Phylogenetic analysis of ovine herpes virus-2 (OHV-2) in malignant catarrhal fever infected cattle in Al-Qadisiyah governorate of Iraq

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Abstract

Ovine herpesvirus-2 is a member of the gammaherpesviruses of the herpesviridae, which is the etiologic agent of malignant catarrhal fever (MCF), a significant fatal disease of cattle. MCF disease was diagnosed in native Iraqi cattle of Al-Qadisiyah governorate of Iraq, during the period from April 2014 and August 2016. Twenty-three blood samples were collected from clinically suspected cattle. The presence of the virus in samples was ascertained based on clinical pictures, postmortem examination and molecular assays. Pansystemic involvement included respiratory, digestive, urinary, nervous systems and ocular lesions were described. A molecular analysis based on a tegument protein gene by specific semi-nested-PCR, DNA sequence and multiple alignments of all PCR products confirmed the Ovine herpesvirus-2 (OHV-2) infection, and revealed a single and double nucleotide deletion, insertion and substitutions. Some of these mutations were non-silent, resulting in changes at the predicted amino acids level into viral tegument protein. The phylogenetic analysis showed the disease was caused by two genovariants of OHV-2 including at one cluster and were related to other sequences from others countries was analyzed. MCF is sporadically occurring in cattle in Iraq, the head and eye form is more pronounced form. It has been concluded that study is provides valuable information about the genetic variation among the OHV-2 genotypes in Iraqi cattle. Based on sequence and phylogenetic analysis of tegument protein gene, this paper elucidated genetic relationship between identified Iraqi OHV-2 with other strains detected in other geographical regions. These results provide new information on the epidemiological and genetically of OHV-2 in Iraq.

Keyword: MCF, Iraq, Ovine herpesvirus-2, genovariants, Al-Qadisiyah governorate, Iraq

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التحليل التطوري لفيروس الهربس (OHV-2) في الأبقار المصابة بالحمى الزنلالية الخبيثة في القادسية في العراق

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الخلاصة

بعد فايروس هربس الأغنام أحد أفراد مجموعة كاما هربس العائدة لعائلة الهربس. مسبب هذا الفيروس مرض الحمى النزلالية الخبيثة وهو مرض قاتل يصيب الماشية. تم تشخيص هذا المرض في هذه الدراسة في محافظة القادسية في العراق خلال الفترة من نيسان ٢٠١٤ لغاية أب ٢٠١٦. تم جمع ثلاث وعشرين عينة دم من الحيوانات المشتبه بصحتها حيث تم تأكيد التشخيص عن طريق العلامات السريرية وفحص ما بعد الموت والتشخيص الجنسي للفايروس. وقد لوحظ انتشار واسع للإصابة في الإجهزة الجسم حيث شملت الجهاز التنفسي والبولي والعصبي والعيون. وتحدث الإصابة بحمى النزلية الخبيثة بحالات فردية في الماشية في العراق وهي شكل العين والعين وبعض من أكثر أشكال المرض حدوث. فيما كان التشخيص الجنسي معهداً على تفاعل اللمحة المستقل عن طريق تشخيص أحد الجينات التركيبية للفايروس يسمى tegument protein gene، أظهرت نتائج التشخيص الجنسي وجود حذف وإدخال واستبدال بعض القواعد النتروجينية
Introduction

Classical malignant catarrhal fever (MCF) is one of highly serious and fatal infectious disease of many species of the families Bovidae and Cervidae, including cattle, buffaloes, other wild living ruminants and various species of Suidae (1-4). MCF caused by about ten members of the MCF viruses grouped within ruminant gamma-herpesviruses of the genus Macavirus (5-7). MCFVs causes a significant economic impact on susceptible hosts (8,9), among these viruses, alcelaphine herpes virus-1 (AlHV-1) and ovine herpesvirus-2 (OvHV-2), are highly related antigenically and genetically and cause the major problem and the ubiquitous prevalent (10). Wildbeest associated-MCF (WA-MCF) can be propagated in vitro and relatively well characterized (11,1). The characterization of sheep association-MCF (SA-MCF) has been restricted because the causative agent has never been successfully propagated in vitro yet (2,12).

