Identification and complete genome characterization of a novel picornavirus in turkey (*Meleagris gallopavo*)

Ákos Boros,1† Csaba Nemes,2 Péter Pankovics,1 Beatrix Kapusinszky,3 Eric Delwart3,4 and Gábor Reuter1,3†

1Regional Laboratory of Virology, National Reference Laboratory of Gastroenteric Viruses, ÁNTSZ Regional Institute of State Public Health Service, Pécs, Hungary
2Veterinary Diagnostic Directorate of the Central Agricultural Office, Kaposvár, Hungary
3Blood Systems Research Institute, San Francisco, CA, USA
4University of California San Francisco, San Francisco, CA, USA

Members of the family *Picornaviridae* are important pathogens of humans and animals, although compared with the thousands of known bird species (>10,000), only a few (n = 11) picornaviruses have been identified from avian sources. This study reports the metagenomic detection and complete genome characterization of a novel turkey picornavirus from faecal samples collected from eight turkey farms in Hungary. Using RT-PCR, both healthy (two of three) and affected (seven of eight) commercial turkeys with enteric and/or stunting syndrome were shown to be shedding viruses in seven (88%) of the eight farms. The viral genome sequence (turkey/M176/2011/HUN; GenBank accession no. JQ691613) shows a high degree of amino acid sequence identity (96%) to the partial P3 genome region of a picornavirus reported recently in turkey and chickens from the USA and probably belongs to the same species. In the P1 and P2 regions, turkey/M176/2011/HUN is related most closely to, but distinct from, the kobuviruses and turdivirus 1. Complete genome analysis revealed the presence of characteristic picornaviral amino acid motifs, a potential type II-like 5′ UTR internal ribosome entry site (first identified among avian-origin picornaviruses) and a conserved, 48 nt long ‘barbell-like’ structure found at the 3′ UTR of turkey/M176/2011/HUN and members of the picornavirus genera *Avihepatovirus* and *Kobuvirus*. The general presence of turkey picornavirus – a novel picornavirus species – in faecal samples from healthy and affected turkeys in Hungary and in the USA suggests the worldwide occurrence and endemic circulation of this virus in turkey farms. Further studies are needed to investigate the aetiological role and pathogenic potential of this picornavirus in food animals.

INTRODUCTION

The family *Picornaviridae* is currently divided into 12 genera: *Aphthovirus*, *Avihepatovirus*, *Cardiovirus*, *Enterovirus*, *Erbovirus*, *Hepatovirus*, *Kobuvirus*, *Parechovirus*, *Sapelovirus*, *Senecavirus*, *Teschovirus* and *Tremovirus* (Knowles et al., 2012). Picornaviruses are small, non-enveloped viruses with single-stranded, positive-sense genomic RNA. In general, the genomes (7.2–9.1 kb long) have a common organizational pattern. The single ORF that encodes the polyprotein is flanked by 5′ and 3′ UTRs. The viral polyprotein has been divided into three regions: P1, P2 and P3. The P1 region encodes the viral capsid proteins (VP4–VP2–VP3–VP1), whilst the P2 and P3 regions encode proteins involved in protein processing (2Apro, 3Cpro and 3CDpro) and genome replication (2B, 2C, 3AB, 3BVPg, 3CDpro, 3DPols) (Racaniello, 2007). In addition, aphthoviruses, cardioviruses, erboviruses, kobuviruses, sapeloviruses, senecaviruses, teschoviruses and ‘turdiviruses’ (proposed) encode a leader (L) protein before the P1 region.

