Polyadenylation of genomic RNA and initiation of antigenomic RNA in a positive-strand RNA virus are controlled by the same cis-element

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ABSTRACT

Genomes and antigenomes of many positive-strand RNA viruses contain 3'-poly(A) and 5'-poly(U) tracts, respectively, serving as mutual templates. Mechanism(s) controlling the length of these homopolymeric stretches are not well understood. Here, we show that in coxsackievirus B3 (CVB3) and three other entroviruses the poly(A) tract is ~80–90 nt and the poly(U) tract is ~20 nt-long. Mutagenesis analysis indicate that the length of the CVB3 3'-poly(A) is determined by the oriR, a cis-element in the 3'-noncoding region of viral RNA. In contrast, while mutations of the oriR inhibit initiation of (-) RNA synthesis, they do not affect the 5'-poly(U) length. Poly(A)-lacking genomes are able to acquire genetically unstable AU-rich poly(A)-terminated 3'-tails, which may be generated by a mechanism distinct from the cognate viral RNA polyadenylation. The aberrant tails ensure only inefficient replication. The possibility of RNA replication independent of oriR and poly(A) demonstrate that highly debilitated viruses are able to survive by utilizing ‘emergence’, perhaps atavistic, mechanisms.

INTRODUCTION

The genomic RNA of many positive-strand RNA viruses is tailed with a 3'-poly(A). In contrast to cellular mRNAs, in which poly(A) is non-template and is added post-transcriptionally, the viral poly(A) is believed to be template-coded (1). Accordingly, the negative strand contains a 5'-poly(U) tract, which is synthesized using the poly(A) of the positive viral RNA as a template. In turn, this poly(U) subsequently serves as a template for the synthesis of the poly(A) of progeny RNA. This alternate usage of the homopolymeric parts of each strand synthesis has not been studied in detail but it was observed that poly(U) in the negative RNA of poliovirus (2,3) and coronavirus (4) may be shorter than poly(A) in the viral RNAs. These observations pose intriguing questions about how a shorter poly(U) is synthesized on a longer poly(A) template and vice versa.

We addressed this conundrum using coxsackievirus B3 (CVB3), an entervoirus belonging to the picornavirus family as a model. Picornaviruses are small icosahedral viruses with a single-stranded 7–8 kb RNA genome of positive (mRNA-like) polarity. This family includes important human and animal pathogens such as poliovirus, hepatitis A virus, foot-and-mouth disease virus and others.

Three cis-acting elements are known to be involved in the replication of entervoirus RNA (5). The 5'-terminal cloverleaf structure (or oriL) is a multifunctional element that interacts, among other ligands, with viral 3CDpro protease and cellular poly(rC) binding proteins PCBP1 and PCBP2 (6–8) to form a ribonucleoprotein (RNP) complex. This complex is thought to be involved in the switch that represses translation and allows replication to commence (9), and is also required for the initiation of both (+)-strand (6) and (−)-strand RNA synthesis. The latter requires genome circularization mediated by an interaction between this RNP and the host poly(A)-binding protein (PABP) bound to 3'-poly(A) (10,11).

Another cis-acting replicative element (cre or ori) is represented by an imperfect stem–loop located within the coding region of the entervoirus RNA (12–14). It ensures the covalent linkage of uridylates to the viral protein VPg via a two-step slide-back mechanism, thereby yielding the
protein primer VPg-pU(pU) (15–18). Both (−)- and (+)-strand RNA 5′-termini have VPg covalently bound (5), but there is no consensus on whether the cre-mediated VPg-uridylylation is required for the initiation of only (+) RNA (19–21) or for the initiation of both strands (14). The former view implies that VPg uridylylation may occur directly on the 3′-poly(A) of (+) RNA, thus providing a primer for elongation into (−)-strands, in line with the finding that polyadenylates indeed support VPg-pU(pU) synthesis in vitro (22).

A third replicative cis-element, the oriR, is located in the 3′-noncoding region (3′NCR) of the viral RNA immediately upstream of the 3′-poly(A). It contains a multi-domain structure maintained by an intramolecular kissing interaction between helical elements X and Y (23–28). Mutations destroying this interaction or changing the mutual orientation of oriR helical elements resulted in a significant loss of fitness, as judged by the acquisition of temperature sensitive and quasi-infectious or lethal phenotypes. It was proposed that oriR is required for an efficient initiation of (−) RNA synthesis (24,26). However, partial and complete deletions of the 3′NCR still generated viable viruses challenging the essential role of 3′NCR and oriR in RNA replication (29–31).

