

A novel passerivirus (family *Picornaviridae*) in an outbreak of enteritis with high mortality in estrildid finches (*Uraeginthus sp.*)

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Abstract

An enteric outbreak with high mortality (34/52, 65.4%) was recorded in 2014 in home-reared estrildid finches (*Estrildidae*) in Hungary. A novel passerivirus was identified in a diseased violet-eared waxbill using viral metagenomics and confirmed by RT-(q)PCR. The complete genome of finch picornavirus strain waxbill/DB01/HUN/2014 (MF977321) showed the highest amino acid sequence identity of 38.9%, 61.6%, 69.6% in P1^{cap}, 2C^{hel} and 3C^{proDPol}, respectively, to passerivirus A1 (GU182406). A high viral load (6.58×10^{10} genomic copies/ml) was measured in a cloacal specimen and in the tissues (spinal cord, lung, and the intestines) of two additional affected finches. In addition to intestinal symptoms (diarrhoea), the presence of extra-intestinal virus suggests a generalized infection in this fatal disease, for which the passerivirus might be a causative agent.

Keywords Passerivirus, Dead bird, Estrildid finch, Picornavirus, Gastroenteritis

The GenBank[EMBL/DDBJ] accession number for the study sequence: MF977321-MF977323.

Picornaviruses are small, non-enveloped viruses of the family *Picornaviridae* that have a positive-sense, single-stranded RNA genome [1]. The RNA genome starts with 5' untranslated region (UTR) containing either an SL-A/B/C cloverleaf fold or the origin of replication (ORI) region, followed by an internal ribosomal entry site (IRES) [2–7]. Generally, with the exception of the dicipiviruses [8, 9], a single polyprotein is translated from the viral RNA genome and is cleaved into smaller viral capsid (structural) proteins and non-structural proteins [1, 10]. In addition, many picornaviruses encode a leader (L) protein before the P1 region, which has diverse functions [11, 12]. Located in different genomic positions, a cis-acting replication element (CRE) is also present in picornavirus genomes [6, 7, 12–15]. The viral RNA genome ends with a 3'UTR, which contains structural-poly(A) tail [16]. The host range of picornaviruses is wide, including both lower and higher vertebrates [<http://www.picornaviridae.com>].

The estrildid finches are very popular hobby birds in Hungary. These birds are members of the family *Estrildidae*, class Aves order Passeriformes [17; <https://www.itis.gov>; <http://www.hbw.com/order/passeriformes>], which currently includes 30 genera with 158 species [18; <http://www.hbw.com/ibc/family/waxbills-estrildidae>]. The estrildid finches naturally originated in India, Australia, and the Indian and South Pacific islands, but the most ancient groups are from Africa [18]. Generally, estrildid finches are small birds that are approximately 8 to 17 centimetres long with an average weight of 7.5 grams. The plumage colour and pattern are varied, but their habitat and eating habits are similar [18, 19]. Their diet consists mostly of seeds (*Panicum miliaceum*, *Setaria italica*, *Phalaris canariensis*) and greens (*Stellaria media*, *Lactuca sativa*, *Plantago major*) or wild furrow-weeds (*Poa annua*, *Poa pratensis*, *Setaria viridis*) [18, 19], but during their reproduction period, high-protein-content foods (*Formica rufa*, *Tenebrio molitor*, *Daphnia magna*) are needed (T. Jantyk personal communication). Frequently, birds for home-rearing are collected from the wild by a stock farmer, often shortening or neglecting the quarantine time and exposing the isolated bird population to external infections.

In this study, using viral metagenomics and molecular methods, a novel passerivirus was identified in cloacal and tissue samples of dead estrildid finches (*Estrildidae*) in Hungary and characterized.

A gastroenteric outbreak with high mortality (34/52, 65.4%) occurred in 2014 in the town of Békés, Hungary, in the finch population, including six violet-eared waxbills (*Uraeginthus granatinus*), six purple grenadiers (*Uraeginthus ianthinogaster*) and 22 red-cheeked cordon-bleus (*Uraeginthus bengalus*) in the genus *Uraeginthus*. Additionally, four red avadavats (*Amandava amandava*) and four orange-breasted waxbills (*Amandava subflava*) in the genus *Amandava*, four Gouldian finches (*Erythrura gouldiae*) in the genus *Erythrura*, four red-billed firefinches (*Lagonosticta*

senegala) in the genus *Lagonostica* and two crimson finches (*Neochmia phaeton*) in the genus *Neochmia* were kept in the aviary. All birds were kept together in a 2.5 × 2.5-metre aviary in a closed building (with walls and a roof) and allowed access to the same water and feed. According to information from the stock farmer, six breeding pairs of violet-eared waxbills were bought from Holland, and six breeding pairs of wild purple grenadiers were bought from Africa. The red-cheeked cordon-bleus were reared locally. All birds of the family *Estrildidae* were breeding pairs of reproductive age except the violet-eared waxbills, purple grenadiers and red-cheeked cordon-bleus, which were younger than 2 years old. An enteric outbreak occurred seven to ten days after the violet-eared waxbills and purple grenadiers were introduced into the farm. The recommended 30-day quarantine was not complied with. At first, all violet-eared waxbills, purple grenadiers and red-cheeked cordon-bleus (members of the same genus) sat languidly with plumped feathers and with their heads bent (nonspecific signs). From this point in time, the appetites of these birds increased, and they mostly showed a preference for mealworms, the larval form of the meal-worm beetle *Tenebrio molitor*, but these birds nonetheless lost weight during several episodes of watery diarrhoea. After the first death, a faecal sample and two dead birds were taken to the local animal hospital for routine bacterial and parasitic examination on November 14, 2014. Hepatosplenomegaly and enteritis were observed during the necropsy of both birds. Bacterial cultivation of liver suspensions in agar slants and modified Drigalszki agar, and STAM staining of liver samples yielded negative results. A flotation assay for parasites in faecal samples from both dead birds gave a negative result for coccidian-like oocyst bodies and cestodes, nematode eggs and adult worms. A total of 34 finches of the genus *Uraeginthus* died during the observation period, but other birds kept in the same aviary remained asymptomatic.