Members of the family *Picornaviridae* are important pathogens associated with several diseases. Picornavirus infections sometimes cause severe disorders of the gastrointestinal tract and the respiratory, neural, hepatocellular and circulatory systems in humans and animals (Racaniello, 2007; Alexandersen et al., 2012). In addition, approximately three-quarters of all emerging infectious disease agents (including...
viruses) in humans are thought to be zoonotic in origin (Woo et al., 2006). Among animal species, birds are a well-known reservoir of emerging infectious diseases such as avian influenza virus (H5N1), West Nile virus and Japanese encephalitis virus in humans (Woo et al., 2006, 2010). Beside the zoonotic agents, duck hepatitis virus (DHAV-1) in the genus Avihepatovirus (Kim et al.; 2006) and the recently identified turkey hepatitis virus (THV) (Honkavuori et al., 2011) circulate among domesticated poultry and may cause significant economical losses. Investigation of novel avian viruses, especially from domestic birds, and comparative analysis of their genomes is required to identify unrecognized pathogens and emerging viruses.

Compared with the thousands of known bird species (>10,000), only a few picornaviruses (n=11) have been described from avian sources. A minority of these picornaviruses have been known for a relatively long time, namely avian encephalomyelitis virus (genus Tremovirus) (Marvil et al., 1999), avian sapelovirus (formerly named duck picornavirus TW90A) in the genus Sapelovirus (Tseng & Tsai, 2007), THV (genus ‘Megrivirus’) (Honkavuori et al., 2011) and DHAV-1 (genus Avihepatovirus). Most of the recently identified avian picornaviruses – two novel species (turdivirus 1 and 2) from wild birds of the family Turdidae (Woo et al., 2010), and pigeon picornavirus A and B from pigeons (Kofstad & Jonassen 2011) – fall in an as-yet-unassigned, but distinct, picornavirus genus. Very recently, a quail picornavirus in common quail (Pankovics et al., 2012) and two novel picornaviruses, tentatively named chicken and turkey galliviruses (ChGV and TuGV, respectively), were discovered in chickens and turkeys (Farkas et al., 2012). The complete genome sequences of pigeon picornavirus A, ChGV and TuGV are not known; for TuGV, only a partial P3 sequence is available.

This study reports the detection and complete genome characterization of a turkey picornavirus, related to the recently identified picornavirus TuGV (Farkas et al., 2012), from faecal samples of healthy and affected commercial turkey (Meleagris gallopavo) collected from different turkey flocks in Hungary. This picornavirus is related distantly to members of the genus Kobuvirus and turdivirus 1 (in the proposed picornavirus genus ‘Orthoturdivirus’) and represents a novel picornavirus species.

**RESULTS**

**Complete genome acquisition and characterization of turkey picornavirus**

In faecal sample M176, following viral metagenomic analysis (Kapoor et al., 2008) we found 15 sequence reads that were assembled into eight contigs (Fig. 1) covering 42% of a picornaviral genome related to turdivirus 1 (GenBank accession no. GU182406) as the closest match using aBLAST search. The complete RNA genome of this turkey picornavirus (turkey/M176/2011/HUN) – which follows the common picornavirus genome organization: 5' UTR-L-P1(VP0-3-1)-P2(2A-B-C)-P3(3A-B-C-D)-3' UTR – consists of 8496 nt, excluding the poly(A) tail (Fig. 1). A large ORF of 7425 nt, which encodes a potential polyprotein precursor of 2474 aa, preceded at the 5' UTR by 761 nt and followed at the 3' UTR by 310 nt and a poly(A) tail, was found. The L protein is 450 nt (150 aa) long. The complete P1 (2562 nt; 854 aa), P2 (2025 nt; 675 aa) and P3 (2391 nt; 796 aa) regions show highest amino acid sequence identity to turdivirus 1 (GenBank accession no. GU182406) and porcine kobuvirus (EU787450) (Table 1). In the P3 region, the study sequence shows 77 and 96% amino acid identity to ChGV (GenBank accession no. JF424824; available sequence length was 3219 nt) and TuGV (GenBank accession numbers JF424828–JF424830; available sequence lengths varying between 1018 and 1020 nt). Possible cleavage
sites of the polyprotein were Q/G (Gln-Gly) in all cases, except for Q_{1271}/N and Q_{1999}/S between 2C/3A and 3C/3D, respectively (Fig. 1), supported by NetPicoRNA prediction (Blom et al., 1996). The base composition of coding region was found to be 22.7 mol% A, 20.1 mol% G, 28.2 mol% C and 29.0 mol% U.

