Molecular Detection, Characterisation and Serological Survey of Chicken Astrovirus from Broiler Flocks in Malaysia

Abdullahi Abdullahi Raji, Aini Ideris, Mohd Hair Bejo and Abdul Rahman Omar

1Laboratory of Vaccine and Biomolecules, Institute of Bioscience, Universiti Putra Malaysia, 43400 UPM, Serdang, Selangor, Malaysia
2Department of Veterinary Pathology, Faculty of Veterinary Medicine, City Campus Complex, Usmanu Danfodiyo University, 840212 Sokoto, Sokoto State, Nigeria
3Department of Veterinary Clinical Studies, Faculty of Veterinary Medicine, Universiti Putra Malaysia, 43400 UPM, Serdang, Selangor, Malaysia
4Department of Veterinary Pathology and Microbiology, Faculty of Veterinary Medicine, Universiti Putra Malaysia, 43400 UPM, Serdang, Selangor, Malaysia

ABSTRACT
Astroviruses have been associated with enteric and extra-intestinal disorders in many animal species, including chickens. Here, we describe the detection and characterisation of chicken astrovirus (CAstV) in broilers and its seroprevalence in broiler breeder flocks. Based on PCR protocol, viral confirmation was carried out on clinical tissue samples from broiler chickens suffering from uneven growth and poor performance. The tissues were molecularly detected for CAstV with differential diagnostic testing against the Newcastle disease virus, infectious bronchitis virus, avian nephritis virus, avian rotavirus, fowl adenovirus and avian reovirus. Polymerase gene-based phylogenetic analyses of the twenty samples detected positive for CAstV indicate they belong to Group I and are related to strains from the US, UK, India and Poland. From these 20 samples, CastV could be isolated from 3 samples upon inoculation in 5-day-old specific-pathogen-free (SPF) embryonated chicken eggs (ECE); virus-infected embryos showed dwarfing, haemorrhages, oedema and gelatinous lesions at harvest. The enzyme-linked immunosorbent assay (ELISA)
results revealed a high prevalence of antibodies against CAstV amongst the broiler breeder flocks tested. It is the first study that describes the detection and prevalence of CAstV in broiler chickens and broiler breeder flocks in Malaysia.

Keywords: Broiler breeder, broilers, chicken astrovirus, Malaysia, PCR, polymerase gene

INTRODUCTION

Enteric poultry viruses of the RNA group such as *Rotaviridae, Reoviridae, Picornaviridae, Astroviridae* and *Coronaviridae* are ubiquitous and could cause varying disease conditions in broiler chickens (Shah et al., 2016). The abundance of these RNA families, especially the *Astroviridae*, has been documented in both healthy and sick broilers, persisting within the ages of 2 to 6 weeks (Pantin-Jackwood et al., 2008). The *Astroviridae* family is divided into 2; the mamastrovirus, which affects mammalian species, and the avastrovirus, which affects avian species (Bosch et al. 2012). In previous studies, two astroviruses belonging to the avastrovirus, viz, avian nephritis virus (ANV) and chicken astrovirus (CAstV), have been detected and characterised (Imada et al., 2000; Baxendale & Mebatsion, 2004).

Chicken astrovirus (CAstV) shares familial characteristics with other astroviruses such as naked, small, round-shaped, positive-sense, single-stranded RNA. The genome is 7.5 kb in length and 38 nm in diameter and consists of three genes encoded in three open reading frames (Koci & Schultz-Cherry 2002; De Benedictis et al. 2011). CAstV is associated with uneven growth, runting stunting syndrome (Baxendale & Mebatsion, 2004; Kang et al., 2018), visceral gout and severe kidney disease (Bulbule et al., 2013), locomotor disorder (de Wit et al., 2011) and white chick syndrome (Sajewicz-Krukowska et al., 2016; Nuñez et al., 2020). Historically, CAstV is known to affect the broiler-type of chickens within the first few days of life, causing significant economic losses due to a decrease in feed conversion, high culling rate and reduced slaughter uniformity, with accompanying increased treatment expenses (Smyth et al., 2009). Conversely, in older chickens, especially breeders, the virus causes a temporary yet significant decrease in the hatch rate, with mid-to-late dead-in-shell embryos (Smyth, 2017).

