Noroviruses subvert the core stress granule component G3BP1 to promote viral VPg-dependent translation.

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Abstract (148)

Knowledge of the host factors required for norovirus replication has been hindered by the challenges associated with culturing human noroviruses. We have combined proteomic analysis of the viral translation and replication complexes with a CRISPR screen, to identify host factors required for norovirus infection. The core stress granule component G3BP1 was identified as a host factor essential for efficient human and murine norovirus infection, demonstrating a conserved function across the Norovirus genus. Furthermore, we show that G3BP1 functions in the novel paradigm of viral VPg-dependent translation initiation, contributing to the assembly of translation complexes on the VPg-linked viral positive sense RNA genome by facilitating 40S recruitment. Our data suggest that G3BP1 functions by providing viral RNA a competitive advantage over capped cellular RNAs, uncovering a novel function for G3BP1 in the life cycle of positive sense RNA viruses and identifying the first host factor with pan-norovirus pro-viral activity.

Keywords (10)
Introduction:

Positive sense RNA viruses rely heavily on host cell factors for all aspects of their life cycle. They replicate on host derived membranous vesicles that are induced following viral infection, the formation of which requires the activity of key membrane bound viral enzymes (Altan-Bonnet, 2017). Within the membrane bound viral replication complex, translation of the viral genome and the synthesis of new viral RNA occurs in a highly coordinated process. Positive sense RNA viruses have evolved novel gene expression mechanisms that enable them to overcome the genome size limitations that accompany error-prone replication and which might restrict their overall coding capacity (Firth and Brierley, 2012). In addition, viral modification of the host cell translation machinery often provides a competitive advantage allowing for the efficient translation of viral RNA in an environment where competing cellular RNAs are in abundance (McCormick and Khaperskyy, 2017). This ability to compete with cellular RNAs is particularly important for the initiation of infection where the incoming viral genome may be present at only a single copy per cell.

We have previously described a novel paradigm of viral translation that relies on the interaction of host translation initiation factors with a virus-encoded protein (VPg), covalently linked to the 5’ end of the genome of members of the Caliciviridae family of positive sense RNA viruses (Chaudhry et al., 2006; Chung et al., 2014; Goodfellow et al., 2005; Hosmillo et al., 2014; Leen et al., 2016). Unlike the 22-amino acid VPg peptides from picornaviruses, the VPg protein linked to the genomes of caliciviruses is significantly larger and is essential for the translation of viral RNA and viral RNA infectivity (Goodfellow, 2011).
Human noroviruses (HuNoV) and sapoviruses (HuSaV) are enteropathogenic members of the *Caliciviridae* family of positive sense RNA viruses, and together cause >20% of all cases of gastroenteritis (GE). They are also a significant cause of morbidity and mortality in the immunocompromised; individuals with genetic immune-deficiencies, cancer patients undergoing treatment and transplant recipients often experience chronic norovirus infections lasting months to years (van Beek et al., 2016). The economic impact of HuNoV is estimated to be at least ~$4.2 billion in direct health care costs, with wider societal costs of ~$60 billion (Bartsch et al., 2016). Despite their socioeconomic impact, we have, until very recently lacked a detailed understanding of much of the norovirus life cycle and many significant questions remain unanswered. HuNoV replicons (Chang et al., 2006), a murine norovirus that replicates in cell culture (Karst et al., 2003; Wobus et al., 2004) and the recent B cell (Jones et al., 2014), stem-cell derived organoid (Ettayebi et al., 2016) and zebrafish larvae infection models (Van et al., 2019), have all provided invaluable tools to dissect the norovirus life cycle. However, due to the technical limitations associated with many of these experimental systems, in comparison to other positive sense RNA viruses, our knowledge of the intracellular life of noroviruses is significantly lacking (reviewed in Thorne and Goodfellow, 2014).

In the current study, we have combined three independent unbiased approaches to identify host factors involved in the norovirus life cycle. Combining experimental systems that incorporated both murine and human noroviruses, allowed the identification of cellular factors for which the function is likely conserved across the *Norovirus* genus. By combining three complimentary approaches, we identify the...
host protein G3BP1 as a critical host factor required for norovirus VPg-dependent translation, identifying a new role for G3BP1 in virus-specific translation.

Results:

Comparative analysis of the norovirus translation initiation complex.

The MNV and the prototype HuNoV Norwalk virus (NV) VPg proteins contain a highly conserved C-terminal domain (Fig 1A) which we have previously shown to be necessary and sufficient for binding to the translation initiation factor eIF4G (Chung et al., 2014; Leen et al., 2016). Using affinity purification on m7-GTP sepharose, we confirmed that the NV VPg protein, as produced during authentic virus replication in a NV replicon bearing cell line, interacts with the cap-binding complex eIF4F (Fig 1B). Components of the eIF4F complex, namely the eIF4E cap-binding protein, the eIF4A helicase and the eIF4GI scaffold protein, along with poly-A binding protein (PABP) and eIF3 subunits, were readily purified on m7-GTP sepharose, whereas GAPDH was not. In NV-replicon containing cells, mature VPg was also enriched on m7-GTP sepharose but the NS3 protein, known to have RNA binding and helicase activity (Li et al., 2018), was not. Furthermore, we demonstrated that transfection of GFP-tagged versions of either the MNV or NV VPg proteins into 293T cells allowed for the affinity purification of eIF4F components and that mutations in the eIF4G binding domain of VPg reduced this association (Fig 1C).

We next used quantitative mass spectrometry of the affinity purified complexes isolated from cells transfected with the GFP-Tagged VPg proteins to identify host factors specifically enriched on the norovirus VPg protein (Fig 1D, Fig S1 and Table
Most of the proteins identified were components of the host cell translation complex including ribosomal proteins, translation initiation factors and host RNA binding proteins. These data agree with but significantly extend our previous observations using a less sensitive multi-step affinity purification approach to characterise host factors associated with the MNV VPg protein only (Chaudhry et al., 2006; Chung et al., 2014). In addition, we identified hnRNPA1 which we have previously shown to act in norovirus genome circularization (López-Manríquez et al., 2013). YBX1, DDX3 and several other proteins that we have previously found to interact with the 5′ end of the viral RNA (Vashist et al., 2012b) were also enriched on VPg (Fig S1). To validate a select number of these interactions and to assess whether the interaction of VPg with eIF4G is required for their association with VPg, we performed western blot analysis of complexes purified from cells transfected with either the WT or eIF4G-binding mutants (Fig 1E). Except for YBX1, the association of all proteins tested were reduced by the introduction of eIF4G-binding site mutations into the MNV VPg protein. Together, these data extend our previous observations and confirm that the norovirus VPg proteins interact with a complex network of host factors, many of which have been implicated in the host cell translation initiation process.

**Determination of the norovirus replication complex proteome.**

To further identify the components of the norovirus translation and replication complex, as formed during authentic viral replication in highly permissive cells, we utilised two recombinant infectious MNV strains that carried epitope purification tags within the NS1/2 or NS4 proteins (McCune et al., 2017) (Fig 2A). The insertion
positions were previously identified using a transposon based mutagenesis screen as sites that tolerate insertions, without compromising virus viability (Thorne et al., 2012). Our approach was somewhat analogous to that recently published for coronaviruses (V’kovski et al., 2019) but instead used stable isotope labelling of permissive cells and the FLAG affinity purification tag rather than proximity labelling. Unlabelled or stable isotope labelled highly permissive BV-2 microglial cells were infected with either wild type MNV or the equivalent virus carrying the FLAG epitope purification tag in either NS1/2 or NS4, and the viral replication complex was purified. The experiment was performed three times by swapping the labelled derivatives of arginine and lysine as described in the materials and methods. Silver stain of the purified complexes confirmed the presence of the bait proteins, with both the uncleaved and cleaved forms of NS1/2 and NS2 being highly enriched (Fig 2B). As expected, complexes purified from NS1/2-Flag virus infected cells co-purified untagged NS4 and vice versa (Fig 2B), as we have previously shown these proteins to interact to form a complex (Thorne et al., 2012). Western blot analysis of the purified complexes confirmed that viral non-structural and structural proteins were specifically enriched in the purified complexes, including NS5 (VPg)-containing precursors (Fig 2C). We noted that anti-NS4 monoclonal antibody was unable to detect protein in the extracts prior to enrichment, which most likely reflected the limited sensitivity of the antibody. Quantitative mass spectrometry of the purified complexes allowed the identified of viral and cellular proteins enriched in the complex (Fig 2D and Table S2).

