**Birnaviridae** Virus Factories Show Features of Liquid-Liquid Phase Separation and Are Distinct from Paracrystalline Arrays of Virions Observed by Electron Microscopy

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**ABSTRACT** To gain more information about the nature of *Birnaviridae* virus factories (VFs), we used a recombinant infectious bursal disease virus (IBDV) expressing split-GFP11 tagged to the polymerase (VP1) that we have previously shown is a marker for VFs in infected cells expressing GFP1-10. We found that VFs colocalized with 5-ethynyl uridine in the presence of actinomycin, demonstrating they contained newly synthesized viral RNA, and VFs were visible in infected cells that were fixed and permeabilized with digitonin, demonstrating that they were not membrane bound. Fluorescence recovery after photobleaching (FRAP) a region of interest within the VFs occurred rapidly, recovering from approximately 25% to 87% the original intensity over 146 s, and VFs were dissolved by 1,6-hexanediol treatment, demonstrating they showed properties consistent with liquid-liquid phase separation. There was a lower colocalization of the VF GFP signal with the capsid protein VP2 (Manders’ coefficient [MC] 0.6), compared to VP3 (MC, 0.9), which prompted us to investigate the VF ultrastructure by transmission electron microscopy (TEM). In infected cells, paracrystalline arrays (PAs) of virions were observed in the cytoplasm, as well as discrete electron dense regions. Using correlative light and electron microscopy (CLEM), we observed that the electron dense regions correlated with the GFP signal of the VFs, which were distinct from the PAs. In summary, *Birnaviridae* VFs contain newly synthesized viral RNA, are not bound by a membrane, show properties consistent with liquid-liquid phase separation, and are distinct from the PAs observed by TEM.

**IMPORTANCE** Members of the *Birnaviridae* infect birds, fish and insects, and are responsible for diseases of significant economic importance to the poultry industry and aquaculture. Despite their importance, how they replicate in cells remains poorly understood. Here, we show that the Birnaviridae virus factories are not membrane bound, demonstrate properties consistent with liquid-liquid phase separation, and are distinct from the paracrystalline arrays of virions observed by transmission electron microscopy, enhancing our fundamental knowledge of virus replication that could be used to develop strategies to control disease, or optimize their therapeutic application.

**KEYWORDS** infectious bursal disease virus, IBDV, *Birnaviridae*, virus factory, replication complex, liquid-liquid phase separation, LLPS, CLEM, FRAPTEM

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The *Birnaviridae* family is divided into 4 genera that infect birds (*Avibirnavirus*), fish (*Aquabirnavirus* and *Blosovirinae*), and insects (*Entomobirnavirus*) (1). Within the genus *Avibirnavirus*, infectious bursal disease virus (IBDV) is of major concern to the global poultry industry due to morbidity, mortality, and immunosuppression in chickens that lead to an increase in secondary infections (2). IBDV is controlled through vaccination (3), and recombinant vaccine strains also show promise as vectors for the...
expression of antigens to control other infections (4, 5). The *Avibirnavirus*, chicken proventricular necrosis virus (CPNV), has also recently been proposed to be the cause of transmissible viral proventriculitis (TVP) (6). Within the genus *Aquabirnavirus*, infectious pancreatic necrosis virus (IPNV) infects salmonid fish, for example salmon and trout, causing high mortality in fry and fingerlings, and many related viruses are associated with disease in a wide variety of fish, mollusks, and crustaceans, resulting in significant economic losses to aquaculture (1, 7). In addition, Blotched Snakehead virus, another birnavirus infecting fish, is the sole member of the genus *Blosnavirus*, and is more closely related to the *avibirnaviruses* than the *aquabirnaviruses* (8). Members of the genus *Entomobirnavirus* infect insects and include Drosophila X virus (DXV) which has played an important role in the study of innate immunity and siRNA pathways (9, 10), Culex Y virus (CYV), and Port Bolivar virus that have been identified in mosquito cells (11, 12).

Despite their abundance and importance, many aspects of the replication of the Birnaviridae remain poorly understood. As the viruses are non-enveloped, and contain a segmented double-stranded (ds)RNA genome, assumptions of their replication strategies have been extrapolated from knowledge of more well-studied dsRNA viruses, for example the Reoviridae family. However, significant differences between the strategies used by the Birnaviridae and the Reoviridae are beginning to be uncovered. For example, birnaviruses do not make use of a core particle during the replication cycle (13). Instead, they possess a single capsid layer and uncoat in the endosome, whereupon the genome, combined in a ribonucleoprotein (RNP) complex with the viral polymerase (VP1) and the nonstructural protein VP3, is extruded into the cytoplasm (14). In the cytoplasm, the VP3 tethers the RNP complex to the cytoplasmic leaflet of endosomal membranes (15) via interacting with phosphatidylinositol-3-phosphate (16), and it is thought that the virus co-opts these cellular membranes to seed the formation of a replication complex, or virus factory (VF), to provide a microenvironment more conducive to replication.