Although establishing the impact of CAstV infections in chickens has been difficult due to the absence of assay, a wide variety of clinical manifestations and gross pathology overlap with other enteric viruses, recent advances in molecular diagnosis have eased its diagnosis. The reverse transcriptase-polymerase chain reaction (RT-PCR) is the gold standard in astrovirus diagnosis (Pérot et al., 2017). Smyth et al. (2009) developed a degenerate primer set capable of amplifying 510 base pairs (bp). The primer pair contains the 24 spacer sequence, a non-coding region between the polymerase genes’ 3’-end and the beginning of *ORF-2* (capsid gene) that could detect all CAstV types (Smyth et al., 2009). The absence of a suitable serological diagnostic tool was addressed with the design of an
enzyme-linked immunosorbent assay (ELISA) kit capable of screening chicken sera for the presence of CAstV-specific antibodies (Skibinska et al., 2015). However, the indirect ELISA kit that is now commercially available can only detect antibodies against members of the Group B CAstV due to a lack of serological cross-reactivity between the capsid genes of Group A and B CAstVs (Smyth, 2017). The kit is valuable for testing breeders before or in-lay for seroconversion against group B CAstVs. Reports have shown that the ELISA kit is useful in longitudinal serosurvey and prevalence studies (Smyth, 2017).

This study described the identification, propagation, and isolation of CAstV from clinical tissue samples submitted by different commercial broiler farms in Peninsular Malaysia to the Avian Diagnostic Unit of the Laboratory of Vaccine and Biomolecules, Institute of Bioscience Universiti Putra Malaysia. Additionally, a serological survey was conducted to determine the prevalence of the virus amongst commercial broiler breeder flocks within Peninsular Malaysia.

**MATERIALS AND METHODS**

**Sample Origin**

Between January 2017 and January 2018, a total of 45 cases from different commercial broiler farms across Peninsular Malaysia with major complaints of uneven growth and poor performance were submitted to the Avian Diagnostic Unit of the Laboratory of Vaccine and Biomolecules, Institute of Bioscience, Universiti Putra Malaysia for molecular screening. History further revealed that the broilers exhibited varying illnesses and lesions on post-mortem examinations. Therefore, chicken astrovirus (CAstV) with extra-intestinal tissue tropism was proposed to cause the outbreak. Initial molecular screening (RT-PCR) with CAstV-specific primers confirmed 22 out of the 45 cases to be CAstV positive. However, an additional investigation on the 22 cases with seven other RNA and DNA enteric viruses’ gene-specific primers (GSPs) confirmed 20 cases to be only CAstV positive. These 20 cases were then considered for further analysis in this study (Table 1).