The carcasses of three violet-eared waxbills (DB01, DB02 and DB03) were also collected by the stock farmer. A cloacal sample from a single bird (DB01) was chosen for viral metagenomics analysis, and corpses were stored at -80 °C. The cloacal sample was diluted in 500 µl of phosphate-buffered saline (PBS) for extraction of total nucleic acid. The suspension was passed through a 0.45-µm sterile filter (Millipore) and centrifuged at 6,000 × g for 5 min. The filtrate was then treated with DNase and RNase [20]. Viral particle-protected nucleic acids were extracted using a QIAamp spin-column kit (QIAGEN), amplified by sequence-independent random RT-PCR, and subjected to viral metagenomics analysis [21]. A viral cDNA library was constructed using a ScriptSeq™ v2 RNA-Seq Library Preparation Kit (Epicentre), and it was sequenced on a MiSeq Illumina platform as described previously [20]. Reads and assembled contigs longer than 100 bp were compared to sequences in the GenBank protein database (BLASTx), using a cutoff value (E-value) of 10⁻¹⁰.

RNA was extracted from a 300- μ l cloacal suspension using TRI Reagent (Molecular Research Centre, Cincinnati, OH, USA) [22]. Sequence-specific primers were designed for the verification of viral metagenomics reads/contigs and to obtain the complete viral RNA genome sequence using various PCR methods [23, 24]. PCR products were sequenced directly and run on an automated sequencer (ABI Prism 310, Applied Biosystems, Stafford, USA). In addition, the organs (midbrain, spinal cord, heart, lung, liver, kidney, pancreas, proventriculus, gizzard, small- and large-intestinal tracts) of two dead violet-eared waxbills – DB02 and DB03 – were also tested by RT-PCR.

The primer pairs used for screening, S13-Passeri-3D-R (5'-CCTCCCAGATCATGTAGT-3', corresponding to nt position 8699-8716 of waxbill/DB01/HUN/2014) and S13-Passeri-3D-F (5'-CCTTATGGTTGGAGGCAA-3', corresponding to nt position 8256-8273 of waxbill/DB01/HUN/2014), were designed based on the 3D^{pol} contig sequence of the isolate from this study. For quantitative RT-PCR, cDNA was synthesized using reverse primer S13-Passeri-3D-R and then amplified using primers (DB01-qPCR-standR (5'-GCATACAGCACATCATCTCCA-3', corresponding to nt position 8391-8411) and DB01-qPCR-standF (5'-AGGCAACCCCTCTGGCTGTGT-3', corresponding to nt position 8268-8288) using a real-time PCR system (LightCycler FastStart DNA Master SYBR Green I, Roche, Mannheim, Germany). For absolute quantification and the generation of a standard curve, a 100-fold dilution series of a single PCR amplicon of the isolate was purified on a silica column (QIAGEN, Hilden, Germany), quantified spectrophotometrically, and used for qPCR analysis. Three technical repeats were performed.

The probable initiation start codon (AUG) was identified using the ATG^{Pr} prediction program (<http://atgpr.dbcls.jp/>), and a probable N-terminal myristoylation signal identified using NMT - The MYR Predictor (<http://mendel.imp.ac.at/myristate/SUPLpredictor.htm>).

Multiple nucleotide (nt) sequences of representative picornaviruses and the isolate from this study were aligned based on codons, using MEGA 6.06 [25], and their predicted amino acid (aa) sequences were compared using BioEdit ver. 7.2.5 [26]. A hypothetical cleavage map of the picornavirus polyprotein was derived from alignments with the closest relative, the picornavirus passerivirus A1 (genus *Passerivirus*) strain thrush/Hong Kong/00356/2007 (GU182406).

All evolutionary analyses were conducted in MEGA 6.06. The nucleotide sequences of representative picornaviruses and the isolate from this study were aligned by the ClustalW codon-based method and pre-tested in a best-nt/ aa-model (maximum likelihood, ML) search. Dendrograms were constructed by the ML method based on the general time-reversible model with gamma distribution (+G) and invariable sites (+I).

The secondary RNA structures of the 5' and 3'UTRs, including the IRES, were predicted using

RNAstructure [27] and ViennaRNA Web Service [28] with manual correction as described previously [4, 29].

The complete sequence of waxbill/DB01/HUN/2014 and the partial 3D^{Pol} sequences of waxbill/DB02-DB03/HUN/2014 were submitted to GenBank under accession numbers MF977321-MF977323.

In 2014, a cloacal sample was collected from a dead DB01 violet-eared waxbill (*Uraeginthus granatinus*) and subjected to viral metagenomics analysis. After *in silico* analysis and *de novo* assembly, 10,393 sequence reads were obtained that showed similarity to viruses (BLASTx cutoff E score $\leq 10^{-10}$). The detected sequences for which more than 15 reads were obtained were from the virus families *Picornaviridae* (N = 9363), *Microviridae* (N = 212), *Circoviridae* (N = 62), *Caliciviridae* (N = 39), *Tombusviridae* (N = 17), “other” (N = 116) and “unclassified” (N = 584). The reads corresponding to the virus family *Picornaviridae* were selected, and approximately 75% of the viral genome was assembled (data not shown). S13-Passeri-3D-R/F primer pairs were used to verify the presence of the viral RNA in the cloacal sample.

The complete RNA genome of strain waxbill/DB01/ HUN/2014 (MF977321) is 9130 nt long, excluding the poly(A) tail. The genome organization is as follows: 5UTR^{IRES-V/L/P1(VP0-VP3-VP1)/P2(2A^{Hbox/NC}-2B-2C)/P3(3A-3B^{VPg}-3C^{Pro}-3D^{Pol})}/3UTR^{'barbell'}-poly(A) (Fig. 1). This isolate has a G+C content of 48.28% and a nucleotide distribution of 26.4% A, 25.3% U, 21.95% G and 26.3% C.