In order to provide more information on the nature of the Birnaviridae VFs, we previously rescued a replication-competent IBDV, strain PBG98, containing a split green fluorescent protein (GFP) tag fused to the C terminus of the VP1 polymerase (PBG98-VP1-GFP11) (17). In infected cells expressing GFP1-10, the GFP11 tag complemented the GFP1-10 molecule and the resultant complete GFP molecule fluoresced green. The GFP signal in PBG-VP1-GFP11-infected cells colocalized with other components of the RNP complex, VP3, and dsRNA, visualized by immunofluorescence microscopy, and with genome segments A and B, visualized by fluorescence in situ hybridization (FISH). We therefore concluded it was a marker for VFs. Using live cell imaging, we found that the VFs moved in the cytoplasm and coalesced together over time (17). Here, we build on this work to further understand the nature of the Birnaviridae VFs, investigating the possibility they are formed from liquid-liquid phase separation using live cell imaging approaches, and defining their ultrastructure by transmission electron microscopy (TEM), and correlative light and electron microscopy (CLEM).

**RESULTS**

IBDV VFs contain newly synthesized RNA. DF-1 cells were transfected with GFP1-10 and infected with the PBG98-VP1-GFP11 virus. Infected cells were treated with actinomycin D to prevent cellular transcription, after which EU was added to the cells that became incorporated into newly synthesized mRNA molecules. After EU treatment, cells were stained with a Click-iT RNA imaging kit to visualize EU and stained with DAPI (Fig. 1). In uninfected control cells lacking actinomycin D treatment, the EU signal, representing cellular mRNAs, was visible in the nucleus, as expected given that transcription is intranuclear. This intranuclear signal was lost in cells treated with actinomycin D, which blocked cellular transcription. VFs were visible in infected cells that were not treated with actinomycin D or EU (Fig. 1A), and in cells that were treated with actinomycin D, but not EU (Fig. 1B), indicating that blocking cellular transcription at this time
point did not prevent VF formation. Infected cells that were not treated with actinomycin D, and were stained with EU had an EU signal in the nucleus, likely representative of cellular mRNA synthesis, and a signal that colocalized with the VFs, indicating that the PBG98-VP1-GFP11 VFs also contained newly synthesized RNA (Fig. 1C). In infected cells treated with actinomycin D, the only EU signal detected colocalized with the VFs in the cytoplasm (Fig. 1D), indicating that the RNA was synthesized independent of the cellular machinery, consistent with the action of the viral polymerase.

**IBDV VFs were not bound by a membrane.** DF-1 cells were transfected with GFP1-10 and infected with the PBG98-VP1-GFP11 virus. Infected cells were fixed and permeabilized with either Triton X-100, which permeabilized the plasma membrane and internal membranes, or digitonin, which permeabilized the plasma membrane but not intracellular membranes. Cells were stained with primary antibodies raised against tubulin (a cytoplasmic marker), PDI (present within the endoplasmic reticulum and therefore a marker of intracellular membrane permeabilization), or VP3, followed by secondary antibodies conjugated to Alexafluor 568 (Fig. 2). As expected, the cytoplasmic tubulin was visible following either permeabilization method, but PDI was not visible when cells were permeabilized with digitonin as it was contained within a membrane-bound organelle (Fig. 2C and D). The PBG98-VP1-GFP11 signal was present in cells permeabilized with either triton X or digitonin, which colocalized with VP3, demonstrating that the VFs were not bound by a membrane.

**FIG 1** IBDV VFs were the sites of de novo RNA synthesis. DF-1 cells were transfected with GFP1-10 and infected with the PBG98-VP1-GFP11 virus 24hpt at an MOI of 1 (A-D). Thirteen hpi, cells were either treated with actinomycin D for 4 h to prevent cellular transcription (B and D), or mock treated with media (A and C), after which cell cultures were either treated with 1 mM EU that became incorporated into newly synthesized RNA molecules (C and D), or mock treated with media alone (A and B). One hour after EU treatment, all cells were stained with a Click-iT RNA imaging kit to visualize EU and stained with DAPI. Scale bars, 20 μm.
IBDV VFs had properties consistent with liquid-liquid phase separation. DF-1 cells were either infected with recombinant PBG98 wt (A and C), or transfected with GFP1-10 and infected with the PBG98-VP1-GFP11 virus (B and D). Infected cells were fixed and permeabilized with either Triton X-100 (A and B), or digitonin (C and D). Cells were then stained with primary antibodies raised against tubulin, PDI, or IBDV VP3, followed by secondary antibodies conjugated to Alexa Fluor 568, and DAPI. Scale bars, 20 μm.

