Gallid Alphaherpesvirus 2 in the Egyptian Turkeys: Molecular Characterization and Establishment of a Universal System for Phylogenetic Classification

Mahmoud Bayoumia,b Mohamed El-Saiedc Basem Ahmeda Magdy El-Mahdyc Haitham Amera

aVirology Department, Faculty of Veterinary Medicine, Cairo University, Giza, Egypt; bDivision of Biomedical and Life Sciences, Faculty of Health and Medicine, Lancaster University, Lancaster, UK; cPathology Department, Faculty of Veterinary Medicine, Cairo University, Giza, Egypt

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Introduction: Gallid alphaherpesvirus 2 (GaHV-2) is a highly contagious oncogenic virus that causes Marek’s disease in chickens and occasionally in turkeys. Among 100 genes identified in GaHV-2 genome, the Meq gene appears to involve viral virulence, oncogenicity, and genetic diversity. Despite the use of Meq gene sequences in phylogenetic classification of GaHV-2 strains circulating in many countries worldwide, no integrated system exists yet. Methods: Turkeys from 2 commercial Egyptian farms were presented with signs of dullness, dehydration, and emaciation. Samples prepared from the internal organs were examined by histopathology and immunohistochemistry. Pools of the internal organs were analyzed by PCR for identification of GaHV-2, avian leucosis virus, and reticuloendotheliosis virus. The Meq gene of an Egyptian strain was sequenced and analyzed in comparison to 40 reference strains for generation of a universal system for phylogenetic classification of GaHV-2 strains. Results: Gross and histopathological examination revealed grayish-white soft masses in the internal organs characterized by diffuse infiltration of pleomorphic neoplastic cells. All lymphoma cells were identified as T-lymphocytes of CD3+ phenotype. Samples of both farms were only positive for GaHV-2 by PCR. Sequence analysis of the Meq gene has classified the current turkey strain as related to the Egyptian strains identified in chicken in 2012. A universal phylogenetic system for classification of GaHV-2 strains into 4 clusters was proposed. The vaccine strains were all grouped in cluster 2, and most of the classical American strains belonged to cluster 4. Cluster 1 was further divided into 3 subclusters (1.1–1.3). Conclusion: GaHV-2 was identified in turkeys for the first time in Africa and the Middle East. Sequence analysis of the Meq gene of the Egyptian strain along with a wide array of the global strains has enabled the construction of a novel phylogenetic classification system.

Introduction

Marek’s disease (MD) is one of the most economically devastating infectious diseases of poultry worldwide. The disease principally affects chickens and infrequently involves turkeys [1–3]. However, reports of MD in turkeys...
are increasing in the last few years [4, 5]. It causes high mortalities among chickens of different age groups, decreased performance, and immune suppression with increasing evidence of vaccination failure [6–8]. MD is caused by Gallid alphaherpesvirus 2 (GaHV-2), which is the prototype member of genus Mardivirus of subfamily Alphaherpesvirinae, family Herpesviridae, and order Herpesvirales [9, 10].

GaHV-2 is an oncogenic highly contagious virus that was classified into 3 serotypes based on antigenic and genetic variations. Serotype 1 (prototyped by CVI988/RISPENS strain) is mostly used as an efficacious live vaccine against MD [10]. Besides, pathogenic strains (mild, virulent, very virulent, and very virulent plus) causing MD are also included [4, 11, 12]. Likewise, serotype 2 (prototyped by SB-1 strain) was used for immunization alone or in combination with MDV serotype 3 [13, 14]. Serotype 3 (turkey herpesvirus; HVT) is nonpathogenic to both chickens and turkeys and is frequently utilized as a heterologous live vaccine against other serotypes [10, 15, 16]. The virus primarily replicates in both B- and T-cells (early cytolysis), and later establishes latent infection in CD4+ T-lymphocytes, inducing T-cell lymphoma in visceral organs, skin, and peripheral nerves [9, 10].

