg
376	Gly	Glu	Glu	Gly	Glu
479	Ser	Gly	Gly	Ser	Gly

Table 5	Pairwise comparisons of amino acids between GA	
vaccines	rom different manufacturers	

Residue position	GA/A	GA/C
3	Gly	Val
32	Arg	Gly
58	Asp	Ser
60	Glu	Asp
68	Gly	Asp
84	Gly	Arg
86	Arg	Ser
90	Gln	Lys
112	Thr	Ser
122	Lys	Gln
141	lle	Arg
144	Asn	Asp
163	Asn	Ser
197	Val	Ser
198	Met	Tyr
199	Arg	Ala
200	Asp	Arg
201	Thr	Ser
257	His	Asp
282	Glu	Asn
283	His	Gln
330	Gly	Arg
389	Asp	Glu
390	Val	Phe
391	Asn	Asp
409	Arg	lle
423	lle	Deletion
479	Lys	Asn
481	Thr	lle

The Conn/B vaccine virus was only reisolated from the swab taken at 3 dpv, from vaccinated birds while the virus was reisolated from all four sampling times from the contact-exposed birds (Table 6). From the birds that were vaccinated with Conn/C vaccine, which also contains the Mass virus, we detected IBV by RT-PCR, but Conn specific primers failed to amplify the Conn virus indicating that the viruses detected in the vaccinated and contact-exposed birds were not Conn virus (data not shown). Sequence analysis confirmed that only the Mass virus was reisolated.

S1 sequence analysis of reisolated vaccine viruses

The S1 amino acid sequences from the reisolated viruses were compared to the sequence from the same vaccine virus that was used for vaccination. Analysis of the virus reisolated from Del/C, GA/C, Mass/A and Mass/C vaccinated and contact-exposed birds showed that the S1 gene amino acid sequence did not change after the virus replicated in birds.

The S1 amino acid sequence analysis of vaccine viruses reisolated from Ark vaccinated and contact birds showed a consistent amino acid change of Tyr326 to Asn326 and a deletion of Asn345 (Table 7) in the vaccine virus obtained from all three manufacturers. Analysis of the original vaccine sequences revealed minor nucleotide peaks in otherwise clean sequence at Tyr326Asn, as shown in Fig. 1 as well as for Tyr43His and Ser213Ala, indicating multiple RNA species in the Ark/A and Ark/C vaccines. The deletion at amino acid residue 345 was not detected in the original vaccines. In addition, amino acid changes Ser130Gly (Ark/A and Ark/C) and Thr323Arg (Ark/C) detected in the reisolated vaccine viruses appeared to result from *in vivo* mutations since minor peaks were not detected in the original vaccine sequences, although selection cannot be excluded.

The S1 amino acid sequence for the virus reisolated from Conn/B vaccinated birds had no changes when compared to the virus in the vaccine vial. The Conn/B virus obtained from contact-exposed birds had an amino acid

			<u> </u>		<u> </u>			
Vaccines ^a	3 dpv ^b Tracheal swab		6 dpv Trache	al swab	9 dpv Trache	al swab	13 or 14 dpv	Tracheal swab
	Vaccinated	Contact	Vaccinated	Contact	Vaccinated	Contact	Vaccinated	Contact
Ark/A	_/+ ^c	d	_/+	_	_/+	_	_/+	_
Ark/B	+	_/+	+	+	+	+	_/+	+
Ark/C	_/+	_	+	_	_/+	_	_/+	_/+
Conn/B	+	_/+	_	+	_	+	_	+
Conn/C	_	_	-	_	_	_	_	_
Del/C	+	_	+	+	+	+	_	_/+
GA/A	+	_	_/+	_/+	+	_/+	_	_/+
GA/C	+	+	_/+	+	_/+	+	_	_
Mass/A	_/+	+	+	+	+	+	_	_/+
Mass/B	+	_/+	+	+	+	+	_/+	+
Mass/C	+	_	+	_/+	+	+	_/+	_/+
MassD/C	+	_	+	+	+	+	+	+

Table 6 RT-PCR and real time RT-PCR analysis of vaccine viruses reisolated from the trachea of vaccinated and contact-exposed birds at 3, 6, 9, 13 or 14 days post-vaccination

^a All samples from negative groups were negative.

^b dpv = days post-vaccination.

 c (-/+) Samples negative by RT-PCR/and positive by real time RT-PCR.

 d (-) Samples negative by RT-PCR and by real time RT-PCR.