**Analysis of 5’ and 3’ UTRs**

The predicted length of the 5’ UTR is 761 nt (Fig. 2). Seventeen AUG motifs were identified in the 5’ UTR. The last in-frame AUG is the predicted initiation codon, at position 762–764. This AUG codon is preceded by a significant polypyrimidine tract [p(Y) tract]. The longest p(Y) tract (UCUUUUCCUCCUUUUUUU) is located upstream in the RNA between nt 640 and 662. Using a BLASTN search, high (89 %) sequence similarity was located upstream in the RNA between nt 640 and 662.

Table 1. Genomic features of turkey picornavirus (turkey/M176/2011/HUN; GenBank accession no. JQ691613) and representatives of existing or proposed picornavirus genera

Amino acid sequence identity is shown as a percentage, based on the P1, P2, P3, 3C^{pro} and 3D^{pol} regions. Bold indicates the two highest levels of amino acid identity.

<table>
<thead>
<tr>
<th>Picornavirus genus</th>
<th>Virus</th>
<th>Genome features</th>
<th>Pairwise amino acid identity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>GenBank accession no.</td>
<td>Size (nt)</td>
<td>G+C content</td>
</tr>
<tr>
<td>Aphthovirus</td>
<td>NC_002554</td>
<td>8115</td>
<td>0.54</td>
</tr>
<tr>
<td>Aviparvovirus</td>
<td>NC_008250</td>
<td>7687</td>
<td>0.43</td>
</tr>
<tr>
<td>Cardiovirus</td>
<td>NC_001479</td>
<td>7835</td>
<td>0.49</td>
</tr>
<tr>
<td>Enterovirus</td>
<td>NC_000258</td>
<td>7440</td>
<td>0.46</td>
</tr>
<tr>
<td>Erbovirus</td>
<td>NC_003077</td>
<td>8821</td>
<td>0.50</td>
</tr>
<tr>
<td>Hepatovirus</td>
<td>NC_001489</td>
<td>7478</td>
<td>0.38</td>
</tr>
<tr>
<td>Kobuvirus</td>
<td>EU787450</td>
<td>8210</td>
<td>0.52</td>
</tr>
<tr>
<td>Parechovirus</td>
<td>NC_001897</td>
<td>7348</td>
<td>0.39</td>
</tr>
<tr>
<td>Sapelovirus</td>
<td>NC_003987</td>
<td>7491</td>
<td>0.41</td>
</tr>
<tr>
<td>Senecavirus</td>
<td>NC_011349</td>
<td>7310</td>
<td>0.50</td>
</tr>
<tr>
<td>Teschovirus</td>
<td>NC_003985</td>
<td>7117</td>
<td>0.45</td>
</tr>
<tr>
<td>Tremovirus</td>
<td>NC_003990</td>
<td>7055</td>
<td>0.45</td>
</tr>
<tr>
<td>‘Salivirus’</td>
<td>GQ179640</td>
<td>7982</td>
<td>0.56</td>
</tr>
<tr>
<td>‘Megivirus’</td>
<td>HM751199</td>
<td>9075</td>
<td>0.46</td>
</tr>
<tr>
<td>‘Orthoturdivirus’</td>
<td>GU182406</td>
<td>8035</td>
<td>0.58</td>
</tr>
<tr>
<td>‘Paratursivirus’</td>
<td>GU182408</td>
<td>7641</td>
<td>0.47</td>
</tr>
</tbody>
</table>