The genetic basis of GaHV-2 pathogenicity, virulence, tumorigenicity, and host range are not yet fully understood [17, 18]. The viral genome encodes for >100 proteins, among which Marek’s EcoQ (Meq) is a basic leucine zipper (b-ZIP) that has the characteristics of a potent transactivator oncoprotein [19, 20]. Meq protein is the strongest candidate in the protein array involved in T-cell transformation caused by serotype 1 viruses [21]. Mutation in the Meq gene has confirmed its role as a potential determinant of virus virulence and oncogenicity [11, 12, 22]. The presence of certain motifs in the transactivation domain of the Meq protein has been reported to associate with the virus pathogenicity in chickens. For instance, the presence of 4 consecutive prolines (PPPP motif) was suggested to correlate with the low pathogenicity of vaccine strains, which have at least 7 PPPP motifs [11, 12]. These evolutionary markers are valuable tools for prediction of the pathogenicity of MDV strains, considering that the in vivo pathotyping of the ever-rising MDV strains in specific-pathogen-free chicks has limited practicality due to several technical and monetary issues.

Meq gene is also associated with considerable genetic diversity and is mostly utilized as a preferred target for phylogenetic studies [23]. Despite the availability of several phylogenetic classifications for GaHV-2 strains circulating in many countries worldwide [8, 9, 18, 23, 24], a number of limitations still exist including (1) lack of representation of turkey isolates (2) use of short inconsistent regions of Meq gene (3) random inclusion of viral strains (4) application of different system for nomenclature of strains used in analysis. These limitations precluded the generation of a universal system for the classification of GaHV-2 strains circulating globally.

MD outbreaks have been documented in both vaccinated and unvaccinated chicken flocks from different regions all over the world [7, 9, 17, 20, 25, 26]. The disease was also incriminated to induce sporadic outbreaks in turkeys in Europe [4, 27–30], USA [5], and Asia [2]. No data are available on the circulation of GaHV-2 in turkey in Africa and the Middle East. The genetic characteristics of the turkey isolates remain to be identified. In this study, GaHV-2 was identified in turkey farms located in 2 Egyptian provinces in 2018. The virus identity was confirmed by histopathology, immunohistochemistry, PCR, and DNA sequencing. Additionally, a universal system for phylogenetic classification of GaHV-2 strains was proposed for further validation.

**Materials and Methods**

**Clinical Specimens**

Turkeys from 2 commercial farms located in Fayoum and Minia governorates in Egypt (95 and 135 days old, respectively) were presented with signs of depression, dehydration, and a decrease in body weight. Guidelines for sample collection and animal use in research were followed according to the Institutional Animal Care and Use Committee, Faculty of Veterinary Medicine, Cairo University. Samples were collected from the liver, kidneys, spleen, proventriculus, pancreas, sciatic nerves, and intestine of dead birds (10 birds for each farm). The tissue specimens were divided into 2 portions; 1 was fixed for histopathological examination and the second was kept frozen at −80°C till use in genetic analyses.

**Histopathological Examination**

Tissue samples were prepared for histopathological examination by fixation in 10% neutral buffered formalin, rinsing in distilled water, dehydration in ascending grades of methanol, clearance in xylene, and finally embedding in paraffin. Tissues were cut into 4-μm-thick sections and stained with hematoxylin and eosin. The images were captured at different magnification powers: 40, 200, 400, and 600 × using an Olympus BX43 microscope equipped with an Olympus digital camera (Olympus, Tokyo, Japan).

**Characterization of Neoplastic Cells**

The neoplastic cells were determined by immune-histochemical technique using an EnVision™ FLEX detection system (Dako, Santa Clara, CA, USA) according to the manufacturer’s guidelines. In brief, the paraffin-embedded sections were dewaxed and rehydrated, and the epitopes were exposed by heat-induced epitope
retrieval through PT-link apparatus as described in [31]. The endogenous peroxidases were blocked by incubation in peroxidase blocking reagent. The tissue sections were incubated overnight with primary mouse antibodies against CD3 and CD20 as markers for T- and B-lymphocytes, respectively [17]. After washing, tissues were incubated for 1 h with dextran coupled with horseradish peroxidase and goat anti-mouse antibodies. The immune reactivity was revealed by the application of 3, 3′-diaminobenzidine tetrahydrochloride and counterstaining with Mayer’s hematoxylin. The positive reaction was indicated as a brown stain under the light microscope.