The synthesis of a (−) RNA strand on the viral RNA template is believed to result in the formation of a genome-length double-stranded ‘replicative form’ (RF) RNA. Positive-strand RNA synthesis is initiated at the 3′-end of the (−) strand but the detailed mechanism is unknown. It has been proposed that destabilization and partial unwinding of this end of the duplex is required for efficient replication (32,33). Transcription of the (−)-strands from the (−)-RNA template results in partially double-stranded ‘replicative intermediate’ RNA molecules consisting of full-length (−) RNA and various numbers of incomplete (+) strands differing in length (5). Newly synthesized virion RNAs are thought to be displaced from the (−) strands by upcoming progeny strands through the duplex-unwinding activity of the viral RNA polymerase 3Dpol (34). However, a number of important issues pertaining to picornavirus genome replication remain contradictory or unsolved (35).

Here, we examined the mechanism for poly(A) and poly(U) synthesis and how their lengths are controlled. We found that the 3′-poly(A) of (+) RNA strands in CVB3 and some other enteroviruses is about 4-fold longer than the 5′-poly(U) of (−) strands and that the poly(A) length is controlled by the oriR. While ensuring efficient negative strand initiation, the oriR does not control the poly(U) length. Removal of the poly(A) (alone or together with the oriR) from transfected viral RNA resulted in regeneration of poly(A)-containing 3′-tails, which, however, ensured only inefficient viral reproduction. These results provide new insights into mechanisms and evolution of genome replication of positive-strand RNA viruses.

**MATERIALS AND METHODS**

**Cells and media**

Buffalo green monkey (BGM) cells were grown in minimal essential medium, MEM (Gibco), supplemented with 10% fetal bovine serum (FBS), 2 mM l-glutamine (Sigma), 100 U penicillin per ml and 100 mg streptomycin per ml (Sigma). HeLa cells were grown in Dulbecco’s MEM (Gibco) with the same supplements (but with streptomycin at 25 mg/ml). HeLa S3 cells (ATCC CCL 2.2) were grown either (i) in tissue culture flasks in Dulbecco’s modified Eagle medium-nutrient mixture F-12 (Ham) (1:1) (Sigma) with the above supplements or (ii) in suspension using suspension MEM (Joklik modified) (Sigma) supplemented with l-glutamine, antibiotics and 10% newborn calf serum (Gibco).
Blocking the 3'-termini of transcript RNA with cordycepin-5'-triphosphate

Cordycepin (3'-deoxyadenosine, Ambion) was added to the 3' end of in vitro transcribed RNA using yeast poly(A) polymerase (USB) according to the manufacturer's recommendations.

RNA transcription and transfection

Sall or MluI linearized cDNA served as templates for in vitro transcription by T7 RNA polymerase (Promega) (26). BGM cells were transfected in duplicate by a DEAE-dextran method (26) and incubated at 36°C and 5% CO2 until complete CPE. Then, the cultures were subjected to three cycles of freezing/thawing prior to storage at –80°C. If 6 days post-transfection no CPE was evident, the cells were lysed by three freeze/thaw cycles and 200 μl of cell suspension was transferred to a fresh culture (post-passage). The 50% tissue culture infective dose (TCID50) was determined by endpoint titration in BGM-cells (36).

Sequencing of recovered virus RNA

Viral RNA extraction was performed using the GenElute™ Mammalian Total RNA Miniprep kit (Sigma). RNA was reverse transcribed using Super RT (HT Biotechnology) and RT1 (Supplementary Data) and PCR-amplified using FPseq and RPseq. RT–PCR products were subcloned into pGEM-T easy vector (Promega). Two independent sequence reactions of selected clones were performed by capillary electrophoresis (3730 DNA Analyzer; Applied Biosystems) and Big Dye Terminator version 3 protocols (Applied Biosystems).

Virus purification

Two T150 flasks of either HeLa cells (for CVA21 strain Coe) or BGM cells (for CVB3 strain Nancy, EV71 strain BrCr and ECHO9 strain Hill) were infected and incubated until complete CPE. After three freeze/thaw cycles, the cell debris was removed by centrifugation at 2500 r.p.m. Supernatants were collected in Beckmann centrifuge tubes containing 5 ml of 30% sucrose. Virions were pelleted at 140,000 × g for 6 h (SW28 rotor, Beckman L8-70M centrifuge). The pellet was resuspended in 200 μl phosphate-buffered saline (PBS) and viral RNA was extracted as described above.