The disease forms range from acute severe inflammatory disease with a short clinical course to a more chronic syndrome occasionally with skin form (13). The summarized symptoms of MCF in cattle are marked with high fever, depression, anorexia, conjunctivitis, corneal opacity, lymphadenopathy, ulceration and erosions of the upper digestive and respiratory systems, and neurological disorders, in chronic cases weight loss, muscle atrophy and bilaterally symmetric alopecia (13,14).

Previously, the disease was diagnosed based on the epidemiological and clinicopathological findings (3). Virus neutralizing antibodies generated against one MCFV cross react with other members of the genus (10). The development of the PCR-assay using specific primers, coupled with sequencing has allowed for the identification and classification of some organisms like viruses that have been impossible or difficult to identify using traditional virological methods (15).

Unfortunately, the knowledge about the current situation of MCF infection of cattle in Iraq is absolutely absent. The main objective of this study was to identify the relative causes of MCF like disease and deaths in farmed cattle. The secondly to describe a molecular identified OHV-2 based semi-nested PCR of tegument protein gene-based sequencing and phylogeny analysis.

Material and methods

Ethics statement

This study, the owners of the animals gave permission for their animals for examination and necropsy with samples collection.

Animals and sample collections

Between April 2014 and August 2016, a total of twenty-three cattle (15 males 1 to 2 years old and 7 females 8 months to 4 years in except one female calf in 4 months of age) from different parts of Al-Qadisiyah governorate show different clinical signs of MCF like disease. Ten milliliters of EDTA blood sample was collected from each animal, and submitted to the diagnostic laboratory at College of Veterinary Medicine, University of Al-Qadisiyah, Iraq.

Post mortem examinations

Two animals were slaughtered and necropsied for detection the main pathological changes of disease.

Sample preparation

Buffy-coat cells from 23 samples were obtained according to (2). 40 ml lysis buffer consist from (10 mM CHK03, 0.1 mM EDTA, 0.15 M NH4Cl pH 7.2) was added to 10 ml of blood, centrifugation at 8500 g for 10 min. The supernatants were discarded. This step was repeated 2–3 times. The cells pellet was resuspended with 40 ml phosphate-buffered saline (PBS), then followed by second time centrifugation and discard the supernatant, and stored at -20 C until use.

DNA extraction

The viral DNA was extracted from frozen peripheral blood leukocytes (PBL) using the virus DNA/RNA extraction kit (Geneaid, New Taipei City, Taiwan), according to the instructions of the manufacturer.

Semi nested-Polymerase chain reaction (Semi nested - PCR)

Potential Semi nested - PCR assay specific for amplified the highly variable region of OHV-2 tegument gene. The primer sets 556 - forward, (5'-AGT CTG GGT ATA TGA ATC CAG ATG CTC-3') (121692–121722) and 755 reverse, (5'-AAG ATA AGC ACC AGT TAT GCA TCT GAT AAA-3') (21312–121340) were used in first step for
amplification of DNA fragments containing 442 bp, a second step used 2µl of first PCR product template with the 555- internal reverse (5’- TTCTGGGTTAGTGCGAGGAGCTTC-3’)(121484–121510), with the same forward primer (16,17). The final reaction mixture contained 20 mM Tris HCl, pH 8.4, 50 mM KCl, 1.5 mM MgCl2, 200 IM dNTPs, 10 pM of each primer, 1 U Taq polymerase (IDT, Belgium), and 4µl DNA template. Briefly, Thermal cycling conditions were carried out with initial denaturation at 95°C / 5 min, (40 cycles of denaturation 94°C /2 min, 52 °C /45 sec. for annealing step, and 72 °C / 60 sec. for extension), followed by 72 °C / 7 min final elongation, then PCR products were loaded on 1.25 % agarose gel and viewed by a BioRad Gel Doc XR image analysis system.

Sequence Determination and accession numbers
PCR amplicons were sequenced in both directions with a sequencing kit in an automated sequencer in Macrogen Korea. Sequencing primers were the same first step PCR reaction (556F and 775R) primers mentioned above. The sequencing has been deposited in the EMBL/GenBank/DDBJ databases with accession numbers are KY635410.1 and KY635409.1.