As expected, all viral proteins, including the VF1 protein product of ORF4, an innate immune antagonist (Bailey et al., 2011), were enriched in the viral replication complex. There was a significant correlation between the relative enrichment of
proteins identified using NS1/2 and NS4 (Spearman correlation of 0.8832), fitting
with our prior knowledge that both proteins form a complex during viral replication
(Thorne et al., 2012). Ontology analysis indicated that proteins involved in vesicle
transport and fatty acid metabolism were significantly enriched (Fig S2 and Table
S3), fitting with previous observations that the viral replication complex is associated
with cytoplasmic membranous structures (Cotton et al., 2017; Hyde and Mackenzie,
2010; Hyde et al., 2009). Several host proteins previously identified in a variety of
biochemical and genetic screens were enriched (Fig S2 and Table S3) providing
additional confidence that the approach identified biologically relevant interactions.
We noted that the VapA and the paralogue VapB, which we have recently identified
as binding to the NS1/2 protein (McCune et al., 2017), were both highly enriched.
Comparison with the data obtained using VPg as a bait protein (Fig 1) showed some
degree of overlap, however it is worth noting that most of the factors that were
identified using VPg were enriched by >2 fold using only the NS1/2 tagged virus and
not the NS4 tagged virus (Fig S2). One of the exceptions to this was the core stress
granule protein G3BP1, which was enriched by both MNV and NV VPg proteins, and
was also enriched in complexes purified using both NS1/2 and NS4-tagged viruses.

Identification of host factors required for norovirus infection using a CRISPR-
knockout screen.

A high density CRISPR library screen was undertaken to identify genes that
contribute to the norovirus life cycle. The Brie library (Doench et al., 2016) was
selected due to the reduced off-target effects relative to previously described
CRISPR libraries used for norovirus studies (Haga et al., 2016; Orchard et al., 2016).
In addition, to minimise the impact of gRNAs that may have deleterious effects on long term cell viability and to increase our ability to detect genes that may be important, but not essential, for norovirus-induced cell death, the infection was reduced to 24 hours as compared to 2-10 days post infection in previous studies. BV-2-Cas9 expressing cells were infected with lentiviruses carrying the Brie gRNA library carrying 78,637 independent guide RNAs to 19,674 genes (Doench et al., 2016). The transduced cells were then infected with two MNV strains, CW3 and CR6, which cause acute and persistent infections in immunocompetent mice respectively (Nice et al., 2012; Thackray et al., 2007), and guide RNA abundance compared to mock infected cells at 24 hours post infection as illustrated in Fig 3A. Genes that were enriched by STARS analysis following MNV infection represent putative pro-viral factors which when disrupted result in slower cell death, whereas those with a negative STARS value represent putative anti-viral factors where virus-induced cell death has occurred quicker, resulting in their underrepresentation in the final pool of cells. MNV-CR6 infection resulted in 212 genes being enriched and 43 being negatively selected (Fig 3B), whereas for MNV-CW3 279 and 19 genes were positively and negatively selected respectively (Fig 3B). In most cases, there was a clear correlation between the datasets obtained using either strain (Fig 3C). STARS analysis was used to ranks genes with positive and negative values (Table S4). In both screens, the MNV receptor Cd300lf was the most highly positively selected gene identified, in agreement with previous reports (Haga et al., 2016; Orchard et al., 2016). The second most highly enriched gene was G3BP1, a gene also identified in one of the two previous CRISPR screens performed on norovirus infected cells (Orchard et al., 2016).
A comparison of the data obtained from all three approaches allowed us to identify several host proteins that were common to all screens (Table S5). G3BP1, the core stress granule component was identified in all three screens as a potential host factor essential for norovirus infection. G3BP1 was found to be associated with the MNV and NV VPg proteins (Fig 1D), enriched in viral replication complexes purified using either NS1/2 or NS4 flag tagged viruses (Fig 2D) and identified in a CRISPR screen using two different MNV strains as a putative pro-viral factor involved in the norovirus life cycle (Fig 3C).

**G3BP1 is essential for murine norovirus replication**

To validate the importance of G3BP1 in the norovirus life cycle we generated G3BP1 deficient BV-2 cell lines (Fig 4A) and examined the impact of G3BP1 ablation on MNV infection. Western blotting confirmed the loss of G3BP1 in the three lines tested and we noted that at in some cases, a concomitant increase in G3BP2 expression was observed as has been previously noted (Kedersha et al., 2016). A clear defect was observed in the ability to replicate to produce infectious virus in three independently selected $\Delta$G3BP1 cell lines (Fig 4B). This effect was mirrored by an inability to induce cytopathic effect leading to virus-induced cell death (Fig 4C). In contrast, the ability of encephalomyocarditis virus (EMCV) to infect and cause cell death was unaffected by the deletion of G3BP1(Fig 4C). These data confirm that cells lacking G3BP1 are highly resistant to norovirus infection.

**G3BP1 is essential for human norovirus replication in cell culture**
To determine if the G3BP1 was also essential for HuNoV, we examined the impact of loss of G3BP1 on human norovirus replication in cell culture using the Norwalk virus replicon. To establish the experimental system, we first confirmed that the presence of VPg on the 5' end of the Norwalk RNA was essential for the replication of the replicon RNA and for the capacity to form G418 resistant colonies. Transfection of replicon RNA, purified from replicon containing cells, into BHK cells readily resulted in the formation of antibiotic resistant cell colonies (Fig 5A). In contrast, RNA that was proteinase K treated prior to transfection was unable to produce replicon containing colonies. Transfection of replicon RNA into wild type U2OS osteosarcoma cells allowed the formation of replicon-containing colonies, although the efficiency of formation was significantly less than that seen in BHK cells (Fig 5B). CRISPR modified U2OS cells that lacked G3BP1 (Kedersha et al., 2016) were unable to support NV replication, as evident by the lack of antibiotic resistant colonies (Fig 5B). To further examine the role of G3BP1 in human Norwalk virus replication, WT or G3BP1 deficient U2OS cells were transfected with NV replicon VPg-linked RNA, and RNA synthesis monitored overtime following the addition of G418. While a significant increase in NV viral RNA levels was seen in WT U2OS cells, those lacking G3BP1 were completely unable to support NV RNA synthesis (Fig 5C). These data indicate that like for MNV, G3BP1 is essential for human Norwalk virus replication.

The RNA-binding domain of G3BP1 is required for its function in the norovirus life cycle.
To confirm the role of G3BP1 in the norovirus life cycle we examined the ability of full length and truncated versions of G3BP1 to restore norovirus replication in G3BP1 knockout cells. A mouse BV-2 G3BP1 knockout cell line was complemented with either full length G3BP1 or variants lacking the RGG or both the RGG and RRM binding domains (Fig 6A) and the impact on viral replication assessed. Complementation with full length murine G3BP1 restored the ability of MNV to induce cell death (Fig 6B) and to produce infectious virus (Fig 6C) back to near wild type levels. In contrast, complementation with a variant carrying a deletion of the RGG domain resulted in limited complementation, and deletion of both the RGG and RRM domains together resulted in complete loss of complementation capacity (Fig 6B&C). These data confirm that the RNA binding domains of G3BP1 are essential for its function in the norovirus life cycle.

G3BP1 is required for a post entry step in the norovirus life cycle.

To further define the role of G3BP1 in the norovirus life cycle we examined whether G3BP1 functioned at the level of viral entry or post-entry. We therefore bypassed the entry phase of the infection process and transfected viral VPg-linked RNA into WT and two independently generated BV-2 ΔG3BP1 cell lines and examined the impact on norovirus replication. Transfection of MNV viral VPg-linked RNA into WT cells resulted in high yields of infectious virus (Fig 7A) and viral proteins (Fig 7C). The levels of infectivity obtained following transfection of ΔG3BP1 cell lines with MNV viral RNA was comparable to that obtained in WT cells in the presence of the nucleoside analogue 2′-C-methylcytidin (2CMC), a known inhibitor of the norovirus RNA polymerase (Rocha-Pereira et al., 2012; 2013) (Fig 7A). No viral proteins were
detected in either of the ΔG3BP1 cell lines suggesting a defect at a very early stage in the viral life cycle (Fig 7C). Transfection of VPg-linked RNA into the ΔG3BP1 cell lines reconstituted with WT G3BP1 restored the ability to produce infectious virus (Fig 7B) and the production of viral proteins (Fig 7D). A minor increase in viral infectivity was observed in the ΔG3BP1 cell line reconstituted with the ΔRGG construct producing viral titres that were higher than those obtained from the WT complemented line in the presence of 2CMC, suggesting low levels of viral replication (Fig 7B). However, the levels of viral proteins produced in this line was below the limit of detection by western blot (Fig 7D). These data confirm that G3BP1 is required for a post entry stage of the norovirus life cycle and that in the absence of G3BP1 only residual norovirus replication is observed.