Tailing of the 3'-end of virion RNA

T4 RNA ligase (Promega) was used to attach a RNA oligonucleotide to virion RNA extracted from purified virus (Figure 1A). A complementary cDNA reverse primer (1) primed the RT reaction. The generated cDNA was amplified using primer 1 and virus-specific forward primers (Supplementary Data). In addition, a semi-nested-PCR was performed using primer 1 and semi-nested (SN) virus specific primers (Supplementary Data) to enhance specificity of the PCR and to increase the quantity of amplified product. The accumulated product was gel-purified and sequenced using the semi-nested PCR forward primers.

5'-RACE-assay

T25 flasks containing, depending on the virus strain (see above), BGM or HeLa cells were infected with an MOI of 10 TCID50/cell. Cells were lysed 6 h post infection (p.i.) and total RNA was obtained as described above. A 5'-rapid amplification of cDNA ends (5'-RACE)-PCR was performed using the 5'-RACE system (Invitrogen) according to the manufacturer's recommendations (Figure 1B). In brief, the RT-reaction on (−) RNA strands was executed by SuperScript™ II (Invitrogen) with virus-specific RT primers (Supplementary Data). After first strand cDNA synthesis, an RNase removed the original RNA template. Single-stranded cDNA was purified and terminal deoxynucleotidyl transferase (Invitrogen) added a 3'-poly(C) stretch. This cDNA was PCR-amplified using an AAP primer (Abridged Anchor Primer; 5'-RACE system, Invitrogen) and virus-specific forward primers (Supplementary Data), followed by a nested-PCR using an AUAP primer (Abridged Universal Amplification Primer) and virus-specific nested primers (Supplementary Data) to enhance specificity of the PCR and to increase the quantity of amplified product. The PCR background did not allow direct sequencing of the PCR-product, although the

![Figure 1](http://nar.oxfordjournals.org/)
band of interest was clearly distinguishable. Therefore, PCR product was excised from the gel and ligated into a pGEM-T easy vector and the 5'-poly(U) size of 10 colonies was determined by sequencing using the nested forward primer (Supplementary Data).

Single-cycle growth analysis
Confluent BGM cell monolayers were infected with an MOI of five TCID50/cell and incubated at 36°C for 2, 4, 6 and 8 h (26). Each growth curve was performed in triplicate and viruses were released by three cycles of freezing/thawing. Virus titers were determined by endpoint titration (36).

Plaque-assay
Six-well dishes with >90% confluent monolayer of BGM cells were incubated with 10-fold virus dilutions in MEM for 1 h and rocked every 15 min. After absorption, the cells were overlaid with 2.5 ml of M199 (Sigma) containing 3% FBS, 1% glutamine, 0.5% gentamycin, 0.5% penicillin, 0.5% Funguzone, 1% MgCl2 and 1.3% Na-bicarbonate in 0.6% agar (1:1 mixture with 2xM199). At 48 or 96 h p.i., cells were fixed by methanol-acetic acid (3:1, vol/vol) for 10 min and stained with a 0.5% crystal violet in 20% ethanol.

Luciferase assay
BGM cells seeded in 6-well plates were transfected using the DEAE-dextran method (26) with 4 µg of replicon RNA. Cells were washed twice with PBS and lysed using 200 µl of CCLR lysis buffer (Promega). The luciferase assay system (Promega) was used to measure the luciferase activity on a BioOrbit 1251 Luminometer.

In vitro translation and replication assay
HeLa cell S10 extract and initiation factors were prepared as described (37). Negative strand RNA synthesis was analyzed as described (38) with minor modifications: 2 µg of CVB3 RNA transcript were mixed with 30 µl of HeLa S10 extract, 2 µl of initiation factors, 5 µl of 10× NTP/energy mix (37) and 1 µl of 100 mM guanidine-HCl in a volume of 50 µl. All other steps were executed as described (38).

RESULTS
The 5'-poly(U) of enterovirus negative strands is significantly shorter than the 3'-poly(A) of the positive RNA strands
Using 3'-tailing and 5'-RACE reactions, we determined the lengths of the 3'-poly(A) of virion RNA and the 5'-poly(U) in (--) RNA for four different enteroviruses; CVB3, ECHO9, enterovirus 71 (EV71) and coxsackievirus A21 (CVA21) (Figure 1A and B). In all cases, the average size of the poly(A) was 80-90 nt, whereas the length of the poly(U) was only ~20 nt (Figure 1C). The poly(A) of CVB3 RNA from purified virions recovered from different cell cultures (BGM, HeLa and COS) was of a similar average size (data not shown).