Table 7 Se	quence of reisolated vaccine v	iruses that showed 51 gene se	equence anterences from the o	original vaccine sequence
Vaccine	Nucleotide position	Nucleotide change	Amino acid change	Amino acid deletion
Ark/A ^a	127 [†]	$T \mathop{\rightarrow} C$	Tyr43His	
	388 [†]	$A \mathop{\rightarrow} G$	Ser130Gly	
	637 [†]	$T \to G$	Ser213Ala	
	976 [†]	$T \rightarrow A$	Tyr326Asn	
	$1033 - 1035^{\dagger}$	AAT deletion		Asn345
Ark/B ^b	976 ^{†,‡}	$T \mathop{\rightarrow} A$	Tyr326Asn	
	1033—1035 ^{†,‡}	AAT deletion		Asn345
Ark/C ^c	127 [†]	$T \! \rightarrow \! C$	Tyr43His	
	388 [†]	$A \rightarrow G$	Ser130Gly	
	637 [†]	$T \rightarrow G$	Ser213Ala	
	968 [†]	$C\toG$	Thr323Arg	
	976 [†]	$T \to A$	Tyr326Asn	
	1033—1035 [†]	AAT deletion		Asn345
Conn/B ^d	172—174 [‡]	AAT deletion		Asn58
	238 [‡]	$T \rightarrow A$	No	
	2 41 [†]	$T \rightarrow A$	No	
	1033 [‡]	$T \mathop{\rightarrow} C$	Ser345Pro	
GA/A ^e	662 [†]	$G {\rightarrow} T$	Ser221Thr	
	988 [†]	$G \to A$	Gly330Arg	
	1152 [†]	$C \rightarrow T$	No	
	1438 [†]	$G \to T$	Lys479Asn	
Mass/B ^f	191 ^{†,‡}	$A\!\rightarrow\! G$	Glu64Gly	
	313 [‡]	$T \rightarrow C$	Ser105Pro	
	1469 [‡]	$C \rightarrow T$	Pro490Leu	

^a Virus isolated at 13 days post-vaccination (dpv).

^b Virus isolated from vaccinated birds at 9 dpv and from contact birds at 13 dpv.

^c Virus isolated at 6 dpv.

^d Virus isolated from vaccinated birds at 3 dpv and from contact birds at 6 and 13 dpv.

^e Virus isolated at 3 and 9 dpv.

 $^{\rm f}$ Virus isolated from vaccinated birds at 3 and 9 dpv and from contact birds at 6 and 13 dpv.

 $^{\dagger}\,$ Vaccinated birds.

[‡] Contact birds.



Figure 1 Chromatograms of vaccine viruses showing major and minor peaks (Tyr326Asn, Gly330Arg, and Glu64Gly) indicating subpopulation selection associated with the reisolated vaccine viruses, and a single peak (Ser221Thr) indicating a mutation was associated with the reisolated vaccine virus.

change of Ser345Pro, a deletion of Asn58 as well as two synonymous nucleotide changes (Table 7). Only one change, Ser345Pro in Conn/B appeared to result from in vivo mutations since no minor peaks were observed in the sequence chromatogram.

The GA/A vaccinated birds had four nucleotide changes that resulted in three amino acid changes, Ser221Thr as shown in Fig. 1, as well as Gly330Arg and Lys479Asn. The

Table 8

Gly330Arg change resulted from selection of a subpopulation since a minor A peak was observed in the chromatogram (Fig. 1), whereas the other changes appear to be the result of *in vivo* mutations. We also observed a synonymous C to T nucleotide change at position 1152.

The S1 sequence in the virus obtained from Mass/B vaccinated and contact birds had a Glu64Gly change. A minor G peak was observed in the chromatogram indicating that the

Reisolated vaccine	Amino acid change	Double peaks ^a
Ark/A (vaccinated 13 dpv) ^c	Tvr43His	Yes

Subpopulations and changes following passage of reisolated virus in embryonating eggs

No No
No
No
No
No
Yes (EP3) ^d
Yes (EP3)
No
Yes (EP4)
No (EP3)
No (EP3)
No (EP3)
No
No
Yes (EP3)

^a Major and minor chromatogram peaks observed in clean sequence data.

^b Viruses were passaged 10 times in 9–11-day old embryonating eggs unless otherwise noted.

