A semi-nested reverse transcription-polymerase chain reaction (Semi-N-RT-PCR) was developed and used to detect the S glycoprotein gene of infectious bronchitis virus (IBV) strains and to discriminate H120 vaccine strain from other strains. Viral RNA was extracted from the allantoic fluid of chicken embryos and from tissues of chickens experimentally infected with different strains of IBV. Amplification and identification of the viral RNA was performed using two sets of primers complementary to a region of the S glycoprotein gene in the Semi-N-RT-PCR assay. The pair of primers used in the first PCR consisted of universal oligonucleotides flanking a more variable region of S1-S2 gene. The second primer pair was used in the Semi-N-RT-PCR and was comprised of one of the primers from the first universal pair together with another universal internal oligonucleotide or a oligonucleotide sequence specific for the H120 strain of IBV. The universal primers detected all reference IBV strains and field isolates tested herein. The Semi-N-RT-PCR had high sensitivity and specificity, and was able to differentiate the H120 vaccine strain from other reference IBV strains; including M41 strain. All tissue samples collected from chickens experimentally infected with H120 or M41 strains were positive in the semi-nested RT-PCR using universal primers, while only the H120-infected tissue samples were amplified by the set of primers containing the H120-oligonucleotide. In conclusion, the ability of Semi-N-RT-PCR to detect distinct IBV strains and preliminarily discriminate the vaccine strain (H120) closes a diagnostic gap and offers the opportunity to use comprehensive PCR procedures for the IBV diagnosis.

INTRODUCTION

The infectious bronchitis virus (IBV) is a coronavirus that causes a highly contagious disease in chickens and belongs to group III of the genus Coronavirus of the Coronaviridae family (Cavanagh & Naqi, 1997). The virus predominantly replicates in the upper respiratory tract followed by viraemic spread to various organs, e.g. kidneys, reproductive tissue and lymphoid tissue (Dhinakar Raj & Jones, 1997). Besides, replication also takes place in the intestinal tract (Dhinakar Raj & Jones, 1996). IBV is an important pathogen characterized by a worldwide distribution and many different variants appear continuously in despite of the use of vaccines. Hence, a fast, sensitive and specific virus detection technique is of interest to the poultry industry. Nevertheless, due to high sequence variability among IBV isolates, it is difficult to develop a reverse transcription-polymerase chain reaction (RT-PCR) technique that can be reliably used to detect and differentiate all IBV isolates.

The IBV contains a genome constituted by a single stranded RNA of positive polarity, which consists of 27 kb and codes for three structural
proteins: the spike glycoprotein (S), the membrane glycoprotein (M) and the nucleocapsid protein (N) (Lai & Cavanagh, 1997).

The N protein of IBV is closely associated with genomic RNA and has a highly conserved amino acid sequence. Therefore, the codifying nucleotide sequences are also conserved and very little variation in the N-gene sequence occurs between various strains of IBV. The S glycoprotein is anchored in the viral envelope and is post-translationally cleaved into two proteins designated S1 and S2. In contrast to the N protein, the S protein is very diverse in terms of both nucleotide sequence and deduced primary protein structure, especially in the upstream part of S1, which contains also hypervariable regions (HVs) distributed among less variable and more conserved sequences of this protein (Kusters et al., 1989; Cavanagh, 1995). The most important epitopes that induce neutralizing antibodies are situated in the S protein (Kant et al., 1992). It is thus essential for the development of protective immunity (Ignjatovic & Galli, 1994), and serotypic differences found in IBV are directly associated this protein.

Therefore, solely the detection of IBV strains by molecular diagnostic techniques can obviously be directed to the nucleoprotein gene sequence, while type differentiation can rely on identification of variable regions of S protein (Handberg et al., 1999). Alternatively to this approach, molecular diagnostic techniques can target more conserved sequences present on S1 and S2 genes of IBV. Universal primers can be designed based on these sequences and then used for the detection of IBV by RT-PCR, whereas specific primers can be designed to be complementary to the more variable regions of these genes and used in a nested or semi-nested PCR to differentiate IBV strains from different serotypes (Keeler et al., 1998; Cavanagh et al., 1999).