**FIG 2** IBDV VFs were not bound by a membrane. DF-1 cells were either infected with recombinant PBG98 wt (A and C), or transfected with GFP1-10 and infected with the PBG98-VP1-GFP11 virus (B and D). Infected cells were fixed and permeabilized with either Triton X-100 (A and B), or digitonin (C and D). Cells were then stained with primary antibodies raised against tubulin, PDI, or IBDV VP3, followed by secondary antibodies conjugated to Alexa Fluor 568, and DAPI. Scale bars, 20 μm.

IBDV VFs had properties consistent with liquid-liquid phase separation. DF-1 cells were transfected with GFP1-10 and infected with the PBG98-VP1-GFP11 virus. ROIs within the VFs were photobleached to approximately 25% of their original intensity, and fluorescence recovery was determined over time. The VFs showed a rapid recovery in fluorescence, to approximately 87% (standard deviation [SD] 3.5) of the original intensity during the course of the experiment (146 s) (Fig. 3A). In a separate experiment, DF-1 cells were transfected with GFP1-10 and infected with the PBG98-VP1-GFP11 virus. Infected cells were treated with 4% 1,6-hexanediol, which has been shown to dissolve the VFs of rotavirus (18). In the presence of 1,6-hexanediol, the VFs of PBG98-VP1-GFP11 dissolved (Fig. 3B and Movie S2), whereas the VFs of mock-treated cells were visible throughout the experiment (Fig. 3B and Movie S1). These data, taken together with our previous study showing VFs rapidly moved in the cytoplasm and coalesced together (17), demonstrated that IBDV VFs had properties consistent with liquid-like behavior.

**Virus Factory like structures (VFLs) formed in the cytoplasm of cells transfected with GFP-VP3, but not other viral proteins.** DF-1 cells were transfected individually with plasmids encoding each of the viral proteins VP1, 2, 3, 4 and 5, tagged to GFP,
and imaged by confocal microscopy. GFP was tagged to the N terminus of the proteins, with the exception of VP2, where GFP was tagged to the C terminus, as the N-terminal tag (GFP-VP2) did not lead to a detectable signal upon imaging. Cells transfected with either GFP-VP1 or VP2-GFP had a fluorescence signature that was present throughout the cytoplasm, whereas cells transfected with GFP-VP3 showed puncta in the cytoplasm reminiscent of the virus factories. We termed these virus-factory-like (VFL) structures. Cells transfected with GFP-VP4 contained wave-shaped structures in the cytoplasm, and cells transfected with GFP-VP5 had a signal that appeared to be near or at the plasma membrane (Fig. 4A). When a small ROI of a VP3-GFP VFL was photobleached to approximately 25% the original intensity with the 488 nm Argon laser, then 170 postbleach images were acquired during the experiment (146 s). The fluorescence intensity was normalized to the background and plotted against time (A). Infected cells were either mock-treated with media, or were treated with a solution of 4% 1,6-hexanediol (hex) 16 hpi and imaged live over a time course of 17 min. For each time course, 50 z-stacks were imaged in total, with each stack being acquired at an interval of 20 s. Images of different time points within the time course indicated by the time stamp (minutes: seconds) are shown (B). Scale bars, 20 μm.

FIG 3 IBDV VFs had properties consistent with liquid-liquid phase separation. DF-1 cells were transfected with GFP1-10 and then infected with the PBG98-VP1-GFP11 virus 24 hpt. At 16 hpi, one ROI with a 0.6 μm diameter was photobleached inside a VF to approximately 25% the original intensity with the 488 nm Argon laser. Then, 170 postbleach images were acquired during the experiment (146 s). The fluorescence intensity was normalized to the background and plotted against time (A). Infected cells were either mock-treated with media, or were treated with a solution of 4% 1,6-hexanediol (hex) 16 hpi and imaged live over a time course of 17 min. For each time course, 50 z-stacks were imaged in total, with each stack being acquired at an interval of 20 s. Images of different time points within the time course indicated by the time stamp (minutes: seconds) are shown (B). Scale bars, 20 μm.

Birnaviridae VFs Show Features of LLPS

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VFs had a lower colocalization with VP2 than VP3 by confocal microscopy. In a previous study, we demonstrated that the VFs had a high colocalization with VP3. In this experiment, we observed the colocalization of the VF GFP signal with VP2, VP3, and VP4 (Fig. 5A) and quantified this using a Manders’ coefficient (MC) (Fig. 5B). We found the MC to be 0.9 for VP3, and 0.2 for VP4, consistent with strong and weak colocalization, respectively. In addition, we observed a lower colocalization with the capsid
protein, VP2 (MC, 0.6) than VP3, that prompted us to investigate the ultrastructure of the VF by TEM.

