Studies on recent IBD virus field variant isolates: Genomic identification and differentiation using RT-PCR-RFLP

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In the present study, 3 pooled proventricular homogenates were collected from 3 broiler flocks, of chicken 15 to 30 days old, from Monofia Governorate. The 3 flocks were suffered from low growth rate, poor feed conversion rate, uneven growth and increased mortalities. Necropsy of dead chickens revealed proventriculitis with increased proventriculus size. IBD viral antigen was detected in the pooled proventricular homogenate of each flock by AGPT using reference antibodies against IBDV and RT-PCR technique. No other viruses were detected; such as Reo virus, CAV, NDV, IBV and ALV-J. Further characterization of the IBDV isolates were conducted by RFLP assay on PCR products using MboI and BstO1 restriction enzymes. Results demonstrate that the 3 IBDV isolates are identical in their RFLP pattern and related to the Del/E variant strain of IBDV.

Infectious bursal disease virus (IBDV) causes an immuno-suppressive disease in poultry leading to mortalities and heavy financial losses to the poultry industry. The virus initiates the destruction of lymphatic tissue in the bursa of fabricious. IBDV has also been implicated as a cause of the transmissible viral proventriculitis (TVP) in broiler chickens (Bayyar et al., 1995; Huff et al., 2001 and Newberry, 1996).

Classical strains of IBDV were found to produce transient lesion in the proventiculus of specific pathogen free (SPF) leghorns (Skeels et al., 1998). Flattening of the papillae, haemorrhage, exudate in the mucosal surface and grossly visible promenances on the serosal surfaces were present. Neither proventricular necrosis nor enlargement was detected. Variant IBDV strains play a role in the TVP, so that vaccination of broiler progeny against these strains could be helpful in reducing the incidence and severity of the disease (Dormitorio et al., 2000). In USA, variant strains of IBDV have been proposed as the possible etiology of proventriculitis (Cookson, 2001; Huff et al., 2001 and Giambrone, 2002). In EGYPT Del/E IBDV variant strain; as detected by an antigen capture ELISA kit; was suspected to be the cause of proventriculitis in 46 broiler flocks suffering from proventriculitis (Hussien et al., 2003). The economical impact of TVP could be summarized in higher than avarage mortality, stunted growth, poor feed conversion rate and passage of the undigested feed in the faeces of the affected chickens (McNulty, 1991 and Goodwin, 1993).

The objective of the present study was to detect, identify and characterize IBDV strain that could be potential causative agent of TVP in broiler flocks in Egypt via using RT-PCR – RFLP assay.

Materials and Methods

IBD viruses.

Field isolates. Proventriculi were collected from 3 affected broiler flocks. Samples of each flock were pooled, homogenized in a sterile phosphat buffer saline (PBS) as 10% (w/v), then exposed to 3 cycles of freezing and thawing, sample suspensions were clarified by centrifugation at 1200 xg for 20 min. at 4°C. The supernatant was collected and passed through 0.45µ then 0.22µ filters. Streptomycin (100mg/ml) and gentamycin (50mg/ml) were added to the filtrate. IBDV was detected in the prepared proventricular homogenate by AGPT using reference serum against IBDV (Nassif, 2001). For confirmation of presence or absence of other viruses in the proventricular homogenate. Each preparation was checked for the presence of other viruses by inoculating the suspension into 20, one day old SPF chicks intra-ocularly and orally (0.2ml/chick). Chicks were observed for 2 weeks, sera were collected at the 14th day post inoculation (PI) and tested for the presence of antibodies for IBDV, NDV, reo virus, CAV, IBV and ALT-
J using commercial ELISA kits (IDEXX and KPL). ELISA results indicated presence of detectable antibodies against IBDV only.

**Classical IBD virus vaccine.** D78 vaccinal strain of IBDV (Intervet, Batch No. 011217D) was used. The lyophilized vaccinal strain was reconstituted in 5ml TNE buffer (10mM Tris-Hcl, 100mM NaCl, 1mM EDTA Amresco PH 8).

**Preparation of viral RNA.** Viral RNA was extracted using the standerd procedures (Jackwood et al., 1996 and Jackwood and Jackwood 1997). viral RNA was extracted from proventricular homogenates and the vaccinal strain of IBDV. One volume of each viral sample was extracted with an equal volume of chloroform, the aqueous layer was collected. SDS (Amresco), and proteinase K (Sigma) were added to the final concentration of 0.5% and 1mg/ml respectively. Following incubation at 37°C for 1h, the samples were extracted with equal volume of acid phenol pH 4.3 (Sigma) and then chloroform: isoamyle alc (24:1). Viral RNA was then precipitated with cold ethanol and centrifugation at 10,000 rpm/min. for 10 min. Viral RNA was resuspended in 100µl DMSO (90%). Proventriculi from non infected SPF chicks were homogenized and exposed to RNA extraction and RT/PCR- RFLP assasy.

**cdNA syntheses.** 5 µl of total RNA was used in cDNA synthesis. Viral RNA was denatured at 95°C for 5 min. followed by addition of RT reaction buffer medium containing 1µg of viral RNA, 2µl of 10x buffer RT buffer, 0.5 mM of each dNTPs, 10µ of RNAsé inhibitor (Fermentas Company), 10 pmol. of each specific primer and 4U reverse transcriptase enzyme (Fermentas Company) in total volume of 20µl. The mixture was incubated at 42 °C for 1h. Primers (700-5’ and 700-3’) used in RT-PCR were designd according to (Jackwood and Sommer 1997). These primers flanked 743 bp fragment of VP2 gene from bp 701 to 1444.

