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DOI:
10.1074/jbc.M114.550657

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Document Version
Publisher's PDF, also known as Version of record

Citation for published version (Harvard):
Chung, L, Bailey, D, Leen, EN, Emmott, EP, Chaudhry, Y, Roberts, LO, Curry, S, Locker, N & Goodfellow, IG 2014, 'Norovirus translation requires an interaction between the C terminus of the genome-linked viral protein VPg and eukaryotic translation initiation factor 4G', Journal of Biological Chemistry, vol. 289, no. 31, pp. 21738-21750. https://doi.org/10.1074/jbc.M114.550657

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J. Biol. Chem. 2014, 289:21738-21750.
doi: 10.1074/jbc.M114.550657 originally published online June 13, 2014

Access the most updated version of this article at doi: 10.1074/jbc.M114.550657

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Norovirus Translation Requires an Interaction between the C Terminus of the Genome-linked Viral Protein VPg and Eukaryotic Translation Initiation Factor 4G*

Received for publication, January 16, 2014, and in revised form, May 27, 2014 Published, JBC Papers in Press, June 13, 2014, DOI 10.1074/jbc.M114.550657

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Background: Noroviruses use a virus-encoded protein, VPg, covalently linked to the viral RNA for translation.

Results: The direct interaction of VPg with the central domain of eIF4G is required for norovirus translation.

Conclusion: eIF4G plays a central role in norovirus VPg-dependent translation initiation.

Significance: The VPg-eIF4G interaction may provide a suitable target for the specific inhibition of norovirus translation.

Viruses have evolved a variety of mechanisms to usurp the host cell translation machinery to enable translation of the viral genome in the presence of high levels of cellular mRNAs. Noroviruses, a major cause of gastroenteritis in man, have evolved a mechanism that relies on the interaction of translation initiation factors with the virus-encoded VPg protein covalently linked to the 5' end of the viral RNA. To further characterize this novel mechanism of translation initiation, we have used proteomics to identify the components of the norovirus translation initiation factor complex. This approach revealed that VPg binds directly to the eIF4F complex, with a high affinity interaction occurring between VPg and eIF4G. Mutational analyses indicated that the C-terminal region of VPg is important for the VPg-eIF4G interaction; viruses with mutations that alter or disrupt this interaction are debilitated or non-viable. Our results shed new light on the unusual mechanisms of protein-directed translation initiation.

As obligate intracellular pathogens, viruses rely exclusively on the host cell for the components of the protein synthesis machinery and have evolved a multitude of mechanisms to enable the translation of viral mRNA into protein in the presence of high concentrations of competing cellular mRNA (1, 2). The translation of mRNA into protein is the final stage in the process of gene expression. The key regulatory step in this process is the initiation phase, a multistep process that results in recruitment of the ribosomal subunits to the start codon on the mRNA. Initiation requires the participation of a number of eukaryotic initiation factors (eIFs)3 to assemble a 43 S preinitiation complex on an mRNA molecule before completion of the 80 S initiation complex at the initiation codon. Cytoplasmic mRNAs are capped at the 5' end, and the first step in translation is the binding of the initiation factor eIF4E, a component of the eIF4F complex, to the 5' cap structure. eIF4F is a complex of three initiation factors: eIF4E is the cap-binding protein, eIF4A functions as an RNA helicase, and eIF4G acts as a scaffold to bridge the mRNA to the 40 S ribosomal subunit via its interaction with eIF3 (3). eIF4G contains three “domains”; the N-terminal one-third contains the eIF4E-binding site (4, 5) and a binding site for poly(A)-binding protein (6), the middle third contains binding sites for eIF3 and eIF4A (7), and the C-terminal third contains another eIF4A-binding site (7) as well as a binding site for the regulatory kinase, Mnk-1. After the 43 S ribosomal complex is bound to the 5' cap of the mRNA, it scans in the 5’ to 3’ direction to find the AUG initiation codon, at which point the 60 S ribosomal subunit joins, forming an 80 S initiation complex that commences protein synthesis.

Noroviruses, members of the Caliciviridae family of small positive-strand RNA viruses, are a major cause of acute gastroenteritis in man (8) but have also been identified in a number of other species including dogs (9, 10), cats (11), sheep (12), and cattle (13). Recent data indicate that noroviruses may also be associated with more significant clinical diseases including necrotizing enterocolitis and benign seizures in infants (14–17). Despite the significant economic impact of noroviruses and

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* This work was supported by funding from the Biotechnology and Biological Sciences Research Council (BBSRC) (BB/I01232X/1) and the Wellcome Trust (WT097997MA) (to I. G.) funding from the BBSRC (BB/I01232X/1; to L. R. and N. L.) as well as BB/J001708/1 (to S. C.).

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§ Author’s Choice—Final version full access.

The abbreviations used are: eIF, eukaryotic initiation factor; MNV, murine norovirus; FCV, feline calicivirus; TAP, tandem affinity purification; hpt, hours post-transfection; HCV, hepatitis C virus; PABP, poly A binding proteins; IRES, internal ribosome entry site; GMP PNP, guanosine 5’-[(β,y-imido)triphosphate].
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numerous ongoing efforts, human noroviruses have yet to be cultivated in the laboratory (18). Our understanding of norovirus biology has been greatly enhanced by the discovery of murine norovirus (MNV) in 2003 (19), which remains the only norovirus that can be cultivated in the laboratory (20). Combined with the availability of various reverse genetics systems (21–23), MNV provides a valuable system with which to dissect the norovirus life cycle and has greatly increased our understanding of the molecular mechanisms of norovirus genome translation and replication (for review, see Ref. 24).