PCR Identification of Tumor Viruses

Total DNA was extracted from the different tissue samples using the GF-1 total genomic DNA extraction kit (Vivantis, Malaysia) according to the manufacturer’s protocol. GaHV-2 was identified in the DNA extract by amplification of the partial ICP4 gene sequence using GaHV2-F and GaHV2-R primers and was confirmed by amplification of the entire Meq gene sequence using GaHV-2 Meq-F and GaHV-2 Meq-R primers (Table 1). The pro-viral DNA of avian leucosis virus (ALV) and reticuloendotheliosis (REV) was identified using ALV-F and ALV-R, and REV LTR-F and REV LTR-R primers, respectively (Table 1). Further confirmation of REV was achieved by amplification of an 855-bp fragment of the env gene using REV env-F and REV env-R (Table 1). The amplification was performed using EmeraldAmp GT PCR Master Mix (Takara, Kusatsu, Japan) in Gene-Amp 9700 thermal cycler (Applied Biosystems, Foster City, CA, USA). PCR was conducted as follows: initial denaturation at 95°C for 5 min, 40 cycles at 95°C for 30 s, 55°C for 30 s (47°C in ALV and 57°C in REV), and 72°C for 1 min, and a final extension at 72°C for 10 min. Specific PCR products were separated by electrophoresis in 1% agarose gel stained with ethidium bromide, visualized by a Gel Doc XR gel documentation system (BioRad Laboratories, Milan, Italy), and purified using the Gel/PCR DNA Fragments Extraction Kit (Geneaid, Taiwan).

DNA Sequencing and Phylogenetic Analysis

The purified PCR product of the GaHV-2 Meq gene of a single sample pool, designated as Fayoum-1 2018, was sequenced on both strands using the specific PCR primers at Macrogen Inc (Seoul, South Korea). Contigs were edited and assembled using Bioedit program, version 7.2.5 (Ibis Biosciences, Carlsbad, CA, USA), and the final sequence was deposited in GenBank with the accession number MK261778. A total of 40 Meq gene sequences were retrieved from GenBank for sequence and phylogenetic studies. Multiple alignment of the corresponding sequences was performed using Clustal W algorithm of the MegaAlign program (DNAStar, Madison, WI) for analysis of percent identity, divergence, identification of mutation hotspots, and prediction of amino acid alterations. The proline (P) content and number of PPPP motifs located in the Meq protein were also evaluated. The phylograms were constructed using the maximum-likelihood method in MEGA 6.0 software. Bootstrapping was calculated at 1,000 replicates to confirm the robustness of the phylogenetic trees.

Table 1. Oligonucleotides used for PCR identification of tumor viruses

| Primers     | Sequence (5′-3′)                                         | Fragment size (bp) | Reference |
|-------------|---------------------------------------------------------|--------------------|-----------|
| GaHV2-F     | GGATCGGCCACCCACGATTACTACC                                  | 313                | [32]      |
| GaHV2-R     | ACTGCGCTCACAAACCCATCTGAC                                  |                    |           |
| GaHV2 Meq-F | ATGTCTCAGGAGCCAGGAGCAC                                     | 1,020              | [7]       |
| GaHV2 Meq-R | GGCTCTCCGTCACCTCTG                                       |                    |           |
| ALV-F       | AATTCTGGTGTGAAATAG                                       | 436                | [5]       |
| ALV-R       | AGTTGTCAAGGAATCGA                                        |                    |           |
| REV LTR-F   | GCGCTGGCTGCTGCAAATG                                      | 200                | [31]      |
| REV LTR-R   | TCCGATCTCGTTTGTGTTGACTG                                   | 855                |           |
| REV env-F   | CCACCGGGTCAATAGATGCAAATG                                  |                    |           |
| REV env-R   | AGTGGCTTTGTGACTGCAGGAGACATA                              |                    |           |

bp, base pair; GaHV2, Gallid alphaherpesvirus 2; Meq, Marek’s EcoQ; ALV, avian leucosis virus; REV, reticuloendotheliosis virus.