**Table 1**

*Samples details with background information*

<table>
<thead>
<tr>
<th>No</th>
<th>Sample ID</th>
<th>Farm Location</th>
<th>Age in days (D)</th>
<th>Clinical Case</th>
<th>Tissue(s) Submitted for Screening</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>IBS395/2017</td>
<td>Kedah</td>
<td>18D</td>
<td>Increased mortality, swollen kidney</td>
<td>Kidney</td>
</tr>
<tr>
<td>No</td>
<td>Sample ID</td>
<td>Farm Location</td>
<td>Age in days (D)</td>
<td>Clinical Case</td>
<td>Tissue(s) Submitted for Screening</td>
</tr>
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</tr>
<tr>
<td>2</td>
<td>IBS404/2017</td>
<td>Kedah</td>
<td>NA</td>
<td>Mortality of about 1% daily for two weeks Fibrous perihepatitis, pericarditis and airsacculitis Swollen kidney, urolithiasis in dehydrated birds</td>
<td>Kidney and caecal tonsils</td>
</tr>
<tr>
<td>3</td>
<td>IBS425/2017</td>
<td>NA</td>
<td>58D</td>
<td>Mortality of about 1% daily Fever, swollen kidneys with moderate urate deposits</td>
<td>Kidney and caecal tonsils</td>
</tr>
<tr>
<td>4</td>
<td>IBS433/2017</td>
<td>Setiawan, Perak</td>
<td>33D</td>
<td>Increased mortality ~1% daily. Swollen kidney</td>
<td>Kidney and caecal tonsils</td>
</tr>
<tr>
<td>5</td>
<td>IBS468/2017</td>
<td>Setiawan, Perak</td>
<td>30D</td>
<td>Swollen kidney and discoloured liver</td>
<td>Kidney and caecal tonsils</td>
</tr>
<tr>
<td>6</td>
<td>IBS489/2017</td>
<td>NA</td>
<td>23D</td>
<td>Visceral gout, swollen kidney, enlarged and swollen gall bladder and mild gizzard erosion</td>
<td>Kidney and caecal tonsils</td>
</tr>
<tr>
<td>7</td>
<td>IBS503/2017</td>
<td>Kedah</td>
<td>30D</td>
<td>Mortality of 0.5-1.2% daily</td>
<td>Kidney and caecal tonsils</td>
</tr>
<tr>
<td>8</td>
<td>IBS508/2017</td>
<td>NA</td>
<td>22D</td>
<td>Swollen kidney with urate deposits</td>
<td>Kidney and caecal tonsils</td>
</tr>
<tr>
<td>9</td>
<td>IBS518/2017</td>
<td>Kedah</td>
<td>26D</td>
<td>Stunted birds and mortality Swollen kidney and enlarged gallbladder</td>
<td>Kidney and Caecal tonsils</td>
</tr>
<tr>
<td>10</td>
<td>IBS543/2017</td>
<td>Yong Peng, Johor</td>
<td>21D</td>
<td>Poor uniformity of the flock with somnolence and culling rate up to 1% on day 21</td>
<td>Proventriculus and caecal tonsils</td>
</tr>
<tr>
<td>11</td>
<td>IBS551/2017</td>
<td>Penang</td>
<td>29D</td>
<td>0.2% mortality, Ruffled feathers, Slow growth/moderate uniformity. PM: swollen kidneys; discoloured and enlarged liver</td>
<td>Kidney</td>
</tr>
<tr>
<td>12</td>
<td>IBS555/2017</td>
<td>Penang</td>
<td>24D</td>
<td>Runtng stuntling. PM: swollen kidney, airsacculitis, pericarditis, thin and loosed elastic intestine</td>
<td>Kidney and caecal tonsils</td>
</tr>
</tbody>
</table>
Detection and Characterisation of CAstV in Malaysia

Molecular Detection

Extraction of Nucleic Acids. Nucleic acids (total RNA and DNA) were extracted from the kidney, liver and intestine homogenates (after three series of freeze-thawing) where the supernatant was harvested after centrifugation of the samples in 50 ml Eppendorf tubes at 1,200 x g for 20 mins at 4°C to pellet the debris in an Eppendorf centrifuge (Eppendorf 5810R refrigerated centrifuge, Germany). The supernatant was then passed through a 0.45 μM Minisart® syringe filter (Sartorius AG, Germany). TRIzol™ Reagent (Invitrogen, California, USA) was then used based on the manufacturer’s instructions for extracting DNA and RNA from the samples.

Conventional PCR and RT-PCR. Primer sets of enteric and extra-intestinal virus used in the differential diagnosis included those of the chicken astrovirus (CAstV) (Smyth et al., 2009), avian rotavirus (AvRV) (Day et al., 2007), chicken parvovirus (ChPV) (Zsak et al. 2013), avian reovirus (ARV) (Pantin-Jackwood et al., 2008), avian nephritis virus (ANV) (Day et al. 2007), infectious bronchitis virus (IB) (Adzhar et al., 1997), Newcastle

Table 1 (Continue)