**OriR** controls the initiation efficiency of negative RNA synthesis and the poly(A) length of virion RNA but not the size of the poly(U) of negative strands
A simple explanation for the smaller size of the 5'-poly(U) tract compared with its 3'-poly(A) template is that (--) strand RNA synthesis is initiated on an internal position of the poly(A) template and there exists a spatial mechanism controlling the choice of this position. One can hypothesize that this mechanism involves the oriR. To examine this, two mutant RNAs, N70/A and N70/B, were generated, in which oriR was replaced by random sequences of 70 nt (Table 1). These RNAs proved to be viable but exhibited a reduced rate of replication [cytopathic effect (CPE) developed 4-6 days after transfection versus 2 days in the case of wild-type (wt) transcripts]. Sequencing the virus pools showed that N70/A had retained the engineered mutations while an adenosine insertion was found in N70/B within an
Figure 2. Characterization of CVB3 3’NCR randomization mutants using single-cycle growth curves (A), in which the 50% tissue culture infective dose (TCID<sub>50</sub>) is set against the time of incubation (time), plaque assay, in which the dilution factor is indicated underneath the depicted well (B), and in vitro replication assay (C). GuHCl, guanidinium hydrochloride. (D) Luciferase activity of wt or mutant RNA determined at 8 and 24 h post-transfection in BGM cells expressed as the fold difference between cells incubated in the presence and absence of GuHCl. (E) 3'-Poly(A) and 5'-poly(U) sizes of the Δ3’NCR-N70/A (N<sub>70A</sub>) and Δ3’NCR-N70/B (N<sub>70B</sub>) mutants. The 3’-poly(A) columns represent the means ±SD of three independent experiments; the 5’-poly(U) columns represent the means ±SD of 10 values. ** corresponds to P < 0.01.

These data confirmed the oriR involvement in the replication of (−) strands and uncovered its role in controlling the poly(A) length of the viral RNA. On the other hand, oriR has little, if any, effect on the poly(U) length in (−) strands.

Structural requirements of oriR that control the poly(A) size

Previous studies indicated that the enterovirus oriR has a conserved structure composed of helical domains joined by a kissing interaction (K) between the loops of the X- and Y-domains (Figure 3A and 3B). Disruption of this interaction by alteration of 6 nt (mutant KD1) or 4 nt (KD2) in the 3’-terminal component of the kissing interaction (Figure 3C) resulted in quasi-infectious transcripts: after a delayed CPE, viable viruses were recovered from the transfected cultures but none of them faithfully retained the engineered structure. The pseudoreversions exhibited by the viruses could be grouped into two types. The representative of one class acquired a mutation restoring potential for a stable, though altered, kissing interaction. Thus, a C insertion at position 7350 could potentially support an alternative 6 nt kissing interaction (Figure 3D). When this mutation was engineered...
Figure 3. Secondary and tertiary structures of wild type and mutant CVB3 3′NCRs. Secondary (A) and tertiary (B) structure of the wt CVB3 3′NCR. The 3′NCR consists of three hairpin domains, X, Y and Z. The structure can be closed by an interaction between the poly(A) with a 4 nt U-stretch overlapping the oriR and the 3D-coding region. The X domain can be stacked to the tertiary ‘kissing’ (K) interaction to form a coaxial helical element, which is connected by single-stranded nucleotide stretches to a second coaxial helical domain Z-Y. (C) Schematic representation of the kissing distortion mutants, KD1 and KD2, in which 6 and 4 bp, respectively, were mutated (underlined). (D) Partial 3′NCR sequence of a virus recovered after transfection with the KD1 mutant. The inserted cytosine residue in the recovered mutant is underlined. (E) The entire 3′NCR sequence of a virus recovered after transfection with the KD2 mutant.

Table 2. Time to CPE after transfection or infection with CVB3 oriR mutants

<table>
<thead>
<tr>
<th>Construct/mutation</th>
<th>Days before CPE Post transfection</th>
<th>Post passage</th>
</tr>
</thead>
<tbody>
<tr>
<td>p53CB3/T7 (wildtype)</td>
<td>1</td>
<td>nd</td>
</tr>
<tr>
<td>p53CB3/T7-ΔX</td>
<td>—</td>
<td>4</td>
</tr>
<tr>
<td>p53CB3/T7-ΔY</td>
<td>—</td>
<td>4</td>
</tr>
<tr>
<td>pRibCB3/T7 (wildtype)</td>
<td>&lt;1</td>
<td>nd</td>
</tr>
<tr>
<td>pRibCB3/T7-ΔXΔY</td>
<td>4/5</td>
<td>nd</td>
</tr>
<tr>
<td>pRibCB3/T7-Δ3′NCR</td>
<td>5</td>
<td>nd</td>
</tr>
</tbody>
</table>

A total of $5 \times 10^6$ cells were transfected with ~4 µg of RNA. ‘—’ means not detectable CPE; nd means not done.
additional mutations were found. All these mutants exhibited a marked deficiency in the virus reproduction (Figure 4A) and a small plaque-phenotype (Figure 4B) as well as inefficient cell-free RF accumulation (Figure 4C) and RNA replication in BGM cells (Figure 4D). The mutations were also accompanied with a severe reduction in poly(A) length, to \(~45\) nt (ΔX), \(~30\) nt (ΔY) and \(~20\) nt (Δ3'NCR) (Figure 4E). The deletion of the Z-domain alone had no significant impact on poly(A) length.