We previously demonstrated that the feline calicivirus (FCV), MNV, and human norovirus VPG proteins can interact directly with the cap-binding protein eIF4E (25, 26). A similar interaction of the plant potyvirus VPG with eIF4E has also been reported and is known to determine the relative susceptibility of plant species to infection (for review, see Ref. 27). Functional analyses of these interactions in calicivirus VPG-dependent translation have indicated that eIF4E is essential for FCV translation as depletion of eIF4E or inhibition of the eIF4E-eIF4G interaction ablates FCV translation. Importantly, however, despite the interaction between the norovirus VPG protein and eIF4E, it appears to be dispensable for MNV translation initiation, at least in vitro (25). Therefore to date, no functional role for a VPG-eIF4E interaction has been described. FCV and MNV belong to the Vesivirus and Norovirus genera of the Caliciviridae family, although a significant degree of similarity is likely to exist in their mechanism of genome translation and replication, subtle differences in the relative requirements for cellular factors between different genera is likely. Previous reports have also highlighted a potential association of the norovirus VPG protein with components of the eIF3 complex (28) and eIF4E, PABP, and eIF4G as well as the ribosomal protein S6 (29), although the functional relevance of these interactions has yet to be demonstrated. Here we describe the proteomic characterization of the murine norovirus translation initiation factor complex, demonstrating that VPG associates directly with the core components of the eIF4F complex and PABP. We further demonstrate that the interaction between eIF4G and VPG is essential for norovirus translation. Furthermore, we demonstrate that eIF4G is required for efficient virus replication in cell culture.

EXPERIMENTAL PROCEDURES

Cells, Plasmids, and Antibodies—Murine macrophage RAW264.7 and microglial BV2 cell lines were cultured in Dulbecco’s modified Eagle’s medium (DMEM) with 10% (v/v) fetal calf serum (FCS), penicillin (100 units/ml), streptomycin (100 μg/ml), and 10 mM HEPES buffer. Baby hamster kidney cells (BHK-21) expressing T7 RNA polymerase (BSRT7 cells) were cultured in similar media lacking HEPES but containing 1 mg/ml G418. Similarly, HEK 293T cells were maintained in media lacking HEPES. HEK 293T cells stably expressing pMEP4-NTAP or pMEP4-NTAP MNV VPG plasmids were supplemented with 50 μg/ml hygromycin B and nonessential amino acids. The HEK293 TREX cells stably expressing pcDNA4/TO-NTAP derivatives of MNV VPG were supplemented with 5 μg/ml Blasticidin and 200 μg/ml Zeocin. All cell lines were maintained at 37 °C and 10% CO2.

The constructs pET28a:HIS N-FAG1–532, pET28a:HIS C-FAG1177–1600, pET28a:HIS M-FAG533–1176, pET28a:HIS p100675–1600, and pET28a:HIS 4GM654–1131, expressing the truncated eIF4GI proteins N-FAG, C-FAG, M-FAG, p100, and 4GM, respectively, were kindly provided by Simon Morley (University of Sussex). Infectious clones of wild type MNV (pT7-MNV 3′RZ) or an MNV mutant with an alanine substitution at position Phe-123 of the VPG protein, F123A (pT7-MNV 3′RZ F123A), were used in various experiments. pCDNA4/TO-FLAG 4GM was generated by PCR amplification of 4GM from pET28a:HIS 4GM using forward (GTAAAGCTTGCACTATGGATTACAAAGGATGAGCCATAGAATTGCTATGAGC) and reverse (TGCGTCGAGTCAACCCTGGTGTTGAAAG) primers to generate a HindIII/XhoI fragment, which was inserted into pCDNA4/TO.

Antiseras to the various MNV non-structural proteins were generated in-house and have been described previously (22, 25). Other antisera was sourced as follows: eIF4E (Santa Cruz; sc-13963), eIF4A (Santa Cruz; sc-14211), PABP (New England Biolabs; 4992), eIF4G (2498; New England Biolabs), GAPDH (AM4300; Ambion), and the histidine tag (HIS; Santa Cruz).

Tandem Affinity Purification—A detailed overview of this protocol is available elsewhere (30). Two expression systems, tetracycline inducible system (pCDNA6/TR and pcDNA4/TO NTAP) and cadmium chloride (CdCl2) inducible system (pMEP4-NTAP), were used for expressing genes of interest, i.e. genes encoding MNV VPG. Both inducible systems expressed a GS tag (containing two protein G units and streptavidin binding peptide). For the CdCl2 inducible system, the transfected 293 cells were selected with 50 μg/ml hygromycin B (Roche Applied Science) and were later induced with 10 μM CdCl2 for 16 h. For the tetracycline inducible system, 293 T-REx cells were selected with 5 μg/ml Blasticidin and 200 μg/ml Zeocin after a 24-h post-transfection of plasmid. A single colony was isolated from the cell population to examine protein expression. ~2 × 106 cells expressing a protein of interest were induced with 1 μg/ml doxycycline for 16 h. Tandem affinity purification (TAP) pulldown assays were then performed on lysates prepared from these cells as described previously (30). Eluted proteins were concentrated using VivaSpin 500™ column (Vivascience). Samples were subject to SDS-PAGE gel electrophoresis and stained with Coomassie Blue. The whole gel lane was divided into either 25 or 9 parts as appropriate, excised, and analyzed by mass spectrometry (Genome Quebec Innovative Center, McGill University).

The McGill University and Genome Quebec Innovation Centre, Canada, performed all mass spectrometry (MS). Proteins present in bands excised from SDS-PAGE gels were identified using a quadrupole TOF micro-mass spectrometer (Waters Micromass) as described previously (31). In this case, however, Mascot was set up to search a Homo sapiens database. Scaffold (Proteome Software Inc.) was used to validate tandem MS (MS/MS)-based peptide and protein identifications. Similar purifications and mass spectrometry analyses were performed with cell lines expressing the GS tag alone to provide a list of nonspecific interactions, which were then removed from the data set. Those remaining were considered VPG-specific interacting proteins.
m7GTP-Sepharose Chromatography—BSRT7 cells were used to overexpress viral proteins from full-length cDNA constructs as described below for the reverse genetics recovery of MNV. At 18 h post-transfection cells were lysed in 120 μl of CAP lysis buffer (1% Triton X-100, 100 mM KCl, 0.1 mM EDTA, 10% glycerol, 2 mM MgCl₂, 20 mM Hepes, pH 7.6, and protease inhibitors mixture) and were incubated on ice for 5 min at 4 °C. Non-lysed cells and debris were subsequently incubated with prewashed m7GTP-Sepharose (25 μl bead volume; GE Healthcare) along with an additional 120 μl of CAP lysis buffer (without 1% Triton X-100) for 2 h at 4 °C to allow binding to take place. Any unbound proteins were later removed by centrifugation at 10,000 × g for 5 min followed by washing 3 times using lysis buffer without Triton X-100.