Results

Gross and Microscopic Lesions

Postmortem examination of dead turkeys has revealed the presence of soft masses in the majority of internal organs. Tumors of kidneys and liver appeared focal in distribution, grayish white in color with variable-sized neoplasms scattered on renal lobes and hepatic surface (Fig. 1a, b). Spleen appeared larger in size (at least twice) with rounded borders and grayish coloration (Fig. 1c). In some examined cases, the spleen showed a mottled appearance. Enlargement in sciatic nerves was remarkable. Histopathological examination showed lymphomatous infiltration of small- to large-sized pleomorphic populations of neoplastic cells with hyperchromatic and mitotic activities in visceral organs and sciatic nerves (Fig. 1d–g). The neoplastic cells appeared multifocal and diffused in distribution with effaced and re-
placed hepatic parenchyma with various necrobiotic changes (Fig. 1d). Furthermore, pleomorphic neoplastic cells were infiltrated between renal tubules and pancreatic islets causing distortion of architecture and atrophy. Moreover, intestinal lamina propria was thickened with diffuse infiltration of heterogenic lymphocytes (Fig. 1e). Additionally, proventricular lobules were infiltrated with a dense population of pleomorphic lymphoid cells between glands (Fig. 1f). Notably, sciatic nerve fibers showed the characteristic type-B nerve lesion that was disrupted by lymphoplasmacytic cell infiltration combined with cell edema and demyelination (Fig. 1g). Immunohistochemical staining identified all lymphoma cells as belonging to the CD3+ phenotype and negative for CD20. CD3-positive staining was distributed over the sections of neoplastic cells infiltrating the liver (Fig. 1h), intestine (Fig. 1i), proventriculus (Fig. 1j), and sciatic nerve (Fig. 1k).

**Fig. 1.** Photomicrographs showing gross and microscopic representation of Marek’s disease lesions in different internal organs of turkeys. Bilateral multifocal variable-sized grayish white foci on renal lobes of the kidneys (a), multifocal discrete white foci in the liver (b), diffuse enlargement in spleen (c), multifocal infiltration of neoplastic lymphoid cells in the liver (black arrows) that replaced the hepatic parenchyma. The inset shows pleomorphic neoplastic cells with mitotic figures (red arrowheads) (d), diffuse infiltration of mixed lymphocyte populations in the lamina propria of the duodenum (e), neoplastic lymphocytic aggregations between the glands of the proventriculus (f), type-B nerve lesion in the sciatic nerve with marked infiltration of pleomorphic neoplastic cells, inflammatory lymphocytes, edema, and demyelination (g), immunohistochemical staining using anti-CD3 antibodies in the liver showing multifocal distribution of neoplastic T-lymphocytes (black arrows) (h), with cytoplasmic reactivity (red arrowheads, insert). Immunohistochemical staining using anti-CD3 antibodies showing positive reactivity of the neoplastic lymphocytic cells in the intestine (f), proventriculus (j), and sciatic nerve (k).

**Identification of the Viral Cause of Neoplastic Lesions**

Pools of the internal organs of both turkey farms were analyzed for 3 possible tumor-causing viruses of poultry, namely, GaHV-2, REV, and ALV, using PCR. GaHV-2 was identified in both sample pools using serotype-specific primers that target a 313-bp fragment of the ICP4 gene. Positive amplification of both samples was verified against serotype I GaHV-2 (Rispens CVI 988 strain) as a positive control and serotype III GaHV-2 (HVT) as a negative control. The entire sequence of the Meq gene was further amplified to differentiate between oncogenic and nononcogenic strains of GaHV-2. The 2 samples produced PCR amplicons of 1,020 bp (an indicator of oncogenic strains), while the nononcogenic Rispens strain generated a longer band of 1,200 bp. Although samples were also positive for REV by amplification of the LTR fragment, no such amplification was demonstrated with PCR targeting the env gene. All samples were negative for ALV.
Phylogenetic Analysis and Genotypic Clustering of GaHV-2 Strains

In a trial to unify the genotypic clustering of GaHV-2 globally, a collection of 40 reference strains identified from chickens and turkeys over a diverse of spatial and temporal ranges was included in the sequence and phylogenetic analysis. These strains were chosen to represent the different phylogenetic clusters and pathotypes described before [8, 9, 17, 18, 23–25]. A large fragment of the Meq gene (924 bp) of the reference strains and Fayoum-1 2018 strain was analyzed using the maximum-likelihood method with bootstrapping of 1,000 replicates. The phylogenetic classification has indicated the separation of GaHV-2 strains into 4 clusters: clusters 1–4 (Fig. 2). The vaccine strains (e.g., CVI988, 3004, 814) were all grouped in cluster 2, whereas the Indian and Japanese strains identified in 2010 and 2011 were members of cluster 3, and most of the classical American strains belonged to cluster 4. Cluster 1 was further divided into 3 subclusters (1.1–1.3). The average homology range between members of the same cluster was 98.5–98.9% and 95.4–97.1% at the nucleotide and amino acid levels, respectively, while the homology within the subclusters ranged from 98.6 to 99.6% (nucleotides) and from 96.7 to 99% (amino acids). All Egyptian strains including Fayoum-1 were grouped within subcluster 1.1 with an overall homology of 99.4–99.6% and 98.7–99% for nucleotides and amino acids, respectively. Another turkey strain recently identified in Italy was grouped within subcluster 1.2.
The sequence Fayoum-1 2018 Meq gene/protein was further studied for identification of distinct characteristics. For technical constraints, complete sequence data of the Meq gene was not obtained. Instead, a large fragment (924 bp) that lacked the first 45 and the last 51 bps of the entire gene was analyzed. Multiple alignment of the partial Meq protein sequence among representative members of the 4 clusters has revealed an overall high degree of sequence conservation. The marked sequence variation was an insertion of a stretch of 60 amino acids ob-