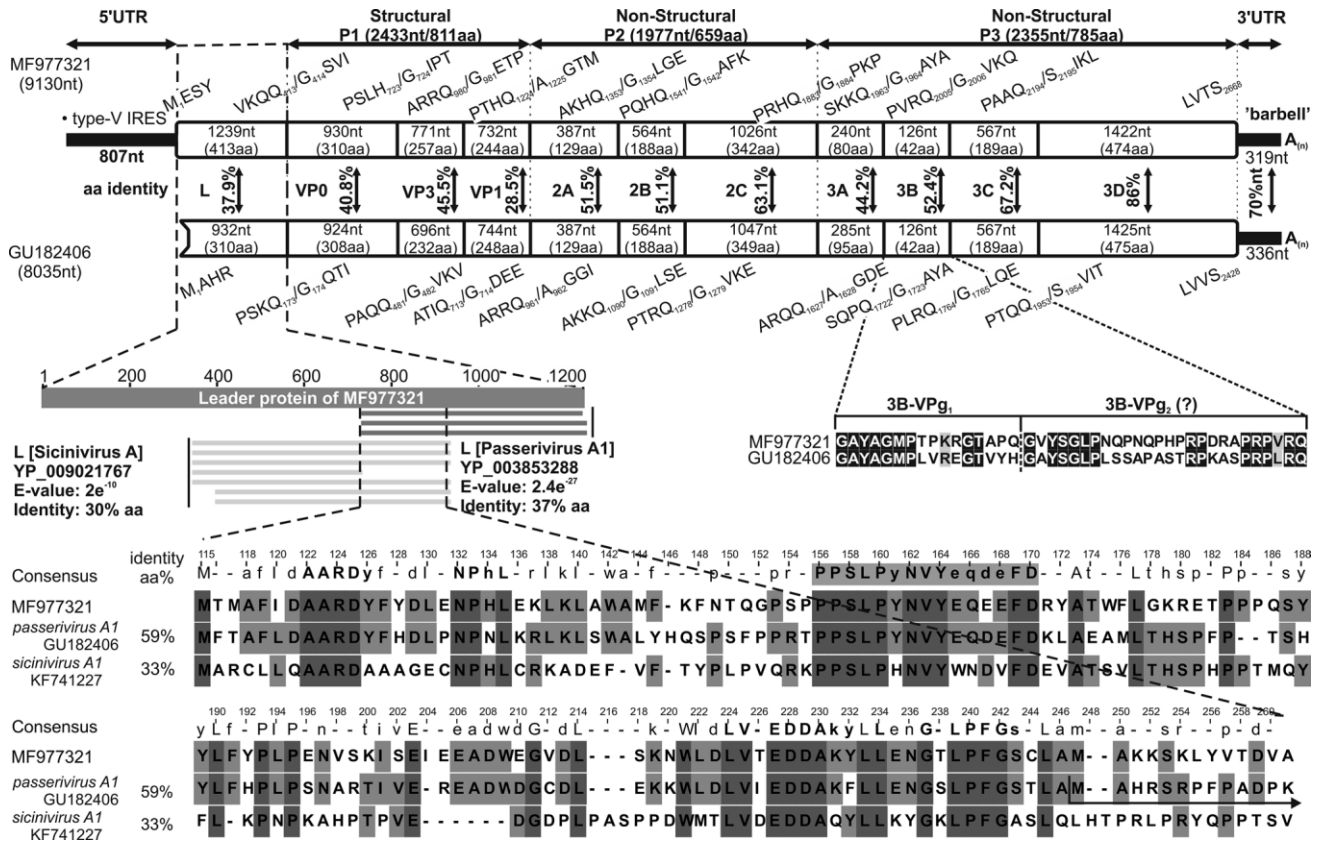


Fig. 1 Comparison of the genome organization of finch picornavirus strain waxbill/DB01/HUN/2014 (MF977321) with that of its closest relative, passerivirus A1 strain thrush/Hong Kong/00356/2007 (GU182406, NC_014411), a member of the genus *Passerivirus*. Genome lengths – excluding the poly(A)-tail – are indicated below the strain name; the nucleotide (nt) and amino acid (aa) lengths of the hypothetical genes are indicated in the gene boxes. The predicted aa cleavage sites are shown at the borders of the regions. Pairwise aa sequence identity scores are shown between the related protein products. The comparative alignment of the leader (L) protein of the study strain and the closest match in based on a GenBank BLASTp search. The L protein of the study strain has 37% (E-value: $2.4e^{-27}$, query cover: 40%) and 30% (E-value: $2.4e^{-10}$, query cover: 40%) amino acid sequence identity to the leader proteins of passerivirus A1 strain thrush/Hong Kong/00356/2007 (GU182406) and scicivirus A1 strain ch/UCC001/Eire (KF741227, NC_023861), respectively. The presumed 5'UTR nt sequence of strain thrush/Hong Kong/00356/2007 (GU182406) [34] encodes a potential L protein product that partially overlaps part with that of scicivirus A1 strain ch/UCC001/Eire (KF741227). The hypothetical initiation site of the prototype passerivirus A1 [34] is indicated by a black arrow. The conserved aa motifs are bolded in the consensus sequence, and the PPSLP amino acids are indicated in grey. The aa sequence alignment of the 3B-VPg region of the isolate from this study and the prototype passerivirus A1 strain indicates the two hypothetical copies of 3B-VPgs

