The immunohistochemistry study of lesions due to avian infectious bronchitis (Serotype 4/91) on different tissues in specific pathogen free chicks

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Abstract: Thirty specific pathogen free (SPF) 20-day-old chicks were inoculated by the intra-tracheal (n=15) or oral (n=15) routes with serotype 793/B of IBV, isolated of Iran. Two groups of 15 SPF chicks which as controls received PBS, either by the intra-tracheal or oral route. All the chicks were observed and examined daily for clinical signs. Each day for 5 days three chicks from each infected and control group were post-mortem and examined pathologically changes in the trachea, lungs, kidneys and intestines. Sections of these tissues were prepared, stained and examined microscopically for histopathological and immunohistochemical changes. Grossly, a small amount of clear mucus and slight congestion were found only in the lumen of the trachea and the lungs of the group chicks which were infected with IBV. The kidneys were pale and slightly enlarge. Immunohistochemical examination revealed similar changes in the kidneys of both groups of chicks infected by the intra-tracheal and oral routes. Viral antigens were detected in the infected cells. The viral antigens were apparent prior to the development of lesions and were detected in the cytoplasm of epithelial cells by 3 days post infection with IBV. The results of this study clearly indicate that IBV (serotype 793/B), isolated in Iran by Momayes et al. (2001), is capable of causing lesions in different tissues most severely in the kidneys of experimentally infected chicks. On the other hand, the serotype has a greater affinity and positive tropism for the kidney than to other tissues.

Key words: infectious bronchitis, serotype 793/B, immunohistochemistry, avian diseases.

Introduction

Infectious bronchitis, a viral disease of great economic importance to the poultry industry, affects the respiratory, urinary and reproductive systems of chickens (14). While IB is considered primarily a disease of respiratory system, different IBV strains may show variable tissue tropisms (8,12,16). Chong and Apostolov (1982) demonstrated with using immunofluorescence, that viral antigen was mainly distributed in many organs and virus replication was also confirmed ultrastructurally by Condon and Marshall (1986) (1,4,5). On the other hand, using streptavidin-biotin immunohistochemical technique and monoclonal antibody to an IBV nucleoprotein, IBV antigen was demonstrated in epithelial cells (Owen et al., 1991; Janse et al., 1994) (10).

The present study describes a sequential immunohistochemical investigation of antigen in the trachea, lung, intestine and kidneys of IBV infected chicks. The aim was to elucidate the localization of viral antigen and its relation to lesions, as well as to determine the main target cells and preferential sites of viral replication and tissue tropism of the serotype.
Materials and Methods

Virus: Serotype 793/B (4/91) IBV was isolated from broiler commercial poultry flocks of 2-18 weeks of age, which were suspected to IBV infection (Momayes et al., 2001). The virus was inoculated (0.1ml) into the allantoic cavity of 10 day-old chicken embryo-yonated SPF eggs. Inoculated eggs were checked twice a day. The allantoic fluid was collected from eggs between 4-5 days post-inoculation (pi). However, 1:4 dilution in 0.01 M PBS (pH=7.4) was used as inoculum at a titre of 105 EID50 /0.1ml for infected groups.

Experimental Design: Sixty 20 day-old SPF chicks were divided into 4 groups. The infected groups (A=15, B=15) were inoculated intra-tracheally (A) and intra-orally (B) with virus (0.3ml), while the control groups (C=15, D=15) were inoculated with 0.3ml of 0.01M PBS (intra-tracheal and orally). The infected and control chicks were housed separately and received food and water ad libitum.

In four groups (infected and control groups), clinical signs were recorded daily for 5 days post-inoculation and three chicks of each were killed at 1, 2, 3, 4 and 5 days p. i.

Immunohistochemistry: At first, the tissues samples sectioned in 3-5μm thickness, and then the slides rinsed in xylole and dilutions of alcohol. The slides rinsed in dionized water and placed in Tris-Hcl solution (0.1mol/l, pH=7.2-7.4) twice for 5 min (13).