G3BP1 is required for viral negative sense RNA synthesis

To define the precise role of G3BP1 in the early stages of the virus life cycle, we used strand-specific RT-qPCR to quantify the levels of viral positive and negative sense RNA in WT and ΔG3BP1 cell lines following infection with MNV. As a control, 2CMC was included following virus inoculation as illustrated in the experimental time line (Fig 8A). The production of viral positive sense RNA was reduced to background levels in the absence of G3BP1, comparable to levels observed when the 2CMC was present during the infection (Fig 8B). Viral negative sense RNA synthesis was also reduced to below the detection limit of the assay in ΔG3BP1 cell lines (Fig 8C). Surprisingly, we were able to detect an ~5 fold increase in viral negative sense RNA production at 6 hours post infection of WT cells in the presence of 2CMC, which,
given that 2CMC was added after the inoculation phase (Fig 8B), likely represents
the first round of viral negative sense RNA synthesis, confirming the sensitivity of the
assay. Addition of 2CMC during the inoculation phase reduced this background
levels (data not shown).

Similar results were obtained following transfection of viral RNA into cells to bypass
the entry phase; viral positive and negative sense RNA synthesis was near (or
below) the sensitivity of the assay following transfection of viral VPg-linked RNA into
two independent ΔG3BP1 cell lines (Fig 8D & E). Complementation with WT G3BP1,
but not the mutant forms lacking the RNA binding domains, also restored viral
positive and negative sense RNA synthesis (Fig 8F & G). We did not detect viral
positive or negative sense RNAs in the ΔRGG compelemented cell line, despite the
presence of low levels of viral infectivity (Fig 7B). This discrepancy likely reflects the
relative sensitivities of the assays and the nature of the strand specific qPCR assay
which requires low levels of RNA input to maintain strand specificity. Together these
data suggest that the function of G3BP1 is prior to, or at the level of viral negative
sense RNA synthesis, with the most logical steps being either viral RNA translation
or the formation of viral replication complexes.

G3BP1 is required for the association of VPg with 40S ribosomal subunits.

We have previously shown that norovirus VPg interacts with eIF4G to recruit
ribosomal subunits and direct viral translation (Chaudhry et al., 2006; Chung et al.,
2014). The interaction between VPg and eIF4G occurs via a direct interaction
between the highly conserved C-terminal region in VPg and the central HEAT
domain of eIF4G (Leen et al., 2016) and does not require any additional cellular cofactors, at least in vitro. The interaction between the eIF4G HEAT domain and the eIF3 complex plays a central role in the recruitment of the 40S ribosomal subunit for translation initiation (Marcotrigiano:2001uq; Kumar et al., 2016; Villa et al., 2013). Our proteomics analysis also confirms that the norovirus VPg proteins form a complex that contains multiple components of the 40S subunit (Fig 1D) and it has been established previously that G3BP1 associates with 40S subunits (Kedersha et al., 2016). To assess a potential role for G3BP1 in the formation of VPg-driven translation complexes in cells, we examined the ability of GFP tagged version of MNV VPg to pull down 40S subunits in the presence and absence of G3BP1. GFP-tagged WT MNV VPg was readily able to pull down eIF4G, G3BP1 and RpS6, a component of the 40S subunit (Fig 9A). However, in the absence of G3BP1, the ability to pull down RpS6 was lost (Fig 9A). Furthermore, we found that disruption of the VPg-eIF4G interaction by the introduction of the F123A mutation into the eIF4G binding domain, also significantly reduced the ability to pull down RpS6 (Fig 9B). These data suggest that the interaction of VPg with eIF4G is important for complex formation with ribosomal proteins and that that G3BP1 contributes in some manner to the formation of this complex.

G3BP1 is require for efficient polysome loading of norovirus VPg-linked RNA

To assess the impact of G3BP on the selective translation of viral VPg-linked RNA following viral infection, we evaluated the impact of loss of G3BP1 on the recruitment of viral RNA to polysomes under conditions where viral RNA synthesis was inhibited, namely in the presence of 2CMC. This approach enabled us to assess only the
capacity of the incoming parental viral RNA to assemble into translationally active complexes, a stage often referred to as the “maiden round” of RNA virus genome translation. To this aim, cells were infected with MNV in the presence of 2CMC and polysomes profiling on extracts prepared from cells at 4 and 9 hours post infection performed (Fig 10A). Quantification of the viral RNA levels in cells in the presence of 2CMC confirmed that the absence of G3BP1 has no impact on the overall levels present at the time points examined (data not shown). We noted that even in the presence of 2CMC, which inhibits viral RNA synthesis, there was a small but measurable increase in free 80S ribosomes over time in WT cells but not in cells lacking G3BP1 (Fig 10A). We have previously found that MNV infection results in translation shut off and that this effect is at least partuialkly due to the activity of the NS6 protease (Emmott et al., 2017). The fact we observed 80S accumulation in WT cells, even in the absence of viral RNA synthesis, but not in cells lacking G3BP1, indirectly lead us to suspect that translation of viral RNA had occurred in WT cells, but was much less efficient in cells lacking G3BP1. Further analyses indicated that while most ribosome-associated norovirus RNA in WT cells was found in polysomes containing fractions, less viral RNA was found in ribosome-containing fractions (1-12 in Fig 10A & B) in the absence of G3BP1 cells and, in comparison to WT cells, very little viral RNA was found in fractions containing polysomes (Fig 10B). Extending the fractionation to include the free RNA and ribonucleoprotein complexes at the top of each gradient confirmed that in the absence of G3BP1 norovirus RNA is less efficient at assembling into polysomal fractions, suggesting a defect at the level of viral protein synthesis (Fig 10C). Together these data support the hypothesis that G3BP1 functions to promote the translation of norovirus VPg-linked RNA, by facilitating the association with 40S subunits.
G3BP1 is required for efficient norovirus VPg-dependent translation in the presence of cellular capped RNAs.

To further examine a potential role of G3BP1 in norovirus VPg-dependent translation, cytoplasmic translationally competent extracts were prepared from WT and ΔG3BP1 cell lines and the translation of highly purified viral VPg-linked viral RNA (Fig S11A&B) examined. Cap-dependent and cricket paralysis virus IRES (CrPV)-dependent translation were comparable in nuclease treated extracts prepared from WT and ΔG3BP1 cell lines (Fig 11A), whereas norovirus VPg-dependent translation was reduced (Fig 11B). Quantification of multiple experiments indicated that translation in nuclease treated extracts was on average reduced by ~40-50% because of G3BP1 ablation (Fig 11C). A similar reduction in in vitro translation was observed across multiple time points (Fig S11C).

The ability of VPg-linked norovirus RNA to be translated in the presence of increasing amounts of total cellular RNAs was then examined to assess the relative role of G3BP1 under conditions where cellular RNA are present. Total RNA isolated from uninfected cells was titrated into the nuclease treated extracts and the impact norovirus VPg-dependent translation examined (Fig 11D). We found that in the presence of cellular RNAs, the translation of norovirus VPg-linked RNA in extracts from ΔG3BP1 cells is reduced by up to 80% in comparison to extracts from WT cells. In agreement with our data using polysome profiling, these data suggest that G3BP1 functions to provide a competitive advantage for norovirus VPg-linked RNA under conditions where cellular RNAs are present.
Discussion

In this study, we have used a combination of biochemical and genetic approaches to identify host factors involved in the norovirus life cycle. Our combined approaches resulted in the identification of the core stress granule component G3BP1 as a host protein critical for the replication of both murine and human noroviruses in cell culture. Furthermore, we determined that G3BP1 plays a key role in the processes of norovirus VPg-dependent protein synthesis, uncovering a new function for G3BP1 in facilitating RNA virus genome translation.

The orthogonal approaches used in the current study provide an unprecedented insight into the identity of host factors with potential roles in the norovirus life cycle. The detailed proteomic analysis of the viral replication and translation complexes formed during MNV infection (Fig 2) resulted in the identification of several host factors with previously identified roles in the MNV life cycle. We focused our efforts on G3BP1 as it was identified in all three approaches and was also identified in a CRISPR screen published during this study (Orchard et al., 2016). Furthermore, we have previously shown that feline calicivirus (FCV), a relative of noroviruses within the Vesivirus genus, cleaves G3BP1 to inhibit stress granule formation (Humoud et al., 2016). In contrast, MNV infection does not result in G3BP1 cleavage and instead forms cytoplasmic foci the composition of which is distinct from canonical stress granules (Brocard et al., 2018).

