Genotypic characterization of infectious bronchitis viruses from India

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Received 4 July 2005; revised 20 February 2006; accepted 5 April 2006

Infectious bronchitis (IB) is an acute, highly contagious disease of chickens characterized by tracheal rales, coughing, sneezing along with excess accumulation of mucus in bronchi. Other disease manifestations such as decline in egg production and quality, kidney damage, enteritis and even pectoral myopathy have also been observed. IB is of economic importance because it causes decrease in weight gain, feed efficiency, egg production and quality. It is caused by a virus belonging to the genus Coronavirus of the Family Coronaviridae. IB virus (IBV) is not a single homogenous type but occurs in different serotypes. Variant serotypes continue to be associated with outbreaks of disease in many countries. In India, IB vaccinations are carried out using vaccines belonging only to the Massachusetts 41 (Ma41) serotype. However, the vaccine strain used may not always be protective and it becomes necessary to continuously monitor the viruses causing disease. There has been anecdotal evidence of IB outbreaks even in vaccinated flocks. Tissue suspensions were inoculated intra-allantoically into 11-day-old embryonated chicken eggs. The allantoic fluids collected at 48 h post-inoculation (PI) were used in reverse transcription-polymerase chain reaction (RT-PCR) to amplify a part of S1 gene. The partial S1 gene products obtained were sequenced. All the seven isolates had sequences with 94.8-98.8% homology with the vaccine strain, H120. The cross-neutralization tests were used to identify the serotype of the field isolates. All the four isolates showed greater than 50% antigenic relatedness value indicating their classification in Massachusetts’s serogroup. The commercial vaccines conferred protection to the viruses isolated.

Keywords: infectious bronchitis virus, chickens, genotyping, antigenic relatedness, protectotyping

IPC Code: Int. Cl. C12N15/11, 15/13, 15/50

Introduction

Infectious bronchitis (IB) is an acute, highly contagious disease of chickens characterized by tracheal rales, coughing, sneezing along with excess accumulation of mucus in bronchi. Other disease manifestations such as decline in egg production and quality, kidney damage, enteritis and even pectoral myopathy have also been observed. IB is of economic importance because it causes decrease in weight gain, feed efficiency, egg production and quality. It is caused by a virus belonging to the genus Coronavirus of the Family Coronaviridae. IB virus (IBV) is not a single homogenous type but occurs in different serotypes. Variant serotypes continue to be associated with outbreaks of disease in many countries. In India, IB vaccinations are carried out using vaccines belonging only to the Massachusetts 41 (Ma41) serotype. However, the vaccine strain used may not always be protective and it becomes necessary to continuously monitor the viruses causing disease. There has been no extensive genomic characterization of IB viruses circulating in India although there has been serological evidence of the presence of an IB variant, 793/B antibodies. There has been anecdotal evidence of IB outbreaks even in vaccinated flocks in India that prompted us to study the nature of IBV involved in field outbreaks. The virus isolates were characterized genotypically, serotypically and protectotypically.

Materials and Methods

Virus Isolation

Trachea, proventriculus, kidney, ileo-caecal junction or oviduct samples were collected from IB suspected birds or received from clinicians, from different parts of India. Pooled 20% tissue suspensions were inoculated intra-allantoically into 11-day-old embryonated chicken eggs (ECE).

Diagnostic RT-PCR

The allantoic fluids of ECR were collected at 48 h post-inoculation (PI) and used in reverse transcription-polymerase chain reaction (RT-PCR) initially for a portion of the nucleoprotein (N) gene using primers and cycling conditions described earlier. Briefly, total RNA extracted by solution D method from the infected allantoic fluids was reverse transcribed in to cDNA using a cDNA synthesis kit (MBI Fermentas, USA). The cDNA was used with ‘N’ gene specific primers for amplification of external
product (402 bp) with the following cycling conditions: initial denaturation at 94°C for 3 min; 35 cycles of 94°C for 45 sec; 60°C for 1 min; 72°C for 2 min, followed by a final extension of 72°C for 7 min. A nested reaction that amplified a 380 bp product was used using 1-2 µL of the first round reaction and the same reaction conditions with an internal set of primers. A total of 47 samples were screened using these primers.

**Phylogenic RT-PCR**

All the samples positive by N gene nested RT-PCR were passaged in ECE thrice in quick succession and the allantoic fluids were concentrated by centrifugation at 35,000 rpm for 2.5 h (Beckmann, 70Ti rotor). RNA was extracted from the pellet and RT-PCR conducted for a portion of the spike protein (S1) gene using primers and cycling condition described earlier. The sensitivity of amplification of the N and S1 genes was estimated by carrying out RT-PCR from dilutions 10^0 EID_{50} to 10^9 EID_{50}.

The partial S1 gene products obtained were sequenced using an automated sequencer (ABI Prism). A commercial vaccine virus (H120) that is commonly used in the field was also sequenced. The nucleotide sequences were aligned using the MegaAlign programme of the LaserGene software (DNASTAR INC., USA) and subjected to phylogenetic analysis. For comparison, published sequences of IB viruses available in GenBank and the vaccine virus sequenced were included.

**Serotyping of Field Isolates**

The cross-neutralization tests used to characterize the field isolates were as described in the OIE manual. Four field isolates (008, 177, S3, U4) and a vaccine strain H120 and their respective sera were used. Hyperimmune sera were raised in chicks as described previously. Each virus was reacted with its homologous serum and also with panels of serially diluted four other heterologous sera. The end points of the homologous and heterologous neutralization titres were determined as the dilution, which showed no signs of dwarfing (complete neutralization) after incubation of inoculated eggs for five days. Each time a control with 10^2.0 EID_{50} of the respective virus (without antisera) was inoculated into 4 eggs to determine the ability of the diluted virus to cause dwarfing. The antigenic relatedness values (R) were calculated by using homologous and heterologous titres. Two viruses having an 'R' value greater than 50% were considered to be related, whereas those with values less than 50% were considered unrelated.