Sections were first treated with a specific monoclonal antibody (1:600) to Matrix (M) and a specific monoclonal antibody (1:600) to Spike (S) antigens of IBV (30 min incubation in room temperature). After washing in Tris-solution (twice), they were treated with peroxidase-labeled Horse Reddish (45min) and washed again in Tris-solution (twice). Sections were then treated with 0.02% diaminobenzidine (DAB)-Hcl in buffered solution at room temperature for 4min, and after washing in Tris-solution (twice), counter-stained with Mayer’s Hematoxylin. In this method, viral antigens in the cytoplasm of infected cells were stained as yellow-brown spots.

Results

Clinical Signs: Some chicks of the infected groups showed tracheal rales, coughing and gasping at 3 days p. i. The signs were less severe by 8 days p. i. Decreased food consumption, ruffled feathers and whitish-green watery diarrhea were noted from 3 to 8 days p. i. Three chicks of infected groups died at 1 to 5 days p. i. The appetite returned to normal at 2 weeks p. i. and almost all signs disappeared after 2 weeks p. i. There were neither clinical signs nor deaths in the control groups.

Gross Findings: In the infected groups, slight hyperemia was at 3 days p. i. and in the tracheal mucosa with a small amount of catarrhal exudate. The kidneys were pale and swollen that specially found in
Figure 3: Intestine from IBV inoculated chick at 5 days post-inoculation having a severely degenerated IBV antigen-positive tubules (×250, ×400).

Intra-oral inoculation from 5 to 10 days post-inoculation. No gross lesions were seen in other organs; also no comparable gross lesions were seen in control groups.

**Immunohistochemistry of the Trachea:** The distribution of viral antigen in tissues is shown in Table 1. Immunohistochemistry of two infected groups

<table>
<thead>
<tr>
<th>Group inoculated</th>
<th>Intra-tracheally</th>
<th>intra-orally</th>
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</thead>
<tbody>
<tr>
<td>Tissue lesion</td>
<td></td>
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<tr>
<td>Trachea</td>
<td>-</td>
<td>b</td>
</tr>
<tr>
<td>Kidney</td>
<td>-</td>
<td>++</td>
</tr>
<tr>
<td>Intestine</td>
<td>-</td>
<td>+</td>
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<tr>
<td>Lung</td>
<td>-</td>
<td>+</td>
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</table>

- a = Days post-inoculation
- b = Mean severity index from three chicks; -, no change; +, mild; ++, moderate; ++++, severe.
- c = Mean no. of antigen-positive epithelial cells per 5 microscopic fields tissue lesion (×400) from three chicks; -, no change; +, 1 to 9; ++, 10 to 20; ++++, over 20

Figure 1. Viral antigen was detected in the tracheal epithelium cells from 2 to 5 days post-inoculation (Table 1). This method demonstrated that viral antigen was primarily distributed in trachea when using intratracheal inoculation. At days 2 of post-inoculation, viral antigen was scattered in a few epithelial cells. Antigen-positive cells often sloughed off into the tracheal lumen. Viral antigen was found increasingly in epithelial cells up to 5 days post-inoculation.

**Immunohistochemistry of the Kidney:** The distribution of viral antigens in tissues is shown in Figure 2. Viral antigen was detected in the epithelial cells of proximal convoluted tubules (PCT), loop of Henle (HL), distal convoluted tubules (DCT) and collecting tubules (CT) and also in few epithelial cells of Bowman’s capsule from 3 days post-inoculation (Table 1). Positive cells increased in number up to 5 days post-inoculation in both infected groups. Viral antigen-positive cells were swollen, degenerated and desquamated into lumina. Inflammatory cells infiltrated in the vicinity of the antigen-positive parts that equalized with the medullary region of kidney in other animals.