Multiple alignment and Phylogenetic Analysis
The consensus sequences were assembled from overlapping fragments of DNA was T-coffee Sequence Alignment version 5.0.9 used for editing, alignment, and translation of the nucleotide sequences. Molecular characterization and phylogenetic analysis of local strains have been undertaken as compare with globally published strains from GenBank. A rooted neighbor-joining tree was constructed by using T-coffee (http://tcoffee.crg.cat/apps/tcoffee/index.html). The robustness of the tree topology was assessed with 1,000 bootstrap replicates as implemented in the program. Percentage values are indicated at nodes of the tree, and translated into predicted protein sequences using the APE-plasmid editor software.

Results
Clinical findings
During 28 months, A total of 20 and 3 animals show severe and relatively mild cases, respectively of MCF were detected until August 2016, all diseased animals was dying after 8 to 20 day were noticed, it exhibits a clinical sign of sever acute to relatively sub-acute (mild) of clinical signs of MCF. The head and eye form was predominant form, keratitis starts from periphery directly to center marked with opacity, sever keratoconjunctivitis, blepharospasm, photophobia, frequently with ocular and bilateral mucopurulent nasal discharge (Figure 1A and B), redness of buccal and nasal mucosa, erosive and necrosis lesions in the mouth papillae especially in chicks and tongue (Figure 2A). Lymph nodes enlargement, high persistent fever (40-41.5°C) were marked signs occasionally with anorexia, digestive system signs seen in 2 males as bloody diarrhea, hematuria revealed in low percentage (one animal). Dullness, shacking of head incoordination and sternal recumbency were a common nervous sign noticed. One adult cow appeared with bilateral corneal opacity with apparent blindness only (Table 1).

Postmortem examination
The necropsy finding revealed ulceration, redness of the tongue and buccal papilla, Petechial to ecchymotic hemorrhages were find within serosal and mucosal surfaces of the mesenteric and internal organs, splenomegaly (Figures 3B), there is sever cystitis and the urinary bladder filled with dark red urine.
Table 1: Clinical manifestations of MCF infected cattle

<table>
<thead>
<tr>
<th>Clinical signs</th>
<th>No. of animals</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fever</td>
<td>23 (100%)</td>
<td></td>
</tr>
<tr>
<td>Keratoconjunctivitis (opacity of corona)</td>
<td>18 (78%)</td>
<td></td>
</tr>
<tr>
<td>Bloody diarrhea</td>
<td>2 (8.7%)</td>
<td></td>
</tr>
<tr>
<td>Ocular and bilateral mucopulent nasal discharge</td>
<td>12 (52.2%)</td>
<td></td>
</tr>
<tr>
<td>Anorexia</td>
<td>23 (100%)</td>
<td></td>
</tr>
<tr>
<td>Blepharospasm</td>
<td>2 (8.7%)</td>
<td></td>
</tr>
<tr>
<td>Blindness</td>
<td>3 (13.03%)</td>
<td></td>
</tr>
<tr>
<td>Erosive and necrosis lesions</td>
<td>5 (21.7%)</td>
<td></td>
</tr>
<tr>
<td>Lymph nodes enlargement</td>
<td>20 (87%)</td>
<td></td>
</tr>
<tr>
<td>Dullness</td>
<td>20 (87%)</td>
<td></td>
</tr>
<tr>
<td>Shaking of head and incoordination</td>
<td>5 (21.7%)</td>
<td></td>
</tr>
<tr>
<td>Sternal recumbency</td>
<td>2 (8.7%)</td>
<td></td>
</tr>
<tr>
<td>Hematuria</td>
<td>1 (4.34%)</td>
<td></td>
</tr>
</tbody>
</table>

PCR

The DNA was sufficiently extracted from PBL of 23 samples were successfully amplified by semi nested-PCR and yielded visible bands as described above, was identified as the cause of MCF disease in cattle (Figure 4).

Sequence analyses of the 442 bp tegument protein gene fragment. Demonstrated the OHV-2 were group under two distanced clusters have highest nucleotide identity (88.24%) among them. In fact, relatively differences were observed in the sequences analyzed from the samples belonging to the same geographical area. The nucleotide sequences that obtained in this study were compared with others viral strains from various countries at the GenBank database using the BLAST program, which revealed the identity ranged from 96.04% to 86.84% (Table 2). The multiple alignments with deduced amino acids were has been amino acids identity 95.7% to 90.1 (Figure 5).