The 5'UTR of the studied strain is 807 nt long (Fig. 1, 2A/B). The first in-frame AUG initiation codon is at nt position 808-810 (AAA₈₀₈UGG) with an optimal Kozak context (GxxAUGG) [30] (Fig. 2A/B). In the GenBank database, 68% nucleotide sequence identity was found from nt 405 to nt 635 between this strain and the type-V IRES of the oscivirus thrush/Hong Kong/10878/2006 (GU182410) (from nt 253 to nt 476) as the closest match (Fig. 2B). Based upon this nt sequence

identity and the results of secondary RNA structure predictions, this isolate was found to have a potential aichivirus-A-like type-V IRES (Fig. 2A/B). Domains H, I, J, K, L and the $Y_{(n)}-X_{(m)}$ -AUG motif are characteristics of type-V IRES elements and were found in the studied sequence [4, 31, 32]. ‘Loop B’, the conserved GNRA tetraloop nucleotide motif in domains Ja and Jb, and the modified eIF4G-binding motifs AGGTACCCG (nt 591 to 599) and CGGATGTGGA (nt 610 to 618) in domain K were also found in the IRES of our isolate [4]. The shape and structural elements of domains D/(D’), E/(E’), G/(G’) were not identifiable [4]. Interestingly, we were unable to identify the origin of replication (ORI), which is usually present at the 5’ terminal end of the 5’UTR in type-V IRES elements [4], but three potential hairpin structures were found. The pyrimidine-rich region (Y: nt 653 to nt 673) and a spacer (X: nt 674 to nt 807) were identified and has 51% nt sequence identity to those of oscivirus A1 (GU182410).

The 3’UTR is 319 nt long and has 70% nucleotide sequence identity to the corresponding region of passerivirus A1 strain thrush/Hong Kong/00356/2007 (GU182406) (Fig. 1). It is also predicted to contain a ‘barbell-like’ secondary RNA structure (Fig. 2C) [16, 29].

The nucleotide sequence motif AAAC/G forms the *cis*-acting replication element (CRE) [7, 15] and is presumed to be active when present in the apical loop of the secondary structure [4, 6, 7, 15]. Based on RNA secondary structure prediction, the thermodynamically most favoured site ($A_{1338}AACG$ or $A_{1346}AACA$ in the loop of the structure with $\Delta G = -21.1$ kcal/mol) of CRE motifs is located between nt 1338 and 1342 or nt 1346 and 1350 in the L protein coding region (Fig. 2D). The genome of the isolate from this study encodes a 2255-aa-long polyprotein that is cleaved into proteins through a post-translational cleavage cascade (Fig. 1).