<table>
<thead>
<tr>
<th>No</th>
<th>Sample ID</th>
<th>Farm Location</th>
<th>Age in days (D)</th>
<th>Clinical Case</th>
<th>Tissue(s) Submitted for Screening</th>
</tr>
</thead>
<tbody>
<tr>
<td>13</td>
<td>IBS574/2017</td>
<td>NA</td>
<td>24D</td>
<td>Pale kidney, foci necrotic liver, pale leg, runting and stunting</td>
<td>Kidney</td>
</tr>
<tr>
<td>14</td>
<td>IBS609/2017</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>Caecal tonsils</td>
</tr>
<tr>
<td>15</td>
<td>IBS667/2017</td>
<td>Penang</td>
<td>27D</td>
<td>Marked enlarged kidney. Urate deposit. Increased mortality (0.5% daily)</td>
<td>Kidney, caecal tonsils and Proventriculus</td>
</tr>
<tr>
<td>16</td>
<td>IBS693/2017</td>
<td>NA</td>
<td>36D</td>
<td>Poor uniformity</td>
<td>Caecal tonsils</td>
</tr>
<tr>
<td>17</td>
<td>UPM713/2018</td>
<td>NA</td>
<td>34D</td>
<td>Poor uniformity. Decreased feed intake. PM: swollen kidney, airsacculitis, pericarditis, perihepatitis</td>
<td>Kidney and caecal tonsils</td>
</tr>
<tr>
<td>18</td>
<td>UPM1007/2018</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>Intestine</td>
</tr>
<tr>
<td>19</td>
<td>UPM1013/2018</td>
<td>NA</td>
<td>27D</td>
<td>NA</td>
<td>Caecal Tonsils</td>
</tr>
<tr>
<td>20</td>
<td>UPM1019/2018</td>
<td>NA</td>
<td>23D</td>
<td>Swollen kidneys. Urate deposits. Visceral gout</td>
<td>Kidney</td>
</tr>
</tbody>
</table>

NA= Not applicable
disease virus (NDV) (Pang et al., 2002) and fowl adenovirus (FAdV) (Meulemans et al., 2001). Conventional PCR and RT-PCR were carried out as described elsewhere (Day & Zsak, 2013; Smyth et al., 2009), with necessary modifications.

**Extraction and Cleaning of RT-PCR Positive Amplicons.** The obtained amplicons were electrophoresed at a voltage of 125 for 45 mins on a 1% agarose gel in 50X TAE Buffer (Thermo Fisher Scientific™, Lithuania). They were visualised by Midori Green Advance DNA stain (NIPPON Genetics Europe, Germany) and viewed using gel viewer GelDoc® (Bio-Rad, California, USA). The 510 bp PCR product of the partial polymerase gene (ORF-1b) was gel excised and purified using the ReliaPrep™ DNA Clean-Up and Concentration System (Promega, Southampton, UK). Purified amplicons were outsourced for sequencing (Macrogen Inc., Seoul, Korea).

**Sequence Data and Phylogenetic Analyses.** The sequences were edited and aligned using BioEdit version 7. The basic local alignment search tool (BLAST) on the National Centre for Bioinformatics Information (NCBI) site was then used to compare all 20 ORF-1b amino acid sequences with those of 28 other CAstV isolates sourced from the GenBank. The GenBank accession numbers of the isolates were FJ476304, FJ476309, KT886453, FJ476307, FJ476308, FJ476296, FJ476298, FJ476297, FJ476294, FJ476295, FJ476300, FJ476301, FJ476302, FJ476305, FJ476306, FJ476293, FJ476303 (Smyth et al., 2009), JF832365 (direct submission), KX397576 (Patel et al., 2017), KX397575, KX397575 (Kang et al., 2018), JF414802 (Kang et al. 2012), MN725025, MN725026 and MK746105 (Xue et al., 2020). The trimmed sequences were then subjected to phylogenetic analysis on MEGA X. The tree was generated using the best scoring model (JTT+G) and Maximum-Likelihood (ML) with 1000 bootstrap replicates.

