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# Independent evolution of tetraloop in enterovirus oriL replicative element and its putative binding partners in virus protein 3C

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**Background.** Enteroviruses are small non-enveloped viruses with (+) ssRNA genome with one open reading frame. Enterovirus protein 3C (or 3CD for some species) binds the replicative element oriL to initiate replication. The replication of enteroviruses features low fidelity, which allows the virus to adapt to the changing environment on the one hand, and requires additional mechanisms to maintain the genome stability on the other. Structural disturbances in the apical region of oriL domain d can be compensated by amino acid substitutions in positions 154 or 156 of 3C (amino acid numeration corresponds to poliovirus 3C), thus suggesting the co-evolution of these interacting sequences in nature. The aim of this work was to understand co-evolution patterns of two interacting replication machinery elements in enteroviruses, the apical region of oriL domain d and its putative binding partners in the 3C protein.

**Methods.** To evaluate the variability of the domain d loop sequence we retrieved all available full enterovirus sequences (>6400 nucleotides), which were present in the NCBI database on February 2017 and analysed the variety and abundance of sequences in domain d of the replicative element oriL and in the protein 3C.

**Results.** A total of 2,842 full genome sequences was analysed. The majority of domain d apical loops were tetraloops, which belonged to consensus YNHG (Y=U/C, N=any nucleotide, H=A/C/U). The putative RNA-binding tripeptide 154-156 (Enterovirus C 3C protein numeration) was less diverse than the apical domain d loop region and, in contrast to it, was species-specific.

**Discussion.** Despite the suggestion that the RNA-binding tripeptide interacts with the apical region of domain d, they evolve independently in nature. Together, our data indicate the plastic evolution of both interplayers of 3C-oriL recognition.

1 Independent evolution of tetraloop in enterovirus oriL  
2 replicative element and its putative binding partners in  
3 virus protein 3C

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## Abstract

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with one open reading frame. Enterovirus protein 3C (or 3CD for some species) binds the

replicative element oriL to initiate replication. The replication of enteroviruses features low

fidelity, which allows the virus to adapt to the changing environment on the one hand, and

requires additional mechanisms to maintain the genome stability on the other. Structural

disturbances in the apical region of oriL domain d can be compensated by amino acid

substitutions in positions 154 or 156 of 3C (amino acid numeration corresponds to poliovirus

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work was to understand co-evolution patterns of two interacting replication machinery elements

in enteroviruses, the apical region of oriL domain d and its putative binding partners in the 3C

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available full enterovirus sequences (>6400 nucleotides), which were present in the NCBI

database on February 2017 and analysed the variety and abundance of sequences in domain d of

the replicative element oriL and in the protein 3C.

**Results.** A total of 2,842 full genome sequences was analysed. The majority of domain d

35 apical loops were tetraloops, which belonged to consensus YNHG (Y=U/C, N=any nucleotide,  
36 H=A/C/U). The putative RNA-binding tripeptide 154-156 (*Enterovirus C* 3C protein  
37 numeration) was less diverse than the apical domain d loop region and, in contrast to it, was  
38 species-specific.

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40 **Discussion.** Despite the suggestion that the RNA-binding tripeptide interacts with the apical  
41 region of domain d, they evolve independently in nature. Together, our data indicate the plastic  
42 evolution of both interplayers of 3C-oriL recognition.

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## Introduction

45 Enteroviruses are small non-enveloped viruses with a plus strand genome about 7500 nt  
46 long, which contains one open reading frame that encodes structural (capsid) and non-structural  
47 proteins, 5' and 3' NTRs (non translated regions), and polyA on the 3' end (Palmenberg,  
48 Neubauer and Skern, 2010). (Figure 1). Most non-structural enterovirus proteins are  
49 polyfunctional. Protease 3CD is a precursor of polymerase 3D and plays a key role in the  
50 initiation of replication (Harris *et al.*, 1992; Gamarnik and Andino, 1998; Thompson and  
51 Peersen, 2004). After translation by host cell ribosomal machinery, the genome is utilized for the  
52 synthesis of the (-) strand RNA, which, in turn, serves as a matrix for the synthesis of multiple

53 daughter (+) strands. Non-translated regions of the genome and a coding sequence within the  
54 genomic region encoding the viral helicase 2C contain replicative elements, which interact with  
55 viral and host proteins. These RNA-protein complexes regulate initiation and further steps of  
56 replication. For poliovirus, the most clinically relevant member of the *Enterovirus* genus, there  
57 are at least three known RNA-protein complexes, which are formed with the replicative elements  
58 oriL, oriR and oriI during replication (Figure 1).

59         Complex oriL with viral protein 3CD and the host protein PCBP2 is crucial for the  
60 transcription initiation (Goodfellow *et al.*, 2000; Vogt and Andino, 2010; Chase, Daijogo and  
61 Semler, 2014). The element oriL has a cloverleaf-like secondary structure with four domains  
62 termed a (stem of the cloverleaf), b, c and d (leaves of the cloverleaf) (Trono, Andino and  
63 Baltimore, 1988; Andino, Rieckhof and Baltimore, 1990) (Figure 1). Previously, it was  
64 demonstrated *in vitro* that 3CD (or 3C) of poliovirus, coxsackievirus B3 and bovine enterovirus  
65 1 interacts with the apical loop and the flanking base pairs of hairpin d in the oriL element  
66 (Andino, Rieckhof and Baltimore, 1990; Du *et al.*, 2003; Ihle *et al.*, 2005) (Figure 1).

67         The apical loops of domain d in genomes belonging to several viruses of *Enterovirus*  
68 genera were shown by NMR experiments to be tetraloops with a specific spatial structure, which  
69 belongs to the UNCG structural class of stable tetraloops (Du *et al.*, 2003, 2004; Ihle *et al.*, 2005;  
70 Melchers *et al.*, 2006). There are several known structural classes of tetraloops, three of which,  
71 named according to consensus sequences, contain tetraloops of extreme stability: UNCG (where

72 N=any nucleotide), GNRA (where R=A/G) and gCUUGc (Uhlenbeck, 1990; Cheong and  
73 Cheong, 2010). Tetraloops of UNCG and GNRA classes are the most widely represented (Woese  
74 *et al.*, 1990; Cheong and Cheong, 2010; Bottaro and Lindorff-Larsen, 2017). Previously, it was  
75 shown that only tetraloops of the UNCG structural class, but not tetraloops of GNRA or  
76 gCUUGc structural classes, can support effective replication of the poliovirus genome (Prostova  
77 *et al.*, 2015). Moreover, the exact sequence of the apical region of poliovirus domain d was of  
78 less importance for effective 3CD-oriL recognition than its spatial structure (Rieder *et al.*, 2003;  
79 Prostova *et al.*, 2015). At the same time, structural disturbance in the apical region of oriL  
80 domain d of poliovirus could be compensated by amino acid substitutions in the tripeptide 154-  
81 156 of the 3C protein (here and hereafter amino acid numeration corresponds to poliovirus 3C  
82 protein) (Andino *et al.*, 1990; Prostova *et al.*, 2015). In addition to triplet 154-156, the conserved  
83 motif <sub>82</sub>KFRDI<sub>86</sub> of the 3C protein also takes part in the oriL recognition (Andino *et al.*, 1990,  
84 1993; Hämmerle, Molla and Wimmer, 1992; Shih, Chen and Wu, 2004). To date a  
85 comprehensive analysis of the diversity in domain d apical region and amino acid tripeptide  
86 sequences in the *Enterovirus* genus has not been conducted.