These results confirmed that efficient viral RNA replication requires an oriR with a sufficiently stable kissing interaction, although the sequence of the K-domain could vary. Also, the efficient initiation of the (−) strand and generation of long poly(A) tails in the progeny (+)RNA could not be supported by its individual domains.

**Effect of poly(A) length on the initiation efficiency of negative-strand RNA synthesis**

As assayed by cordycepin-treated transcript RNA (see Materials and Methods), the minimal poly(A) size ensuring efficient cell-free RF RNA synthesis was \(~20–30\) nt (Figure 5). These results are in line with previous reports showing that a poly(A) of \(~20\) nt is sufficient for the efficient replication of poliovirus (also an enterovirus) (39), whereas
shorter poly(A) resulted in a dead or highly debilitated virus (2,10,31,40).

Thus, the minimal length of the poly(A) in the (+) RNA ensuring efficient replication is ~20 nt, coinciding with the poly(U) length in the (−) RNA.

**Regeneration of RNA 3′-tails in viruses recovered from poly(A)-lacking genomes**

To better understand the role of poly(A) in RNA replication, mutant viral cDNAs were generated, in which poly(A) was either missing [Δpoly(A)] or replaced by other sequences, such as G23 or C9G [Δpoly(A)-G23 and Δpoly(A)-C9G]. The Δpoly(A) construct contained 3′-terminal TCGA derived from a Sall site at the linearization locus. Transcripts of all these constructs exhibited a CPE delayed for 2–3 days and generated viruses with newly acquired 3′-poly(A), as evidenced by the ability of oligo(dT)18 to serve as a reverse primer in PCR-based sequencing. The RNA of the recovered Δpoly(A) mutant corresponded to its DNA construct, whereas the poly(G) moiety of the G23 mutant lost 21–22 G-residues and the C9G moiety of the respective mutant was replaced by a CCUU sequence (Table 1). No alterations within the heteropolymeric part of 3′NCR was detected. The recovered viruses exhibited a nearly wt plaque phenotype at 36°C but formed somewhat smaller plaques at 39°C (Table 3). In BGM-cells, the poly(A)-lacking RNA of Δpoly(A) construct generated, at 10 h post-transfection, luciferase levels equal to ~10% of that of wt RNA, whereas both G23 or C9G constructs produced luciferase levels similar to that found in the cells transfected with wt RNA in the presence of a potent inhibitor of viral reproduction guanidine-HCl (data not shown).

In view of oriR involvement in polyadenylation of viral RNA, the effect of simultaneous deletion of heteropolymeric 3′NCR and poly(A) was investigated. When applied to the 53CB3/17-based RNA, such a damage resulted in a nonviable genome. However, the equivalent ribozyme-containing transcript with a genuine 5′-end, Δ3′NCR-Δpoly(A), was viable, though exhibited a delayed CPE of 5 days after transfection. The viral genome acquired a 3′-poly(A) but no other modifications in the 3′-terminal portion of the molecule (Table 1).

To better understand the requirements for polyadenylation of poly(A)-lacking RNA, viral genomes were generated in which the 3′NCR was replaced by various structures. The poly(A)-lacking RNA with the oriR replaced by 111 random nucleotides, Δ3′NCR-N111-Δpoly(A), acquired a 3′-poly(A) preceded by an AU-rich stretch containing a U13 element and an AAUAAA hexamer corresponding to the cellular polyadenylation signal (Table 1). A ribozyme-lacking N70-Δpoly(A) mutant was dead. However, insertion of an AAUAAA-containing oligonucleotide into the randomized N70 sequence produced a viable virus with 3′-poly(A) preceded by a long AU-rich region containing multiple AAUAAA hexamers (Table 1). The oriR-lacking RNAs terminated with G15 or C9G (together with a Sall site-derived UCGA) also generated viable genomes in which 3′-poly(A) preceded by relatively long poly(U) tracts and an oligo(A) stretch interspersed with several single U residues, again generating AAUAAA polyadenylation-like signals (Table 1). All the recovered poly(A)-lacking viruses with 3′NCR deletion/substitution generate minute plaques (Table 3), suggesting that they had a severe defect in viral replication.