^c Virus reisolated from vaccinated or contact-exposed birds on the day indicated (dpv = day post-vaccination).

^d Each egg passage was sequenced. (EP#) = Egg passage number where the change was observed.

change resulted from selection of a subpopulation (Fig. 1). The virus isolated at necropsy from the contact-exposed birds showed additional changes of amino acid Ser105Pro and Pro490Leu (Table 7). Those changes apparently resulted from *in vivo* mutations.

The S1 amino acid sequence analysis of MassD/C vaccinated and contact-exposed birds were not consistent. The virus obtained from the first swabs from the vaccinated birds at 3 dpv and the contact-exposed birds at 6 dpv had amino acid changes that were not observed in the viruses isolated from subsequent swabs (data not shown).

Sequence analysis of reisolated vaccines passaged in eggs

To determine if genetic changes observed in the reisolated vaccine viruses would revert to the original vaccine sequence we passaged them a maximum of ten times in embryonating eggs and examined selected passages for genetic reversion (Table 8). No reversion was observed in the Ark/A, Ark/B, and Ark/C viruses reisolated from vaccinated birds following 10 passages. The Ark/B viruses reisolated from contact-exposed birds reverted from Asn to Tyr at position 326 and an additional Asn at position 345 was observed at pass 3 in embryonating eggs, which was the same as the original vaccine. The Conn/B virus reisolated from contactexposed birds reverted to the original vaccine sequence at position Pro345Ser after 4 passages in eggs but the deletion of Asp58 in the reisolated vaccine was still present following 10 passages. None of the GA/A reisolated vaccine viruses reverted following 3 egg passages. The GA/A virus was only passaged 3 times in eggs due to contamination. The Mass/B detected by RT-PCR from vaccinated birds at 9dpv could not be rescued in embryonating eggs; thus, the 3dpv swabs were passaged in eggs. Reisolated Mass/B from vaccinated and contact-exposed birds did not revert to the original vaccine sequence at position Glu64Gly following 10 passages in eggs. Mass/B virus reisolated at 13 dpv from contact-exposed birds reverted to the original vaccine sequence at position Pro105Ser and Leu490Pro following 3 passages in eggs.

Discussion

In this study, we examined the S1 sequence of 12 vaccines for IBV and compared that sequence data to the same vaccines reisolated from vaccinated and contact-exposed chickens. We focused on the S1 gene for this analysis because it is widely known that the S1 subunit of spike contains serotype specific epitopes, as well as epitopes that induce virus-neutralizing antibodies and S1 is directly involved in attachment to host cells. In addition, it was shown that by far, most genetic changes occur in the S1 gene during adaptation to the host [15–17]. Until now, it was not clear if mutations or selection of a more fit subpopulation was responsible for the changes observed when coronaviruses are attenuated or adapted to a particular host system [16]. Our data clearly shows that selection of a subpopulation of intra-vaccine guasispecies results from infection and replication of IBV vaccine viruses in chickens and that genetic mutation among IBV vaccines also occurs.

With a couple of exceptions where it was necessary to sequence cloned amplicons due to sequencing difficulties, we sequenced RT-PCR products because that data represents all of the genetic diversity within the population of viruses in a given isolate. Sequence ambiguities in otherwise clean sequence data can be used to identify diverse viral RNA genomes within an isolate [5]. Furthermore, the area under the peaks when two or more peaks are present at any single nucleotide position in the sequence chromatogram can be used to identify the major and minor subpopulations. Our data show differences in S1 sequence within the same vaccine serotype from different manufacturers, differences in S1 sequence between different vaccine serials from the same manufacturer, and intra-vaccine S1 differences. Differences in the S1 gene between the same vaccine serotype from different manufacturers likely reflects the source of the original virus strain used to produce the vaccine and the methods and number of passages in embryonating chicken eggs used to attenuate the virus. S1 gene differences between vaccine serials from the same manufacturer are most likely due to the generation of mutations through replication of the viral genome. The World Organization for Animal Health (OIE) guidelines (http://www.oie.int/eng/normes/mmanual/A_summry.htm) for production of IBV vaccines stipulates that no more than 5 passages beyond the master seed can be used to produce the vaccine. IBV has been reported to have a 1.5% mutation rate per year, and SARS-CoV was reported to have a non-synonymous substitution rate of $1.16{-}3.30{\,\times\,}10^{-3}$ per site per year [7,8]. In addition, it has been reported that IBV can adapt to cell culture following as few as 2-5 passages and as many as 25 amino acid substitutions have been documented in the S1 gene after only 7 passages in cell culture [18,19]. Clearly, coronaviruses have the capacity for rapid genetic change, which likely accounts for sequence differences between different serials of the same vaccine.