87         The replication of an enterovirus is a low-fidelity process, generating, on average, one  
88 mutation per genome (Sanjuán *et al.*, 2010; Acevedo, Brodsky and Andino, 2013). The high  
89 probability of mutation allows the virus to adapt to a constantly changing environment on the  
90 one hand, but requires additional mechanisms to maintain genome stability on the other (Wagner

91 and Stadler, 1999; Lauring, Frydman and Andino, 2013). The aim of the present study was to  
92 understand co-evolution patterns of two interacting replication machinery elements in  
93 enteroviruses, the apical domain d of oriL and the 3C protein.

94

## 95 **Materials and methods**

### 96 **Formation and filtration of sets of full genomes**

97 All available nucleotide sequences (as of February 2017) containing the *Enterovirus*  
98 genus with length 8000>n>6800 were extracted from the NCBI database. For every species, a  
99 multiple sequence alignment was conducted using MAFFT version 7 with default settings (Katoh  
100 and Standley, 2013). Sequences that contained more than 50 N characters in succession and  
101 sequences that were annotated as “Modified\_Microbial\_Nucleic\_Acid”, were removed from  
102 alignments. All sequences that differed from any other sequence in the dataset by less than 1% of  
103 the nucleotide sequence were omitted in order to reduce the bias caused by over-represented  
104 sequences.

### 105 **Analysis of tetraloop and amino acid variety in the sets of genomes**

106 For analysis of domain d sequence variety, the multiple sequence alignments were used.

107 The relevant region of multiple sequence alignment and the respective names of sequences were



108 analysed in Microsoft Excel. To analyse correlation of the domain d loop and tripeptide of 3C  
109 sequences the same alignments were translated in the protein 3C coding region. The resulting  
110 amino acid sequences that corresponded to tripeptides 154-156 (poliovirus 3C numerations) were  
111 analysed using Microsoft Excel. An amino acid frequency plot was created via the WebLogo  
112 server using the set of filtered genomes for every species (Crooks *et al.*, 2004). To do this, the  
113 multiple sequence alignment of filtered genomes of every species was translated in the region  
114 that codes protein 3C, while positions 71-89 and 147-160 were saved in separate MAS files,  
115 which were then used to produce logos.

## 116 **Domain d secondary structure**

117 The domain d secondary structure was folded using the Vienna RNA Websuite server  
118 with subsequent manual editing (Gruber *et al.*, 2008; Lorenz *et al.*, 2011). Algorithm accounting  
119 for minimum free energy and partition function was used.

## 120 **Results**

### 121 **Sample characteristics**

122 To evaluate the variability of the domain d loop sequence we retrieved all available  
123 complete genome (8000>n>6800 nucleotides) enterovirus (EV) sequences that were present in  
124 the NCBI database on February 2017. Representatives of *Enterovirus A* (1173 sequences in

125 total), *Enterovirus B* (414), *Enterovirus C* (773), *Enterovirus D* (462), *Enterovirus E* (12),  
126 *Enterovirus F* (13), *Enterovirus G* (15), *Enterovirus H* (3), *Enterovirus J* (7), *Rhinovirus A* (202),  
127 *Rhinovirus B* (76) and *Rhinovirus C* (51) species were analysed. As expected, genomes of  
128 epidemiologically significant viruses were the most represented in the database. For example,  
129 66% of *Enterovirus A* species genomes belonged to the EV71 type, the causative agent of hand,  
130 foot and mouth disease (Solomon *et al.*, 2010); most *Enterovirus C* species sequences (78%)  
131 belonged to poliovirus; and most *Enterovirus D* species sequences (98.7%) represented EV68, an  
132 aetiological agent of severe respiratory illness (Oermann *et al.*, 2015). The number of genome  
133 sequences of each species that contained the oriL region is shown in Table 1.

134 Sequences of apical regions in oriL domain d and the amino acids involved in RNA  
135 recognition in 3C protein were analysed. In genomes of *Enterovirus E* and *Enterovirus F* species  
136 with two oriLs (Pilipenko, Blinov and Agol, 1990; Zell *et al.*, 1999) sequences of both oriLs  
137 were analysed (Table 1). To reduce the bias towards particular loop sequences present in a large  
138 set of closely related genomes, which, for example, belonged to one outbreak, all sequences that  
139 differed from any other sequence in the dataset by less than 1% of the nucleotide sequence were  
140 omitted. After curation, the sizes of the largest data sets decreased dramatically, but the number  
141 of unique loop sequences in every set did not change significantly (Table 1). Unique tetraloop  
142 variants were lost for *Enterovirus A* (tetraloop UGUG), *Enterovirus C* (tetraloops CCCG, CAUG  
143 and UGUG) and *Enterovirus D* (tetraloop UUGG). This indicates that, even among closely

144 related genomes, the tetraloop sequence can vary. Indeed, in several outbreaks caused by EV71  
145 or PV1, closely related genomes contained different apical domain d sequences (not shown). It  
146 should be noted that the filtration of the dataset using a 95% sequence identity threshold resulted  
147 in a dramatic loss of unique tetraloop variants (107 genomes out of 1052 were left after filtration,  
148 while 13 unique tetraloop variants out of total 17 variants were detected in the filtered set).

### 149 **Variability in the oriL domain d apical loop sequence**

150 The secondary structure of domain d was conserved in all species, except *Enterovirus G*,  
151 in which an elongated domain d was observed (Figure 2) (Krumbholz *et al.*, 2002).

152 The variety and occurrence of various loops in the apical region of domain d in all  
153 species of the *Enterovirus* genera were analysed in filtered sets of full genome sequences. Most  
154 domain d apical loops were tetraloops (i.e., they consisted of four nucleotides) (Table 2).  
155 However, triloops (3-nucleotide loop) could be found in genomes of *Enterovirus C* and  
156 *Rhinovirus A* and *B* species, whereas pentaloops (5-nucleotide loop) were detected in genomes of  
157 *Enterovirus E* species (Table 2).