The results indicated that poly(A)-lacking viral RNA can be viable but that 3′-poly(A) is readily regenerated. This poly(A) is usually, but not always, preceded by newly emerged AU-rich sequences.

**Genetic instability of aberrant 3′NCR in viral RNA**

To understand whether the aberrant 3′NCRs could be functionally ‘improved’ by natural evolution, viruses containing such 3′NCR were subjected to additional passages. The 3′NCR of Δ3′NCR-N70/A RNA remained unchanged after five passages, although the plaque size was somewhat increased (data not shown), possibly due to alterations elsewhere in the viral RNA. Four clonal derivatives of Δ3′NCR-N70/B RNA exhibited several differences compared with the original recovered viruses and underwent further changes upon passages (Table 4). Modifications included a marked increase in the length of internal oligo(A) blocks (e.g. from 3 to 23 nt in Δ3′NCR-N70/B4) and deletions of 13 or 29 nt-long heteropolymeric segments. The virus recovered after transfection with Δ3′NCR-N111-Δpoly(A) as well as its four clonal derivatives exhibited, upon passaging, in addition to lengthening of internal poly(A), also lengthening of poly(U) and acquisition of relatively extended AU-rich sequences containing the AAUAAA hexamers (Table 4). None of these changes were accompanied by a significant increase in the plaque size.

**DISCUSSION**

Evidence is presented here that the mutual templating of the 3′-poly(A) of (+) RNA strands and 5′-poly(U) of (−) strands in viruses with positive RNA genomes, exemplified by enteroviruses, is not straightforward. In CVB3 (and three other enteroviruses), poly(U) is ~20 nt-long, whereas poly(A) is ~4 times longer. Destruction of the oriR, a cis-element in the 3′NCR of genomic RNA, significantly reduces the poly(A) size and decreases the efficiency of (−) RNA synthesis but does not affect the poly(U) length. Even though efficient genome replication requires the presence of a poly(A) of at least ~20 nt, its complete removal did not kill the virus because of the poly(A) capacity to regenerate.

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**Table 3. Plaque phenotypes of CVB3 oriR mutants**

<table>
<thead>
<tr>
<th>Virus</th>
<th>Plaque size (mm)</th>
<th>36°C</th>
<th>39°C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild-type CVB3</td>
<td>4.1 ± 0.9</td>
<td>4.3 ± 1.4</td>
<td></td>
</tr>
<tr>
<td>G23</td>
<td>3.6 ± 0.7</td>
<td>3.1 ± 0.9</td>
<td></td>
</tr>
<tr>
<td>C9G</td>
<td>4.0 ± 1.0</td>
<td>2.8 ± 0.4</td>
<td></td>
</tr>
<tr>
<td>Δ3′NCR+Δpoly(A) G15</td>
<td>&lt;0.5</td>
<td>&lt;0.5</td>
<td></td>
</tr>
<tr>
<td>Δ3′NCR+Δpoly(A)C14G</td>
<td>&lt;0.5</td>
<td>&lt;0.5</td>
<td></td>
</tr>
<tr>
<td>Δ3′NCR+Δpoly(A)</td>
<td>&lt;0.5</td>
<td>&lt;0.5</td>
<td></td>
</tr>
<tr>
<td>Δ3′NCR-N111+Δpoly(A) P1</td>
<td>&lt;0.5</td>
<td>&lt;0.5</td>
<td></td>
</tr>
<tr>
<td>Δ3′NCR-N111+Δpoly(A) P5</td>
<td>&lt;0.5</td>
<td>&lt;0.5</td>
<td></td>
</tr>
<tr>
<td>N70+Δpoly(A)+AAUAAA</td>
<td>&lt;0.5</td>
<td>&lt;0.5</td>
<td></td>
</tr>
</tbody>
</table>

*The size was measured on the second day p.i., except for plaques with a diameter <0.5 mm, which became discernible only on the third day p.i.*

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[Image of page with table and text]
The stop codon is shown in italics and given as subscripts in parenthesis, the sequences acquired after transfection are in bold, the cellular polyadenylation signal, AAUAAA, is underlined, Δ means deletion and the partial SalI site is double underlined. For the mutant N70/B cDNA and N111 sequences see Table 1.

*Recovered virus.

**Passage 10.

### Mechanism and control of viral RNA polyadenylation

The existence of a longer poly(A) compared with its poly(U) template indicates that polyadenylation of viral RNA is not limited to the faithful copying of the template. Theoretically, additional polyadenylation may be because of either a reiterative usage of the poly(U) template or a non-template reaction catalyzed by a terminal nucleotide transferase (TNT)-like enzyme or poly(A) polymerase. Since the poly(A) is markedly shortened in the absence of oriR, the mechanism should involve this cis-element.