**The PCR.** PCR was conducted according to (Nassif, 2001). The optimal reaction mixture contained 1µg RT product, 5 µl of 10x buffer, 200µM of each dNTPs, 30 pmol of each primer, MgCl2 (2-4 mM) and 2.5U Taq polymerase (Fermentas Company). in a reaction containing and. Primers were used at a concentration of. The PCR was performed using thermal cycler (T-gradient PCR, Biometra) in a program that involved initial denature at 94°C for 3 min. followed by 25 cycles of denature at 94°C for 1 min., annealing at 55°C for 1min., extension at 72°C for 1min and final extension at 72°C for 10min. The amplicon was visualized on 1.5% agarose under U.V illumination.

**RFLP assay.** RFLP assay was performed according to (Jackwood and Sommer, 1997). The amplicon was 1st purified using PCR product purification kit, then digested using the restriction enzyme BstO1 and Mbo1 separatly according to the manufacturer's instructions. Briefly, 1µg of amplicon was digested for 1h at 60°C for BstO1 and 37°C for Mbo1, following the incubation the restriction fragment was separated on 1.5% agarose gel utilizing 100bp DNA ladder and 123bp ladder. Purification kit and restriction enzymes were obtained from Fermentas Company.

**Results**

The RT-PCR products from the 3 proventricular homogenates and the D78 vaccinal strain of IBDV were identical to

| **Table (1): RFLP profile of the IBDV field isolates and D78 vaccinal strain.** |
|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Code No. of the strain | BstO1 | BstO1 | BstO1 | BstO1 | BstO1 | BstO1 | BstO1 | BstO1 |
| Field isolate (1) | 424 | 209 | 172 | 154 | 119 | 403 | 362 | 280 |
| Field isolate (2) | | | | | | | | |
| Field isolate (3) | | | | | | | | |
| D78 | | | | | | | | |

* The length of the yielded fragments was in base pairs.

* Shaded boxes designate the presence of a restriction fragment after digestion with BstO1 or Mbo1 enzymes.
each other and were of 743 bp length (Fig. 1). No RT/PCR product was detected from the proventricular homogenate of the negative control SPF chicks (Fig. 1). The results of RFLP assay for the 3 field isolates of IBDV and the D78 vaccinal strain are summarized in Table (1). The RFLP patterns of the 3 IBDV field isolates were similar to each other when digested with Bst01 enzyme and yielded 3 fragments of 424, 172 and 119 bp in length (Fig. 2). On the other hand, the RFLP pattern of the D78 vaccinal strain was differed and 4 fragments of 209, 172, 154 and 119 bp were produced (Fig. 2). The RFLP profiles of the 3 IBDV field isolates; when Mbo1 enzyme was used, were also identical to each other and 2 fragments of 403 bp and 229 bp length were generated (Fig. 3), while 2 fragments of 362 and 229 bp length were produced when the PCR product of the D78 strain was digested with Mbo1 enzyme (Fig. 3).

Results of RT/PCR-RFLP assay denoted that the 3 IBDV field isolates were related to the same IBDV strain which is identical to Del/E variant strain of IBDV.

**Discussion**

The present study reports the detection and identification of IBDV associated with TVP in broiler chicken in Egypt. The detection and identification of IBDV in the present study using RT-PCR/RFLP from the proventricular homogenates confirms that it could be the a potential cause of the TVP. Our results are in agreement with that reported by (Bayyari et al., 1995, Goodwin et al., 1995 and 1996 and Hussein et al., 2003).

In the present study, RT/PCR-RFLP assay was used for detection and identification of the IBDVs from 3 proventricular homogenates in addition to D78 vaccinal strain. The primers were designated to amplify a 743 bp fragment of the VP2 gene from bp 701 to bp 1444. The RT/PCR products of the 3 field isolates of IBDVs and D78 vaccinal strain were of the same size (743 bp length). Similar results were obtained by Jackwood and Sommer (1998).

RT-PCR/RFLP assay is used to differentiate the IBDV strains into 6 molecular groups, IBDV strains within a group were genitically and antigenically related molecular group 1 and 2 contained variant viruses; groups 3, 4 and 6 contained classic viruses, the molecular group 5 contained the Leukert strain viruses, (Jackwood and Jackwood, 1997 a,b and Jackwood, 1998 and Jackwood and Sommer, 1999).
In our study, the results of RFLP assay on 3 field isolates of IBDV; using MboI and BstO1 enzymes; confirms that the 3 isolates are variant strains of IBDV they were identical to each other and related to the molecular group 2 of IBDVs that containing the Del/E variant strain (Fig 2, 3). On the other hand, the RFLP assay of D78 vaccinal strains affirms; as expected; that is related to molecular group 4 that contains the classical strains. Our results are in agreement with that of (Hussein et al., 2003) who detected the same variant strain of IBDV; from broiler flocks that were suffered from proventriculitis in Egypt, using antigen capture ELISA. Many Egyptian authors reported the presence of variant strains of BDV in Egypt; (El-Sanousi et al., 1994, Sultan, 1995 and Metwally et al., 2003) who detected IBDV variant strains in broiler flocks in 58.3% of the tested bursae. The majority of the detected variants, in such study were related to Del/E strain of IBDV. This variant strain is identified at the molecular level for the first time in Egypt. Further investigations are required to investigate pathogenesis, pathogenicity, the immunosuppressive potentiality of this variant strain on molecular level for the first time in Egypt. Trials for preparation of inactivated vaccine from this variant strain is highly commandable.

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