**Propagation of the CAstV Positive Samples in SPF-ECE.** All the 20 positive tissue samples were processed and blindly passaged four times by inoculation into 5-day old specific-pathogen-free (SPF) embryonated chicken eggs (ECE) via the yolk-sac route as described by Erica & Stephens (2016). Healthy SPF-ECE (Malaysian Vaccine Pharmaceuticals) were selected by candling five days post-lay, transferred into a class 2 biosafety cabinet (ESCO Class II BSC, Singapore), disinfected with ethanol on the surface and allowed to dry. A hole that penetrated the shell and egg membrane at the top of the egg in the centre of the air cell was made using an egg punch. Slowly, 0.2 ml of the homogenate suspension was inoculated using a 1cc syringe with a 22 G, 1” needle (Terumo, Philippines). The hole was then glued with wax, and eggs were placed back into the incubator and candled daily for ten days. The death of an embryo within 24 hours was considered to be non-specific. Dead embryos were chilled at 4°C before harvesting and observation of lesions.
Detection and Characterisation of CAstV in Malaysia

Characterisation of Gross Lesions on the Inoculated SPF-ECE. The embryos were observed for lesions associated with viral infections such as haemorrhages, oedema, reddening of the tips of extremities of the limbs, deformities, dwarfin, gelatinous aspects on the embryo and liver discoulouration.

Extraction of Viral RNA from Chicken Astrovirus Inoculated SPF-ECE. Harvested SPF embryos (whole) if embryo mortality occurred within 2 to 5 days post-inoculation (dpi) or liver and intestines if embryo mortality occurred between 8 to 10dpi were homogenised, and all other procedures were conducted as described earlier.

Molecular Detection of CAstV RNA by RT-PCR. The detection was conducted as described previously by Smyth et al. (2009), with some modifications. Briefly, primer pairs targeting the polymerase gene were used in the RT-PCR, and a reaction was carried out using Superscript IV One-Step RT-PCR System. A reaction mix comprising 12.5 μL 2 x Platinum SuperFi RT-PCR Master Mix, 0.2 μL SuperScript IV RT Mix, 2 μL of each of the RNA template, forward and reverse primers to a final concentration of 0.5 μM each, and diethylpyrocarbonate (DEPC) H₂O to a final volume of 25 μL were used. The thermal cycler (Bio-Rad C1000 Touch Thermal Cycler (Bio-Rad, California, USA) was set on the following conditions: reverse transcription at 50°C for 10 min and initial denaturation 98°C for 2 min, followed by 40 cycles of denaturation at 98°C for 10 sec, annealing for 10 sec at 65.5°C, extension 72°C for 30 sec and a final extension at 72°C for 5 min.

Serological Survey Against CAstV Antibodies

Collection of Serum Samples. A total of 420 serum samples (30 sera per house) were collected for serological screening of antibodies against CAstV from 14 breeder flocks representing four commercial broiler breeder farms in Peninsular Malaysia. The age range of the breeders was between 30 and 62 weeks age. Further details regarding history, medication, vaccination and location remain confidential. Serum samples were collected in-line with FAO protocol and stored at 4°C until the analysis.

Serological Screening Using ELISA Kit. A CAstV Group B Antibody test kit based on enzyme-linked immunosorbent assay (ELISA) (BioChek, the Netherlands) was used in determining the broiler breeder antibody titre based on the manufacturer’s protocol. A microplate reader (Biotek EL800, Biotek Instruments, USA) with a 405 nm filter reader was used in reading the plates. A sample to positive (S/P) ratio above 0.70 (titre of 967) was considered positive.
RESULTS AND DISCUSSION

Enteric viruses and their associated diseases cause significant economic losses in poultry worldwide, yet knowledge about them is still lacking. Their effect on chicken and turkey production lines has been reported in several countries, indicating their widespread presence in both healthy and sick young chicks, especially those of the broiler-type and poults (Shah et al., 2016). Here, we described the identification, propagation, isolation and partial (ORF-1b) sequence characterisation of CAstV from tissue samples of broiler chickens suffering from different illnesses in Malaysia between January 2017 to January 2018. This study also followed these analyses with a serosurvey of four commercial broiler breeder farms in Peninsular Malaysia to ascertain the presence and extent of the spread of the virus.

Molecular screening of the total nucleic acids (DNA and RNA) extracted from the 20 tissue samples using virus-specific conventional RT-PCR and PCR assays revealed that all the 20 samples were positive for CAstV but not for other agents screened. The expected amplicon size of 510bp could be obtained from all the positive samples (Figure 1). The primers, which were designed from a conserved region (polymerase gene encoded in ORF-1b), contained the promoter site that aids in the synthesis of CAstV genome subgenomic RNA cleaved in the ORF-2 capsid gene (Smyth et al., 2009), thus, enabling high degrees of specificity and sensitivity.