G3BP1 is one member of a group of G3BP proteins (Ras-GTPase-activating protein (SH3 domain)-binding proteins), referred to as Rasputin in insects, that possess
RNA binding activity and have multiple cellular functions including the regulation of RNA stability and translation in response to stress. Originally identified as a protein that interacted with Ras-GTPase activating protein (RasGAP), more than two decades of research have significantly expanded our knowledge of the multifunctional role in cellular processes. It is now well accepted that G3BPs play a role in cancer cell survival, cancer metastasis and invasion, processing of specific miRNAs and stress granule formation (Reviewed in (Alam and Kennedy, 2019). Stress granules are dynamic cytoplasmic ribonucleoprotein complexes that form rapidly under stress conditions and within which cellular RNAs are stored in stalled translation complexes (Protter and Parker, 2016). In the context of viral infection, numerous studies have suggested that many, if not all, viruses must interact in some manner with stress granules as there is growing evidence that the formation of cytoplasmic stress granules is part of the antiviral defence mechanism (Reviewed in (McCormick and Khaperskyy, 2017). Some viruses interact with stress granules to promote viral replication (Cristea et al., 2010; Kim et al., 2016; Panas et al., 2014; 2012) whereas some do so to counteract the inhibitory effect of stress granules on the translation of viral RNA (Panas et al., 2015; White et al., 2007).

Our data suggests that G3BP1 plays a key role in promoting the translation of norovirus VPg-linked viral RNA. Positive sense RNA viruses have evolved mechanisms to ensure the efficient translation of their viral genomic RNAs in the presence of high concentrations of competing cellular RNAs. These mechanisms include the use of internal ribosome entry site elements (IRES), modified cap-dependent mechanisms (Firth and Brierley, 2012; Jaafar and Kieft, 2019) and the ability to target the host cell translation machinery to generate an environment where viral RNA translation is favoured over cellular capped RNAs (Walsh et al., 2013).
G3BP1 is known to associate primarily with free 40S subunits and not 80S monosomes (Kedersha et al., 2016). Our data supports a hypothesis whereby the association of G3BP1 with 40S ribosomal subunits provides a selective advantage for norovirus VPg-dependent translation, thereby uncovering a new function in virus-specific translation. The mechanism by which G3BP1 contributes to this process has yet to be fully explored but our data supports the hypothesis that G3BP1 directly or indirectly promotes the recruitment of 40S ribosomal subunits to VPg-driven translation complexes. The RGG motif of G3BP1 is known to be essential for the association between G3BP1 and 40S subunits as well as the ability to from stress granules, whereas data would suggest that the RRM may play a regulatory role (Kedersha et al., 2016). These domains were also required for the function of G3BP1 in the norovirus life cycle (Fig 6) confirming that the G3BP1 association with 40S is important for its role in promoting norovirus VPg-dependent translation. Importantly, RGG domains are known to have many functions (Thandapani et al., 2013) and therefore in the context of G3BP1 function in the norovirus life cycle, may also contribute to unknown interactions that promote norovirus translation. Previous work on alphaviruses have shown that G3BP1 is sequestered by binding to the nsP3 protein (Panas et al., 2012; 2014; 2015). Furthermore, this interaction occurs via an FGDF motif also found in other viral proteins including the ICP8 protein of herpes simplex virus (Panas et al., 2015). While the MNV VPg protein has a similar motif FGDGF (Fig 1A), this motif is not conserved in the GI Norwalk virus VPg protein. Therefore our data suggest that the interaction of VPg with G3BP1 is not direct, fitting with our observation that this interaction is reduced by mutations in the eIF4G binding domain (Fig 1A and Fig 9B.) While our data fit with a primary role for G3BP1 in norovirus translation, we are unable to exclude the possibility that G3BP1 plays
other roles in the viral life cycle. Recent studies have confirmed that G3BP1 is
enriched at sites of viral RNA synthesis (Brocard et al., 2018; Fritzlar et al., 2019) so
it is possible that G3BP1 makes multiple contacts between the 40S subunit and viral
RNA genome directly.

The technical challenges associated with studying human norovirus replication in cell
culture have limited the experimental approaches we could use to validate the role of
G3BP1 in human norovirus translation. However, our results have clearly
demonstrated that in the absence of G3BP1, human Norwalk virus is unable to
replicate or form replicon-containing colonies. The presence of G3BP1 in the NV
VPg-containing complexes again fits with our hypothesis that G3BP1 plays a role in
promoting viral VPg-dependent protein synthesis.

We have previously found that norovirus infection leads to a translation bias whereby
acellular mRNAs induced in response to infection are inefficiently translated (Emmott
et al., 2017). Our data suggested that this modification of host cell translation was at
least partially driven by the ability of the viral NS6 protease to cleave PABP and the
induction of apoptosis which results in cleavage of cellular translation initiation
factors (Emmott et al., 2017). Furthermore, we have more recently shown that the
ability of the protease to cleave PABP and other substrates is controlled by
polyprotein processing and interactions with other viral proteins (Emmott et al.,
2019). Recent work would agree with our observations and confirms that the
translational bias is not driven by phosphorylation of eIF2α and that activation of
GCN2 leads to the phosphorylation of eIF2α (Brocard et al., 2018). We note however
that others have suggested that NS3 may contribute to translational shut off (Fritzlar
et al., 2019), with the caveat that this observation was made outside of the context of
infected cells and using overexpressed tagged proteins. We suspect that noroviruses use multiple mechanisms that work co-operatively to enable the control of host gene expression and the subsequent translation of the cellular mRNAs. The relative contribution of these processes in any given cell type may also differ depending on the degree to which the cells can sense and respond to viral infection through the induction of innate and apoptotic responses.

The observation that many of the factors enriched using the VPg protein were only enriched on complexes purified with NS1/2 tagged infectious MNV, could suggest that the viral proteins present in the viral translation complex are distinct from those present in complexes active for viral RNA synthesis. However, we cannot formally rule out other possible explanations including the possibility that the specific enrichment of translation factors on NS1/2 occurs because NS1/2 is the first protein to be translated from ORF1, therefore unprocessed NS1/2 at the N-terminus of the ORF1 polyprotein being actively translated could function as an anchor, facilitating the enrichment of ribosomes and the associated factors. In addition, we have previously seen that VPg-containing precursors may bind the translation initiation factor eIF4G less well (Leen et al., 2016), which could prevent some VPg (NS5) containing precursors associating with translation initiation complexes.

In conclusion, our data adds significantly to the growing body of literature on the role of G3BP proteins in the life cycle of viruses and further extends the functional roles of G3BP1 to include the promotion of viral translation processes. We identify G3BP1 as a host protein that has a critical role in the life cycle of murine and human noroviruses, identifying the first cellular pro-viral protein with pan-norovirus activity. Furthermore, given the apparent importance of G3BP1 to an early stage of the
norovirus life cycle, this work suggests that targeting G3BP1 may hold future therapeutic potential.

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Figure 1. The norovirus VPg proteins interacts with ribosome associated translation initiation factors. A) Amino acid sequence alignment of the GV murine norovirus VPg sequences with VPg from representative human noroviruses from GI Norwalk virus (NV), GII, and GIV. The position of the site of RNA linkage to the highly conserved tyrosine residue is highlighted in green. The eIF4G binding motif is boxed and the position of the C-terminal single amino acid change known to interfere with eIF4G binding highlighted in orange. B) m7-GTP sepharose was used to affinity purify eIF4F containing complexes from either wild-type BHK cells (BHK) or BHK cells containing the Norwalk virus (NV) replicon (BHK-NV). Samples of the lysate (L) or the affinity purified complexes (m7) were separated by SDS-PAGE then analysed by western blot for the indicate proteins. Molecular mass shown on the left of the gels represent the positions of molecular weight markers. C) GFP fusion proteins to either the wild type (WT) or C-terminal eIF4G binding domain mutants of the MNV and NV VPg proteins (F123A, F137A) were transfected into human 293T cells and subjected to immunoaffinity purification using anti-GFP. Samples of the input lysates (Input) and the purified complexes (GFP-IP) were then separated by SDS-PAGE and analysed by western blot analysis for the indicated proteins. Mock transfected cells served as a specificity control. The approximate expected molecular mass of each protein is shown to the left. D) Quantitative proteomics was used as described in the text to identify host factors that were affinity purified following transfection of GFP-tagged derivative of either the NV or MNV VPg proteins. Proteins specifically enriched in comparison to the GFP control are shown. Data visualisation was performed using Cystoscape (Shannon et al., 2003). E) Western blot analysis of cell
extracts (Input) or immunoprecipitated (GFP IP) complexes isolated from cells transfected as described in panel C. For clarity, the molecular masses shown in this panel refer to the expected mass of the protein being examined.