**Protectotyping of a Field Isolate**

Two groups of six chicks each were inoculated with H120 or Massachusetts serotype (Ma5) vaccines at day old at the recommended vaccine dose of 10^3.0 EID_{50} per bird oculo nasally. One group (of six chicks) was left unvaccinated to serve as challenge controls. At 3 weeks of age all the three groups were challenged with a representative field isolate 008 (10^3.0 EID_{50}) oculo nasally. Four days post-challenge the birds were killed humanely and assessed for protection ensured by the vaccines against the field virus. In each group, trachea from 3 birds was used to assess ciliostasis. An individual chick was recorded as protected against challenge if the ciliostasis score for that trachea was less than 20. For each group, a ‘protection score’ was calculated by the formula:

\[
\text{Mean ciliostasis score of vaccinated/infected group} = \frac{1 - \text{Mean ciliostasis score of challenge controls}}{100}
\]

The maximum obtainable protection score is 100. The higher the score reaching towards 100, higher the level of protection provided by that vaccine. Tracheas from the remaining three birds from each group were used for virus isolation in ECE.

**Results**

**Diagnostic and Phylogenic RT-PCR**

A total of 47 samples were screened using primers specific to amplify the part of N-gene of IBV. Ten samples were found positive by N gene nested PCR and all were amplified using S1 gene primers after consecutive passages and concentration. The sensitivity of the N-gene RT-PCR was found to be 10^3.0 EID_{50} while that of the nested PCR was 10^9 EID_{50} and that of the S gene PCR was 10^14.0 EID_{50} (Fig. 1). Seven isolates and one vaccine virus belonging to H120 strain were used in sequence analysis. The sequence data of other 3 isolates were not of good quality to be used for analysis. The nucleotide sequence of the H120 vaccine virus sequenced in this study was 100% similar to the already available sequence for that virus in GenBank (Accession No. M21970). The other 7 sequences had nucleotide percent similarity with H120 strain ranging from 94.8 to 99.8%. The phylogenetic tree constructed based on aligned sequence data is shown in Fig. 2.
Serotyping

In the cross-neutralization tests, the end point titres of all the 4 isolates tested had less than 2-fold differences with the H120 vaccine sera titres (Table 1). All the four isolates showed greater than 50% antigenic relatedness value indicating their classification in Massachusetts’s serogroup.

Protectotyping

Since all the isolates tested belonged to the same serogroup Mass 41, one representative isolate (008) was selected for use in the in vivo protection experiment. The ciliostasis score of the unvaccinated, challenged group was 22 out of maximum score of 40 obtainable. However, the vaccinated challenged birds had a score of 4.6. The protection scores for both the H120 and Ma5 vaccinated groups were 79%. The tracheal homogenates of the unvaccinated challenged chickens were all positive for virus while those of chickens vaccinated with IBV vaccines were negative.

Discussion

In India, Newcastle disease (ND) has always been the most prevalent respiratory pathogen in chickens. The incidence and isolation of IBV has been rare, probably due to the ubiquitous presence of ND viruses that masked IBV infections. Now extensive vaccination with several types of ND vaccines is routinely being used, bringing the disease largely under control. In this context, IB has started causing great economic losses. In this study, no distinct variant genotype of IBV could be detected. All the seven isolates had sequences of 94.8-98.8% homology with the vaccine strain, H120. The least homology (high percent divergence) was with the isolate IBV 186 (94.8%). All the isolates sequenced had 1-4 nucleotide differences from that of the H120 vaccine virus. Thus it appears that the vaccine viruses are acquiring point mutations and insertions during its spread in the field. Although speculative, it is possible that in due course, the accumulation of these mutations may lead to viruses that may escape protection by existing vaccines. Serotype analysis by cross-neutralization tests also confirmed them as Mass serotype. The commercial vaccines conferred protection to the viruses isolated. Mass serotype based vaccines are used worldwide, though they are protective against the homologous serotype they spread in field, creating a confusing epizootiological problem. The exact discrimination of wild type and Mass vaccine strains is very difficult.

The sequencing results demonstrated the co-circulation of vaccine and wild type infectious bronchitis viruses belonging to Mass 41 serogroup and are further justification for continuous monitoring of circulating strains in order to rationally modify

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Table 1—Antigenic relatedness (R) values of IBV isolates

<table>
<thead>
<tr>
<th>Isolates</th>
<th>Relatedness (%)</th>
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<tbody>
<tr>
<td>H120</td>
<td>100</td>
</tr>
<tr>
<td>008</td>
<td>70.7</td>
</tr>
<tr>
<td>177</td>
<td>100</td>
</tr>
<tr>
<td>S3</td>
<td>50</td>
</tr>
<tr>
<td>U4</td>
<td>70.7</td>
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<td>50</td>
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Fig. 1—Sensitivity of S1 gene PCR: Lanes1-5, 10^2 to 10^5 EID<sub>50</sub>; Lane 6, Negative control; M, 100 bp Marker lane

Fig. 2—Phylogenetic tree based on nucleotide sequence alignment of the S1 gene variable region of different IBV isolates. (Branched distance correspond to sequence divergence).
vaccination strategies to make them appropriate to the field situation. However, there was no evidence of any variant viruses present at least among the few IBV sequenced in this study.

Acknowledgement
The authors thank the Tamil Nadu Veterinary and Animal Sciences University, Chennai for providing all the facilities to carry out this research work.

References
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