**Immunohistochemistry of the Intestine:** The distribution of viral antigen in tissues is shown in Figure 3. Viral antigen was detected in the epithelium and in a few epithelial cells of crypts from 3 days post-inoculation. Viral antigen-positive cells were frequently degenerated and desquamated into lumina (Table 1).

**Immunohistochemistry of the Lung:** The distribution of viral antigen is shown in Figure 4. Viral antigen was detected in the epithelium and in a
few epithelial cells of the alveolar mucous glands from 2 days p.i. (Table I).

**Discussion**

This study was conducted to demonstrate the viral antigen, frequency and severity of antigen distribution, tissue tropism with using of immunohistochemistry method for the serotype 793/B (4/91), which recently has been reported in the Iran (Momayes et al., 2001). In the present study, it is interesting to note that the infected chicks showed almost similar patterns of clinical signs, gross findings, and distribution of tissues viral antigens. Of course while the chicks infected by the intra-tracheal route had an earlier (from 2 days p.i.) distribution of antigen. Thus the suggestion by Purcell et al., (1976) that the route of infection with IBV may affect the incidence of disease is not supported (11).

The serotype 793/B (4/91) isolated in Iran (Momayes et al., 2001), has a broad tissue distribution that included respiratory, digestive, and urinary tract tissues. The frequent finding of viral-antigen in the epithelium of kidney in the present study, suggests that the virus has a tropism to epithelium, specially in collecting tubules of kidney.

The immunohistochemical findings in the respiratory, digestive and urinary tracts in this study are similar to those described previously on other strains of IBV (Purcell and McFerran 1972, ChenandItakura 1997, ChenandHosi 1996) (2, 4, 6, 7, 10, 13, 17).

In trachea infected by the serotype 4/91, viral antigen was localized in the epithelial layer that deciliated. Also, distribution of viral antigen in epithelium of crypts is shown in intestine. In lung infected by virus, antigen was detected in epithelial cells of the alveolar mucous glands. In kidney, viral antigen was localized mainly in the epithelial cells. Antigen was prominently distributed in the collecting tubules (CT), distal convoluted tubules (DCT) and proximal convoluted tubules (PCT), which showed marked damage of their epithelial cells at the peak of lesions; of course antigen-positive PCT epithelial cells were few. Thus, IBV 4/91 seemed to attack mainly the ducto-tubular tissues in kidney (Figure 5).

In the present study, IBV antigen appeared in renal epithelial cells prior to the development of renal lesions. The virus induced renal lesions which can be considered to be a ducto-tubular interstitial nephritis.

The sequential observations by immunohistochemistry (from 1st-5th day p.i.) support previous evidence that the virus first infects the tracheal and lung mucosa, then replicates in intestine and specially in renal epithelial cells (ChongandApostolov, 1982; Owen et al., 1991; Janse et al., 1994) (3, 4, 9, 10, 15). Presence of the virus antigen seemed to be must correlated with the pathogenesis of histological changes. The findings from this study support the view that IBV 793/B has a greater affinity and pathogenicity for the kidney than to other tissues.

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مقاله ایمیتونو هیستو شمیمیایی ضایعات بیماری برونشت عفونی طیور سویه بر روی بافت‌های مختلف جوجه‌های سالم

سعید مهدی‌ی/ عباس‌تولی/ سید علی پوری‌خس/ رضا متی/ مهردادکش‌الدینی

ویژه تحقیقات سرمو/ ورد ورز/ رازی تهران/ ایران

دانا محمدی/ حمیدرضاکیشانی/ تهران/ ایران

نام/ موسسه‌های تحقیقاتی کرمان/ کرمان/ ایران

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نتیجه‌گیری: نتایج نشان داد که فاکتور SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. اثر SPF بر بافت‌های مختلف جوجه‌های سالم مطرح شده است. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان داد که SPF می‌تواند عامل مؤثری در حفظ سلامتی بیماری برونشت عفونی طیور سویه باشد. این تحقیق نشان D