The 310 nt long 3’ UTR sequence of turkey/M176/2011/HUN was similar in length (309 and 311 nt) and nucleotide sequence (73 and 96 % nucleotide identity) to ChGV strain CHK1/USA/2010 (GenBank accession no. JF424824) and TuGV strain TRK90/USA/2010 (JF424829). The predicted secondary RNA structure of the 3’ UTR shows the presence of multiple stem–loops (Fig. 3a) and a 48 nt long ‘barbell-like’ structure (Fig. 3a, b) of the 3’ UTRs with a relatively short p(Y) tract at the upper loop region of the turkey/M176/2011/HUN, which was highly similar in position and nucleotide sequence (UUUCUUU, identical nucleotides) to the core binding site of PTB located at the 3’ UTR of murine norovirus 1 isolate Mu/NoV/GV/MNV1/2002/USA (GenBank accession no. AY228235.2) (Bailey et al., 2010). As well as being found in turkey/M176/2011/HUN, the ‘barbell-like’ structure was also recognizable in members of the genera Aviparvovirus and Kobuvirus, with a 9 + 6 nt long identical nucleotide motif in the lower loop regions (Fig. 3b–d). The position of the conserved ‘barbell-like’ structure at the 3’ UTRs of different picornaviruses shows significant variability. The first nucleotide of this structure was 227 nt from the stop codon in DHAV-1 (GenBank accession no. DQ249299; 3’
UTR length 317 nt), 56, 70, 72 nt in Aichi virus (FJ890523; 3’ UTR length 218 nt), mouse kobuvirus (JF755427; 3’ UTR length 241 nt) and canine kobuvirus (JN088541; 3’ UTR length 243 nt), respectively, and 167, 171, 171 nt in ChGV (JF424824; 3’ UTR length 310 nt), TuGV (JF424829; 3’ UTR length 314 nt) and the study sequence, respectively (Fig. 3d).

**Analysis of coding regions (L, P1, P2 and P3)**

Turkey/M176/2011/HUN encodes a 150 aa L protein upstream of the capsid-encoding genome region; it exhibits no significant sequence similarity to picornaviruses, including aphtho-, cardio-, erbo-, sapelo-, tescho- and turdivirus. Homologous sequences were also not found using
a conserved domains search in CDD (Marchler-Bauer et al., 2011). Neither the catalytic dyad (Cys and His) of papain-like thiol proteases found in the foot-and-mouth disease virus L protein (Gorbalenya et al., 1991) nor a presumed zinc-binding motif, Cys-His-Cys-Cys, found in Theiler’s murine encephalomyelitis virus and quail picornavirus (Chen et al., 1995; Pankovics et al., 2012) was present.

The P1 genome region is located downstream of the L protein-encoding site. As in kobu-, avihepato-, parecho- and turdiviruses, the potential cleavage site between the VP4 and VP2 is not recognizable and turkey/M176/2011/HUN virions are probably built from only three capsid monomers (VP0-VP3-VP1). Multiple potential myristylation motifs (GxxxS, where x is a non-conserved amino acid) (Chow et al., 1987) were recognized in the N-terminal end of VP4 (Fig. 1).

The P2 genome region encodes the non-structural proteins 2A-2B-2C. The 2A protein of turkey/M176/2011/HUN contains no sequence repeats. The conserved tyrosine residue at the third position of VPg proteins of all known picornaviruses was present (e.g. Aichi virus; Yamashita et al., 1998). The conserved catalytic triad (H, E/D, C) of the 3C viral cysteine-active-centre protease was seen in turkey/M176/2011/HUN with a modification (I instead of E/D) (Bazan & Fletterick, 1988); the active-site cysteine in motif GxCG (x, variable) was also present (Gorbalenya et al., 1989) (Fig. 1). The RNA-dependent RNA polymerase (3D pol) contained highly conserved motifs (KDEL, GGxPSG, YGDD and FLKR) (Fig. 1).