IBV is conventionally diagnosed by virus isolation in embryonated eggs, followed by immunological identification of isolates. Since two or three blind passages are often required for successful primary isolation of IBV, this procedure can be time-consuming. Alternatively, IBV may be isolated by inoculation in chicken tracheal organ cultures. This method is sensitive (Cook et al., 1976), but is also laborious. Additionally, IBV may be detected directly in tissue of infected birds by in situ hybridization (Collisson et al., 1990) or by immunohistochemistry (IHC) (Yagyu & Ohta, 1990; Nakamura et al., 1991; Chen et al., 1996; Dhinakar Raj & Jones, 1996).

RT-PCR has proved useful in detecting several RNA viruses (Cavanagh, 1993). This technique performed on IBV extracted from allantoic fluid is promising as a diagnostic tool and has been used by Jackwood et al. (1992) and by Adzhar et al. (1996). The fragments generated by RT-PCR were successfully used to differentiate IBV strains by restriction fragment length polymorphism (RFLP, Kwon et al., 1993b), sequencing (Zwaagstra et al., 1992) and hybridization (Kwon et al., 1993a). The development of RT-PCR methods using directly avian tissues is in progress (Kwon et al., 1993a; Cavanagh et al., 1997; Falcone et al., 1997; Jackwood et al., 1997; Cavanagh et al., 1999; Handberg et al., 1999), but these procedures still need to be improved. Although they do detect IBVs properly, there are some drawbacks related to differentiation between virus strains. This problem is particularly important when it is necessary to discriminate among genetically different strains that are classified within the same serotype.

Bird flocks are routinely vaccinated against IB with an attenuated live vaccine containing the strain H120 of IBV, which belongs to the Massachusetts serotype. This results in an additional problem concerning the discrimination of virulent field strains from the vaccine serotype using RT-PCR or even RFLP.

The present study describes the development and evaluation of a general and a strain-specific semi-nested RT-PCR for rapid detection and discrimination between the H120 vaccine strain and other IBV strains, including the strain M41, using organ samples harvested from infected birds or allantoic fluid (AF) of specific pathogen free (SPF) chicken eggs inoculated with IBV.

**MATERIALS AND METHODS**

**Viruses**

Three different reference IBV strains (H120, M41 and Arkansas 99) and one field isolate (A034) were propagated in 10-day-old specific-pathogen-free (SPF) embryonated chicken eggs. The eggs were inoculated by allantoic sac route, as described Owen et al. (1991). Virus strains were kindly provided by Empresa Brasileira de Pesquisa Agropecuária -Embrapa (Concordia, SC, Brazil).

**Oligonucleotides**

The oligonucleotides used to amplify gene S fragments are listed in Table 1. Primers SYU (+) and SYU (-) were described by Yu et al. (2001), whereas two new oligonucleotides, VBIS- and Oligo-H120, were
designed using Gene Runner (version 3.0.5). The anti-sense VBIS- oligonucleotide is complementary to a highly conserved sequence located at the beginning of the S2 gene. Oligo-H120 is complementary to a specific sequence of the H120 strain of IBV and was designed based on the nucleotide sequences published by Kusters *et al.* (1989), as well as on alignment results from Clustal X, followed by analysis using Gene Runner (version 3.0.5).

The primers used to amplify gene S fragments are listed in Table 1, whereas Figure 1 shows the annealing position of each oligonucleotide on S1 and S2. There was no similarity between the used oligonucleotides and other nucleotide sequences present in GenBank database using the BLAST tool of the National Center for Biotechnology Information (www.ncbi.nlm.nih.gov).

**RNA extraction**

Total viral RNA was extracted from 500 µl of allantoic fluid (AF) collected from IBV-inoculated 10-day-old embryonated chicken eggs using Trizol (Invitrogen, Carlsbad, CA). The extracted RNA was resuspended in 12 µl of RNase-free water, treated with diethylpyrocarbonate (DEPC).

**RT-PCR**

A general RT-PCR procedure was carried out according to the instructions provided with SuperScript II H-reverse transcriptase (Invitrogen). The synthesis of the cDNA first strand was performed using 5 µl total viral RNA primed with random hexamers.