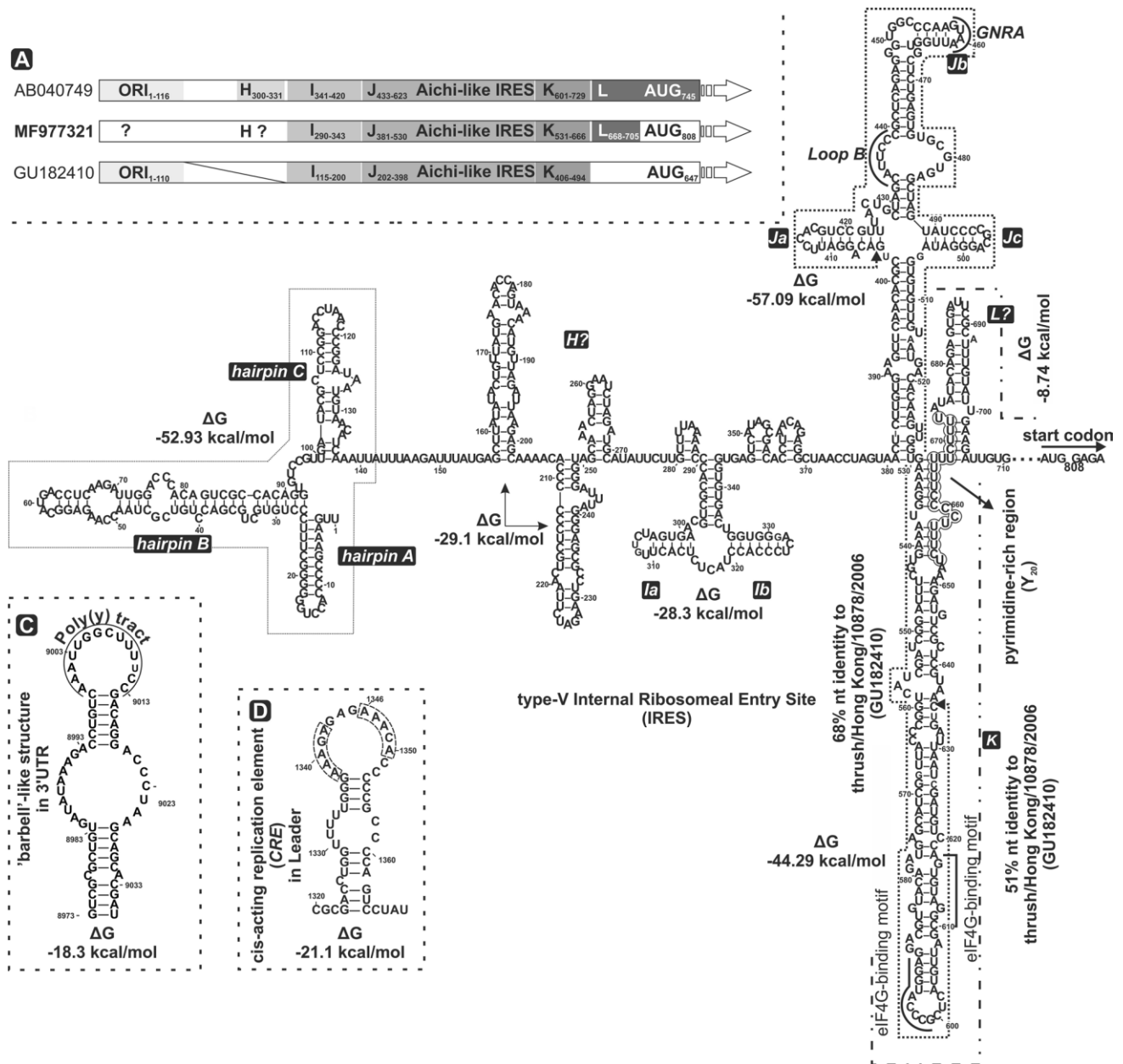


Fig. 2 Predicted RNA secondary structures of finch picornavirus strain waxbill/DB01/HUN/2014 (MF977321). **A**) Schematic representation of the 5'UTR of the study strain based on similarities to the corresponding parts of the closest relative oscivirus A2 (genus *Oscivirus*) strain thrush/Hong Kong/10878/2006 (GU182410) and aichivirus A (genus *Kobuvirus*) strain A846/88 (AB040749). Domains H, I, J, K and L are labelled, indicating the starting and ending nucleotide positions, and the initiation codon of each polyprotein is shaded. **B**) Secondary structure model of the finch picornavirus 5'UTR, including a type-V internal ribosomal entry site (IRES) [4, 29]. The shape and structural elements of domains I, J and K were identified, and domains H and L were hypothesized. The structural elements D, E, F and G were not identified in the studied strain. The conserved 'loop B' and nucleotides GNRA (GUAA) were identified in domain Ja and Jb, respectively. Nucleotides indicated by dotted lines showed 68% and 51% nt sequence identity to *oscivirus A2* (genus *Oscivirus*) strain thrush/Hong Kong/10878/2006 (GU182410). The essential pyrimidine-rich $Y_{(n)}$ motif between domains K and L and the putative in-frame AUG start codon (start from nt 808) are indicated. The predicted 'barbell-like' RNA secondary structure in the 3'UTR (**C**) and the cis-acting replication element (CRE) in the leader (L) protein (**D**) were

modelled *in silico*. The position of the poly(Y) tract in the ‘barbell-like’ structure and the two alternate CREs are indicated by dotted lines in the apical loops of the structures

Potential polyprotein cleavage sites were predicted based on a comparison with the prototype passerivirus A1 strain thrush/Hong Kong/00356/2007 (GU182406) as the closest relative (Fig. 1). The putative L protein is 1239 nt (413 aa) long, and a BLASTx search showed 37% aa sequence identity to the closest match, passerivirus A1 strain thrush/Hong Kong/00356/2007 (GU182406) and 30% aa sequence identity to sicinivirus A1 strain ch/UCC001/Eire (KF741227) in the corresponding region (Fig. 1). The putative 5’UTR of strain thrush/Hong Kong/00356/2007 [8] is in fact a protein-encoding nucleotide sequence based on our analysis, and the predicted protein has 59% aa sequence identity to the L protein of the isolate from this study. Several conserved amino acid motifs, including A₁₂₂ARD, P₁₅₆₉PSLP (**PxLPxNxxxxxxFD**), E₂₂₇DDA and L₂₃₉PFG, were detected in both picornaviruses, which could indicate a homologous function of this protein and suggest that the published genome sequence of the prototype passerivirus A1 strain is not complete (Fig. 1) [33]. The P1 structural and P2 and P3 non-structural regions are 2433 nt (811 aa), 1977 nt (659 aa) and 2355 nt (785 aa) long, respectively. A potential N-terminal myristoylation motif (**GxxxT/S**, **G₄₁₄SVIYN**) was present at the N-terminal end of the hypothetical VP0 protein. The conserved **H**-box/**NC** amino acid motifs (**H₁₂₄₉WAI** and **N₁₃₀₈CTNW**) in 2A; the conserved Walker A (**GxxGxGKS**, **G₁₆₇₃KPGSGKS**), Walker B (**DDLxQ**, **D₁₇₂₄DIGQ**) and Walker C (**KGxxxxSxxxxx(S/T)(S/T) N**, **K₁₇₅₇GMTYTSKVIIMTSN**) motifs in 2C were also identified. Interestingly, the 3B protein is 42 amino acids long, suggesting that there are potentially two copies of unequally long VPgs encoded likewise in the genome of the prototype passerivirus A1 strain (Fig. 1). The RNA-binding domain (**KFRDI**, **Q₂₀₈₆CRDI**), the modified picornavirus H-D-C catalytic triad (**TDV₂₁₃₅Y** and **GLC₂₁₄₇G**), the histidine (**H₂₁₆₇VAG**) residue in the trypsin-like protease domain in 3C, and well-conserved aa motifs (**K₂₃₅₄DELR**, **G₂₄₈₈NPSG**, **Y₂₅₂₈GDD** and **F₂₅₇₇LKR**) in 3D [34] are also present in the isolate from this study.

Comparing the P1^{cap}/2C^{hel}/3C^{pro}/3D^{pol} protein products of waxbill/DB01/HUN/2014 to the closest relative, passerivirus A1 strain thrush/Hong Kong/00356/2007 (GU182406) and the corresponding proteins of representative picornaviruses (Supplementary Table 1), these proteins showed 38.9%, 60.9% and 69.3% aa sequence identity, respectively. Phylogenetic analysis of the predicted protein products these coding regions showed that the strain from this study clustered together with passeriviruses of the family *Picornaviridae* (Fig. 3A).