158 The most common loop sequences belonged to consensuses YNMG (Y=C/U, N=any,  
159 M=A/C; tetraloops with UNCG class spatial structure belong to this consensus) and YNUG  
160 (tetraloops with UNCG class and gCUUGc class spatial structures belong to this consensus)  
161 (Table 2). Consensus YNMG and consensus YNUG together corresponded to 24 unique  
162 sequence variants. Interestingly, in our dataset of 2,842 full genomes, four tetraloops out of 24

163 possible variants were never found in the domain d apical region: UUAG, UCAG, CUAG and  
164 CCAG (Table 2). Thus, dinucleotides UA and CA are likely to be avoided at positions 2 and 3 of  
165 the tetraloop in enterovirus genomes.

166 In *Enterovirus A* species, 17 out of 24 possible unique tetraloop sequences were  
167 identified (Table 2, Table S2). Twelve unique loops of *Enterovirus A* belonged to consensus  
168 YNMG, while the other five belonged to consensus YNUG. The most abundant tetraloop in  
169 *Enterovirus A* genomes, in contrast to other species, was CUCG (Table 2, Table S1). This is  
170 explained by the prevalence of this tetraloop in the EV71 C4 genotype (Table 2, Table S1). The  
171 frequency of other tetraloop sequences varied significantly (Table 2, Table S2). One tetraloop  
172 (UGUG) was lost upon filtration. Such sequences here and below were manually added to the  
173 final data set to maintain diversity of loop sequences within the species, as well as provide  
174 comprehensive information about sequences in apical domain d in viable viruses (Table 2).  
175 Interestingly, EV71 sequences contained 13 out of 17 tetraloop variants, which were detected in  
176 the *Enterovirus A* genus (Table 2). In other words, the diversity of tetraloops in one discrete  
177 lineage in general resembles its diversity in the unification of different discrete lineages.

178 In *Enterovirus B* genomes, 18 unique tetraloops out of 24 possible were found. Twelve  
179 of these tetraloops belonged to consensus YNMG and six to consensus YNUG (Table 2, Table  
180 S3). The most abundant tetraloops were CACG (98 genomes), UACG (51 genomes) and UGCG  
181 (31 genomes), which were also present among the most abundant tetraloops of *Enterovirus A*

182 species.

183 In genomes of the *Enterovirus C* species, nine unique tetraloops belonged to the YNMG  
184 consensus and four to the YNUG consensus. Three unique tetraloops were lost upon filtration  
185 and added to the final data set (CCCG, UGUG, CAUG) (Table 2, Table S4). Two genomes  
186 annotated in the NCBI data base as Human coxsackievirus A21, strain Coe, (accession number  
187 D00538) and Human coxsackievirus A21, strain BAN00-10467, (accession number EF015031)  
188 contained triloops CAG and CCG, respectively. The most abundant tetraloops in *EV-C* species  
189 were UACG (106), CACG (101), UGCG (43) (Table 2, Table S1, Table S4), which corresponds  
190 to the Sabin vaccine strains of poliovirus serotypes 2, 3 and 1, respectively. To evaluate bias  
191 caused by the redundant number of vaccine strain sequences in the data set, we subtracted  
192 genomes of vaccine/vaccine derived poliovirus strains from the analysed set. As a result,  
193 tetraloops UACG, CACG and UGCG were still the most frequent variants (Table 2, Table S1).

194 Only 57 *Enterovirus D* genomes out of 419 were left after 1% identity filtration. Fifty  
195 genomes belonged to Human enterovirus 68, the aetiological agent of respiratory illness. All  
196 genomes of this type contained loop UUCG in the domain d apical region. Other tetraloops were  
197 UUUG (1), CUCG (2), CCCG (1), CUUG (2) and CACG (1) (Table 2, Table S5). One tetraloop  
198 (UUGG) was lost upon filtration and manually added to the final data set.

199 Species *Enterovirus E* and *F* have two oriLs in the 5'NTR, generally with similar  
200 sequences in the apical region of domain d (Pilipenko, Blinov and Agol, 1990; Zell *et al.*, 1999)

201 (Table S6). As such, we united sequences from the first and the second oriL of these viruses in  
202 the heat map (Table 2). Domain d loops in 10 genomes of *Enterovirus F* were tetraloops, while,  
203 in 10 *Enterovirus E* genomes, there were both tetraloops (first oriL) and pentaloops (first and  
204 second oriL) (Table 2, Table S6). There were four diverse tetraloop sequences in oriLs of  
205 Enterovirus E and F with no obvious preference between these species. These sequences were  
206 GCUA, GUUA, GCCA, AUUA (Table 2, Table S6). Tetraloop AUUA was found once in the  
207 first oriL domain d of EV-F (strain PS87/Belfast, accession number DQ092794) (Table 2, Table  
208 S6). There were six diverse pentaloop sequences in domain d of Enterovirus E genomes –  
209 GCUUA, GUUUA, GCCUA, GCGUA, GAUUA, GUCUA (Table 2, Table S6).

210 All domain d loops in genomes of Enterovirus G, H and J species were tetraloops; all  
211 except one tetraloop variant belonged to consensus YNMG (Table 2, Table S7). One Enterovirus  
212 G representative had a GUUA tetraloop sequence (strain LP 54, accession number AF363455),  
213 similar to loops of Enterovirus E and F species (Table 2). This genome had only one oriL with  
214 the same domain d length as that of Enterovirus G genomes (Krumbholz *et al.*, 2002).

215 All except one (isolate V38\_URT-6.3m, accession number JF285329) of the full  
216 genomes of *Rhinovirus A* species and all full genomes of *Rhinovirus C* species had tetraloops in  
217 the apical regions of domain d (Table 2). Tetraloops of these viruses in almost all cases belonged  
218 to consensus YNMG, with one exception found in *Rhinovirus C* (tetraloop CUUC, isolate JAL-1,  
219 accession number JX291115) (Table 2, Table S8). All loops in the apical region of *Rhinovirus B*

220 domain d were triloops (Table 2, Table S8).

221 Thus, the secondary structure of domain d was very similar among species of the  
222 *Enterovirus* genus, with the exception of *Enterovirus G* species (Figure 2). The apical region of  
223 domain d has a high diversity of sequences; however, in species of *Enterovirus A, B, C, D, G, H*  
224 and *J* and *Rhinovirus A* and *C*, it mostly corresponds to the same consensus, that is, YNHG  
225 (Y=C/U, N=any, H=A/C/U).