The reiterative hypothesis (Figure 6A) assumes that the synthesis of a longer poly(A) is due to a kind of ‘stuttering’ of the viral polymerase, common to negative-strand RNA viruses (41,42). The oriR role may be envisioned as follows. The nascent oriR, just released from the RNA-dependent RNA polymerase (RdRP) molecule engaged in the 5′-poly(U) copying, interacts with this RdRP in cis (43) and ‘encourages’ it to repeatedly use the poly(U) template. The RdRP/oriR interaction may be aided by other cellular (44) or viral proteins (45). Cases of increase in the length of internal oligo(A) in picornovirus (23,46) and foot-and-mouth disease virus (47) RNAs support the possibility of reiterative copying during picornavirus RNA replication. In this model, additional polyadenylation of viral RNA is replication-coupled, which is compatible with relative inefficiency of the RdRP complementation in trans (48,49).

On the other hand, poly(A) elongation may result from 3′-terminal adenylate additions. In this case, the nascent viral RNAs has only ~20 nt-long poly(A). Further polyadenylation is accomplished by a TNT-like enzyme activated by binding to oriR. This job may be performed by a host poly(A) polymerase (50) or the terminal adenylate transferase activity may be an intrinsic property of the viral RdRP, as reported for the poliovirus enzyme by Neufeld et al. (51). A TNT activity has also been detected in RdRP of hepatitis C virus (52).

The oriR-dependent poly(A) additions to the ‘immature’ viral RNA may be post-replicative (i.e. occurring after completion of nascent positive strand transcription) (Figure 6B) or post-translational (i.e. accomplished by the nascent RdRP molecule or TNT-like enzyme) (Figure 6C). The post-translational hypothesis would explain why translation of viral RNA is a prerequisite for its replication (53). The ‘maturaion’ of viral RNA through its further polyadenylation is required for its usage as template for the next round of RNA replication and also contributes to its stabilization.

Also it cannot be excluded that the oriR itself controls the poly(A) length just by its stabilization.

Although the above hypotheses are speculative, each of them makes specific predictions amenable for experimental testing.

### Control of initiation of negative strands

Although oriR is not essential for the (−) strand synthesis (29,31; this study), it is important for its efficiency. Again, several mechanisms of such a control can be considered. One may suppose that binding of the viral RdRP (within an RNP complex or otherwise) to the oriR is a prerequisite for efficient initiation of (−) RNA synthesis or that the very presence of oriR makes the poly(A) tail more accessible for the initiation complex. Either of these notions can easily be combined with the models of poly(A) elongation considered above.

Remarkably, transcripts harboring certain oriR mutations produced progeny in which these mutations were not retained but either a wt-like oriR structure was regained or this structure was completely destroyed (see KD1 and KD2). Hence, it could be more harmful for the virus to possess certain oriR modifications than to lack this element altogether. This may indicate that some oriR modifications do not permit formation of a functional initiation complex, resulting in a dead genome.

Even though oriR controls the efficiency of (−) RNA synthesis, it does not appear to affect the site at which this synthesis is initiated. The fixed location of this site could be independent of the poly(A) size, if it is merely determined by any sort
of template masking. Rather the results seem to suggest the existence of a tool able to measure the appropriate distance (~20 nt) downstream of the heteropolymeric part of the viral RNA regardless of its sequence or structure. The nature of this tool is yet to be elucidated, but a possible role of the CRE(2C), the site of VPgpUpU synthesis utilized for (−) strand RNA synthesis, cannot be excluded (14). One may also speculate that a putative cis-element in the RdRP-coding part of the viral RNA (31) may be involved in such measurements.

There is still another possibility for maintaining a constant poly(U) length. Following the nidovirus example, where the (−) RNA strand is synthesized discontinuously (55,56), one may contemplate that Vpg-dependent synthesis of enterovirus (−) strands starts at or near the 3′-end of the poly(A) template but after synthesizing ~20 nt-long poly(U), the viral RdRP together with the nascent poly(U) jumps to the heteropolymer/homopolymer boundary to continue the elongation. As yet, such a mechanism for picornaviruses is not supported by any evidence.

**Emergency tailing of positive strands**

Efficient enterovirus replication requires the presence of a poly(A) tail in viral RNA with a minimal length of ~20 nt. Nevertheless, most poly(A)-lacking constructs proved to be (quasi)-infectious, generating genomes with newly-acquired AU-rich tails terminating with poly(A). Such nucleotide preference might reflect peculiarities of a yet-to-be-defined tailing reaction or merely resulted from selection because other types of tails might confer a lesser fitness. Mechanisms of elongation of the RNAs lacking poly(A) (or poly(A) and
3’NCR) are not necessarily the same as those operating in the case of wt viruses.