Finally, proteins associated with the cap structure were eluted with SDS-PAGE reducing sample buffer and analyzed by Western blot after subjected to SDS-PAGE electrophoresis.

RNA Affinity Chromatography—Ribosome-bound MNV RNA translation complexes were purified as described previously (32). Briefly, 25 μg of purified Vpg-linked MNV RNA or in vitro transcribed HCV IRES (as described in Ref. 33) were pre-annealed to biotinylated DNA oligonucleotides targeting the 3′ end of the MNV genome (5′-GCATCTAACTACCAC-AAAGAAAAAGAAC-3′; 5′-GCCCTGCTACTCCCGATCC-TTGGG-3′) or the HCV IRES construct (5′-GGGATTTCTGTATCTGGC-3′; 5′-TTTCTGATCTGGCTTCA-3′) and translated for 15 min at 30 °C in rabbit reticulocyte lysates pretreated with 5 μM GMP-PNP and 0.5 μM puromycin before the addition of 100 μg/ml cycloheximide. Translation complexes were subsequently immobilized onto Streptavidin Dynabeads M-280 (Invitrogen). After extensive washing in THEMK buffer (34 mM Tris, 66 mM HEPES, 0.1 mM EDTA, 2.5 mM MgCl₂, 75 mM KCl, pH 7.8), the complexes assembled onto MNV RNA or HCV IRES were released by adding RNase H. The presence of individual eIFs was then assayed by immunoblotting as described above and compared with a control lane containing rabbit reticulocyte lysate.

Reverse Genetics Recovery of Mutant Viruses—Reverse genetics recovery of recombinant MNV was performed by infecting baby hamster kidney cells with foxplow virus expressing T7 RNA polymerase followed by transfection of MNV cDNA expression constructs as described previously (22, 34). To analyze protein expression, cells were harvested 24 h post-transfection for Western blot analysis. Virus yield was determined by TCID₅₀.

Identification of the Minimal Vpg Binding Domain within elf4GI—Conducts expressing the truncated elf4GI proteins, N-FAG, C-FAG, M-FAG, p100, and 4GM (described above) were co-transfected with either an infectious clone of wild type MNV or MNV containing an alanine substitution at position Phe-123 of the VPg protein (F123A) into FPV-infected BSRT7 cells using Lipofectamine 2000 (Invitrogen). pcDNA3.1+ was used as a negative control. At 24 h post-transfection, the cells were lysed in 120 μl of CAP lysis buffer (1% Triton X-100 mM KCl, 0.1 mM EDTA, 10% glycerol, 2 mM MgCl₂, 20 mM Hepes, pH 7.6, and protease inhibitor mixture) and centrifuged at 13,000 × g in a benchtop microcentrifuge to pellet insoluble debris.

The cleared lysate was then added to prewashed Ni-CAM beads and an additional 120 μl of CAP lysis buffer lacking Triton X-100 was added. The reaction mix was incubated on a rotary mixer at 4 °C overnight. The beads were then washed 5 times using a more stringent CAP buffer (200 mM KCl, 0.1 mM EDTA, 10% glycerol, 2 mM MgCl₂, 20 mM Hepes, pH 7.6) in the absence of Triton X-100. Proteins were eluted by resuspending the beads in 45 μl of SDS sample buffer and boiling the samples for 5 min. Purified protein was resolved in an SDS-PAGE gel and analyzed by Western blotting.

His-Vpg-elf4GI Pulldown Assay—The pETM11 plasmid encoding MNV Vpg 1–124 is as described elsewhere (35). QuikChange site-directed mutagenesis (Stratagene) was used to produce the MNV Vpg 1–124 F123A mutant. elf4GI 4GM (652–1132) was amplified by PCR from a human elf4GI clone (NCBI accession number AAM69365.1) and cloned into a modified pGEX2T vector using BamHI and EcoRI restriction enzymes (primer details available upon request). GST and GST-elf4GI 4GM were expressed in Escherichia coli (DE3) Rosetta and E. coli (DE3) CodonPlus cells, respectively. Expression was induced during mid-log phase by the addition of 1 mM final isopropyl 1-thio-D-galactopyranoside in the case of GST for 4 h at 37 °C and for 3.5 h at 30 °C for GST-elf4GI 4GM. Wild type and F123A MNV Vpg were described as previously (35).

GST and GST-elf4GI 4GM were purified by affinity chromatography using Glutathione-Sepharose 4B resin (GE Healthcare). GST was dialyzed against 50 mM Tris buffer, pH 7.6, 200 mM NaCl, and 2 mM β-mercaptoethanol. GST-elf4GI 4GM was dialyzed against the same buffer but with 150 mM NaCl and then further purified by size exclusion chromatography using a Superdex S200 10/300 column (GE healthcare). Both proteins were finally dialyzed against 50 mM Tris buffer, pH 7.6, containing 150 mM NaCl (binding buffer). His-tagged MNV Vpg proteins (wild type and the F123A mutant) were purified by affinity chromatography to TALON resin (Clontech) and subsequently dialyzed against 50 mM sodium phosphate, pH 6.5, 300 mM NaCl, and 1 mM DTT.

In pulldown experiments 25 μl of TALON beads were used to capture the bait protein (His₆-tagged MNV Vpg WT or F123A at a final concentration of 5.5 μM) and GST or GST-elf4GI were used as prey proteins as a final concentration of 1.1 μM. The proteins were mixed together with 450 μl of binding buffer in SigmaPrep spin columns (Sigma) and incubated by rotation at 4 °C for 90 min before collection of the flow through fraction. The resin was washed with 750 μl of binding buffer then twice with 10 mM imidazole in 750 μl of binding buffer. The bait was eluted with 100 μl of 500 mM imidazole in binding buffer. Captured proteins were analyzed by SDS-PAGE.

siRNA Knockdown and Reconstitution Experiments—elf4GI-siRNA-mediated reduction of protein expression was performed in HEK 293T cells. Cells were transfected with either control nonspecific siRNA (GGCAGUUCUUUGUAG-GAUUCG) or elf4GI-specific siRNAs (CCCAUACUGGAGGAG). At 24 h post-transfection the cells were retransfected with the appropriate siRNA and in the case
of reconstitution experiments co-transfected with either pCDNA4/TO or pCDNA4/TO FLAG 4GM. 24 h after this second transfection cells were transfected with VPg RNA isolated from MNV-infected BV-2 cells. Cells were then harvested at the appropriate times for Western blotting. Percentage reconstitution of viral translation was determined using Li-Cor Odyssey imaging whereby the levels of viral translation observed in the eIF4G siRNA- and empty vector (pCDNA)-transfected cells was set as background, and the values obtained from 4GM-expressing cells were expressed relative to the nonspecific siRNA-transfected cells. siRNA-mediated reduction of eIF4E was performed in BV-2 cells using a pool of eIF4E specific siRNAs (Santa Crux, sc-35285) transfected as described previously (31).