The non-structural proteins of the P3 genome region comprise 3A-3BVPg-3Cpro-3Dpol. The short 3B protein (small genome-linked protein or VPg) of turkey/M176/2011/HUN contains no sequence repeats. The conserved tyrosine residue at the third position of VPg proteins of all known picornaviruses was present (e.g. Aichi virus; Yamashita et al., 1998). The conserved catalytic triad (H, E/D, C) of the 3C viral cysteine-active-centre protease was seen in turkey/M176/2011/HUN with a modification (I instead of E/D) (Bazan & Fletterick, 1988); the active-site cysteine in motif GxCG (x, variable) was also present (Gorbalenya et al., 1989) (Fig. 1). The RNA-dependent RNA polymerase (3D pol) contained highly conserved motifs (KDEL, GGxPSG, YGDD and FLKR) (Fig. 1).

Phylogenetic analysis of turkey picornavirus

The relatively high nucleotide and amino acid sequence identity at the P3 genome region of turkey/M176/2011/HUN to TuGV and ChGV is also observable in the P3
Fig. 4. Phylogenetic relationship between turkey picornavirus (turkey/M176/2011/HUN; GenBank accession no. JQ691613; indicated in bold), representative members of the 12 picornavirus genera and unassigned picornaviruses, based on amino acid sequences of the different picornavirus coding regions: P1 (a), P2 (b) and P3 (c).
Table 2. Epidemiological and clinical background of tested turkeys from eight turkey farms in Hungary, and diagnostic results of the collected faecal samples for turkey picornavirus

<table>
<thead>
<tr>
<th>Farm location</th>
<th>Sample ID</th>
<th>Age (days)</th>
<th>Symptoms</th>
<th>Routine diagnostic tests*</th>
<th>VP1 RT-PCR†</th>
<th>GenBank accession no.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bogyiszlo´</td>
<td>B-177</td>
<td>8</td>
<td>Catarrhal enteritis, foamy/watery caecum content</td>
<td>par, – bact, –</td>
<td>Positive</td>
<td>JQ691615</td>
</tr>
<tr>
<td></td>
<td>B-415</td>
<td>14</td>
<td>Rickets</td>
<td>par, – bact, –</td>
<td>Positive</td>
<td></td>
</tr>
<tr>
<td>Szalánta</td>
<td>B-195</td>
<td>7</td>
<td>Catarrhal enteritis, foamy/watery caecum content</td>
<td>par, – bact, –</td>
<td>Negative</td>
<td></td>
</tr>
<tr>
<td>Karaközöröcsök</td>
<td>B-232</td>
<td>21</td>
<td>Growth depression</td>
<td>par, – bact, –</td>
<td>Positive</td>
<td></td>
</tr>
<tr>
<td>Nagyalásony</td>
<td>B-308</td>
<td>47</td>
<td>Growth depression, skeletal disorders</td>
<td>par, – bact, –</td>
<td>Positive</td>
<td></td>
</tr>
<tr>
<td>Bakonyőléske</td>
<td>B-356</td>
<td>14</td>
<td>Uneven growth, catarrhal enteritis, foamy/watery caecum content</td>
<td>par, – bact, + (E. coli bacteraemia)</td>
<td>Positive</td>
<td></td>
</tr>
<tr>
<td>Bikal</td>
<td>B-407</td>
<td>42</td>
<td>Skeletal disorders</td>
<td>par, – bact, –</td>
<td>Positive</td>
<td></td>
</tr>
<tr>
<td>Nemesböd</td>
<td>PP-1</td>
<td>7</td>
<td>Healthy</td>
<td>None</td>
<td>Negative</td>
<td></td>
</tr>
<tr>
<td></td>
<td>PP-2</td>
<td>14</td>
<td>Healthy</td>
<td>None</td>
<td>Positive</td>
<td></td>
</tr>
<tr>
<td></td>
<td>PP-3</td>
<td>21</td>
<td>Healthy</td>
<td>None</td>
<td>Positive</td>
<td>JQ691614</td>
</tr>
<tr>
<td>Bögöte</td>
<td>M176</td>
<td>20</td>
<td>Growth depression</td>
<td>par, – bact, –</td>
<td>Positive</td>
<td>JQ691613§</td>
</tr>
</tbody>
</table>

*par, Pathogen parasite diagnostics based on microscopic observation using native faecal samples; bact, routine bacterial culture of heart blood, and liver or bone marrow.
†For the primer sequences, see Table S1 (available in JGV Online).
§Complete genome.
phylogenetic tree, where the gallviruses and the study sequence cluster together and form a distinct lineage among the other representative members of the family Picornaviridae.