The following mix was used in the first PCR: 5 µl of the first strand of cDNA 20 pmol of each primer (SYU+ and VBIS-), 10 X PCR buffer, 0.2 mM each dNTP, 1.0 mM MgCl₂, and 2.0 U Taq DNA polymerase (Invitrogen), and water to a final volume of 50 µl. Cycling conditions were as follows: one cycle of initial denaturation (95°C, 10min); 25 cycles of denaturation (94°C, 1min), annealing (51°C, 2min), and extension (72°C, 3min); and final extension (72°C, 10min).

**Semi-nested PCR with universal oligonucleotides**

The semi-nested PCR was performed with 5 µl of the cDNA produced in the RT-PCR diluted 1:20 in DEPC-treated water, 20 pmol of each primer (SYU+ and SYU), 10 X PCR buffer, 0.2 mM each dNTP, 1.0 mM MgCl₂, and 2.0 U Taq DNA polymerase (Invitrogen) and water to a final volume of 50 µl. The following cycling conditions were used: initial denaturation (95°C, 8min); 25 cycles of denaturation (95°C, 1min), annealing (45°C, 2min), and extension (72°C, 3min); and final extension (72°C, 10min).

**Semi-nested PCR with H120 strain-specific primer**

The ideal annealing temperature was previously determined using a gradient thermal cycler and reactions were prepared as follows.

The semi-nested PCR was performed with 5 µl of the cDNA produced in the RT-PCR diluted 1:20 in DEPC-treated water, 20 pmol of each primer (SYU+ and O-H120), 10 X PCR buffer, 0.2 mM each dNTP, 1.0 mM MgCl₂, and 2.0 U Taq DNA polymerase (Invitrogen), and water to a final solution of 50 µl, using the following conditions: initial denaturation (95°C, 8min); 25 cycles of denaturation (94°C, 30s), annealing (51°C, 2min), and extension (72°C, 3min); and final extension (72°C, 10min).

**Experimental infection and evaluation of diagnostic sensitivity and specificity**

Three groups of SPF White Leghorn chickens were kept in positive pressure isolator units. At three weeks of age, one group was inoculated with a negative virus suspension (SPF-AF) and two groups were inoculated by ocular and intra-nasal routes with 10⁻⁶ (EID₅₀/ml) of the H120 or M41 strains of IBV. Six trachea samples were collected of each group at 5 days after experimental infection. Tissue samples were submitted to virus isolation test (VI) and passed thrice in embryonated chicken eggs. Samples were considered negative or positive based on the absence or presence of virus.

### Table 1 - Oligonucleotides used in Semi-N-RT-PCR.

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence¹</th>
<th>Position²</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>SYU+</td>
<td>5'-TAYTAYTACCARAGYGCYTT-3' 144-163/S1</td>
<td>X04722</td>
<td>YU <em>et al.</em> (2001)</td>
</tr>
<tr>
<td>VBIS-</td>
<td>5'-GGACCTTATCCATACGC-3' 1881-1865/S2</td>
<td>X04722</td>
<td>Designed</td>
</tr>
<tr>
<td>SYU -</td>
<td>5'-TTAGADGTRWAVACAAGRTCACCATT-3' 508-533/S1</td>
<td>X04722</td>
<td>YU <em>et al.</em> (2001)</td>
</tr>
<tr>
<td>Oligo-H120</td>
<td>5'-GAAACACGTATAGAATGCTG-3' 401-382/S1</td>
<td>M21970</td>
<td>Designed</td>
</tr>
</tbody>
</table>

* + = sense. - = antisense. Y, R, D and W represent mixtures of nucleotides C/T, A/G, A/G/T and A/T respectively, for degenerated primers.² Annealing site on S1-S2 sequence, which is indicated by the accession number from GenBank.
of typical embryonic lesions induced by IBV, such as stunting, curling or embryo death (Gough et al., 1988).

Analytical specificity

The analytical specificity of the semi-nested RT-PCR was assessed using different antigen suspensions of homologous IBV strains (M41 and H120), a heterologous reference strain (Arkansas 99) and a heterologous field isolate (A034), as well as non-related viruses such as Newcastle disease virus (NDV/La Sota vaccine strain), infectious bursal disease virus (IBDV/Lukert vaccine strain) and avian pneumovirus (APV/PL-21 vaccine strain).