Supplementary Figure 1: Host factors binding to the norovirus VPg.
Quantitative proteomics was used as described in the text to identify host factors that were affinity purified following transfection of GFP-tagged derivative of either the NV or MNV VPg proteins. Proteins specifically enriched in comparison to the GFP control are shown in Panels A and B respectively and where protein binding was reduced. Panel C illustrates the proteins previously found to interact with the 5′ or 3′ termini of the MNV genome (Bailey et al., 2010; Vashist et al., 2012b) or to associate with MNV VPg using tandem affinity purification (Chung et al., 2014). Data visualisation was performed using Cystoscape (Shannon et al., 2003)

Figure 2: Proteomic characterisation of the norovirus replication complex using infectious epitope tagged MNV. A) Schematic representation of NS1/2-FLAG and NS4-FLAG viruses contain insertions of nucleotide sequences encoding the FLAG peptide DYKDDDDK (in yellow) in their coding sequences. The NS1/2-FLAG virus FLAG peptide was inserted between 2 of the 3 caspase-3 cleavage sites present in NS1/2 (underlined). B) BV-2 cells labelled with stable derivatives of arginine and lysine were infected with either wild type MNV (WT) or recombinant epitope-tagged MNV as described in the materials and methods. 12 hours post infection samples were lysed, samples pooled and immunoaffinity purifications performed as described in the text. Samples of the cell lysates (Input) and the affinity purified complexes (IP:Flag) were analysed by SDS-PAGE on a 4-12% gradient gel
prior to silver staining. The positions of the NS1/2, NS2 and NS4 proteins is shown.

C) Western blot analysis of lysates purified from cells infected as described in panel B, for various viral proteins, confirming the specific enrichment of viral replicase components. Plot showing detailing overlapping proteins identified in pulldowns from cells infected with FLAG-tagged NS1/2 or NS4 expressing viruses. All MNV proteins were identified in association with NS1/2 and NS4 (light blue) including the viral polymerase NS7, demonstrating enrichment of the MNV replication complex. Proteins previously identified as host factors potentially involved in some aspect of the norovirus life cycle through various biochemical or genetic screens are shown in red. Selected highly enriched proteins are highlighted in black. The NS1/2 binding partner VapA (McCune et al., 2017) and paralog VapB were both enriched by NS1/2 and NS4.

Supplementary Figure 2: Additional analyses of NS1/2 and NS4-associated proteins. MNV proteins highlighted in light blue, and G31 in gold. A) Gene ontology analysis of proteins copurifying with NS1/2 or NS4 (Log2 SILAC ratio >2 for either protein). Proteins in selected, mutually exclusive gene ontology categories were identified using PANTHER overrepresentation analysis and are plotted in different colors. B) A number of factors previously identified as MNV host factors using proteomics or CRISPR approaches (Orchard et al., 2016, Chung et al., 2014, Vashist et al., 2012) copurified in pulldowns of NS1/2 and NS4 (highlighted red). C) Novel putative MNV host factors highly enriched (Log2 SILAC ratio >4 for either protein) by NS1/2 and NS4 are plotted in black. D) Proteins identified in pulldowns of NS1/2 and NS4 which were also identified in pulldowns of MNV VPg are plotted in red. E) Proteins enriched in pulldowns of NS1/2 and NS4 which were also identified
using CRISPR screening as MNV host factors are plotted in red (positive host
factors) and yellow (negative host factors), along with G3bp1 in gold.

Figure 3: CRISPR screen identifies host genes positively and negatively
selected upon MNV infection. A) Schematic overview of the infection CRISPR
screen workflow. B) Volcano plot identifying candidate genes enriched upon MNV-
CW3 (red) or MNV-CR6 (blue); red or blue labelled genes correspond to the top-ten
positive or negatively selected genes ranked by the STARS algorithm.

Figure 4: CRISPR knockout of G3BP1 renders cells non permissive for MNV
replication. A) Western blot analysis of three independent ΔG3BP1 clones for
GAPDH, G3BP1 and G3BP2. B) High multiplicity, single cycle growth curve analysis
of the impact of G3BP1 ablation on MNV replication. BV-2 ΔG3BP1 clone C cells
were infected at a MOI of 10 TCID50/cell, samples were collected at the time points
illustrated, the samples then processed and titrated by TCID50 as described in the
text. The error bars represent standard errors of three biological repeats and the
data are representative of at least three independent experiments. C) Wild type (WT)
or ΔG3BP1 clone C BV-2 cells were plated in a 96 well plate and subsequently
infected using a serial dilution of either EMCV or MNV. Cells were fixed in
paraformaldehyde and stained with crystal violet 5 days post infection. D) Light
micrographs of WT or ΔG3BP1 cells either mock infected (-) or infected with EMCV
or MNV and visualised 5 days post infection.

Figure 5. G3BP1 is required for human Norwalk virus replication in cell culture.
A) Colony formation ability of human norovirus VPg-linked RNA isolated from BHK-
NV replicon containing cells is dependent on the presence of VPg. NV VPg-linked RNA isolated from BHK-NV cells was either mock treated or treated with proteinase K prior to transfection into BHK cells. Wells were transfected with either 1.5 μg or 0.75 μg of total RNA isolated from NV replicon containing BHK cells. Following 2 weeks of antibiotic selection with G418, surviving replicon containing colonies were fixed and stained with crystal violet in paraformaldehyde. B) NV replicon colony forming assay in WT and G3BP1−/− U2OS cells performed as described in panel A, with the exception that colonies were stained 12 days post transfection. C) Quantification of NV replication in WT or ΔG3BP1 U2OS cells following transfection of viral VPg-linked RNA. Viral RNA was quantified by RT-qPCR following transfection and antibiotic selection. The error bars represent the standard error of three biological repeats and are representative of three independent experiments.

**Figure 6: MNV replication in BV-2 cells requires the RNA binding activity of G3BP1.** Western blot analysis of wild type BV-2 cells (WT) or a ΔG3BP1 cells (clone 1B2) and the respective complemented lines expressing either WT or the various G3BP1 truncations. Cells were lysed prior to separation by 12% SDS-PAGE. B) WT or ΔG3BP1 cells complemented with the indicated constructs were plated in a 96 well plate then infected with a serial dilution of MNV, before being fixed and stained 5 days post infection as described in the text. C) WT or ΔG3BP1 cells complemented with the indicated constructs were infected with MNV at an MOI of 10 TCID50 per cell. After 24 hours the virus yield was determined by TCID50. The error bars represent the standard error of three independent repeats. The data are representative of at least two independently repeated experiments.
**Figure 7:** Loss of G3BP1 results in a defect following transfection of viral VPg-linked RNA into ΔG3BP1 cells. A) The indicated cell lines were transfected with MNV viral RNA and harvested at 9 hours post transfection for TCID50 to assess the virus yield. In some instances, the nucleoside analogue 2CMC was included to inhibit viral replication. The dotted line indicates the limit of detection (LOD) and the error bars represent the standard error from three biological repeats. B) Infectious virus yield from ΔG3BP1 and reconstituted cell lines performed as described in panel A. C) and D) illustrate the accompanying western blots for samples prepared in panel A and B respectively. Samples were prepared at 24 hours post transfection, prior to harvesting, separation by SDS-PAGE on a 4-12% gradient gel prior to western blotting for the indicated proteins.

**Figure 8:** The Lack of G3BP1 results in a failure to produce viral negative sense RNA. The experimental design is illustrated in A. Wild type or ΔG3BP1 (1B2) cells were infected prior to the addition of the nucleoside analogue 2CMC to prevent viral RNA synthesis. Samples were harvest at the indicated time post infection and viral positive (B) and negative sense RNA quantified by stand specific RT-qPCR (C). Error bars represent standard error of three biological repeats. LOD refers to the limit of detection of the assay. D) and E) Viral RNA synthesis following transfection of viral VPg-linked RNA into WT or two ΔG3BP1 cell lines. F) and G) Viral RNA synthesis following transfection of viral VPg-linked RNA into ΔG3BP1 (1B2) complemented with full length G3BP1 or truncated derivatives.

**Figure 9:** G3BP1 is required for the association of VPg with 40S ribosomal subunits. A) GFP-Trap immunoprecipitation of complexes isolated on with GFP
alone or GFP tagged wild type MNV-VPg demonstrating the pull down of eIF4G1, G3BP1 and 40S subunits (Rps6). BVv2 cells were transfected with the relevant constructs, lysates prepared and GFP-Trap pull downs performed as detailed in the text. Samples were separated by SDS-PAGE and western blotted for the proteins as shown. B) Mutations in the eIF4G binding domain ablate the association of VPg with G3BP1 and 40S subunits. GFP-Trap pull downs were performed as described in panel A with the addition of the MNV VPG Fq123A mutation known to reduce the association with eIF4G.

Figure 10: G3BP1 is required for polysome association of viral RNA association. A) Polysome profiles of the ribosome containing fractions from mock or MNV infected wild type (WT) or ΔG3BP1 (1B2) BV-2 cells at 4 and 9 hours post infection (moi 3 TCID50/cell). B) Relative viral RNA levels present in ribosome containing fractions expressed relative to WT infected BV-2 cells. C) Extended gradient fractionation of WT or ΔG3BP1 cells infected with MNV and harvested 9 hours post infection. Viral RNA levels across the gradient are expressed as described in panel B.