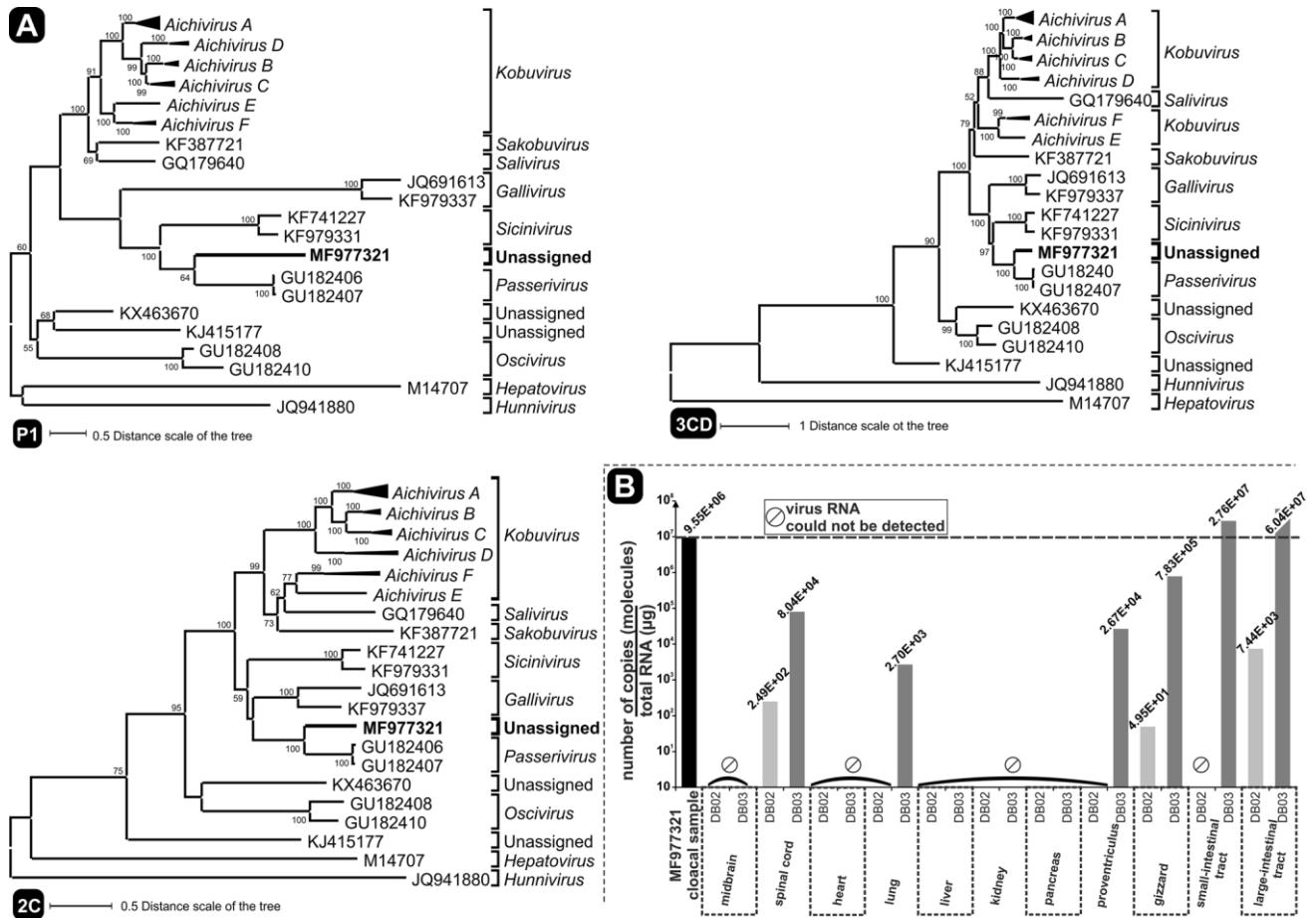


Fig. 3 **A**) Phylogenetic analysis of finch picornavirus strain waxbill/ DB01/HUN/2014 (MF977321 in bold) and representative picorna- viruses, based on the complete P1^{cap}, 2C^{hel} and 3C^{pro}/3D^{pol} nucleotide sequences. Two strains, BHUV1/2008/HUN (JQ941880, hunnivirus A1, genus *Hunnivirus*) and HM-175 (M14707, hepatitis A virus, genus *Hepatovirus*) were used as outgroups in each phylogenetic tree. Evolutionary analysis was conducted in MEGA6 [25], and evolutionary history was inferred using the ML method based on the GTR model with discrete gamma distribution (+G), allowing evolutionarily invariable sites (+I), and bootstrap values were determined for 1000 replicates. The tree is drawn to scale, with branch lengths measured in the number of substitutions per site. Species within the genus *Kobuvirus* were grouped, and representatives of the type species were used in each phylogenetic tree (*Aichivirus A*, KF831027, KJ958930, KM091960, JN387133, JF755427, JQ898342, AB040749, and KJ934637; *Aichivirus B*, GU245693, KF006985, and AB084788; *Aichivirus C*, KF793927, EU787450, and JX177612; *Aichivirus D*, LC055960 and LC055961; *Aichivirus E*, KT325852; *Aichivirus F*, KJ641686 and KJ641691; FFUP1/Portugal/2012, KF387721; hu/NG-J1/Nigeria/2007, GQ179640; turkey/ HUN/M176/2011, JQ691613; ch/518C/Hong Kong/2010, KF979337; newt/II-5-Pilis/2014/HUN, KX463670; tortoise/UF4/USA/2009, KJ415177; robin/Hong Kong/10717/2006, GU182408; thrush/Hong Kong/10878/2006, GU182410; thrush/Hong Kong/00356/2007, GU182406; thrush/Hong Kong/00805/2007, GU182407; ch/ UCC001/Eire, KF741227; ch/55C/Hong Kong/2008, KF979331. **B**) Quantitative analysis of novel finch picornavirus RNA in a violet- eared waxbill DB01 cloacal sample and DB02, DB03 organs and intestines by real-time RT-PCR using SYBR Green I chemistry. The y-axis represents the number of genome copies per total RNA (µg).