Amplicons from the 20 positive tissue samples were sequenced to confirm the detection of CAstV. The 20 CAstV positive samples demonstrated the similarity of 76 to 98% and 88 to 99% at the levels of nucleotide and amino acids sequences, respectively. Analysis of the

Figure 1. Amplification of CAstV specific products. Lane 1 and 23, molecular weight markers; Lane 2, negative control; Lanes 3, IBS395/2017; 4, IBS404/2017; 5, IBS425/2017; 6, IBS433/2017; 7, IBS468/2017; 8, IBS489/2017; 9, IBS503/2017; 10, IBS508/2017; 11, IBS518/2017; 12, IBS53/2017; 13, IBS551/2017; 14, IBS555/2017; 15, IBS574/2017; 16, IBS609/2017; 17, IBS667/2017; 18, IBS693/2017; 19, UPM713/2018; 20, UPM1007/2018; 21, UPM1013/2018; 22, UPM1019/2018.
453 bp nucleotide sequence of each of the 20 samples corroborated the findings of earlier reports by Pantin-Jackwood et al. (2008) and Todd et al. (2009a). Phylogenetic analysis displayed two distinct groups containing subgroups. In contrast, the first group (Group I) comprised all the Malaysian isolates and many subgroups; the second group (Group II) contained many subgroups and comprised all the three available Chinese and four of the UK isolates. Hence, the 20 CAstV sequences were sub-grouped in Group I, with isolates from the US, India, South Africa, the UK, and Germany (Figure 2), sharing an amino acid sequence similarity of 95.36 to 99.33% within the sequenced partial polymerase gene. On the other hand, the Malaysian isolates shared a similarity of 88.74 to 90.06%, with Group II CAstV consisting of isolates from the UK and China. Although this classification of CAstV based on groups I and II are no longer considered in CAstV classification, it appears from this study and a previous study by Smyth et al. (2009) that Group I CAstVs could be highly prevalent. Some observations worthy of note in this study were the absence of prior sampling and/or serological screening of the flocks as early as day 0. The 100% detection rate from the tissue samples showed that the infection could have set in at an early age. It seems likely that this infection could have been a carryover in the poultry house that was either shed by earlier broilers (horizontally acquired) or spread vertically by infected chicks. Serological evidence has shown that infections with CAstV are highly prevalent in broiler breeders and their progeny (Todd et al., 2009b).