The analysis of these sequences revealed OHV-2 in geographical area of study grouped at two distinctive clusters (called Iraqi1 and Iraqi2). The identity among detected virus in this study was 88.24%, a mild number of nucleotide substitutions in both strains, and deletions at Iraqi1 strain located between were observed in both clusters. And an insertions or frame shift mutations were detected. No other variations like deletions were tack place. The identity between sequences and reference sequences obtained from GenBank database varies from 86.84 to 96.04.

Table 2: demonstrate the nucleotides identities and predicted amino acids identities of partial sequence of tegument protein gene among Iraqi strains and between them and others published references

<table>
<thead>
<tr>
<th>Strains</th>
<th>Nucleotide alignment</th>
<th>Deduced amino acid alignment</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Iraq2</td>
<td>Iraq1</td>
</tr>
<tr>
<td>Iraq2</td>
<td>88.24</td>
<td>90.64</td>
</tr>
<tr>
<td>Iraq1</td>
<td>86.84</td>
<td>90.10</td>
</tr>
<tr>
<td>Canada</td>
<td>92.11</td>
<td>95.05</td>
</tr>
<tr>
<td>Brazil</td>
<td>92.11</td>
<td>96.04</td>
</tr>
<tr>
<td>India</td>
<td>92.11</td>
<td>96.04</td>
</tr>
<tr>
<td>Egypt</td>
<td>92.11</td>
<td>96.04</td>
</tr>
<tr>
<td>Germany</td>
<td>92.11</td>
<td>96.04</td>
</tr>
<tr>
<td>South Africa</td>
<td>92.11</td>
<td>95.70</td>
</tr>
<tr>
<td>Turkey</td>
<td>90.48</td>
<td>92.73</td>
</tr>
<tr>
<td>Magnolia</td>
<td>90.48</td>
<td>92.73</td>
</tr>
<tr>
<td>Italy</td>
<td>90.48</td>
<td>92.73</td>
</tr>
<tr>
<td>United state</td>
<td>88.89</td>
<td>91.30</td>
</tr>
</tbody>
</table>
Figure 3: (A) petechial and ecchymotic hemorrhages of omentum and serosa of abomasum of the right ocular globe fibrinopurulent bronchopneumonia, (B) petechial hemorrhages within the mesenteric, thoracic and pleural surfaces, ulcerative stomatitis and rhinitis, pulmonary edema, lymphadenomegaly and bilateral multifocal hemorrhagic.

Figure 4: PCR amplicons obtained with OHV-2 specific primers from the PBL of MCF like disease infected cattle. Lane Ladder, molecular size markers; Lanes (1 to 5) represent the five Agarose gel electrophoresis of ethidium bromide-stained PCR products amplified from different DNA samples with primers specific for OHV-2 (A) first step PCR of OHV-2 DNA of tegument protein gene, the 422bp corresponding to ladder (B) second step of semi nested- PCR for the OHV-2 of the same gene shows 223bp.

Figure 5: Alignment of predicted amino acid sequences from the OHV-2 DNA tegument protein PCR from MCF infected Iraqi cattle, with ten known MCF viruses. Unsimilar residues between aligned sequences for a given position are colored (green). The additional residue (asparagine) colored (red) in iraq2 strain, Identity between the OHV-2 predicted amino acid sequence and other OHV-2 viruses from various countries is shown in bold adjacent to the ends of the sequences. GenBank numbers (and references) for Canada, Brazil, India, Egypt, Germany, South Africa, Turkey, Mongolia, Italy and United states, KJ705001.1_USA, KX060582.1, KJ658293.1, HM216475.1, KJ420947.1, JX442930.1, KP015737.1, LC203437.1, JN084009.1, HM216481.1, JN084011.1 respectively.
All OHV-2 tegument protein sequences were with mild diversity. The similarity ranged between 95.7 and 86.84% at the nucleotide sequence level and between 95.7 and 90.35% at the deduced amino acid sequence level. Iraq2 strain have three nucleotide bases for coding additional residue of asparagine which are located at 163–165 position of tegument protein gene of one of Iraqi strains, and two amino acid substitutions.