To our knowledge, this is the first report where both selection of a more fit virus subpopulation and mutations are associated with replication of modified live IBV vaccine viruses in chickens. Vaccine virus subpopulation selection following vaccination, as evidenced by major and minor peaks in otherwise clean sequence data, were observed for Tyr43His, Ser213Ala, and Tyr326Asn changes in reisolated Ark/A and Ark/C vaccine, Gly330Arg in reisolated GA/A vaccine and Glu64Gly in reisolated Mass/B vaccine. Genetic mutations as evidenced by single peaks in clean sequence data and no reversion following egg passage of reisolated viruses, were observed for Ark/A at Ser130Gly, and Ark/C at Ser130Gly and Thr323Arg. Mutations were also detected for GA/A at positions Ser221Thr and Lys479Asn. However, it should be noted that existing viral RNA species could be responsible for apparent genetic mutations if that viral RNA subpopulation was below our level of detection. The level of detection for the RT-PCR test used herein is approximately 500 template copies/ml (unpublished data). No genetic subpopulations for Conn/B at position Ser345Pro were detected in the original vaccine, however; the reisolated vaccine reverted to Ser after only 4 egg passages. This was also observed for Mass/B at positions Ser105Pro and Pro490Leu, which reverted after only 3 egg passages. Reversion after only

a few egg passages suggests that the genetic change is important for growth in eggs but it is not clear if selection of a subpopulation or mutations was responsible for those changes.

A deletion of 3 nucleotides (AAT) coding for Asn345 was observed in all of the reisolated Ark vaccines indicating that that genetic change is important for *in vivo* replication of Ark type vaccines. However, based on sequence data from cloned RT-PCR product, the reisolated Ark/B vaccine reverted following embryonating egg passage indicating that a low-level virus population retaining the 3 nucleotide sequence was being maintained *in vivo*, since it is unlikely, but not impossible, that the virus could acquire an AAT insertion through a recombination event or other mechanism after only a few back passages in eggs. It was necessary to clone the RT-PCR products from the Ark/B vaccine to obtain sequence data of single fragments, presumably because viruses with and without the deletion were being maintained in the population.

Amino acid sequence differences in the IBV spike glycoprotein are responsible for diverse serotypes of the virus, which do not cross-protect [20]. Some IBV vaccines, so called protectotypes, have been shown to provide more crossprotection than other similar vaccines [21]. It is likely that these protector-type vaccines have a more diverse population of viruses capable of simulating a broader immune response in the host. In addition, it has been shown that viral subpopulations can work together to invade and replicate in the host [22,23]. Thus, it seems logical that a vaccine having a more dynamic quasispecies population ought to be more efficacious than one producing fewer mutant sequences. However, molecular evolution can lead to the selection of virulent viruses as well as the emergence of new viral pathogens [9]. A largely undocumented but widely accepted phenomenon of rolling vaccine reactions in commercial chicken flocks occurs when less than 100% of the flock receives the vaccine allowing repeated transmission of the vaccine virus from vaccinated birds to unvaccinated flock-mates. The highly infectious nature of IBV combined with vaccination and re-infection of vaccine viruses in a commercial chicken flock that can contain over 20,000 birds, likely allows the virus to undergo molecular evolution. The evidence for this can be found in the multitude of variant viruses with sequences extremely similar to the vaccines used in the flocks [24].

In summary, it appears that high-titered IBV vaccines contain a mixture of genetic variants and vaccination of chickens results in selection of the most fit viral RNA subpopulations as well as generation of mutations. This allows rapid adaptation to the host ensuring replication and stimulation of an immune response, however; accumulating evidence also indicates that it can lead to persistence and the emergence of new viral pathogens [5,9]. Recognizing selection and mutation of vaccine viruses in the host is important for our understanding of how vaccine viruses can potentially replicate, persist and evolve to cause disease, and applies to all modified live coronavirus vaccines including potential vaccines against the SARS-CoV. This information can also be used to develop safer, higher quality, more targeted modified live vaccines containing viral subpopulations that successfully induce an efficacious immune response.

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