Gross examinations of the embryos revealed haemorrhages, runting, oedema and gelatinous lesions in the initial passages, with little to no discoloration at the tips of the upper and lower extremities. Dwarfing was conspicuously observed in the fourth passage in chickens that died on 10 dpi (Figure 3). These lesions were mostly observed in IBS503/2017, IBS543/2017 and UPM1019/2018 isolates and were consistent from the first to the fourth passage compared to the remaining 17 samples. The negative control embryos were devoid of lesions through all four passages. The CAstV inocula from the tissue samples were not able to be propagated in-vivo in the SPF-ECE yolk sac, except for three isolates (IBS503/2017, IBS543/2017 and UPM1019/2018) out of the 20 samples screened. The choice of the yolk sac as the route for inoculation was chosen because it was a nutrient-providing centre to the chicken embryo and with a link to the embryo moments after fertilisation (Nuñez et al., 2015). Four passages in SPF-ECE also failed to amplify the CAstV up to detectable levels, which led to the conclusion that the 17 CAstV positive samples could not establish a productive infection except the three isolates. These undetectable samples were not inhibited competitively by other infective enteric viruses such as ARV, AvRV, FAdv-1 and ChPV because they were initially ruled out (based on RT-PCR and PCR screening). Similar challenges have been observed by other researchers in growing isolates of astroviruses in SPF-ECE or cell culture (Smyth et al., 2009; Chamings, 2016; Palomino-Tapia et al., 2020). Although some CAstV strains are readily
propagated in embryonic eggs or cell culture (de Wit et al., 2011; Smyth et al., 2013), the explanations for why these isolates develop while others do not is yet to be established. Strikingly, with embryo mortality observed as early as 4dpi up to 10dpi, macroscopic lesions were characterised by haemorrhages and oedematous embryo with a conspicuous dwarfing of the embryo, especially at 10 dpi (Figure 3). No characteristic lesions were observed on the embryonic membranes. This early embryo mortality pattern and lethality of the Malaysian isolates differ from earlier reports on CAstV isolates but is likely similar to turkey astrovirus (TAstV), which produces an early lethal effect on TAstV-inoculated embryos (Tang et al. 2006).
All the 420 serum samples were positive for CAstV specific antibodies in the serological study. Results from a one-way ANOVA indicated that the means of the antibodies were unequal, \( F(13,406) = 4.798, p = 0.000 \). Similarly, Tukey’s honest significance test showed a statistically significant difference \( (p = <0.05) \) between the age groups. The highest antibody titres were observed at 30, 60 and 62 weeks of age, with titres of 4180±1835.8, 4248±3006.5 and 4020±3516.5, respectively. As a whole, the study has presented that CAstV infection is common amongst broilers and broiler breeder flocks in Malaysia. These seropositivities in the broiler breeder flocks investigated suggest that the virus can occur in adult laying hens, consequently providing the basis of transmission from breeder hens to day-old chicks. The lower levels of seropositivity observed in some flocks tested can result from decreased virus spread on account of low replication and excretion of the virus to the environment or development of protective immunity against the virus as recorded at ages 54 to 58 weeks. Interestingly, instead of a decreased seropositivity at ages 60 and 62 weeks, an increase in titre was observed, thus suggesting that the birds were probably re-infected and not protected against the CAstV infection. The high seropositivity observed here is similar to the findings of Todd et al. (2009b) and Xue et al. (2017), where broiler parent, great-parent, and great-grandparent flocks of older ages exhibited high CAstV specific-antibody titres. Hence, in the absence of a vaccine against CAstV infection, the detection of high levels of antibodies against CAstV in the broiler breeder flocks could be actively acquired as a result of exposure to natural infection.

The identification of CAstV in the broiler breeder flocks demonstrates the potential challenges in eliminating the virus. As such, reports have indicated that some strains of CAstV can be transmitted vertically, with a possible pathogenic effect in chicks (Baxendale 2019).
& Mebatsion, 2004; Smyth, 2017). Consequently, SPF-ECE producers must adopt measures to eradicate CAstV in their flocks to prevent possible contamination of vaccines derived from chick embryos. In addition, serologic screening as a tool aids in demonstrating the flock’s infection and its freedom from infection. Therefore, screening as described here is useful in an extensive testing exercise.

Based on the history of the birds (Table 1) and the prior identification of CAstV from tissues submitted for screening, it is clear that CAstV contributed to the clinical manifestations and lesions observed. The antibody titres recorded in this study indicates that CAstV is present in commercial broilers and broiler breeder flocks in Malaysia. The fact that this was the first study to be conducted, coupled with the lack of a vaccine against the virus, eradicating the virus may not be easy, but vaccinating the breeder flocks that will then transfer maternal antibodies (mAb) to their progeny will be necessary. Although mAb cannot fully protect chicks against CAstV, based on earlier work on ANV (Todd et al., 2009a), the first identified astrovirus in chickens, a vaccine can limit the spread of CAstV infection if developed.

In conclusion, in this study, we detected CAstV based on an RT-PCR assay and characterised the partial segment of the polymerase gene (ORF-1b) from 20 tissue samples of broiler-type chickens belonging to different broiler flocks suffering from varying illnesses in Peninsular Malaysia. These characterised partial polymerase genes were in Group I based on the earlier grouping of CAstV into two groups. Serological screening of broiler breeder flocks suggests high prevalent of CAstV infection based on Group B type CAstV specific ELISA. To the best of our knowledge, this is the first paper that documents the presence of CAstV in Malaysia. Further study on genome sequencing and pathogenicity is currently underway to better understand the characteristics of the CAstV strain circulating in Malaysia.

ACKNOWLEDGEMENTS
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Detection and Characterisation of CAstV in Malaysia


Detection and Characterisation of CAstV in Malaysia