Figure 11: G3BP1 is required for efficient norovirus VPg-dependent translation.
Translation competent extracts from ΔG3BP1 cells are fully competent for cap-dependent and CrPV IRES dependent translation (A). B) Translation of MNV VPg-linked viral RNA is diminished in extracts prepared from ΔG3BP1 cells. Translation of viral RNA in rabbit reticulocyte lysates (RRL) are used as a side by side comparison. The positions of the viral proteins are indicated (1-5). C) Quantification of protein
products and total translation levels across multiple experiments. The error bars represent the standard error of three independent experiments. D) Norovirus VPg-dependent translation in extracts from WT or ΔG3BP1 cells in the presence of increasing concentrations of total cellular RNA isolated from uninfected cells. Translation of viral RNA in rabbit reticulocyte lysates (RRL) are used as a side by side comparison. E) Quantification of viral proteins produced in panel (D).

**Supplementary Figure 11:** A) Characterisation of viral VPg-linked RNA. The sensitivity of purified MNV VPg-linked RNA to various nucleases was compared to *in vitro* transcribed capped MNV gRNA (cap-gRNA) and the MNV1 full length cDNA construct. RppH was included as a decapping enzyme require for Xrn1-mediated cleavage of capped RNAs. Following digestion, the samples were analysed on a 1% native agarose gel. B) *In vitro* translation of viral VPg linked RNA in rabbit reticulocyte lysates demonstrated robust translation and the production of a protein profile indistinguishable from *in vitro* transcribed capped genomic RNA (cap-g). Capped sub-genomic (cap-sg) RNA was included to demonstrate the location of the VP1 and VP2 proteins. C) The translation of MNV VPg-linked RNA in extracts prepared from ΔG3BP1 cells is reduced across multiple time points. *In vitro* translations prepared in rabbit reticulocyte lysates (RRL) using *in vitro* transcribed capped genomic RNA (cap-g) or capped sub-genomic (cap-sg), along with viral VPg-linked RNA, was used as a reference for the expected mass of the viral proteins.
Materials and methods.

Cells. The murine microglial BV-2 cell line (Blasi et al., 1990) was provided by Jennifer Pocock (University College London). BV-2 cells were maintained in DMEM supplemented with 10% FCS (Biosera), 2 mM L-glutamine, 0.075% sodium bicarbonate (Gibco) and the antibiotics penicillin and streptomycin. BHK cells engineered to express T7 RNA polymerase (BSR-T7 cells, obtained from Karl-Klaus Conzelmann, Ludwid Maximillians University, Munich, Germany) were maintained in DMEM containing 10% FCS, penicillin (100 SI units/ml) and streptomycin (100 μg/ml), and 0.5 mg/ml G418.

Generation of G3BP1 KO cells. BV-2 cells were cultured in DMEM containing 10% FBS and 1% HEPES. G3BP1 knockout BV-2 cells were generated using two approaches. The clone 1B2 was generated by transiently transfected with Cas9 and a sgRNA (5TTCCCCGGCCCCGGCTGATGNGG) targeting exon 7 of G3BP1. BV-2 cells were then single cell cloned and G3BP1 was sequenced by Illumina HiSeq. BV-2 cells are polyploid at the G3BP1 locus as described previously (Orchard et al., 2016). Clone 1B2 also had three independent deletions at the sgRNA binding site resulting in deletions of 1, 2, and 5 base pairs respectively. The mutations introduced into the IB2 BV-2 cell clone resulted in frame shifts at nucleotide positions 253, 254 and 244 and the absence of detectable G3BP1 protein as measured by western blot. G3BP1 knockout BV-2 cell clones A, C and F were generated using an independent approach that relied on first generating a pool of three lentiviruses carrying guide RNAs TGTGCAACATGTCCGGGGC, CAAACTCCCGCCCGACCAGC and TAGTCCCCTGCTGGTCGGGC targeting the first 100bp of the coding sequence, cloned into pLentiCRISPRv2 (Sanjana et al., 2014). BV-2 cells were then transduced
with the pool of 3 lentiviruses, selected by puromycin treatment for 72 hours, prior to cloning by limiting dilution. Individual clones were then screened by western blot for the absence of G3BP1.

G3BP1 complementation. Mouse G3BP1 cDNAs were subcloned into pCDH-MCS-T2A-puro-MSCV lentiviral vector (System Biosciences) by NEBuilder HiFi DNA assembly (New England Biolabs). Mouse G3BP1 was subcloned from pCM6-G3BP1 (MR207441; Origene). Mouse G3BP1 lentiviral constructs deficient in the C-terminal RGG domain (mG3BP1ΔRGG) and the RGG and RRM domains (mG3BP1ΔRGGRRM) were generated by Gibson cloning from the pCMV6-G3BP1 vector. Lentivirus was generated by co-transfecting pCDH-G3BP1-T2A-puro-MSCV with pCMV-VSV-G and pSPAX2 into 293T cells with Trans-IT LT1 (Mirus Biosciences) per manufacture instructions. Two days post-transfection, supernatants were harvested, filtered through a 0.22 micron filter, and stored at -80°C. Lentivirus encoding G3BP1 or an empty control was then used to transduce G3BP1 KO 1B2 BV-2 cells. Two days post-transduction BV-2 cells were selected with puromycin (2.5ug/ml) for six days.

MNV growth curves
To determine the effects of G3BP1 disruption on MNV replication G3BP1 WT, KO, or complemented cells were plated in each well of a flat bottom 96-well plate and then infected with either MNV strains CW1, CW3, or CR6 as described in the text. Infected cells were flash frozen at -80°C at the times post infection indicated in the text. Viral replication was then assessed by plaque assay or TCID50 in BV-2 cells as described in the text. In cases where the appearance of virus-induced cytopathic
effect was examined, infected monolayers were either visualized by light microscopy
directly or fixed with crystal violet in formalin, prior to washing and imaging.

**CRISPR screens.** The CRISPR screen was performed similarly to that described
previously (Orchard et al., 2016) with a number of modifications that included the use
of the Brie gRNA library to reduce off target effects (Doench et al., 2016) and shorter
infection times to improve the recovery of gRNAs that may also compromise cell
viability. BV-2 cells stably expressing Cas9 nuclease (Orchard et al., 2016) were
transduced with the Brie library using previously described protocols (Doench et al.,
2016). MNV strains CW3 and CR6 were used to infect BV-2 CRISPR library at MOI
5 pfu/cell and cells were isolated 24 hours post infection and preparation of gDNA
for sequencing as described previously (Orchard et al., 2016). The screen relies on
the premise that guide RNAs targeting genes that are overrepresented following
infection represent genes that when disrupted are protected against infection and
therefore likely represent factors with pro-viral activity. Likewise, genes for which
guide RNA are underrepresented suggest that infection had proceeded faster and
the gene is antiviral. Following sequencing, the data was analyzed by STARS
method as previously described (Doench et al., 2016; Orchard et al., 2016).
Visualization of candidate genes was accomplished using R (RStudio, Inc., Boston,
MA).

**Maintenance of SILAC cell lines.** Stable isotope labelling of amino acids in cell
culture of BV-2 cells (SILAC, Ong et al., 2002), was carried out in high-glucose
DMEM lacking arginine and lysine (Sigma-Aldrich), supplemented with dialyzed fetal
bovine serum, 1% L-glutamine, 1X nonessential amino acids, 10 mM HEPES, and
1X penicillin/streptomycin. SILAC media were supplemented with Light (R0K0), Medium (R6K4) or Heavy (R10K8) Arginine and Lysine (Cambridge Isotope Laboratories). BV-2 cells were maintained in SILAC medium for 2 weeks to ensure complete metabolic labelling of proteins. Labelling of HEK-293T cells was performed essentially as described for BV-2 cells, with the omission of 10 mM HEPES and 1X non-essential amino acids from the cell culture media.

DNA based recovery of murine norovirus. Experiments were performed according to previously published protocols (Chaudhry et al., 2007). Briefly, BSRT7 cells were infected to an MOI of 0.5-1 PFU/cell with fowlpox virus expressing T7 RNA polymerase. Cells were then transfected with a plasmid encoding the MNV full length clone, or a derivative thereof (e.g. pT7 MNV 383FLAG 3'Rz or pT7 MNV 2600FLAG 3'Rz, our FLAG-tagged virus constructs containing FLAG tags in either NS1/2 or NS4 respectively). MNV was harvested by freeze-thaw at 24h post-transfection.