Real-time PCR quantification of waxbill/DB01/ HUN/2014 revealed the presence of an average

of 6.58×10^{10} genomic copies per ml of faeces (SD: $\pm 3.37 \times 10^9$) in the cloacal sample. The organs of the other two dead violet-eared waxbills – DB02 and DB03 – were also tested by RT-PCR using the screening primer pair S13-Passeri- 3D-R/F. Nine out of 24 tissue samples were RT-PCR positive (Fig. 3B), and all were confirmed by direct nucleotide sequencing to contain passerivirus sequences. The nucleotide sequence of the viral RNA from organs of DB02 was identical to that of waxbill/DB01/HUN/2014, but a sequence with four nucleotide differences with 98% nt identity was detected in tissue samples of DB3. These mutations resulted in one amino acid change, isoleucine to threonine at aa position 2587 in the 3D^{pol} region. Passerivirus RNA was also detected by qPCR in spinal cord (DB02 and DB03), lung (DB03), gizzard (DB02 and DB03) and proventriculus (DB03), small intestine (DB03) and large intestine (DB02 and DB03) specimens. The viral copy number relative to the amount of total RNA (μg) was the highest in the large intestine of DB03 (higher than in the cloacal sample of DB01), but a high viral load was also detected in other parts of the enteric tract, lung and spinal cord (Fig. 3B).

The Aves class is considered to be the largest class of vertebrates, with over 10,000 currently surviving species. For that very reason, live wild birds could be the key factors for transmission of agents of infectious diseases and could carry and spread potentially unrecognized pathogens to become an emerging risk (<http://www.onehealthinitiative.com/>). On the other hand, our knowledge of viral bird diseases is still fragmentary. The first comprehensive screening of viral agents in birds was published by Woo et al. in 2010 [34]. In that study, 201 different wild bird species of 50 avian families were analysed to investigate the presence of viral agents. The prototype passeriviruses (formerly turdivirus 1) were identified in dead robins and thrushes (*Cosychus saularis* and *Turdus sp.* in avian family *Turdidae*), but among the tested respiratory and alimentary specimens collected from 515 different dead estrildid finches (family *Estrildidae*), passeriviruses were not found using RT-PCR [34]. In the present study, we report the detection and complete genome characterization of a novel passerivirus from an enteric outbreak with high mortality in an estrildid finch population.

To date, strains thrush/Hong Kong/00356/2007 (GU182406; NC_014411) and thrush/Hong Kong/00805/2007 (GU182407) have been the only known members of the genus *Passerivirus*, family *Picornaviridae* [1, 8]. However, the 5' end of both viral genomes could be incomplete based on our analysis. The finch picornavirus strain waxbill/DB01/HUN/2014 showed a passerivirus-like genome organization with high amino acid sequence identity 38.9%, 61.6%, 69.6% in P1^{cap}, 2C^{hel} and 3C^{proD^{pol}}, respectively, to passerivirus A1 (GU182406). According to the taxonomic genus demarcation criteria of the International Committee on Taxonomy of Viruses (ICTV) (http://www.picornastudygroup.com/definitions/genus_definition.htm), novel picornavirus genera are

defined if internal ribosomal entry site (IRES) structures are not homologues and if the amino acid (aa) sequence identity of the orthologous proteins P1^{cap}, 2C^{hel}, 3C^{pro} and 3D^{pol} is less than 34%, 36%, 36% and 36%, respectively. The novel finch picornavirus strain waxbill/DB01/HUN/2014 can therefore be considered a member of the genus *Passerivirus*, and it appears to be the first member of this genus for which a complete genome sequence has been obtained, including the type-V IRES in the 5'UTR and the complete leader (L) sequence between the 5'UTR and the P1 regions.

An origin of replication (ORI) structure with a pseudoknot is common in picornaviruses with type-V IRES elements [4], but we were not able to identify such a structure in the isolate described here. It instead contains three hairpin structures at the beginning of the 5'UTR region, as has been found in murine kobuvirus 1 strain M-5/USA/2010 (JF755427) and rabbit kobuvirus 1 strain rabbit01/2013/HUN (KT325852) [35, 36]. A converse example is the porcine kobuvirus 1 strain sw/S-1-HUN/2007 (EU787450), in which the ORI secondary structure with the pseudoknot is followed by type-IV IRES structures [37]. This observation strengthens the hypothesis that structural elements/modules can be transferred or exchanged between (picorna)virus genomes by recombination, maintaining genetic diversity and virus viability in a new host species [4, 38–40]. The domains I, J, K and the Y_n-X_m-tract between domain K and the initiation AUG motif are believed to be indispensable elements of type-V IRES elements [4, 31, 32]. Focusing on the Y_n-X_m-tract, the pyrimidine-rich region (Y_n) in the finch picornavirus is exceptionally long and is followed by a long spacer (X_m) similar to that in the oscivirus A2 strain thrush/ Hong Kong/10878/2006 (GU182410) (data not shown).

Among picornavirus proteins, the leader (L) protein shows the greatest diversity, both in function and in size [11, 12]. More than half of the known members of picornavirus genera encode a leader (L) protein at the 5' end of the ORF (<http://www.picornaviridae.com>), but only a few of them have a known function [41–45]. The **PPSLP (PxLPxNxxxxxxFD)** aa motif in the L protein of waxbill/DB01/HUN/2014 is present in other picornaviruses of different genera, suggesting a common origin and analogous function of this gene [33]. Despite the amino acid sequence similarity between the leader proteins of the waxbill strain and two other bird-origin picornaviruses, passerivirus A1 strain thrush/Hong Kong/00356/2007 (GU182406) and sicinivirus A1 strain ch/UCC001/Eire (KF741227), the function or the biological activity of this L protein is still unknown.

The order Passeriformes is the largest of all orders in the Aves class, including more than 50% of all bird species. The family *Estrildidae* includes many finch species. These small birds are natively found throughout the old world tropics and Australasia [18]. Estrildid finches are popular as exotic pets, for home-rearing, and for exhibition in Hungary and throughout Europe. Stock farmers in Belgium, the

Netherlands, and Germany successfully breed estrildids, often with no documented information about the origin of the first few generations. Ideally, incoming birds should be isolated in a separate cage/building for 30-days or longer. With this quarantine procedure, many unexpected contagion sources can be excluded. In this case, the incubation period appeared to be no more than 7-10 days [46] between suspected exposure to the virus and onset of clinical signs.