### 226 **Variety of RNA-recognition tripeptide of 3C**

227 Two motifs of protein 3C are involved in RNA recognition and interact with oriL: the  
228 conservative motif KFRDI (positions 82-86 of poliovirus 3C) and the putative RNA-binding  
229 tripeptide (positions 154-156 in poliovirus 3C) (Andino *et al.*, 1990, 1993; Hämmerle, Hellen  
230 and Wimmer, 1991; Shih, Chen and Wu, 2004). Substitutions in the putative RNA-binding  
231 tripeptide are known to compensate for disturbance in the apical region of domain d, such that  
232 the RNA-binding tripeptide is a putative candidate to co-evolve with the domain d loop (Andino  
233 *et al.*, 1990; Prostova *et al.*, 2015). There are other amino acids that have been found to affect  
234 oriL-3CD interaction, but tripeptide 154-156 (*Enterovirus C* 3C protein numbering here and  
235 below) is the only one that is proven to compensate structural disturbance in the domain d apical  
236 region (Andino *et al.*, 1990, 1993). To evaluate the possible co-evolution between the domain d  
237 tetraloop and its putative interaction partners in protein 3C, relevant sequences in the filtered full  
238 genome data sets were analysed.

239 Motif  $_{82}\text{KFRDI}_{86}$  was conserved in all species, as well as amino acids Glu 71 and Cys  
240 147 of the protease catalytic triad (Figure 3). Always in second position of the putative RNA-  
241 binding tripeptide (position 155) was Gly.

242 No mutual dependence between loop sequences and tripeptide sequences was found  
243 within enterovirus genomes of the same species. For example, *Enterovirus A* genomes contained  
244 17 unique variants of the tetraloop sequence, whereas the predominant fraction of 3C sequences  
245 (548 out of 564) contained the conservative tripeptide VGK at positions 154-156 (Figure 3,  
246 Table S9). It is noteworthy that, this tripeptide was not found exclusively only in genomes of the  
247 EV71 serotype, although genomes of this serotype prevailed in the data set. Other *Enterovirus A*  
248 genomes contained tripeptides VGR (seven out of 564), TGK (four out of 564), IGK (three out  
249 of 564), VGE (one out of 564) and SRK (one out of 564) (Figure 3, Table S9). Genomes with  
250 tripeptides other than VGK contained no peculiarities of the domain d loop sequence (Table S9).  
251 This observation confirms that the specific loop sequence is not likely to be the main subject for  
252 recognition by the RNA-binding tripeptide. Similarly, all or almost all genomes of *Enterovirus B*  
253 (242 out of 244), *Enterovirus C* (272 out of 274), *Enterovirus D* (all), *Enterovirus G* (seven out  
254 of 8), *Enterovirus H* (a total of two: genomes – one with TGK, one with TGR), *Enterovirus J*  
255 (all) and *Rhinovirus B* (36 out of 37) species contained tripeptide TGK at positions 154-156 of  
256 the 3C protein (Figure 3, Table S7, S9, S10). Alternative tripeptides were TGR in two genomes  
257 of *Enterovirus B* and one genome of *Enterovirus H*; IGK in one genome of *Enterovirus C*



258 species and in one genome of *Rhinovirus B* species; PGK in one genome of *Enterovirus C*  
259 species; and MGK in one genome of *Enterovirus G* species (Table S7, S9, S10).

260 Genomes of *Enterovirus E* and *F* species contained two oriLs with tetraloops in domain d  
261 mostly of consensus GYYA or pentaloops of consensus GHBUA, where H = A/C/U and  
262 B=U/C/G. All genomes contained tripeptide MGK at positions 154-156 of protein 3C (Figure 3,  
263 Table S7). Interestingly, a similar loop-tripeptide pair was found in one genome of *Enterovirus G*  
264 species (strain LP 54, accession number AF363455). It contained tetraloop GUUA in domain d  
265 of its single oriL and tripeptide MGK in 3C. Unlike this unique genome, other genomes of  
266 *Enterovirus G* species contained tetraloops of YNMG consensus and tripeptide TGK in the  
267 protein 3C.

268 Rhinovirus genomes contained tetraloops, mostly of consensus YNMG (*Rhinovirus A*  
269 and *C* species) or triloops (*Rhinovirus A* and *B*) (Table 2). *Rhinovirus A* genomes with tetraloops  
270 in domain d contained tripeptides in 3C with positively charged amino acid before the tripeptide,  
271 but not in its final position, as in case of genomes of *Enterovirus A-C* species (Figure 3, Table  
272 S11, S12). The sequence of tripeptides, which did not depend on the tetraloop sequence, was, in  
273 descending order, IGQ (the most abundant, 65 genomes out of 119), IGL (20 genomes out of  
274 119), IGS, VGS, IGN, VGQ, IGV and VGH (Table S11). In the case of the *Rhinovirus A*  
275 genome with triloop UCU in domain d (isolate V38\_URT-6.3m, accession number JF285329),  
276 protein 3C contained tripeptide TGK without positively-charged amino acid before it (Table

277 S11). All genomes of *Rhinovirus B* species contained triloops in domain d, with all but one (with  
278 IGK) containing tripeptide TKG in 3C. Genomes of *Rhinovirus C* contained tetraloops mostly of  
279 consensus YHCG (H=all but G) and tripeptides in 3C without a positively charged amino acid at  
280 the last position (TGN, VGN, TGH) or outside of the tripeptide (Table S12). One genome  
281 contained tetraloop CUUC paired with most abundant tripeptide, that is, TGN (23 out of 37  
282 genomes)(Table S12).

283 Thus, dependence between apical domain d sequences and tripeptides in protein 3C  
284 within a species was not detected (Figure 3). We can state that the tripeptide and motif KFRDI  
285 are almost non-variable within a species compared to the domain d loop sequence, but there is a  
286 specifically preferred tripeptide sequence for each species. Hence, tripeptide sequences are  
287 species-specific, while the domain d loop sequences are almost universal among *Enterovirus A*,  
288 *B*, *C* and *D* and *Rhinovirus A* and *C* species.

## 289 Discussion

290 Most of the domain d apical loops in enterovirus genomes were represented by tetraloops.  
291 The most common variants of tetraloop sequences corresponded to consensus YNHG (Y=C/U,  
292 N=any, H=A/C/U) (Table 2). Similar results were obtained in our previous experimental work,  
293 where eight apical nucleotides of domain d of the poliovirus genome were randomized and  
294 viable variants were selected *in vitro*, with the majority of selected tetraloops belonging to

295 consensus YNHG (Prostova *et al.*, 2015). Some tetraloops of consensus YNHG were found in  
296 genomes in the NCBI database, but not among the variants selected *in vitro*, namely tetraloops  
297 CACG, CUCG, UAAG, UGAG, CAAG, CGAG, UGUG and CUUG (Prostova *et al.*, 2015).  
298 Tetraloops UGAG, UGUG and CUUG were reconstructed with a U\*\*\*\*G flanking base pair in  
299 the context of the poliovirus genome strain Mahoney, which supported effective virus replication  
300 (Prostova *et al.*, 2015).