Analytical sensitivity

A series of 10-fold dilutions (10⁰ to 10⁻¹⁰) of RT products (cDNA) from H120 and M41 strains were used as a substrate in PCR and semi-nested PCR to evaluate the analytical sensitivity of both reactions. The amplified products of each dilution and technique were visualized by ethidium bromide staining after agarose gel electrophoresis.

RESULTS

Universal oligonucleotides evaluation in the semi-nested RT-PCR of the IBV S1 gene

Two degenerate primers (SYU+, SYU-) complementary to S1 regions and an anti-sense universal primer complementary to the beginning of S2 gene were used as “universal” primers in the semi-nested RT-PCR. This procedure allowed for a “general” amplification of the major part of S1 gene (1,737 bp) in the first PCR, followed by the amplification of a smaller fragment of S1 (451 bp) in the semi-nested PCR. Figure 2 shows that a fragment with the expected size (451 bp) was amplified when all four IBV strains were tested (H120, M41, Arkansas and A034).

Use of a specific primer to differentiate H120 strain by semi-nested RT-PCR

Based on differences in the nucleotide sequence, a specific primer was designed for the vaccine strain (H120). This specific primer was used together with one of the two degenerate primers (Table 1) in the semi-nested PCR of the amplified product obtained in the first PCR. An amplified product of 320 bp was
detected only for the H120 strain, differentiating this virus from the other IBV strains, including the M41 strain (Figure 3).

**Analytical Specificity**

The specificity of this procedure is also supported by the observation that only IB viruses were amplified, whilst non-related avian viral pathogens such as Newcastle, Pneumovirus and Gumboro were not detected in the PCR or semi-nested PCR techniques (Figure 4).

**Analytical Sensitivity**

The sensitivity of the PCR and semi-nested PCR was evaluated by using 10-fold dilutions of the cDNA obtained by RT from H120 and M41 strains of IBV. The detection limit of the PCR was $10^9$ and $10^7$ of cDNA, for H120 and M41 strains, respectively, which showed a band of approximately 1.7 kb. In the semi-nested PCR, the detection limit of the 451-bp fragment was $10^{-4}$ and $10^{-5}$ of cDNA for H120 and M41, respectively (Table 2).

<table>
<thead>
<tr>
<th>VBI strain</th>
<th>PCR detection limit</th>
<th>Semi-Nested PCR detection limit</th>
</tr>
</thead>
<tbody>
<tr>
<td>H120</td>
<td>$10^7$</td>
<td>$10^4$</td>
</tr>
<tr>
<td>M41</td>
<td>$10^1$</td>
<td>$10^1$</td>
</tr>
</tbody>
</table>

**Diagnostic Sensitivity and Specificity**

To evaluate the efficacy of the semi-nested RT-PCR, 12 tracheal samples collected from chickens infected with the strains H120 (6 samples) and M41 (6 samples) of IBV were tested. Tissue samples collected from H120 infected chickens were positive in the semi-nested RT-PCR performed with either the “universal” primers or the primer specific for the H120 strain. However, tracheal samples collected from M41 strain were positive only in the semi-nested RT-PCR using “universal” primers, whereas no amplified product was detected in these samples when the semi-nested RT-PCR was performed with specific H120 strain primer. IBV isolation in SPF embryonated chickens was possible from all tracheal tissue samples infected with H120.
and M41 strains. On the contrary, no IBV was detected by semi-nested RT-PCR, nor virus was isolated in tissue samples collected from the negative control birds.

**DISCUSSION**

The frequent emergence of new variants of IBV is one of the major obstacles to the effective control of the infection caused by this pathogen (King, 1988, Wang *et al*., 1994, Wang & Tsai, 1996). Relevant changes in IBV antigenicity and virulence have been reported in several countries (King, 1988, Wang & Tsai, 1996, Wang *et al*., 1997, Di Fabio & Rossini, 2000).