RESULTS

Proteomic Analysis of the Murine Norovirus Translation Initiation Factor Complex—Given the lack of functional data on the initiation factor requirements for norovirus translation initiation, we aimed first to identify the components within the VPg-initiation factor complex and then to determine which of these were functionally important. We, therefore, established a method for purifying VPg from transfected cells with any associated initiation factors. This approach is based on the well known TAP method originally described in yeast (36) but subsequently adapted for use in mammalian cells (30, 37). Murine norovirus VPg was fused at the Nterminus to an affinity purification (TAP) tag consisting of two protein G domains, a tobacco etch virus protease cleavage site, and a streptavidin binding peptide (Fig. 1B). This expression cassette was cloned into two plasmid-based expression systems, the first an episomally maintained plasmid containing the inducible metallothionein promoter along with the Epstein-Barr virus EBNA1 protein and origin of replication (pMEP4) and the second based on pcDNA4:TO containing a CMV promoter controllable by the tetracycline repressor protein (Fig. 1B). Human 293T cells were used for this approach as we previously observed that, although these cells are not permissive to MNV infection due to the lack of a suitable receptor, robust MNV translation and replication occurs upon transfection of viral RNA into the cytoplasm (22). The MNV permissive cell line RAW264.7 proved to be unsuitable for this approach due to poor yields of tagged VPg recovered from these cells (data not shown).

Cell lines expressing either the tag alone (NTAP) or with TAP-VPg were generated, and expression was shown to be stimulated by the addition of cadmium chloride for the pMEP4-based system or doxycycline for the pcDNA4:TO-based expression system (data not shown). Protein expression was readily detectable by virtue of the protein G binding domain present in the TAP tag.

FIGURE 1. Generation of cell lines expressing TAP-VPg. A, schematic representation of the norovirus genome. The positions of the three major open reading frames (ORF1, -2, and -3) and the murine norovirus-specific open reading frame 4 (ORF4). The position and subgenomic RNA start site and the ORF1 polyprotein cleavage sites are highlighted. The NS1-NS7 nomenclature as proposed by Sosnovtsev (50) has been used to illustrate the identities of the mature cleavage products along with the previously described nomenclature (N-term, helicase, RdRp etc.). B, the domain layout of the TAP-tag is illustrated graphically highlighting the positions of the two protein G IgG binding domains, the tobacco etch virus (TEV) protease cleavage site, and the streptavidin-binding peptide along with the streptavidin binding peptide. C, m7GTP-Sepharose chromatography of cells expressing the NTAP tag or TAP-VPg fusion protein. Lysates from induced cells were incubated with m7GTP-Sepharose to enrich the eIF4F complex. Both lysates (L) and the elutions (m7) from the m7GTP-Sepharose were analyzed by Western blot for eIF4E and the presence of the protein G binding domain present in the TAP tag. An asterisk is used to highlight a TAP-MNV VPg degradation product present in the cell lysate.
A VPg-eIF4G Interaction Is Involved in Norovirus Translation

Figure 2. Proteomic analysis of the norovirus translation initiation factor complex. Final elutions after the tandem affinity purification procedure are detailed under “Experimental Procedures,” prepared from either cells expressing the TAP tag alone or the NTAP-VPg fusion protein. Panel A contains the final elutions obtained from the pMEP4-based cadmium chloride inducible expression system, whereas panel B displays the results obtained using the pCDNA4:TO tetracycline inducible system. In the latter case, the purification was performed in the presence and absence of ribonuclease (RNase). An asterisk is used to highlight the position of the bait protein after tobacco etch virus protease digestion and biotin-mediated elution.

To identify the host factors associated with VPg, large scale affinity purifications were performed with extracts from cells expressing the TAP tag alone or TAP-MNV VPg. SDS-PAGE analysis of the purified complexes demonstrated that a number of host cell factors were co-purified with VPg that were not present in complexes isolated using the TAP tag alone (Fig. 2). The identities of the proteins isolated using the TAP method were determined by the excision from the SDS-PAGE gel, digestion with trypsin followed by mass spectrometry. Specific associations with VPg were determined by the exclusion of proteins also identified in the purification using the TAP tag alone. To eliminate the possibility of pulling down proteins due to nonspecific RNA binding ribonuclease was included in purifications performed using the doxycycline-inducible expression system as sequence analysis and our unpublished observations indicated that the norovirus VPg protein has RNA binding activity (data not shown). The inclusion of ribonuclease did not appear to affect the profile of proteins co-purified with VPg (Fig. 2B).

Mass spectrometry was used to identify proteins present within the samples with a minimum of 2 unique peptides and >90% identification probability. Proteins present in the complexes isolated from cells expressing the TAP tag alone were excluded as nonspecific interacting proteins. Analysis of the proteins isolated only on VPg reproducibly identified the core eIF4F components, eIF4A, -4E, and -4G, along with PABP as factors that associate with VPg (Table 1). eIF4G1 and 4GII were reproducibly identified using both experimental systems as were eIF4E, eIF4AI, and -4AII and PABP1 and -4. In addition, components of the eIF3 complex were found associated with VPg; the episomally maintained cadmium chloride-inducible system also identified the eIF3 complex (subunits A–I and K–M), whereas the tetracycline inducible system found subunits A–C, E, and K only. In addition to the canonical translation initiation factors, the episomally maintained expression system also identified a number of cellular factors including ribosomal proteins (P2, S25, S12, and S15a), LARP1, DDX9, IGF-BP1, tubulin, eEF1a, and ABCE1. As these proteins were not identified in both experimental systems and the focus of this study was to determine the role of canonical initiation factors in VPg-dependent translation, these additional host cell proteins were not examined in more detail. However, it is worth noting that some of these proteins have previously been identified as playing a role in the life cycles of other RNA viruses (discussed in more detail below).