301         Conversely, tetraloops UUAG, UCAG and CCAG, found in domain d of selected *in vitro*  
302 viable poliovirus variants, were able to support virus reproduction; however, they were not found  
303 in naturally circulating viruses (Prostova *et al.*, 2015). One tetraloop of the YNHG consensus  
304 (CUAG) was neither found in genomes from the NCBI database (n=2842), nor in the  
305 randomized poliovirus genomes selected *in vitro* (n=62) (Table 2). Thermodynamic stability is  
306 unlikely to be the reason why this and other tetraloops were unrepresented as the melting  
307 temperature of stem loops with avoided tetraloops is within range of the melting temperature of  
308 YNHG tetraloops, which supported replication (Proctor *et al.*, 2002). Moreover, tetraloops  
309 UUAG and UCAG are common in rRNA (Woese *et al.*, 1990). Sample insufficiency cannot be  
310 excluded for both database and *in vitro* selected sets of genomes, but it is safe to conclude that  
311 these tetraloop variants are at least extremely rare. In any case, the fact that the incidence of  
312 these tetraloops is much less than for other tetraloops indicates that such variants are possibly  
313 less fit.

314 The most abundant tetraloops in the domain apical region of genomes from the NCBI  
315 database and variants selected *in vitro* could be compiled into consensus UNCG and CNCG  
316 (Table 2, Table S1). At the same time, these tetraloops are most abundant in rRNA, and, with  
317 certain closing base pairs, among the most thermodynamically stable tetraloops (Woese *et al.*,  
318 1990; Proctor *et al.*, 2002). Tetraloops of these consensus and some other found tetraloops of  
319 the YNHG consensus form a specific spatial structure of the UNCG structural class of stable  
320 tetraloops (Cheong, Varani and Tinoco, 1990; Varani, Cheong and Tinoco, 1991; Du *et al.*,  
321 2003, 2004).

322 Another set of tetraloops, which correspond to GNYA consensus, was found both in  
323 genomes of *Enterovirus E* and *F* and in genomes of viable polioviruses selected *in vitro*  
324 (Prostova *et al.*, 2015). Tetraloop GCUA was able to support the effective replication of  
325 poliovirus and, together with tetraloop GUUA, is known to assume an UNCG fold (Ihle *et al.*,  
326 2005; Melchers *et al.*, 2006; Prostova *et al.*, 2015). In sum, these data suggest that the spatial  
327 structure, rather than the exact sequence, is the main subject for recognition by virus protein 3C.  
328 Structure-based recognition of tetraloops occurs in several known RNA-protein complexes. For  
329 example, tetraloops with a GNRA class structure in the context of bacteriophages P22 and  $\lambda$   
330 genome transcription antitermination element boxB are specifically recognized by the  
331 bacteriophage N-protein arginine-rich motif (ARM) (Cai *et al.*, 1998; Legault *et al.*, 1998;  
332 Schärpf *et al.*, 2000). Arginines and lysines of the ARM recognize the shape of the negatively

333 charged phosphodiester backbone of the stem-loop and positions N-peptide for hydrophobic or  
334 stacking interaction with a non-conserved nucleotide of the loop (Cai *et al.*, 1998; Legault *et al.*,  
335 1998; Schärpf *et al.*, 2000; Thapar, Denmon and Nikonowicz, 2013). Another example of  
336 structure-specific recognition is the complex of the double-stranded RNA-binding domain  
337 (dsRBD) of RNase Rnt1p and AGNN class tetraloop (Chanfreau, Buckle and Jacquier, 2000;  
338 Lebars *et al.*, 2001; Wu *et al.*, 2001, 2004; Wang *et al.*, 2011; Thapar, Denmon and Nikonowicz,  
339 2013). Motif dsRBD recognizes the phosphodiester backbone at the 3' side of the tetraloop and  
340 its non-conserved third and fourth nucleotides (Wu *et al.*, 2004; Wang *et al.*, 2011; Thapar,  
341 Denmon and Nikonowicz, 2013).

342         The sequence to structure degeneracy (different RNA sequences are able to form similar  
343 spatial structure) is the known phenomenon (Petrov, Zirbel and Leontis, 2013; Bottaro and  
344 Lindorff-Larsen, 2017). Moreover, it is suggested to refrain from associating sequences with a  
345 particular fold (D'Ascenzo *et al.*, 2016, 2017). Together with the literature data our result let us  
346 assume that sequence-structure degeneracy is an universal way in which RNA tetraloops are  
347 used in nature (Lebars *et al.*, 2001; Wu *et al.*, 2004; Ihle *et al.*, 2005; Petrov, Zirbel and Leontis,  
348 2013; D'Ascenzo *et al.*, 2016, 2017; Bottaro and Lindorff-Larsen, 2017).

349         It can be speculated that pentaloops in domain d of the *Enterovirus E* genome and  
350 triloops of domain d of rhinoviruses have the potential to comprise the same UNCG fold as some  
351 YNHG and GNYA tetraloops. For HRV14 domain d, it was shown that its triloop resembles the

352 structure of the first and last two nucleotides of UNCG structural class tetraloops (Headey *et al.*,  
353 2007). There are pentaloops with four nucleotides that belong to consensus UNCG, GNRA or  
354 gCUUGc, which are able to form spatial structures of corresponding structural classes with the  
355 fifth bulged nucleotide (Cai *et al.*, 1998; Schärpf *et al.*, 2000; Theimer, Finger and Feigon, 2003;  
356 Oberstrass *et al.*, 2006; Liu *et al.*, 2009). It is possible that four nucleotides of the pentaloops in  
357 domain d of Enterovirus E species have a UNCG fold with one bulged nucleotide.

358 Tetraloops that did not belong to the YNHG or GNYA consensus were found in both sets  
359 of natural and *in vitro* selected genomes. However, in an experiment such variants were found to  
360 evolve towards the YNHG or GNYA consensus (Prostova *et al.*, 2015). Apparently, tetraloops  
361 that do not belong to the YNHG or GNYA consensus are less fit in most settings and under  
362 experimental conditions. However, as these variants may still be found in a few naturally  
363 circulating viruses (consequently, they have emerged and been fixed), we speculate that they  
364 may be beneficial under specific replication conditions.