Molecular studies of IBV have shown that genetic and antigenic variants of this virus can emerge, mainly as a result of a few alterations or mutations in nucleotide sequences of the S1 gene, while the majority of the IBV genome remains unaltered (Lai & Cavanagh, 1997). The genetic variability in the S1 subunit of the envelope spike glycoprotein gene represents an adaptive mechanism of the virus to immune selective pressures associated with intensive IBV vaccination and other management practices (Gelb *et al*., 1991). As a consequence, several serotypes of the virus are recognized and additional variant serotypes continue to evolve and cause disease episodes. Moreover, the vaccination of many broiler and layer flocks has been routinely made with a live attenuated vaccine against IB containing the strain H120. This strain belongs to the Massachusetts serotype and represents an additional problem for the discrimination of virulent field strains from this serotype, using RT-PCR or even RFLP techniques. Thus, accurate and rapid serotype determination has a great relevance in controlling infectious bronchitis.

As genetic variation is a quite common event among IBV isolates, the design of PCR primers to detect all IBV isolates is very difficult. Therefore, the selection of appropriate primers to be used in RT-PCR is essential for the final results in terms of sensitivity and specificity of this diagnostic technique. An additional difficulty of such approach is to conciliate, in one basic procedure of RT-PCR, the detection and identification of a given serotype or strain of IBV (Cavanagh *et al*., 1999).

However, comparative sequence analysis of the S1 gene of IBV revealed that there are few conserved regions flanking the variable and hypervariable regions (HVs) of this gene (Adzhar *et al*., 1996, Keeler *et al*., 1998), allowing the construction of universal primers. In the present study, primers complementary to the beginning of S1 gene (SYU+) and to the beginning of S2 gene (VBIS-) were used in RT-PCR in order to detect a significant number of different IBV strains. A similar approach has been reported, in which a pair of universal primers was designed and successfully used in the amplification of the S1 gene of IBV by RT-PCR (Kwon *et al*., 1993, Adzhar *et al*., 1996, Keeler *et al*., 1998). The primers flanked the HV1 and HV2 regions of S1 gene (Keeler *et al*., 1998), or annealed to the beginning of S1 (positive-sense) and S2 (negative-sense) (Kwon *et al*., 1993, Adzhar *et al*., 1996), in a design similar to our RT-PCR, but using different annealing sites.

Our test was designed specially for the detection of IBV, followed by the discrimination of H120 vaccine strain from other IBV strains, including the M41 strain, which is genetic and antigenically closely related to H120. The semi-nested RT-PCR performed with the three universal S1-S2 primers was able to detect the four IBV strains assayed here with high analytical specificity and sensitivity. Two of these strains were originally classified in the same serotype, e.g., Massachusetts, but one is attenuated and used in the live vaccine formulation (H120), whereas the other is highly virulent, particularly for the chicken respiratory tract (M41). Other two IBV strains belong to different serotypes, Arkansas 99 (serotype Arkansas) and a Brazilian field isolate (A034). Conversely, the semi-nested RT-PCR using a combination of the universal S1-S2 primers and the primer H120-OLIGO-generated amplified products only when the H120 strain was assessed and was able to discriminate it from the two heterologous IBV strains, as well as from the M41 strain.

Such results proved that the comparative sequence analysis of IBV S1/S2 genes have, in fact, regions that were conserved among different IBV serotypes, because three of these conserved regions were used to develop “universal” primers for amplifying IBV genomic RNA by semi-nested RT-PCR. The “universal” primers amplified all IBV strains tested. Furthermore, the “universal” primers used here were found to be IBV-specific because they did not amplify genetic material from other avian respiratory pathogens, e.g., Pneumovirus, Gumboro disease virus and Newcastle disease virus.

Although the nucleocapsid and membrane genes provide good targets for the design of “universal” primers that can be used for coronavirus identification, the co-existence of some conserved regions along with several variable regions of the S1 gene nucleotide sequence also permit their use for the design of both “universal” and “strain-specific” RT-PCR primers, as
explained above and performed successfully in this study.

CONCLUSION

The method reported here was successfully used to detect IBV and efficiently discriminated the vaccine attenuated H120 strain from other IBV strains, including the M41 strain.

REFERENCES


King DJ. Identification of recent infectious bronchitis virus isolated that are serologically different from current vaccine strains. Avian Diseases 1988; 32:362-364.


Owen RL. Detection of viral antigen following exposure of one-day-old chicken to the Holland-52-strain of IBV. Avian Pathology 1991; 20:663-673.