VPg Binds to the eIF4F Complex via an Interaction with eIF4G—To identify the interactions within the VPg-dependent translation initiation complex that are required for viral translation, we attempted to identify which components of the complex that interacted with VPg directly and contribute to viral translation. Although direct affinity measurements were not possible using the TAP purification approach, we reasoned that the interaction of VPg with the protein with the highest affinity would be resistant to increasing ionic strength. Tandem affinity purification was again performed using the murine norovirus NTAP-VPg bait protein; however, during the final washing steps increasing concentrations of sodium chloride, ranging from 125 mM to 1 M, were included to increase the stringency of the washes. Our results indicated that although the levels of eIF4E, eIF4A, and PABP isolated using NTAP-VPg were readily reduced as the ionic strength increased, the levels of eIF4G remained largely unchanged (Fig. 3A). These data would suggest that the norovirus VPg associates with the eIF4F complex and PABP, yet it can form a stable high affinity interaction with eIF4G. It is worth noting, however, that in addition to disrupting electrostatic interactions, increasing sodium chloride concentrations stabilize hydrophobic interactions. Therefore, these data alone do not confirm eIF4G as a direct binding partner; however, they confirm, using the number of unique peptides identified for each protein (Table 1) as a semiquantitative indirect measure of protein abundance in the purified complex, it was also apparent that eIF4G was enriched in the complex with respect to the other proteins isolated.

The presence of eIF4G in the norovirus translation initiation complex was confirmed by affinity purification of assembled translation complexes from rabbit reticulocyte lysates programmed with either HCV IRES containing RNA or MNV VPg-linked RNA (Fig. 3B). Western blot analysis of the purified complex confirmed the presence of eIF4G and eIF4E on MNV VPg-linked RNA, both of which were absent from complexes purified on HCV IRES containing RNA as expected. In contrast, eIF3f was found associated with both MNV VPg-linked RNA and the HCV IRES containing RNA.

Mutations in the C Terminus of VPg Affect Initiation Factor Binding and Virus Viability—We have recently determined the structure of the MNV VPg protein (35) and demonstrated that
although the central domain consists of two α helices, the N and C termini are largely disordered. To identify the regions within the norovirus VPg protein that are required for the interaction with initiation factors, we targeted several conserved amino acids for alanine mutagenesis. 14 individual point mutations in VPg were introduced into the MNV infectious clone, and protein expression was driven by coinfection of the cells with a fowlpox virus expressing T7 RNA polymerase as previously described (22). This approach was used to circumvent problems associated with reduced protein expression from non-viable viral genomes.

The effect of each mutation on virus viability was assayed using virus yield 24 h post transfection, and the effect of mutations on the ability of VPg to associate with initiation factors was assessed by the enrichment of the eIF4F complex using m7GTP-Sepharose followed by Western blotting for VPg. The mutation K3A ablated virus recovery, but this effect was mostly due to an effect on polyprotein processing as large VPg-containing precursors, not typically observed during infection, were readily detected (highlighted with an asterisk in Fig. 4).

The mutations K5A and K7A, which reduced virus yield by 10-fold, had no appreciable effect on initiation factor binding or, as shown during our previous work, the ability of the VPg to function as a template for covalent attachment of RNA to VPg by polymerase-mediated nucleotidylation (35). Our previous work has highlighted that mutations in the region Asp-23 to

![A VPg-eIF4G Interaction Is Involved in Norovirus Translation](http://www.jbc.org/content/289/31/21743)