Molecular Characterization of the Turkey GaHV-2 Meq Protein

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served in cluster 2 strains, which brings the Meq protein longer in vaccine than in virulent strains. Cluster-specific sites were not observed except for residue 80 (tyrosine in cluster 1 and aspartic acid in other clusters). Likewise, subcluster-specific sites were only evident in subcluster 1.2 that share 2 amino acid substitutions: A88T and Q93R. All Egyptian strains possess a unique amino acid substitution E263D, whereas the turkey strain Fayoum-1 2018 is the only one that shows the mutation record E54K (Fig. 3). To predict the pathogenicity of the identified strain, the PPPP motifs were counted, and only 3 motifs were observed in the Egyptian MDV strains in both chickens and turkeys. Additionally, one interrupted motif from PPPP to PAPP was recorded at position 177. This interruption characterizes subclusters 1.1 and 1.3 (Fig. 3).

Discussion

GaHV-2 is an oncogenic herpesvirus that replicates predominantly in T-lymphocytes inducing lymphomatous lesions in chickens and less frequently in turkeys. Despite intensive vaccination and adoption of control measures, outbreaks of MD still occur globally with a significant impact on the economy of the poultry industry [7, 17, 18]. Understanding the genetic diversity among the clinical isolates of GaHV-2 is a key factor that helps determine the causes of vaccination insufficiency to date. Furthermore, the role of turkeys in the epidemiology and evolution of GaHV-2 should be elucidated. In the current study, GaHV-2 strains of turkey origin were identified in 2 commercial farms in Egypt. The polymorphism of the Meq gene as a determinant of virus virulence and tumorigenesis was studied in different GaHV-2 strains identified worldwide over an extended period of time, including the current Egyptian turkey strains.

The major complaint of the 2 turkey flocks was emaciation and uneven growth with a suspect of an oncogenic viral cause. Necropsy of the emaciated turkeys has revealed distinct enlargement of visceral organs with the development of grayish-yellow nodules. These lesions were commonly reported in GaHV-2-infected chickens [9, 17] and turkeys [27, 30]. Enlargement of the sciatic nerve was peculiar in the dissected turkeys. Previous reports only documented infiltration of pleomorphic neoplastic cells in sciatic nerves of turkeys with no evidence of neural lesions [27, 30]. It seems that the Egyptian strain(s) may have greater pathogenicity and/or tumorigenicity. Further pathological and molecular characterization of the Egyptian turkey isolates is required to identify the basis of enhanced neuropathogenicity.

In addition to GaHV-2, 2 avian retroviruses including ALV and REV were reported to infect turkeys displaying similar histopathological alterations [6, 27, 29]. Therefore, a stepwise diagnostic approach was followed to confirm that GaHV-2 is the primary cause of disease syndrome, including histopathological examination, immunohistochemistry, PCR, and DNA sequencing. B-cell lymphomas are caused by ALV [32–34], while GaHV-2 recruits T-cells [34, 35]. Both types of lymphocytes are infiltrated in neoplastic cell formation by REV [30]. Immunohistochemical analysis of tissue samples in the current study has revealed infiltration of lymphocytes with CD3+ and CD20-phenotype, which is specific for T- but not B-cells (Fig. 1h–k). This result augmented that the observed histopathological findings principally referred to GaHV-2 infection [5, 17, 35].