The position of turkey/M176/2011/HUN in the phylogenetic trees based on deduced amino acid sequences constructed from the other genome regions (P1, P2) shows a consistent relationship to turdivirus 1 (GenBank accession no. GU182406) of the proposed genus ‘Orthoturdivirus’ and a distant relationship to viruses of the genus Kobuvirus and to klasseviruses/saliviruses (Fig. 4).

Identification and phylogenetic analysis of additional turkey picornavirus sequences

A specific primer pair was used to amplify the complete VP1 of turkey picornavirus to screen faecal samples from healthy and affected commercial meat turkeys collected from an additional seven turkey farms (Table 2). All tested faecal samples were RT-PCR-positive for turkey picornavirus except for samples B195 and PP1, which were collected from 7-day-old turkeys (Table 2). Two VP1-positive samples (from one affected and one healthy turkey) were selected randomly for sequence and phylogenetic analysis. Pairwise comparisons of the three acquired VP1 sequences show 95–96 % nucleotide and 98 % amino acid sequence identity. This high degree of nucleotide and amino acid sequence similarity was also noticeable in the phylogenetic tree (Fig. 5), where all three turkey picornavirus sequences from Hungary grouped together and formed a distinct cluster among picornaviruses. The distinction of the study sequences from the closest available relative, turdivirus 1, was based on the high sequence difference, where the VP1 study sequences (265 aa) show only 18 % amino acid sequence identity to turdivirus 1 (GenBank accession no. GU182406).

DISCUSSION

Using a metagenomics and RT-PCR approach, we report the first complete genome characterization and an initial prevalence study of a turkey picornavirus in Hungary that showed a high degree of sequence similarity at the partial P3 genome region to picornaviruses reported by Farkas et al. (2012), tentatively named galliviruses. Galliviruses are present in chicken and turkey flocks in the USA (Farkas et al., 2012), although the complete genome sequence of both ChGV (GenBank accession no. JF424824; longest available sequence length 3219 nt) and TuGV (JF424828–JF424830; longest available sequence length 1020 nt) remain unknown. Because of the high nucleotide and amino acid sequence identity of turkey/M176/2011/HUN to TuGV at the available partial P3 genome region, we refer to this group of novel picornaviruses as galliviruses.

Taxonomic classification of turkey picornavirus

Complete genome analysis of the turkey picornavirus, turkey/M176/2011/HUN, showed that the closest relatives were turdivirus 1 (proposed genus ‘Orthoturdivirus’) and porcine kobuvirus (genus Kobuvirus). According to the current criteria of new genus definition from the Picornaviridae Study Group of the International Committee on Taxonomy of Viruses (http://www.picornastudygroup.org/definitions/...
genus_definition.htm), novel picornavirus genera are defined by amino acid identity in the P1, P2 and P3 regions of <40, <40 and <50%, respectively. Based upon this theoretical agreement and the supporting result of the phylogenetic analysis, turkey picornavirus, together with galliviruses (Farkas et al., 2012), probably represents a novel picornavirus genus.

Analysis of the 5’ UTR structure of turkey picornavirus

Analysis of the 5’ UTR revealed that turkey picornavirus has a type II-like IRES – first identified among avian origin picornviruses – based upon sequence and secondary structural similarities to coronavirus, and particularly to EMCV. Initiation of translation on these IRESs begins with specific binding of the eIF4G to J–K domains, which is stimulated by eIF4A (Yu et al., 2011). In addition to canonical translation factors, type II IRESs also require IRES trans-acting factors (ITAFs), such as PTB, that are hypothesized to promote local conformational changes of domain J and to bring it into closer proximity to the base of domain I (Yu et al., 2011). Structural analogies to the EMCV IRES predict similar translation initiation in turkey picornavirus. On the other hand, the picornavirus 5’ UTR contains a poly(C) tract between domains C and D of the 5’ UTR that varies in length (from 80 to 250 nt) and is associated with virulence (Racaniello, 2007). Whether turkey picornavirus contains a poly(C) tract – which we could not determine by 5’ RACE – currently remains unknown.