**TABLE 1**

| Protein name | Abbreviation | Protein molecular weight (Da) | Protein accession numbers | Number of unique peptides/Sequence coverage | Protein accession number | Number of unique peptides/Sequence coverage |
|--------------|--------------|------------------------------|---------------------------|------------------------------------------|--------------------------|------------------------------------------|
| Fas tyrosine translation initiation factor 4 γ, 3 variant | eIF4GIII | 195,843 | Q05060 | 47 | 32.1% | Q05060 | 35 | 26.6% |
| Fas tyrosine translation initiation factor 4 γ, 1 variant | eIF4GII | 178,140 | Q47E8 | 40 | 40.1% | Q47E8 | 49 | 31.8% |
| Fas tyrosine translation initiation factor 3, subunit A | eIF3a | 166,557 | B1AMV5.Q41452.20014 | 9 | 9.4% | B1AMV5.Q41452.20014 | 12 | 9.4% |
| Fas tyrosine translation initiation factor 3, subunit B | eIF3b | 92,475 | Q05211 | 2 | 1.7% | Q05211 | 2 | 1.7% |
| ATP-dependent RNA helicase A | eIF4A | 103,329 | A1qvY7.099613 | 10 | 13.4% | A1qvY7.099613 | 10 | 13.4% |
| La-related protein | eIF4A | 92,475 | A40584.A40584 | 14 | 21.6% | A40584.A40584 | 14 | 21.6% |
| Polyadenylate-binding protein 4 | eIF5A | 70,766 | B1AQ09.014310.0562 | 11 | 34.8% | B1AQ09.014310.0562 | 11 | 34.8% |
| Polyadenylate-binding protein 2 | eIF5B | 70,653 | P19990 | 24 | 40.6% | P19990 | 24 | 40.6% |
| ATP-binding cassette sub-family E member 1 | eIF2C | 67,298 | P16221 | 6 | 9.9% | P16221 | 6 | 9.9% |
| Fas tyrosine translation initiation factor 5, subunit L | eIF5L | 66,711 | B0Q9P9.Q0709262 | 9 | 16.2% | B0Q9P9.Q0709262 | 9 | 16.2% |
| Fas tyrosine translation initiation factor 5, subunit D | eIF5D | 63,956 | A5g8a0.Q0728565 | 10 | 22.1% | A5g8a0.Q0728565 | 10 | 22.1% |
| Insulin-like growth factor 2 mRNA-binding protein 1 | eIF6 | 53,408 | P60228.098633.0962XX5.Q0BW4K | 21 | 47.3% | P60228.098633.0962XX5.Q0BW4K | 21 | 47.3% |
| Fas tyrosine translation initiation factor 3, subunit E | eIF6 | 52,200 | A29BF3.Q08701.098229.Q0NF5 | 3 | 6.7% | A29BF3.Q08701.098229.Q0NF5 | 3 | 6.7% |
| Elongation factor 1-alpha | eEF1A | 56,123 | Q3YR52.093326 | 2 | 33.5% | Q3YR52.093326 | 2 | 33.5% |
| Tubulin, beta 2C | TUB2C | 49,813 | Q47F23.093326 | 5 | 47.7% | Q47F23.093326 | 5 | 47.7% |
| Fas tyrosine translation initiation factor 4A, II | eIF4A | 46,386 | ARK76Q6.P0684 | 15 | 70.4% | ARK76Q6.P0684 | 15 | 70.4% |
| Fas tyrosine translation initiation factor 4A, I | eIF4AM | 46,137 | ARK76Q6.P0685 | 15 | 70.4% | ARK76Q6.P0685 | 15 | 70.4% |
| Fas tyrosine translation initiation factor 1, subunit M | eIF5M | 47,787 | AKR76Q6.P0686 | 18 | 45.4% | AKR76Q6.P0686 | 18 | 45.4% |
| Acetyl coenzyme A synthetase | ACS2A | 41,720 | AKR76Q6.P0687 | 24 | 58.6% | AKR76Q6.P0687 | 24 | 58.6% |
| Fas tyrosine translation initiation factor 3, subunit H | eIF3H | 39,913 | Q1S372.093607.093800.Q0AM9B | 3 | 8.8% | Q1S372.093607.093800.Q0AM9B | 3 | 8.8% |
| Fas tyrosine translation initiation factor 3, subunit F | eIF3F | 37,546 | ARK90.093607 | 6 | 24.6% | ARK90.093607 | 6 | 24.6% |
| Fas tyrosine translation initiation factor 3, subunit I | eIF3I | 36,484 | Q13457.Q0148S | 5 | 15.7% | Q13457.Q0148S | 5 | 15.7% |
| Fas tyrosine translation initiation factor 3, subunit G | eIF3g | 35,994 | Q57S21.093608 | 2 | 9.1% | Q57S21.093608 | 2 | 9.1% |
| Fas tyrosine translation initiation factor 4E | eIF4E | 25,080 | ARK90.093607.Q025Q0 | 2 | 11.3% | ARK90.093607.Q025Q0 | 2 | 11.3% |
| Fas tyrosine translation initiation factor 5, subunit K | eIF5K | 23,042 | ARK76Q6.P0684 | 2 | 11.3% | ARK76Q6.P0684 | 2 | 11.3% |
| 40S ribosomal protein S14 | RPS14A | 14,822 | N55-MYN1.04000 | 12 | 65.3% | N55-MYN1.04000 | 12 | 65.3% |
| 40S ribosomal protein S12 | RPS12 | 14,497 | N55-MYN1.04000.P0683 | 12 | 65.3% | N55-MYN1.04000.P0683 | 12 | 65.3% |
| MNV VPg | VPg | 14,281 | P60251 | 3 | 22.4% | P60251 | 3 | 22.4% |
| 40S acidic ribosomal protein P2 | RPS25 | 13,723 | N55-MYN1.04000.P0686 | 3 | 40.9% | N55-MYN1.04000.P0686 | 3 | 40.9% |

**FIGURE 3. VPg interacts with eIF4G.** A, cells expressing either the NTAP fusion tag or NTP-MNV VPg were used to perform tandem affinity purification as described under “Experimental Procedures”; however, before biotin elution the samples were washed with buffer containing 125 mM, 500 mM, or 1 M sodium chloride. After biotin elution, samples were concentrated, separated by SDS-PAGE, and then analyzed by Western blotting. Both lysates (L) and the elutions (E) were analyzed for the presence of eIF4G, eIF4A, eIF4F, and PABP. Asterisks are used to highlight the positions of the TAP tag and the TAP-VPg proteins detectable in the lysate due to the presence of protein G binding domains. B, translation initiation complexes were purified from rabbit reticulocyte lysates (RRL) programmed with either the HCV IRES or VPg-linked MNV RNA by RNA-affinity purification as described under “Experimental Procedures.” Samples were subsequently analyzed by Western blot for the presence of various translation initiation factors with a sample of the input rabbit reticulocyte lysates used as a control.
Glu-28, most of which are lethal for virus viability, alter the ability of VPg to function as a template for nucleotidylation (35), but none of these was found to affect the ability of VPg to bind initiation factors. In contrast, the mutations V115A, D116A, and F123A altered the levels of infectious virus produced; a reduced virus yield was observed in the case of V115A and D116A, whereas the mutation F123A completely ablated infectious virus production. Although the majority of the mutants had little or no reproducible effect on the ability of VPg to associate with eIF4F compared with wild type VPg, the mutants V115A, D116A, and F123A showed a consistently reduced interaction with eIF4F (Fig. 4). The mutation F123A clearly resulted in the greater reduction in the amount of VPg recovered using m7GTP-Sepharose; low levels of VPg F123A were reproducibly detected only upon long exposure of the Western blots (data not shown). The degree to which the mutation affected virus recovery appeared to correlate with the ability of VPg to be co-purified with initiation factors (Fig. 4). In addition, sequence analysis indicated that the V115A mutant was unstable and readily reverted back to WT after repeated passage in cell culture (data not shown). These data would indicate that the disordered C terminus of the norovirus VPg protein contains amino acids involved in the interaction with eIF4G.