Oncogenic GaHV-2 infection was further confirmed by PCR amplification of the partial sequence of the ICP4 gene and the entire Meq gene. PCR testing for REV using primers specific for the long terminal repeats (LTR) has generated positive amplification results. Although co-infection between GaHV-2 serotype I and REV has been documented in turkeys [6, 36], the use of LTR for the identification of REV is mostly misleading. LTR of REV can be inserted in the fowl-pox genome from vaccine and field strains [37]. Therefore, another PCR was performed to prove the absence of full integration of REV proviral DNA in the turkey genome using a primer set specific for the env gene. The results have proved that GaHV-2 is the sole cause of the disease syndrome.

Cell transformation caused by GaHV-2 is principally linked to the transactivation of the Meq gene [17, 21]. The length of the Meq gene is variable according to the virulence and pathogenicity of the involved strain. Vaccine strains (CVI988, 814, and 3004) and those that induce mild symptoms (e.g., CU-2) have 60 amino acid (aa) longer than virulent strains. Insertion of a stretch of 60 aa residues may have a suppressive effect on Meq gene expression in vaccine and mild strains [9]. The presence of PPPP motifs in the transactivation domain was also linked to the virus pathogenicity. The higher number of the PPPP motif in a given strain is associated with the reduced virulence in affected chicken [11, 12, 25]. In this report, a shorter Meq gene of 1,020 aa length was identified in Fayoum-1 2018 strain. In addition, 3 PPPP motifs in the transactivation domain were recognized with proline content of 21.1% (data not shown). When comparing these figures with vaccine strains that display 7 PPPP motifs and 23.6% proline
content, it may be predicted that Fayoum-1 2018 strain is a virulent to very virulent field strain as proposed before with similar strains [12, 26]. Sequence analysis has revealed specific aa residues (T88, R93, D263; Fig. 3) similar to those identified in the Egyptian very virulent strains prevalent in chicken in 2012 [9]. This specific pattern may reflect the potential interspecies transmission of the new turkey strain from chicken, and may explain the enhanced neuropathogenicity; however, further molecular and pathological studies are required to support this hypothesis.

Phylogenetic classification of GaHV-2 based on Meq gene sequences has been described before by many groups. However, no consistent grouping of the strains was accomplished till now. The phylograms either presented unclustered strains [18], clustered strains according to the country of origin [23], or clustered strains in 2 or 3 groups identified by Latin numbers I, II, and III [8, 9, 17, 24]. Many strains (e.g., Indian Tn-n strains, Chinese strains GX070060, WS03, and YA) were allocated in different clusters according to the criteria used in the phylogenetic analysis [8, 9]. To unify the system used for phylogenetic clustering of GaHV-2 strains, all the previously constructed phylograms were analyzed and strains that represent every single cluster were chosen. Special care was taken to cover the entire array of geographical regions, temporal chronology of outbreaks, and different host systems. We also proposed the use of almost the entire Meq gene in the generation of the unified phylogram. It is expected that the phylogenetic classification proposed in this study will provide a platform that will help to understand the epidemiology and evolution of GaHV-2 in both chicken and turkeys.

In conclusion, GaHV-2 was recorded in turkeys for the first time in Egypt (and possibly in Africa and the Middle East). The emergence of MD in a new species might pose threats to the poultry industry, particularly in the context of close proximity between the commercial farms of chicken and other avian species in Egypt. The inefficient biosecurity practices adopted in turkey farms compared to that of chicken counterparts provides ample opportunities for multispecies transmission and development of outbreaks. Comprehensive epidemiologic and molecular studies are essentially required to explore the prevalence and significance of GaHV-2 in turkeys worldwide.

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Statement of Ethics

Guidelines for sample collection and animal use in research were followed according to the Institutional Animal Care and Use Committee, Faculty of Veterinary Medicine, Cairo University.

Conflicting Interest Statement

The authors declare that they have no conflicting interests.

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Author Contributions

Conceptualization: Haitham Amer and Magdy El-Mahdy. Methodology: Mahmoud Bayoumi and Mohamed El-Saied. Formal analysis and investigation: Haitham Amer, Basem Ahmed, and Magdy El-Mahdy. Writing – original draft preparation: Mahmoud Bayoumi and Mohamed El-Saied. Writing – review and editing: Haitham Amer, Basem Ahmed, and Magdy El-Mahdy. Resources: Mohamed El-Saied. Supervision: Haitham Amer and Magdy El-Mahdy.

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