365 A similar structure of domain d and its apical region suggests the free exchange of this  
366 region between genomes of the same and different species of Enterovirus genera. Indeed, viable  
367 intra and inter species recombinants for this region could be obtained *in vitro* (Muslin *et al.*,  
368 2015; Bessaud *et al.*, 2016). To evaluate the relative impact of the high mutation rate and  
369 recombination on domain d apical loop variability, sequences of EV71 C4 genotype viruses were  
370 analysed. The natural recombination in EV71 genotype C4 is much less frequent than other

371 *Enterovirus A* types (Lukashev *et al.*, 2014); meanwhile only one recombinant genome  
372 (accession number HQ423143) was detected in our data set. Therefore, the variability of its  
373 domain d loop sequence reflects changes that were only accumulated via mutations. The  
374 diversity of the domain d loop sequence of EV-71 C4 viruses was far less prominent than among  
375 *Enterovirus A* genomes and represented only by five tetraloop sequence variants (Table 2). As  
376 the most recent common ancestor of EV71 genotype C4 dates back about 20 years (McWilliam  
377 Leitch *et al.*, 2012), this diversity, although limited, has only emerged very recently. On the other  
378 hand, the high sequence variability of the domain d apical region in all enterovirus genomes was  
379 possibly assisted by inter- and intra-species recombination events.

380 Interestingly, in contrast to the similar structure of domain d and the very similar  
381 distribution of its apical sequences in genomes of different enterovirus species, its putative RNA-  
382 recognition tripeptide of 3C is diverse (Figure 3). Most *Enterovirus A* genomes contain tripeptide  
383 VGK in 3C, while there is a prevalence of the TGK tripeptide among genomes of *Enterovirus B*,  
384 *C* and *D* species (Figure 3). Genomes of *Rhinovirus A* and *C* also contain common enterovirus  
385 tetraloops in the domain d apical region, but, in 3C, unlike other species, they contain tripeptides  
386 without positively charged amino acids (Figure 3, Table S11, Table S12). Positively charged  
387 amino acids are often involved in the interaction with RNA, in particular, with phosphates of the  
388 RNA backbone. As such, they are of importance to RNA-protein recognition (Jones *et al.*, 2001;  
389 Bahadur, Zacharias and Janin, 2008). In *Rhinovirus A* genomes, positively charged amino acid

390 “jumped” from the last position of the tripeptide (position 156) to the position that precedes the  
391 tripeptide (position 153) (Figure 3, shown by an arrow). The residue at position 153 starts and  
392 the residue at position 156 ends the reverse turn between beta strands dII and eII of protein 3C  
393 (Mosimann *et al.*, 1997; Matthews *et al.*, 1999; Cui *et al.*, 2011). In a crystal structure of the  
394 Rhinovirus A2 protein 3C, the side chain of Lys153 (preceding the tripeptide) is positioned in a  
395 region similar to that of the side chain of Lys156 (in last position of the tripeptide) in the crystal  
396 structure of Enterovirus 71 and Poliovirus 1 proteins 3C (Mosimann *et al.*, 1997; Matthews *et*  
397 *al.*, 1999; Cui *et al.*, 2011). Thus, Lys at position 153 of 3C has almost the same potential to  
398 interact with the RNA-ligand as Lys at position 156 (Mosimann *et al.*, 1997; Matthews *et al.*,  
399 1999; Cui *et al.*, 2011). Genomes of *Rhinovirus C* species do not contain a positively charged  
400 amino acid, either inside the tripeptide of the 3C protein, or in the neighbouring positions,  
401 possibly indicating that tripeptide 154-156 in the protein 3C of *Rhinovirus C* genome does not  
402 interact directly with RNA. Thus, 3C is able to recognize domain d of the oriL with tripeptides of  
403 a different sequence. In contrast to the domain d structure and its apical sequence, the tripeptide  
404 is species-specific. The diversity of the tripeptide, which is expected to recognize domain d, has  
405 several compatible explanations. Residue 154 of the tripeptide possibly does not interact with  
406 domain d directly. The tripeptide may be involved into a species-specific cooperative amino acid  
407 network (amino acid “epistasis”). Moreover, different tripeptides could reflect slightly different  
408 molecular mechanisms for domain d recognition.



409           The complexity of the tripeptide's role in domain d recognition can be shown in several  
410 examples. The 3C protein of different species with the same RNA-binding tripeptide is not  
411 guaranteed to bind the same structured domain d. Genomes of the *Rhinovirus B* contain triloops  
412 in the apical region of domain d, which are paired with tripeptide TGK in 3C, common for  
413 genomes with tetraloops. In contrast, protein 3C of the Coxsackie virus B3 (*Enterovirus B*  
414 species, containing tripeptide TGK) cannot recognize oriL sufficiently well when domain d is  
415 capped with a triloop (Zell *et al.*, 2002). This indicates that the sequence of the RNA-binding  
416 tripeptide is probably not the exclusive participant in oriL-3C recognition. In other words,  
417 different molecular mechanisms of oriL-3C recognition have evolved in every enterovirus  
418 species independently. For example, it was shown for Rhinovirus 14 (*Rhinovirus B* species) that  
419 protein 3C recognizes the stem region of domain d, rather than its apical loop (Leong *et al.*,  
420 1993). Another oriL-3C recognition mechanism is seemingly employed by *Enterovirus E* and *F*  
421 species, two oriLs of which play the same role as the single oriL in genomes of other  
422 enteroviruses (Pilipenko, Blinov and Agol, 1990; Zell *et al.*, 1999). The apical loop of their  
423 domain d is a tetra- or pentaloop with a sequence that differs from the loop consensus of other  
424 enteroviruses. The RNA-binding tripeptide in 3C is species-specific as well, and is always MGK  
425 (Table S6). Interestingly, one genome of *Enterovirus G* species had the same pair domain d loop:  
426 tripeptide of 3C, i.e., GUUA MGK. Domain d of *Enterovirus G* species is prolonged in  
427 comparison to the length of domain d in genomes of other species (Krumbholz *et al.*, 2002)

428 (Figure 2). Tripeptide MGK in the 3C of *Enterovirus E, F* and *G* possibly indicates another  
429 molecular mechanism of oriL-3C recognition (Krumbholz *et al.*, 2002). Therefore, we assume  
430 that, though putative RNA-binding, the tripeptide, in most cases, possibly interacts with the  
431 domain d apical region (since amino acid substitutions in it are known to compensate for  
432 structural disturbance in domain d); however, this interaction is not the only one that determines  
433 the evolution of oriL-3C interaction. Altogether, the data suggest that the independent evolution  
434 of the putative RNA-binding tripeptide of 3C and domain d of oriL occurs.