Conserved motif analysis in the viral polyprotein of turkey picornavirus

During the genome analysis, characteristic picornaviral motifs (e.g. Rhv-like capsid domains, 2C, 3C and 3D functional domains) were recognized in the viral polyprotein, with minor variance observed in some motifs (e.g. isoleucine instead of the common glutamic acid or aspartic acid in the 3C catalytic triad or more than one potential myristylation motif at the N-terminal end of VP0). The role and functionality of these variations remain to be elucidated.

Analysis of the 3’ UTR ‘barbell-like’ structure

A series of stem–loops, similar to that described by Farkas et al. (2012), was identified at the 3’ UTR of turkey/M176/2011/HUN. The highly structured heteropolymeric regions of picornaviral 3’ UTRs are very diverse and their functions are not yet completely understood, although some evidence indicates that the 3’ UTRs could contain specific binding sites for viral and/or cellular proteins (Rohll et al., 1995). The presence of a nearly identical p(Y) tract located at the same position (upper loop region of the 3’ UTR) in murine noroviruses (MNVs) and in turkey/M176/2011/HUN indicates that it could serve as a core binding site for the cellular poly(C)-binding proteins (PCBP1 and 2) and/or PTB (Makeyev & Liebhaber, 2002; Sawicka et al., 2008), as demonstrated previously for MNVs (Bailey et al., 2010). Further studies with functional analyses will be required to identify cellular or viral factors that might interact with this region.

A 48 nt long ‘barbell-like’ structure with two identical motifs at the lower loop was identified in galliviruses, in turkey/M176/2011/HUN and in members of two other picornavirus genera. Studies of the picornaviral 3’ UTR revealed the presence of a ‘kissing’ interaction between different 3’ UTR stem–loops (Pilipenko et al., 1996), and of 3’ UTR–5’ UTR IRES interactions (Liu et al., 2009) with complementary base pairing. A genome-wide complement sequence search to find sections of sequence corresponding to the two identical motifs of the ‘barbell-like’ structure was unsuccessful. The possible function of this conserved structure is unknown, although the highly conserved nature of this ‘barbell-like’ motif suggests a pivotal role in the viral reproduction process.

Turkey picornavirus in healthy and affected turkeys

The general presence of turkey picornavirus in faecal samples of healthy and affected turkeys in Hungary in the USA (Farkas et al., 2012) strongly suggests the worldwide occurrence and endemic circulation of this virus. The presence of this virus in healthy turkeys does not rule out its pathogenic potential related to stunting syndrome and/or poult enteritis complex, similar to other enteric viral pathogens (e.g. turkey astro-, rota- and reoviruses), as these viruses are also detectable in apparently healthy turkey flocks (Pantin-Jackwood et al., 2007, 2008; Day et al., 2010). Furthermore, it is also possible that this virus could contribute to the development of symptoms as part of coinfections with other pathogens (Day et al., 2010). A clear connection between turkey picornavirus infection and clinical outcomes will require more detailed investigation, including experimental animal infection studies and periodic surveys.