Although there is some evidence that the picornavirus was related to the disease, i) passerivirus sequence reads from the cloacal specimen represents up to the 90% of the total viral metagenomic sequence reads, suggesting an enterically transmitted agent, ii) samples from more than one affected bird (three of the three tested dead estrildid finches) with similar symptoms from the same outbreak were positive for this novel passerivirus, iii) the passerivirus genome sequence is present in different organs, indicating an extra-intestinal and generalized infection, and iv) a high passerivirus viral load was measured in a cloacal specimen, a direct etiological connection and confirmation is lacking in this retrospective study. FFPE specimens were not available for *in situ* hybridization, and there was no inoculation study of naïve animals or additional testing of healthy and sick populations, which could confirm that the virus might be correlated with disease. Passeriviruses in the family *Picornaviridae* might be responsible for sporadic disease in wild birds and outbreaks in home-kept birds with fatal outcome. Further molecular epidemiological studies are needed to explore the diversity, biology, and detailed pathogenesis of passerivirus in domestic and wild bird species.

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Compliance with ethical standards

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Conflict of interest The authors declare that they have no conflict of interest.

Ethical approval All applicable international, national, and/or institutional guidelines for the care and use of animals were followed.

Supplementary Table 1. The amino acid (aa) identity of a finch picornavirus strain waxbill/DB01/HUN/2014 (MF977321) to the corresponding regions of representative picornaviruses. The highest aa values are bolded. The identity calculation was carried out using BioEdit [26], based on the codon-based aligned nucleotide sequences by MEGA6 [25].

Genus	Type	Strain	Acc. num.	waxbill/DB01/HUN/2014 (MF977321)		
				P1 ^{cap}	2C ^{hel}	3C ^{proD} pol
<i>Kobuvirus</i>	Feline kobuvirus 1 (FeKV-1)	FK-13/South Korea/2011	KF831027	31.9	45.7	51.7
		12D240/South Korea/2012	KJ958930	33	46.3	51.1
		TE/52/IT/13	KM091960	32.9	46.1	51.6
	Murine kobuvirus 1 (MuKV-1)	M-5/USA/2010	JF755427	32.8	46.8	50.6
		KoV-SewKTM (Nepal/2009)	JQ898342	30.1	46.1	49.9
	Canine kobuvirus 1 (CaKV-1)	dog/AN211D/USA/2009	JN387133	31.2	45.4	50
	Aichi virus 1 (AiV-1)	A846/88	AB040749	32.2	45.9	51.5
	European roller kobuvirus 1 (ERKV-1)	SZAL6-KoV/2011/HUN	KJ934637	32.9	45.2	49.2
	Bovine kobuvirus 1 (BKV-1)	bov/U-1/Japan	AB084788	33	46	50.1
	Ferret kobuvirus (FKV)	ferret/MpKoV38/NL/2010	KF006985	32.4	45.9	51.5
	Ovine kobuvirus 1 (OKV-1)	sheep/TB3/HUN/2009	GU245693	32.8	46.9	50.8
	Porcine kobuvirus 1 (PKV-1)	sw/S-1-HUN/2007	EU787450	32.4	44.9	49.8
		wb/WB-1-HUN/2011	JX177612	32.7	45	48.7
	Caprine kobuvirus (CapKV)	black goat/12Q108/KOR/2012	KF793927	32	45.5	49.7
	Kagovirus 1 (KV-1)	cattle/Kagoshima-1-22-KoV/2014/JPN	LC055961	32	44.2	49.6
	Kagovirus 2 (KV-2)	cattle/Kagoshima-2-24-KoV/2015/JPN	LC055960	30.8	43.9	49.2
Rabbit kobuvirus 1 (RKV-1)	Rabbit01/2013/HUN	KT325852	31.8	42.7	49.8	
Aichivirus F1 (AiV-F1)	BtMr-PicoV/JX2010	KJ641686	32.6	46.1	48.8	
Aichivirus F2 (AiV-F2)	BtMf-PicoV-2/GD2012	KJ641691	32.5	44	51.2	
<i>Salivirus</i>	Salivirus A1 (SaV-A1)	hu/NG-J1/Nigeria/2007	GQ179640	31.9	46	46
<i>Sakobuvirus</i>	Sakobuvirus A1 (SakV-A1)	FFUP1/Portugal/2012	KF387721	30.7	44.6	49.9
<i>Gallivirus</i>	Chicken gallivirus 1 (ChGV-1)	chicken/518C/Hong Kong/2010	KF979337	29.4	46.5	54.3
	Gallivirus A1 (GV-A1)	turkey/M176/2011/HUN	JQ691613	29	43.9	55
<i>Sicinivirus</i>	Sicinivirus A1 (SiV-1)	chicken/UCC001/Eire	KF741227	34.7	50.3	57.6
	unassigned siciniviruses	chicken/55C/Hong Kong/2008	KF979331	34	49.3	58.1

<i>Passerivirus</i>	Passerivirus A1 (PasV-A1)	thrush/Hong Kong/00356/2007	GU182406	38.9	60.9	69.3
		thrush/Hong Kong/00805/2007	GU182407	38.9	61.6	69.6
Unassigned	Livupivirus A1 (LiV-A1)	newt/II-5-Pilis/2014/HUN	KX463670	33.6	41.6	45.5
Unassigned	Tortoise rafivirus A1	tortoise/UF4/USA/2009	KJ415177	31	38.7	43.9
<i>Oscivirus</i>	Oscivirus A2	thrush/Hong Kong/10878/2006	GU182410	30.7	42.1	45.9
	Oscivirus A1	robin/Hong Kong/10717/2006	GU182408	30.9	41.3	48.8
<i>Hepatovirus</i>	Human hepatitis A virus (HHAV)	HM-175	M14707	24.6	33.9	34.5
<i>Hunnivirus</i>	Hunnivirus A1 (HuV-A1)	BHUV1/2008/HUN	JQ941880	29.1	35.5	34.7

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