435

436

## Conclusions

437 We analysed the variety and occurrence of the replication element oriL's functional loop  
438 and its protein ligand virus protease 3C. RNA-binding motifs of 3C are species-specific, in  
439 contrast to domain d loop sequences: the sequence variety of domain d loop is almost the same  
440 for *Enterovirus A, B, C* and *D* and *Rhinovirus A* and *C* species, whereas tripeptide sequence  
441 variety differs. The conservation of the tripeptide sequence within species, together with the  
442 almost universal diversity of tetraloop sequences among species, indicates the occurrence of the  
443 independent evolution of these two elements. Our result suggest the structure-based, rather than  
444 sequence-based, recognition of domain d by virus protein 3CD. These, together with the data  
445 reported in the literature, let us assume that the sequence-structure degeneracy is a universal way

446 RNA in which tetraloops are used in nature.

447

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665

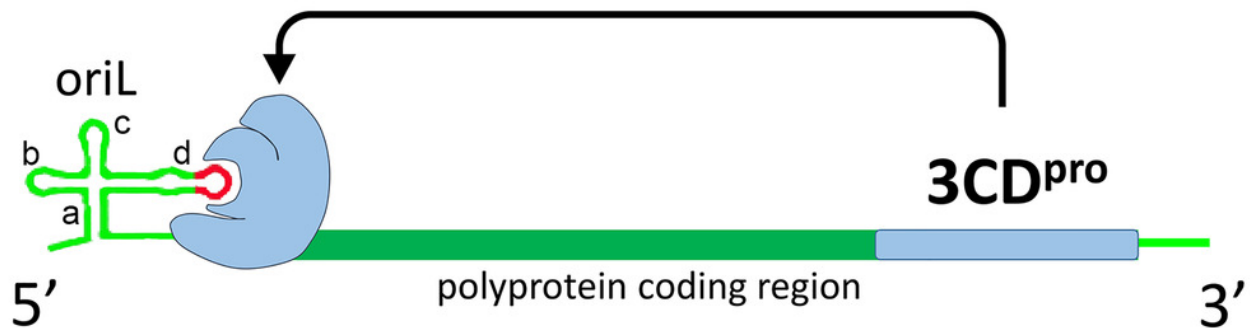
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# Figure 1

Schematic representation of poliovirus genome and detailed representation of secondary structure of poliovirus replicative element oriL.

Protein 3C and its RNA-binding motifs  $_{82}\text{KFRDI}_{86}$  and  $_{154}\text{TGK}_{156}$  are shown (here and below amino acid numeration corresponds to poliovirus protein 3C) (Prostova et al., 2015 with modifications).

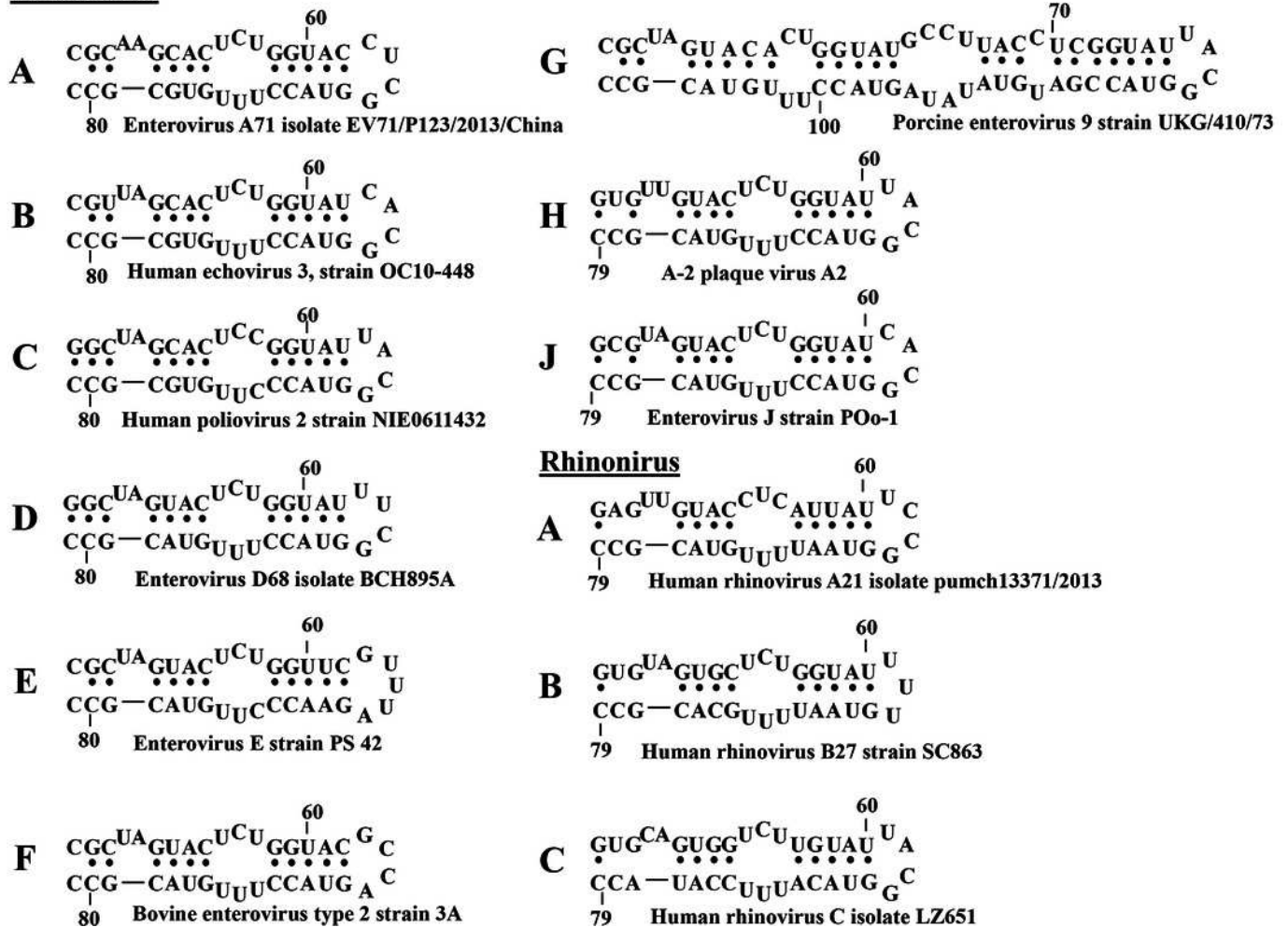


## Figure 2

Secondary structure of oriL domain d of distinct enterovirus species.

For *Enterovirus E* and *F* domain d of the first oriL is shown. Secondary structure of domain d of Porcine enterovirus 9 strain UKG/410/73 was folded with use as reference Krumbholtz et al., 2002.