Sequences and phylogenetic analyses of nucleotides and protein of tegument protein gene showed that the strains from Iraq had some diversity from each other by 95% and 94% respectively, diverging by 10 out of the 309 nucleotides and 9 diverging out of 105 amino acids analyzed (Figure 6).

**Discussion**

In Iraq, despite the clinically presence of MCF, there is no any study concerning with cases of the disease in cattle, therefore the evaluation of suspected MCF cases by clinical and molecular methods for detect the causes of the disease which was likely to have contributed directly to the death of cattle in Iraq, so this study is the first report has established the presence of the disease in native cattle of the country. And the use of a diagnostic assay directly targeting a viral component would be of great relevance to confirm the diagnosis (18).

The major clinical signs observed wide fluctuation in severity, almost presence head and eye form and other forms were similar to previous reports of MCF in cattle and other domestic and wild ruminants like buffaloes, bison and deer (19-22). Some cases were no developing the erosive or ulcerative lesions in mouth and oral cavity, these results were agreeing with (23). Animals represent with corneal opacity, blandness and anorexia were also noted by (24).

Analyses of genetic variation of OHV-2 obtained from clinical case samples by looking at important region to address the hypothesis that the highly polymorphic tegument protein gene is a useful epidemiological marker of OHV-2 strain variation, in addition, the use of highly specific primers for conserved regions of viral DNA that flanked the variable region is useful to differentiate among MCF causative members (22,25). These primers have been used previously for detection of OHV-2 DNA (16,17,26,27).

Amplification of the segment of tegument protein gene was attempted in DNA samples from 23 cattle with MCF sporadic presentations were yielded a clear single band of the expected size for each reaction this agreement with (16,24). Others members of the MCFVs are currently detected by the presence of the tegument protein gene (22).

The novel OHV-2- genotypes were identified as the etiology of disease in the Iraqi cattle, and the broad spectrum of clinical signs that may relate to these polymorphisms of virus. This data indicated that the virus present in cattle has some diversity compared to reference OHV-2. The nucleotide identity was similar to previous study of (18) which finds that the sequence identity ranged from 90.9 to 100% by used the same sets of primers.

The results of the partial sequence comparisons and phylogenetic analysis revealed that the strains of OVH-2 circulating within Al-Qadisiyah governorate is at least two new sub genotypes, these some differed from each other genetically by several nucleotides, these differences based a tegument protein gene we analyzed, with four nucleotide substitutions, four nucleotide deletions in Iraq1 strain and
three additions in Iraq2 strain, like this genetic diversity were detected (26). Nevertheless, these Iraqi OVH-2 tegument protein gene sequences derived from cattle are 95.5 % - 88.83% homologous to those derived from cattle in other geographical regions.

The asymptomatic sheep and goat (OHV-2) detected in healthy sheep and goat (unpublished study) would have been the most likely source of infections. In other hand, samples (blood and nasal swabs) were collected from other species of domestic animals. Furthermore, that, the most infections occur by close contact between the carrier animals and susceptible host (26). Several authors reported the transmission of OHV-2 has been occurred when cattle have been separated at least 70 m to5 km from a lamb (26,28,29).

The phylogenetic analysis and the genetic identity among Iraqi genotypes and with those from other countries showed some genomic differences, suggesting that the OHV-2 diagnosed in Iraqi cattle was genetically different from that found in other regions of the world.

By sequencing the tegument protein gene of the OHV-2 isolates, it showed that the isolates were in the clade with the unidentified strains from Germany and Egypt as well as with the OHV-2 from India. Depend on predicted protein sequence comparisons (105 amino acids) of OHV-2 revealed that the strain Iraq2 differed clearly from all known OHV-2 strains detected in cattle but shared 95% similarity with the strains obtained from Germany, Italy and Egypt.

Conclusions

This study considerate a first report about occurrence of MCF in Iraq, and there are at least two different genetic variants of OHV-2 clustered in one group responsible of disease occurrence in Iraq.

References


