LARYNGOTRACHEITIS: REPRODUCIBILITY OF THE DISEASE AND COMPARISON OF DIAGNOSTIC METHODS

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This paper corresponds to an “extended abstract” selected for oral presentation in the 22nd Brazilian Congress of Microbiology, held in Florianópolis, SC, Brazil, in November 17-20, 2003

ABSTRACT

Infectious laryngotracheitis virus (ILTV) cause mild to severe respiratory disease in chickens, the purpose of our study being to use Brazilian isolate of ILTV to reproduce ILTV disease in chickens by experimental infection and to compare three diagnostic methods (nested polymerase chain reaction (PCR), virus isolation, histopathology) for detection of ILTV. Forty-eight chickens intratracheally inoculated with ILTV and a further 48 with PBS, showing mild respiratory signs 48 hours post infection (PI) but no signs of infection after day 10 PI. Every 2 days PI, six birds were arbitrarily selected from the control and infected groups, sacrificed and the trachea collected. Both the nested PCR and virus isolation detected the virus from day 2 until day 12 PI. However, at day 12 PI, PCR detected ILTV DNA in 100% of the samples while the virus isolation method detected ILTV in only 33% of the samples. Tracheal histopathology showed intranuclear inclusion bodies on days 8 and 10 PI. The results indicate that the field-isolate of ILTV studied by us is of low pathogenicity and that our nested PCR protocol was able to detect positive samples over a longer infection period than many ILTV diagnostic test already described.

Key words: infectious laryngotracheitis virus, avian pathology, diagnosis.

INTRODUCTION

Infectious laryngotracheitis virus (ILTV) is a member of the Herpesviridae family that cause severe or mild respiratory disease in chickens. Various techniques have been described for the detection of ILTV, including the polymerase chain reaction (PCR) (1,2), virus isolation (3,4) and histopathology (4). The distribution of ILTV is worldwide (4). The virus was detected in Brazil in 1974 but since then there have been no outbreaks in Brazil. We have previously described the isolation of some field-virus strains (5), the purpose of the present study being to reproduce the disease with one of the Brazilian ILTV isolates through the experimental infection of chickens and to compare three diagnostic methods (nested PCR, virus isolation and histopathology) for their ability to detect ILTV.

MATERIALS AND METHODS

Viruses

The ILTV positive control was the vaccine strain from Laryngvac of Solvay Animal Health (Charles City, Illinois, USA). The field-isolate used was obtained from layer-hens with respiratory illness collected in the southeast region of Brazil in 2002 (5). The virus was grown and titrated in 9 day-old embryonated eggs.

Experimental infection

Ninety-six broilers from a commercial lineage were reared in an isolation facility. At 6 weeks of age they were divided into two groups of 48 birds (infected and control group). On day zero the chickens were intratracheally inoculated with 0.2 mL the ILTV field-strain containing 0.2 x 10⁶EID₅₀ (median embryo

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infected dose), the control group being inoculated with the same volume of PBS. On days 2, 4, 6, 8, 10 and 12 post infection (PI), six birds were arbitrarily selected from each group, sacrificed and their tracheas collected separately and tested.

**Virus isolation**

Each trachea was ground in a mortar and pestle and suspended in PBS which was then centrifuged to remove cell debris. For each trachea, 0.2 mL of the supernatant was inoculated into the chorioallantoic membrane (CAM) of a 9-day-old embryonated egg which was then incubated for 7 days and the EID$_{50}$ estimated from the number of opaque plaques on the CAM.

**Histopathology**

The CAMs and the tracheas were fixed in 10% formalin. The fixed samples were processed through graded alcohol, embedded in paraffin, sectioned, and stained with haematoxylin and eosin.

**PCR**

The DNA was extracted by the phenol-chloroform method (1). Two sets of nested primers were selected from the published sequence data for the ILTV thymidine kinase (TK) gene (1), the internal primers ILTV/PCR 5’ and ILTV/PCR 3’ (p1 and p2) were as previously described (1) while the external primers (p3 and p4) were selected to increase sensitivity. The first PCR reaction with primers p3 and p4 was performed for 30 cycles, followed by another 30 cycles with the p1 and p2 primers. PCR products were analyzed by electrophoresis in a 1.2% agarose gel stained with ethidium bromide.

**RESULTS AND DISCUSSION**

Experimentally infected chickens showed mild respiratory signs 48 hours PI, including depression and dyspnea, no respiratory signs being observed after day 10 PI. Two birds died between days 8 and 10 PI displaying hemorrhagic tracheal exudates. The results indicate that the ILTV field-isolate is a sample of low pathogenicity. Virus isolation detected ILTV from day 2 to day 12 PI, all tracheas positive by the virus infection assay also producing the expected 647 bp PCR amplification product and thus being positive by the nested PCR test (Fig. 1). On day 12 PI, the PCR was positive in 100% of the samples but the virus infection test was positive only in 33.33% of the samples, syncytia and inclusion bodies being histopathologically detected in the tracheal sections from day 6 and 8 PI. Comparison of the results of the three diagnostic tests performed on poultry infected with ILTV suggested that PCR was the more sensitive. Some authors have described experimental studies which indicate that ILTV is not detected, or is inconsistently detected, 6 days PI by different diagnostic tests (2,3,4). Our results indicate that the nested PCR protocol was able to detect positive samples over a wider infection period than many ILTV diagnostic test already described.

**REFERENCES**