To generate higher titre stocks, WT MNV, NS1/2-FLAG MNV, and NS4-FLAG MNV (Thorne et al., 2012) generated using the DNA based recovery method described above were passaged once in BV-2 cells. After 80-90% of cells displayed visible cytopathic effects (CPE) of viral infection, flasks containing infected cells were frozen at -80°C. Flasks were frozen and thawed twice before cell debris was removed by centrifugation at 4000 rpm for 10 minutes in a benchtop centrifuge. Viruses were pelleted by centrifuging over a 30% sucrose cushion at 76,755xg in a SW32ti rotor for 2 hours at 4°C. Virus pellets were resuspended overnight in PBS to achieve 100-fold concentration. Concentrated virus was then passed through a 23-gauge blunt
needle 15 times, and clarified by centrifugation at maximum speed in a benchtop microcentrifuge for 10 minutes. Supernatant aliquoted, and titrated prior to use.

**Infection of SILAC labelled BV-2 cells with FLAG-tagged viruses.** SILAC-labelled BV-2 cells were infected with WT MNV, NS1/2-FLAG or NS4-FLAG viruses at an MOI of 10 TCID50 cell. Infections were performed in triplicate, using different combinations of SILAC-labeled BV-2 cells each time to control for any impact of the SILAC labelling. Infected cells were then plated in the appropriate SILAC media. At 10 hours post-infection, cells were harvested by scraping, and pelleted at 500xg for 5 minutes. Cells were then washed 3 times with ice-cold PBS, and were lysed in (0.5% Nonidet-P40 substitute, 10 mM Tris-HCl pH 7.5, 150 mM NaCl, 0.5 mM EDTA, 2 mM MgCl). Benzonase nuclease (Sigma-Aldrich) was added to lysis buffer to a concentration of 5 μl/ml to prevent nonspecific interactions mediated by RNA or DNA.

**Transfection of SILAC labelled HEK-293T cells with GFP-tagged VPg.** SILAC-labelled HEK-293T cells were transfected with pEGFP-C1 (control) or derivatives thereof containing either human or murine norovirus VPg protein as described in Emmott & Goodfellow, (2014). GFP fusions of both wild-type protein or mutant VPg containing a mutation to inhibit initiation factor binding (MNV: F123A, HuNoV: F137A) were used. Cells were transfected using Lipofectamine 2000 (Life Technologies) according to the manufacturers protocol, using antibiotic-free SILAC media in place of Opti-mem. The experiment was performed in triplicate and SILAC labels switched in one of the replicates.
FLAG and GFP-TRAP immunoprecipitation. FLAG immunoprecipitations were performed following the manufacturer’s protocol (FLAG M2 beads, Sigma Aldrich) as described (Thorne et al., 2012). In brief, protein concentration in lysates was normalized using BCA. Lysates were then diluted with 1 volume of wash buffer (10 mM Tris-Cl pH 7.5, 150 mM NaCl, 0.5 mM EDTA). Equal volumes of anti-FLAG affinity gel were dispensed into either WT infected cell lysates, or lysates of cells infected with NS1/2-FLAG or NS4-FLAG. Binding was carried out overnight at 4°C with rotation. After binding, beads were washed 3 times with wash buffer. All liquid was carefully removed from each tube, before boiling in SDS-PAGE loading buffer for 10 minutes. GFP-trap immunoprecipitation of GFP-tagged VPg was accomplished using GFP-trap beads (Chromotek) per the manufacturer’s protocol, as described (Emmott and Goodfellow, 2014). RNase cocktail (Ambion) was also included in the lysis buffer at a concentration of 5 μl/ml to prevent non-specific interactions mediated by RNA. In all cases, light, medium, and heavy-labelled proteins eluted from the beads for each experimental replicate were pooled together in a ratio of 1:1:1 before submission for mass spectrometry analysis at the University of Bristol Proteomics Facility.

Mass spectrometry analysis. Mass spectrometry analysis was performed at the University of Bristol Proteomics Facility. In brief, samples were run into precast SDS-PAGE gels for 5 minutes, the entire sample cut from the gel as a single band, and then subjected to in-gel tryptic digestion including reduction and alkylation using a ProGest automated digestion unit. The resulting peptides were fractionated using a Dionex Ultimate 3000 nanoHPLC system in line with an LTQ-Orbitrap Velos or Orbitrap Tribrid Fusion mass spectrometer.
Interpretation of SILAC Proteomics data. Raw data files were processed and quantified using Maxquant v1.5.5.1 or 1.6.0.16 (Tyanova et al. 2016). The GFP-VPg experiments were searched against the Uniprot human database (70,550 entries, downloaded September 19th 2016) plus a custom FASTA file containing the wild-type and mutant VPg sequences. The raw data, search results and FASTA files can be found as part of PRIDE submission PXD007585 (Reviewer username: reviewer75984@ebi.ac.uk, Password: BH2pTctW). The FLAG-virus experiments were searched against the Uniprot mouse database (Swiss-prot only, 16,966 entries, downloaded May 19th 2018) plus a custom FASTA file containing the various Murine norovirus proteins. The raw data, search results and FASTA files can be found as part of PRIDE submission PXD011779 (Reviewer username: reviewer49419@ebi.ac.uk, Password: eLYwivNP). Data were searched with default Maxquant parameters including upto 2 missed tryptic cleavages, oxidation of methionine and N-terminal acetylation as variable modifications, and carbamidomethylation of cysteine as a fixed modification. The data were searched against a reverse database and PSM and Protein FDR were set to 0.01. The requantify option was not selected.

GFP-VPg data were analysed as described previously (Emmott and Goodfellow, 2014). FLAG-virus experiments were analysed by computing the pairwise ratios of samples infected with NS1/2-FLAG or NS4-FLAG relative to WT MNV-infected controls. Log₂ SILAC ratios for proteins identified in at least 2/3 replicates were averaged, and ratios for NS1/2-FLAG:WT and NS4-FLAG:WT were plotted for comparison of host cell proteins by viral replication complex-associated proteins.
Assessment of virus-induced cytopathic effect. BV-2 WT, KO G3BP1 or respective G3BP1 complemented cells as described in the text, were seeded onto 96 well plates and infected with serial 10-fold dilutions (starting at MOI=10 TCID50/cell) of MNV (CW1) or EMCV. At 48h post-infection, cells were fixed in ice-cold methanol and stained with toluidine blue prior to washing and imaging.

Cap-Sepharose purification for eIF4F complex. Cell lysates were prepared from BHK parental cells or BHK containing GI Norwalk virus (BHK-NV) replicon cells in cap-Sepharose lysis buffer (100 mM KCl, 0.1 mM EDTA, 10% glycerol, 2 mM MgCl$_2$, 20 mM HEPES, pH 7.6 in KOH) with 1% TX-100, proteinase and phosphatase inhibitor cocktails (Calbiochem). Cytoplasmic extracts were centrifuged and RNase treated for 15 min at room temperature. At least 1000 µg of the cell lysates were incubated with Sepharose beads coupled to 7-methylguanosine (m$^7$GTP, Jena Biosciences). Input cell lysates were collected for western blot analysis while the remaining were incubated overnight with continuous rotation at 4°C. The eIF4F-enriched complex was precipitated and washed 2 times with cap-sepharose lysis buffer. Bound proteins were eluted in 2x reducing SDS-PAGE samples buffer and resolved by SDS-PAGE prior to western blot.

Human Norwalk virus colony formation assay. Total RNAs extracts from BHK or BHK-NV replicon-containing cells (Kitano et al., 2018) were pretreated with and without proteinase K (10 µg/ml) in 10 mM Tris, pH 8.0, 1.0 mM EDTA, 0.1 M NaCl, and 0.5% SDS. Pretreated RNAs were immediately purified using GenElute RNA purification columns (Sigma). Serial 10-fold dilutions of mock or proteinase K-treated
RNAs were transfected in BHK cells and 24 h post transfection, cells were passaged and maintained in growth media containing 0.5 mg/ml G418. Colonies began to form after 5 d and were allowed to grow until 14 d. All plates were harvested at day 14 and well-formed colonies were fixed in 10% formaldehyde and stained with toluidine blue. A similar protocol was followed to assess colony formation in U2OS cells with the exception that selections were maintained for up to 12 days post transfection. Where indicated, cell aliquots from each time point were collected for qRT-PCR analysis to assess viral RNA synthesis over time.