### Enterovirus



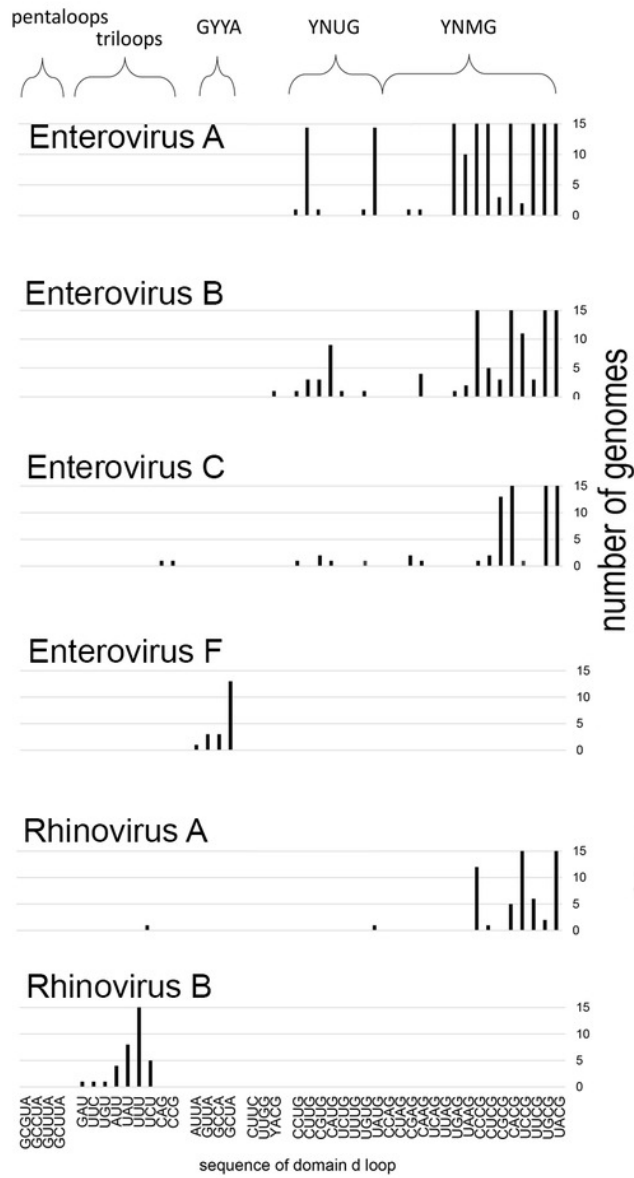


## Figure 3

Distribution of domain d loop sequence and amino acid motifs in the 3C protein.

**A** - Distribution of domain d loop sequences. The regions corresponding to tetraloop consensus, triloops and pentaloops are shown. Number of genomes cut off at 15 for clear view of sequence distribution. **B** - The frequency plot of amino acid sequence of 3C in species of genus *Enterovirus*. The amino acid sequence logo was done with WebLogo server (Crooks *et al.*, 2004). Arrows indicate amino acids of the proteolytic triade (Glu71 and Cys 147), the first and the last amino acids of motif  $_{82}\text{KFRDI}_{86}$ , the putative RNA-binding tripeptide 154-156 of 3C and Lys153 in the protein 3C of *Rhinovirus A*.

A



B



**Table 1** (on next page)

Number of full genome sequences that contained oriL region and number of unique domain d sequences before and after filtration.

For *Enterovirus E* and *F* number of unique tetraloops is shown separately for first and the second oriL.

Species	Number of full genome sequences	Number of full genome sequences after 1% nucleic identity filtration	Number of unique tetraloops before filtration		Number of unique tetraloops after filtration	
Enterovirus A	1052	564	17		16	
Enterovirus B	339	244	18		18	
Enterovirus C	747	274	15		12	
Enterovirus D	419	57	7		6	
Enterovirus E	12	10	6	5	6	5
Enterovirus F	13	10	4	3	4	3
Enterovirus G	10	8	6		6	
Enterovirus H	3	2	2		2	
Enterovirus J	8	5	3		3	
Rhinovirus A	151	118	8		8	
Rhinovirus B	50	37	7		7	
Rhinovirus C	38	37	6		6	

1

**Table 2** (on next page)

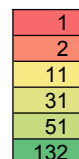
Occurrence of domain d apical sequences in filtered sets of full genomes of different enterovirus species.

Tetraloops CCCG, UGUG, CAUG and UUGG that were unique for species *Enterovirus A, C* and *D* and were lost upon filtration, were added and are shown in blue. The gradient coloring from red to green represents abundance heat map for the genomes with different domain d sequence.

1 Table 2 Occurrence of domain d apical sequences in filtered sets of full genomes of different  
 2 enterovirus species and serotypes. Tetraloops CCCG, UGUG, CAUG and UUGG that were unique  
 3 for species *Enterovirus A, B, C* and *D* and were lost upon filtration, were added to maintain the  
 4 diversity of loop sequence and are shown in blue. The gradient coloring from red to green  
 5 represents abundance heat map for the genomes with different domain d sequence.

6

Loop sequence	Enterovirus													Rhinovirus			
	A				B	C			D	E	F	G	H	J	A	B	C
	all	EV71	EV71 C4 genotype	non EV71		all	PV	non PV									
Triloops																	
CCG						1		1									
CAG						1		1									
UCU														1	5		
UUU															17		
UAU															8		
AUU															4		
UGU															1		
UUC															1		
GAU															1		
YNMG Tetraloops																	
UACG	85	28		57	51	106	64	42				3	1	2	38	15	
UGCG	114	2		112	31	43	31	12				1	1		2		
UUCG	16	16	14		3				50						6	6	
UCCG	2			2	11	1	1								53	10	
CACG	48	28		20	98	101	54	47	1			1		2	5		
CGCG	3	2		1	3	13	6	7				1					
CUCG	132	127	126	5	5	2		2	2						1	3	
CCCG	40	39	28	1	16	1		1	1						12	2	
UAAG	10	10			2												
UGAG	22	22			1												
UUAG																	
UCAG																	
CAAG	1	1			4	1		1						1			
CGAG	1			1		2		2									
CUAG																	
CCAG																	
YACG					1												
YNUG Tetraloops																	
UAUG	54	1		53											1		
UGUG	1			1	1	1		1									
UUUG									1								
UCUG					1												
CAUG					9	1		1									
CGUG	1			1	3	2		2									
CUUG	34	34	35		3				2								
CCUG	1	1	1		1	1		1									
GYYA Tetraloops																	
GCUA									2	13							
GCCA										3							
GUUA									2	3	1						
Other tetraloops																	



UUGG									1							
CUUC																1
AUUA											1					
Pentaloops																
GCUUA											7					
GUUUA											2					
GCCUA											4					
GCGUA											1					
GCGUA											1					
GAUUA											1					
GUCUA											1					

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