The family Picornaviridae is currently divided into 12 genera; two (Avihepatovirus and Tremovirus) include only avian picornaviruses. Due to the increasing number of novel picornaviruses of different (e.g. avian) origins, the taxonomy of the family Picornaviridae is evolving rapidly. Among the four novel picornavirus genera proposed in 2011, one includes avian picornavirus(es) (THV in the proposed genus ‘Megrivirus’) (Knowles et al., 2012). Furthermore, two of the three recently identified avian picornaviruses (turdviruses 1 and 2) are probably the type species of two novel genera, ‘Orthoturdivirus’ and ‘Paraturdivirus’ (Woo et al., 2010) and the third, quail picornavirus (Pankovics et al., 2012), stands at the borderline of a potential novel picornavirus genus. Turkey picornavirus, together with galliviruses (Farkas et al., 2012), is a member of another potential genus of avian picornaviruses. The accelerating pace of novel avian picornavirus characterization indicates that there are still many unidentified
and highly divergent avian picornaviruses circulating among domestic and wild birds.

METHODS

High-throughput pyrosequencing and analysis. A faecal sample (M176) collected in April 2011 from a commercial meat turkey (M. gallopavo) showing stunting syndrome in a turkey farm located in north-west Hungary was selected for viral metagenomic analysis. PBS-diluted specimens were passed through a 0.45 μm sterile filter and centrifuged at 6000g for 5 min. The pellet was mixed with a mixture of nucleases to enrich for particle-protected nucleic acids (Victoria et al., 2009). Nucleic acids were extracted using a QIAamp Viral RNA Mini kit (Qiagen) according to the manufacturer’s instructions. Viral nucleic acid libraries containing both RNA and DNA molecules were constructed by sequence-independent random RT-PCR amplification, as described by Victoria et al. (2009). 454 pyrosequencing using 454 GS FLX technology was then performed as described previously (Kapoor et al., 2008; Victoria et al., 2009). The pyrosequencing reads and assembled sequence contigs were compared with the GenBank nucleotide and protein databases using BLASTX and BLASTN, respectively.

Complete genome acquisition of turkey picornavirus. Specific primer pairs (Table S1, available in JGV Online) were designed based on the sequence contigs from the pyrosequencing reads to determine the complete nucleotide sequence of the turkey picornavirus using RT-PCR and long-range RT-PCR. RNA was extracted from 150 μl faecal suspension [35–40 % (v/v) in 0.1 M PBS] using TRizol reagent (Invitrogen) according to the manufacturer’s instructions. RT-PCR and long-range RT-PCR were performed as described previously (Reuter et al., 2002; Boros et al., 2012). The 5' and 3' ends of the genome were determined by using a 5'/3' RACE PCR kit (Roche) as described previously (Boros et al., 2011). PCR products were sequenced directly with a BigDye Terminator Cycle Sequencing Ready Reaction kit (PE Applied Biosystems) using the primer-walking method and run on an automated sequencer (ABI PRISM 310 Genetic Analyzer; Applied Biosystems).

Detection of turkey picornavirus in different turkey flocks. Additional faecal samples from seven different turkey flocks located in western Hungary were collected in 2011 from turkeys of different ages and showing different symptoms, and three samples (PP1–3) were collected from a healthy turkey for three consecutive weeks after hatching (Table 2). Native faecal samples were initially tested by microscopic observation for pathogen parasites and by routine bacterial culture of heart blood, and liver or bone marrow. RNA samples were tested for turkey picornavirus by RT-PCR as described above and using specific primer pairs targeting the VP1 region (Table S1).

Sequence and phylogenetic analysis. Sequences from representative members of different picornavirus genera were obtained from GenBank and the study sequences were aligned using CLUSTAL_X software (version 2.0.3) (Thompson et al., 1997); similarity calculations were performed by using GeneDoc software (version 2.7) (Nicholas & Nicholas, 1997). Phylogenetic trees of the amino acid alignments were created using the neighbour-joining method based on the Jones–Taylor–Thornton matrix-based model of MEGA software (version 5) (Tamura et al., 2011). Bootstrap values (based on 1000 replicates) for each node are given when >50 %. The secondary structures of 5′ and 3′ UTRs were predicted using the Mfold program (Zuker, 2003) and the two-dimensional model was drawn by using the Corel Draw Graphics Suite (version 12). The complete genome sequence of novel turkey picornavirus was submitted to GenBank with accession numbers JQ691613–JQ691615.

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