To further validate our observation that the C terminus of VPg contributes to the interaction with eIF4G, we performed small scale tandem affinity purification with both the WT and the F123A mutant form of MNV VPg and examined the puri-

![Figure 4](https://www.jbc.org/)

**FIGURE 4.** Mutations in the C terminus of VPg affect initiation factor binding and virus viability. BSRT7 cells infected with a fowlpox virus expressing T7 RNA polymerase were transfected with full-length cDNA clones of MNV containing either WT or various VPg mutants. 24 h post transfection cells were lysed, and the resulting lysate (L) was then subjected to m7GTP-Sepharose affinity chromatography. After purification, the proteins associated with m7GTP-Sepharose beads (m7) were separated by SDS-PAGE and analyzed by Western blotting with antisera to MNV VPg and eIF4E. Asterisks highlight the position of high molecular mass VPg-containing precursors formed as a result of incorrect polypeptide processing. Note that all Western blots were performed at the same time, and identical exposures were used to generate the figure shown. The m7GTP-Sepharose data presented is a single representative dataset from at least three independent repeats. The effect of VPg mutations on virus replication was also summarized in this figure. Virus viability is expressed as virus yield 24 h post transfection relative to wild type (+ + +) as assayed by >5 independent experiments. Typical yields of wild type virus were 1–5 x 10^4 TCID50 units. —, no virus detected; +, up to 100 TCID50 detected; ++, up to 1000 TCID50; ++++, up to WT levels of virus detected, typically >10,000 TCID50 per ml.

![Figure 5](https://www.jbc.org/)

**FIGURE 5.** The F123A mutation abolishes the VPg-eIF4G interaction. Tandem affinity purification was performed on human 293T cells transiently transfected with plasmids containing the TAP tag alone, TAP-wild type MNV VPg (WT), or TAP-MNV VPg containing the mutation F123A. Samples of the cell lysate (L) or the eluted purified complex (E) were analyzed by Western blot for eIF4A, eIF4G, PABP, and eIF4E. Asterisks are used to highlight the position of the TAP-VPg fusion proteins detected by the binding of the primary antibody to the protein G domains present in the TAP tag.
fied complex for eIF4E, eIF4A, eIF4G, and PABP. The mutation F123A ablated the ability of VPg to co-purify eIF4G, eIF4A, and PABP but did not affect the ability of NTAP-MNV VPg to co-purify eIF4E (Fig. 5), suggesting that the eIF4E and eIF4G binding regions on VPg are distinct. The ability of the F123A VPg mutant to interact with eIF4E is in agreement with the low levels of this mutant isolated using m7GTP-Sepharose (Fig. 4) and the ability of recombinant F123A VPg to bind eIF4E using a capture ELISA assay (data not shown). These data would suggest that the direct interaction of VPg with eIF4G largely determines the eIF4F binding capacity of the norovirus VPg protein.

VPg Interacts with the Central Domain of eIF4G via a Direct Protein-Protein Interaction—To determine which portions of eIF4G interact with the MNV VPg protein, we examined the ability of various previously described domains of eIF4GI to co-purify VPg when expressed in mammalian cells. Based on our previous observation that foot and mouth disease virus L protease-mediated cleavage of eIF4G had no effect on norovirus translation in vitro (25), we hypothesized that the interacting domain would be within the p100 region of eIF4G (amino acids 675–1600). Mammalian cells, previously infected with a fowlpox virus expressing T7 RNA polymerase were co-transfected with the various eIF4GI expression constructs (or empty vector) and either a wild type murine norovirus cDNA clone (WT) or one containing the F123A mutation in VPg (F123A). Lysates (L) or the purified complex (P) were prepared and used for nickel-affinity purification followed by Western blotting for the presence of either VPg or the recombinant protein (His).

FIGURE 6. VPg interacts with the central domain of eIF4G. A, cleavage map of eIF4GI highlighting the positions for translation factors. The arrow and the corresponding numbers illustrate the locations of the individual protease cleavage sites mediated by the cellular proteins caspase 3 or the viral protease Lpro. The expression constructs used in the assay are also illustrated along with the specific amino acids residues encompassed within the construct. B, nickel-affinity purification of His-eIF4G fragments. Cells previously infected with a foxpox virus expressing T7 RNA polymerase were co-transfected with the various eIF4GI expression constructs (or empty vector) and either a wild type murine norovirus cDNA clone (WT) or one containing the F123A mutation in VPg (F123A). Lysates (L) or the purified complex (P) were prepared and used for nickel-affinity purification followed by Western blotting for the presence of either VPg or the recombinant protein (His).
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Although GST-4GM readily co-purified with WT MNV VPg, GST did not. Moreover, the F123A mutation in the MNV VPg protein significantly reduced the interaction of VPg with eIF4G in this assay (Fig. 7). These data confirm that the VPg-eIF4G interaction occurs via a direct protein-protein interaction and does not require additional cellular proteins.

Neither the eIF4E-eIF4G Interaction nor High Levels of eIF4E Are Required for Norovirus Translation and Replication—We previously reported that the norovirus VPg protein can interact with eIF4E directly but have also shown that eIF4E is dispensable for norovirus translation in vitro (25). To explore whether the eIF4E-eIF4G interaction is required for norovirus translation during an authentic viral life cycle, we examined the effect of overexpression of 4E-BP1 on virus translation and replication in permissive cells. 4E-BP1 interferes with the eIF4E-eIF4G interaction, and 4E-BP1 overexpression results in the inhibition of eIF4G recruitment to the 5' end of mRNA. This process is regulated by phosphorylation of 4E-BP1 as only the phosphorylated form of 4E-BP1 interacts with eIF4E (38). Wild type or a non-phosphorylatable mutant version of 4E-BP1 were overexpressed in cells, and the effect on MNV replication was examined (Fig. 8, A and B). The overexpression of 4E-BP1 resulted in inhibition of the eIF4E-eIF4G interaction and had no effect on MNV viral protein expression (Fig. 8A), virus titer (Fig. 8B), or viral RNA levels (not shown), whereas the ability of eIF4G to be co-purified with eIF4E on m7GTP-Sepharose was significantly reduced by ~80 and ~60% for the WT and mutant forms of 4E-BP1, respectively (data not shown). Further evidence that eIF4E does not play a direct role in norovirus translation was obtained by examining the effect of siRNA-mediated reduction of eIF4E on norovirus replication (Fig. 8C). Reduction of cellular eIF4E levels by >90% (Fig. 8C) had no appreciable effect on MNV virus titer under the conditions used (Fig. 8D), suggesting that high levels of eIF4E are not required for efficient virus replication in immortalized microglial cells.