At least for the cases, where the engineered RNA did not possess an appropriate oligoadenylate to bind, or to generate, the VPg-pUpU primer [e.g. Δ3’NCR-N70-Apoly(A) and Δ3’NCR-N111-Apoly(A)], it is likely that some tailing of the viral RNA occurred prior to its use as a template for the (−) strand. Such tailing may involve a TNT-like enzyme of viral or host origin or non-replicative recombination (46,57,58), or both. Participation of recombination is supported by the observed inclusion of apparently host-derived sequences identical to a part of the human heterogeneous nuclear RNP U mRNA (residues 2408–2437) into the acquired 3’-tails upon passages (data not shown).

Passages of viruses with altered 3’-tails resulted in further modifications, e.g. elongation of internal poly(U) and poly(A) stretches. One may hypothesize that in addition to the terminal nucleotide transfer and recombination, reiterative synthesis using either 3’-end-adjacent sequences of (+) strand or 5’-end-adjacent sequences of (−) strand as templates may be involved. It seems likely that similar aberrant tailing can occur during the wt virus infection as well, but low-fitness viruses thereby generated are readily outcompeted and become undetectable.

Not surprisingly, the AU-rich sequences often contain one or more canonical cellular nuclear polyadenylation signals, AAUAAA. It is unknown whether they appear just by chance or are selected because they promote recruitment of the host polyadenylation machinery. A possible contribution of the AAUAAA signals to the appearance of oligo(A) stretches in the aberrant 3’-tails of the viral RNA is supported by the finding that the addition of this signal restored viability of Δ3’NCR-N70-Apoly(A) RNA (Table 2). Also in Bamboo Mosaic Virus it was shown that this hexameric sequence is involved in genomic RNA polyadenylation (59). Furthermore, treatment of human cells with siRNAs specific for the cellular polA polymerase resulted in downregulation of the enzyme and inhibition of poliovirus replication (C. Polacek and R. Andino, unpublished data). The origin of the poly(U) blocks in the (+) RNA molecules is enigmatic. It cannot be excluded that they are also due to a TNT-like activity. Remarkably, these segments are often located upstream of the 3’-poly(A). Hence, if tailing occurs at the level of viral RNA, the 3’-poly(A) may be added by snap-back synthesis on poly(U) blocks. However, the scarcity of experimental data makes such notion rather speculative.

Mechanisms of the (−) RNA initiation on the abnormal 3’-tails of the genomic RNA are yet to be determined, but our data suggest that viral RNA undergoes polyadenylation prior to replication.

Robustness of the viral genome

Illuminating examples of the robustness of the enteroviral genomes are accumulating. The oriR, a highly conserved cis-element, controls polyadenylation of the progeny RNA and initiation of the (−) strands but is nevertheless dispensable for viral viability ([29–31]; this study). Removal of poly(A), an element required for efficient genome replication, especially in combination with oriR inactivation, inflicted a severe damage to the viral fitness, though again usually not fatal. The poly(A) is functionally replaced by newly acquired AU-rich poly(A)-terminated 3’-tails potentially able to accommodate the VPg primer. Such a relatively unspecific regeneration of damaged 3’-tails of viral RNAs is not unique for enteroviruses (60).

Several mechanisms may be responsible for the viral robustness. First, the error-prone viral RdRP generates a variety of neutral or slightly detrimental mutants, not accumulating under normal conditions. However, ‘at emergency’, when the viral genome is severely invalidated, some of such mutations may result in a fitness increase, as exemplified by apparent restoration of the functional oriR structure after destruction of the kissing interaction [(24); this study] or changing mutual orientation of the oriR helical elements (25). Second, oriR belongs to a class of cis-elements, which are important but not essential. It is tempting to assume that such elements could have been acquired at some steps of enterovirus evolution in order to increase viral fitness. If so, the oriR-lacking mutants could be regarded as relying on ancient, ‘atavistic’ replicative tools. Regeneration of poly(A) after its complete deletion suggests the existence of unspecific mechanisms, which may ‘revive’ nearly dead viruses. Some of these mechanisms perhaps involve host enzymes.

Although certain RNA damages may generate rather crippled viruses, their very survival gives them chances to acquire by recombination either their lost element(s) or structurally unrelated but functionally active element from other viruses or hosts. Such loss of an element and acquisition of its functional substitute from an outside source may be an important path of viral evolution.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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