**Polyribosome fractionation analysis.** BV2 WT and BV2 ΔG3BP1 cells were seeded at a density of 7.5 x 10⁶ cells per T-75 flask, and then either mock infected or infected with MNV1 (CW1) at MOI 3 TCID50 per cell in the presence of 2-CMC (400µM) for each set of infection. After 1h, the inoculum was then removed; the cells were washed and maintained in growth media containing 2-CMC accordingly until the cells were harvested at 4h and 9h p.i. Prior to harvesting, cells were treated with cycloheximide (CHX) for 10 mins at 37°C (Sigma-Aldrich; 100 µg/ml) and were rinsed with 5 ml of ice-cold PBS supplemented with CHX 100 µg/ml. Polysome lysis buffer [20 mM Tris-HCl pH 7.5, 150 mM NaCl, 5mM MgCl, 1 mM DTT, 1% Triton X-100, 100 µg/ml cycloheximide, 25 U/ml TURBO DNase (Life Technologies)] was used to lyse the cells. Lysates were clarified by centrifugation for 20 min at 13,000 g at 4°C. Aliquots of the lysates were collected for BSA assay and qPCR analysis against MNV1 RNA before proceeding with fractionation. Input lysates were normalized to total protein concentration and RT-qPCR was used to confirm the levels of viral RNA in samples were comparable. Lysates were subjected next to 10-50% sucrose gradient centrifugation for 90 mins SW41Ti rotor at 190,000 x g at 4°C.
The gradients were fractionated at 0.5 ml/min and the levels of RNA in each sample measured using an in line-254 nm spectrophotometer connected to a chart recorder. RNAs were extracted from each fraction, converted to cDNA and immediately used for qPCR. The distribution of viral RNA across the gradient was then calculated as percentage (%) of the viral RNA seen in WT BV-2 cells using the reference gene (GAPDH) to obtain normalized values across the gradient. Samples were performed in duplicates on the same qPCR plate, and the observations were robust across three independent experiments. Data were collected using a ViiA 7 Real-Time PCR System (Applied Biosystems).

Transfection of VPg-linked MNV RNA into BV-2 cells. VPg-linked RNA purified from MNV-1 virus particles was transfected in BV-2 cells using NEON™ as previously described (Yunus et al., 2010). Total cell lysates were harvested at 3 and 9 hours post transfection with RIPA buffer. 10 μg total lysates were analysed by 4-12% SDS-PAGE (Invitrogen) and antibodies against MNV, VPg, G3BP1 and GAPDH were used for detection using LI-COR® Odyssey® CLx. Virus yield was determined by TCID₅₀. For strand-specific qPCR detection of MNV RNA, total cellular RNA was extracted using GeneElute Mammalian Total RNA Miniprep kit (Sigma). RT-qPCR was performed as described previously (Vashist et al., 2012a).

Purification of MNV VPg-linked RNA. BV-2 cells were infected at an MOI=0.01 TCID₅₀ per cell and harvested after ~30 post infection. Cell debris was removed by low speed centrifugation for 10 minutes and supernatant loaded onto 5 ml of 30% sucrose solution in PBS. MNV particles were pelleted using a SW32Ti rotor at 25,000 RPM for 4 hours at 4°C. Virus was then resuspended in PBS and total RNA
extracted from soluble fraction. Where detailed, the authenticity of the viral RNA was examined by nuclease digestion. 500 ng of viral RNA or plasmid DNA was treated with DNase I (10U, Roche), XrnI+RppH (1U XrnI + 5U RppH, both from NEB) or RNase cocktail (0.5U RNase A + 20U RNase T1, ThermoFisher) at 37 °C for 10 minutes. Then analysed on 1% agarose gel.

**Preparation of BV2 S10 cytoplasmic extracts.** Preparation of BV-2 S10 extracts was based on a previously published protocol (Rakotondrafara and Hentze, 2011; 2006). BV-2 cells were harvested, washed with PBS, and lysed with 1x packed volume of hypotonic buffer containing 10 mM HEPES pH7.6, 10 mM potassium acetate, 0.5 mM magnesium acetate, 5 mM DTT, 1x protease inhibitors cocktail (EDTA-free, Roche). Cells were lysed on ice for 45 minutes, then passed through 25G and 27G needles to achieve >95% lysis. Cell lysates were then centrifuged at 10,000 x g for 10 minutes at 4 °C twice and the supernatant collected. The total protein concentration was measured by Bradford assay and normalised to 20 mg/ml before freezing at -80 °C until use. For micrococcal nuclease treatment, S10 extracts were thawed on ice, 1 mM calcium chloride and 200 unit/ml final concentrations of micrococcal nuclease (NEB). Cell lysates were incubated at 25 °C for 15 minutes before adding 3 mM final concentration of EGTA was added.

**In vitro translation of BV2 S10 lysates.** *In vitro* translation assays were set up based on a previous protocol (Favre and Trepo, 2001). Translation reactions were set up in 12.5 μl total volume containing 5 μl BV2 S10 lysate, 2.5 μl 5X translation buffer, 0.25 μl of 5 mg/ml creatine kinase, 1.25 μl RRL, 0.225 μl of 5 M potassium acetate, 0.25 mM of 100 mM magnesium acetate, 5.13 μCi 35S-labelled methionine...
(PerkinElmer) and 10-100 ng/μl RNA as detailed in the text. 5X translation buffer contains 350 mM HEPES, 75 mM creatine phosphate, 10 mM ATP, 3.75 mM GTP, 100 μM amino acid minus methionine, 3.75 mM spermidine and 0.375 mM S-adenosyl-methionine. For control experiments using RRL (Promega), the reactions were set up according to manufacturer’s instructions using 0.5-1 ng/μl RNA. Reactions were incubated at 30 °C for 90 minutes before addition of 12.5 μl trans-stop buffer containing 10 mM EDTA and 0.1 mg/ml RNase A and incubated at room temperature for 10 minutes, then 25 μl 5X loading buffer was added to the reaction and heated at 95 °C for 5 minutes. 10 μl lysates were resolved in 15% SDS-PAGE and exposed to a phosphorimag screen and visualised using a TyphoonFLA7000 machine. For non-radioactive translation, 1.25 μl of 1 mM methionine was used instead of 35S-labelled methionine, and the reactions were stopped with 100 μl 1x passive lysis buffer (Promega) and the luminescence read using a GloMax luminometer (Promega).
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Figure 1

A. CLUSTAL alignment of murine and human eIF4G binding domains.

B. Western blot analysis of protein expression in BHK and BHK-NV cells.

C. Co-immunoprecipitation of VPg-GFP and eIF4G.

D. Heat map of protein expression levels.

E. Dual-labeling experiment with GFP and VPg-GFP.
Supplementary Figure 1

A. Host proteins binding to the murine norovirus VPg protein

B. Host proteins binding to the Norwalk virus VPg

C. Host proteins previously identified as interacting with the MNV VPg protein or the termini of the MNV genomic RNA

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1. Infect at MOI 5 pfu/cell for 24 hours
2. Collect surviving cells
3. Determine gRNA abundance relative to mock infected

Figure 3
Figure 5

A. Mock

VPg-linked RNA

-Pk

+Pk

B. WT ΔG3BP1

C. Viral RNA fold change vs D0

|         | d0  | d4  | d12 |
|---------|-----|-----|-----|
| WT      | 100 | 1000| 1000|
| ΔG3BP1  | 100 | 1000| 1000|
Figure 6

A.

kDa

WT ΔG3BP1 Empty G3BP1 ΔRGG ΔRRM

G3BP1

ΔG3BP1

G3BP2

ΔG3BP1 + empty

ΔG3BP1 + G3BP1

ΔG3BP1 + ΔRGG

ΔG3BP1 + ΔRGG/RRM

B.

MNV

Mock MNV EMCV

Mock MNV EMCV

Mock MNV EMCV

Mock MNV EMCV

Mock MNV EMCV

Mock MNV EMCV

Mock MNV EMCV

Mock MNV EMCV

Mock MNV EMCV

Mock MNV EMCV

C.

TCID50/ml (Log10)

WT ΔG3BP1 + empty

ΔG3BP1 + G3BP1

ΔG3BP1 + ΔRGG

ΔG3BP1 + ΔRGG/RRM

ΔG3BP1

LOD 0Hr
Figure 8

A. Time post-infection (h)

B. Viral positive sense RNA

C. Viral negative sense RNA

D. Viral positive sense RNA

E. Viral negative sense RNA

F. Viral positive sense RNA

G. Viral negative sense RNA
Figure 9

A.

| Input | Pull-down |
|-------|-----------|
| GFP   | GFP       |
| GFP   | eIF4G     |
| GFP   | G3BP1     |
| GFP   | RpS6      |

B.

| Input | Pull-down |
|-------|-----------|
| GFP   | GFP       |
| GFP   | eIF4G     |
| GFP   | G3BP1     |
| GFP   | RpS6      |
Figure 10

A.

Mock 4 Hr

Mock 9 Hr

Polysomes

B.

WT

ΔG3BP1

4 Hr

9 Hr

Polysomes

C.

WT

ΔG3BP1

Free RNA & RNP

Free subunits

Polysomes

Fractions

Fractions
Figure 11

A. 

B. 

C. 

D. 

E. 

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