The eIF4G Central Domain Is Required for Norovirus VPg-dependent Translation—To determine if eIF4G plays a role in the translation of the VPg-linked norovirus RNA, we examined the effect of eIF4G siRNAs on norovirus replication (Fig. 9). Due to difficulties in reproducibly reducing eIF4G levels in mouse macrophage or microglial cells permissive to MNV infection, we used human cells instead. Transfection with eIF4G siRNAs was followed by transfection with viral VPg-linked RNA. This

FIGURE 7. VPg binds the central domain of eIF4G via a direct protein-protein interaction. The ability of recombinant eIF4G central domain (4GM) to interact with either wild type MNV VPg (WT) or the F123A MNV VPg mutant was examined using a His-tag pulldown assay. Recombinant GST or GST-4GM was mixed with purified His-tagged WT or F123A MNV VPg, and the resulting complex was purified using cobalt affinity chromatography. Samples of the purified proteins (Input), the mixtures of the proteins before purification (Reaction mix), and the final bound fraction were separated by SDS-PAGE and analyzed by Coomassie Blue staining.

FIGURE 8. Reduced levels of eIF4E and inhibition of the eIF4E-eIF4G interaction are not required for efficient norovirus replication. A, MNV permissive microglial cells BV2 were transiently transfected with either an empty plasmid vector (EMP) or plasmids expressing HA-tagged derivatives of wild type (WT) or a non-phosphorylatable mutant of 4E-BP1 (MUT). Cells were then infected with MNV1 at a multiplicity of infection of 3 TCID50/cell, and viral protein expression as well as virus titer (B) was determined at various times post infection. C, BV2 cells transfected with either non-targeting siRNA or an siRNA directed against eIF4E were infected with MNV1 at a multiplicity of infection of 10 TCID50/cell, and the effect on viral titer examined at various times post infection (D).
approach was based on the observation that MNV VPg-linked RNA is infectious when transfected in human cells (22), which suggest that all the cellular machinery required for efficient viral RNA translation and replication is conserved. At 11 and 16 h post transfection samples were analyzed for viral protein by Western blot and the production of infectious virus (Fig. 9). A significant reduction in both viral protein expression and virus replication was observed at both time points; the virus yield was reduced 8.9-fold (∼89%) after 11 h and 5.6-fold (∼82%) after 16 h. To confirm that the observed effect on viral translation was specific and that the central domain of eIF4G was sufficient for viral translation, we examined the ability of the 4GM domain alone to complement eIF4G siRNA-treated cells. Fluorescence-based Western blotting and quantification of multiple independent experiments confirmed that the expression of the 4GM domain of eIF4G was sufficient to restore viral translation in eIF4G siRNA-transfected cells (Fig. 10).

**DISCUSSION**

Until recently the novel mechanism of calicivirus genome translation has been poorly characterized, largely due to the inability to culture many members of the virus family in immortalized cells. FCV, a member of the *Vesivirus* genus, has until very recently provided the only tractable experimental system with which to characterize this novel mechanism of calicivirus VPg-primed viral genome translation. However, with the discovery of a cultivatable member of the *Norovirus* genus, namely MNV, it has now been possible to use genetic and biochemical assays to examine the role of protein-protein interactions in the norovirus life cycle. Our previous work on norovirus translation using MNV as a model system demonstrated a direct interaction with eIF4E, which at least in vitro was not required for viral translation, but also implicated a functional role for the helicase eIF4A in viral translation initiation (25, 26). In agreement with other earlier observations from our laboratory (25), we have now used a proteomics approach to observe that the MNV VPg protein can be used to affinity purify the entire eIF4F complex from cells, including eIF4G. In addition to eIF4F, we identified PABP, components of the eIF3 complex, and a number of cellular proteins previously implicated in protein synthe-
troscopy and found that the MNV VPg protein consists of a domain of VPg using nuclear magnetic resonance (NMR) spectroscopy. The current study would suggest that VPg is likely to make multiple contacts with the eIF4F complex, including interactions with eIF4G independently. Our previous data and that presented in (Fig. 5) would suggest that either the mutation blocks the interaction of eIF4G on the RNA helicase eIF4A (44). Therefore, recruitment of eIF4E to the 5′ end of norovirus RNA would be predicted to enhance the helicase activity of the eIF4A that has formed a complex with the VPg-associated eIF4G. The interaction of VPg with eIF4E may also play a role in regulating eIF4E phosphorylation, as eIF4E is phosphorylated with MNK1 via an interaction with the eIF4G scaffold protein (45). Phosphorylated forms of eIF4E also play a role in the translation of a subset of cellular mRNAs, including those involved in the response to viral infection (46). Our ongoing studies have suggested that the VPg-eIF4E interaction may simply function to recruit eIF4E to facilitate the formation of polysome-associated phosphorylated eIF4E. The role of the VPg-eIF4E interaction is, therefore, most likely involved in controlling the host response to infection, but further studies are required.

eIF4G is also known to be phosphorylated (47), although the physiological relevance of eIF4G phosphorylation has become apparent only recently; data have highlighted that phosphorylation of eIF4G by protein kinase Ca (PKCa) increases an interaction with MNK1 (48). Independent observations indicate that eIF4G phosphorylation is stimulated during norovirus replication in cell culture and that this phosphorylated form of eIF4G is associated with MNV VPg in infected cells (29). The impact and functional relevance of initiation factor phosphorylation on the norovirus life cycle is currently under study.

The novel mechanism of VPg-primed translation initiation is not unique to caliciviruses and has been widely characterized in a number of plant viruses where it contributes to susceptibility to infection (for review, see Ref. 27). In the case of the potyvirus tobacco etch virus, VPg interacts with both eIF4E and eIFiso4E, increasing the affinity of the eIF4 complex for tobacco etch virus RNA (49). The effect of the norovirus VPg-eIF4G interaction on the affinity of the complex for the norovirus RNA genome is as yet unknown, but it is also possible that similar allosteric effects of the VPg-eIF4G interaction on other compo-

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components of the translation initiation factor complex contribute to the stabilization of the initiation factor complex and promote ribosome recruitment.

Overall our work has provided additional insight into the novel mechanism of VPg-dependent viral translation initiation. Given the key role of translation initiation in the life cycle of noroviruses, we would propose that the interaction of VPg with eIF4G might provide a suitable therapeutic target for these pathogens.

Acknowledgment—We particularly thank Peter Cherepanov (London Research Institute) for helpful suggestions during the early stages of the project.

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