Paracrystalline arrays (PAs) of virions and electron-dense regions of the cytoplasm were observed in infected cells by TEM. DF-1 cells were either transfected with GFP1-10 and then infected with PBG98-VP1-GFP11 (Fig. 6A), or were infected with the avian reovirus (ARV) strain S1133 (Fig. 6B), or the recombinant wild-type (wt) PBG98 (Fig. 6C), and imaged by TEM. We observed PAs of virions in the cytoplasm of cells infected with IBDV strain PBG98-VP1-GFP11 (Fig. 6A). Upon further observation, we were unable to discern any other structures within the PAs, even by electron tomography (ET) (Movie S3). In contrast, DF-1 cells infected with ARV strain S1133 contained VFs that had a more complex ultrastructure, including an electron-dense area, as well as PAs of virions (Fig. 6B). In IBDV-infected cells, we observed discrete electron-dense regions in the cytoplasm that were distinct from the PAs. These were observed in cells transfected with GFP1-10 and infected with PBG98-VP1-GFP11 (data not shown), as well as DF-1 cells that were not expressing GFP1-10 and were infected with wt PBG98 (Fig. 6C).

VFs correlated with discrete electron-dense regions of the cytoplasm that were distinct from the PAs. In order to determine what structures the VFs we observed by confocal microscopy correlated with by TEM, DF-1 cells were seeded onto gridded glass coverslips, transfected with GFP-VP3, and infected with the PBG98-VP1-GFP11
virus. Infected cells were stained with Hoechst 33342 and imaged live by confocal microscopy (Fig. 7A and B). After imaging by confocal microscopy, cells were fixed and prepared for TEM. The same cell that was imaged by confocal microscopy was imaged by TEM (Fig. 7C), and the two images overlaid (Fig. 7D and E). We observed that the VFs correlated with the electron dense regions of the cytoplasm and were distinct from the PAs of virions (Fig. 7).

**DISCUSSION**

We previously demonstrated that in cells transfected with GFP1-10 and infected with an IBDV containing GFP11 fused to the C terminus of the VP1 polymerase, the GFP1-10 and GFP11 molecules came together to form the full-length GFP molecule, which fluoresced green. In addition, we demonstrated that the GFP signal colocalized with viral ribonucleoprotein (vRNP) components VP3 and dsRNA (17), indicating that the GFP-positive structures were VFs. In the present study, we extended these

**FIG 5** VFs had a lower colocalization with VP2 than VP3 by confocal microscopy. DF-1 cells were transfected with GFP1-10 and infected with the PBG98-VP1-GFP11 virus 24hpt at an MOI of 1. At 24 hpi, infected cells were fixed, permeabilized, and stained with anti-VP2, anti-VP3, or anti-VP4, followed by secondary antibodies conjugated to Alexa Fluor 568, and DAPI. Scale bars, 20 μm (A). An overlap coefficient, (an alternative to the Pearson’s correlation coefficient, created by Manders et al.) was used to quantify the colocalization of the VP1-GFP signal of the VFs with the VP2, VP3, and VP4 signals (B).
observations, confirming that the GFP-positive structures also colocalized with EU that was incorporated into newly synthesized RNA, even when cellular transcription was blocked by actinomycin D, demonstrating that the VFs contained newly synthesized RNA. As the GFP-positive structures contain the viral polymerase, it is possible that they are the sites of de novo RNA synthesis. However, it is also possible that viral RNA may be synthesized elsewhere in the cell, with transcripts subsequently accumulating in the VFs. Further experiments are needed to test these hypotheses.

Next, we sought to understand more about the nature of IBDV VFs. We demonstrated that the VFs were not bound by a membrane, which is consistent with previous literature demonstrating that structures containing VP3 were visible following cellular permeabilization with digitonin (15). It has previously been documented that upon entering the cytoplasm, VP3 within the vRNP complex binds to phosphatidylinositol-3-phosphate on the cytoplasmic leaflet of endosomal membranes to seed the formation of the VF (16). Interestingly, the Birnaviridae are thought to form a phylogenetic link with single-stranded positive (ss+) sense RNA viruses (1, 19), and the finding that the virus manipulates cellular membranes, presumably to provide a microenvironment more favorable for replication, is more akin to strategies used by ss+ RNA viruses in replication. Our finding that the VF is not bound by a membrane does not preclude the fact that there may be membrane structures within the VF, and Delgui et al. have demonstrated endosomal membrane markers colocalize with VP3–positive structures in QM7 cells infected with IBDV, even at 36 hpi (20). We did not observe any membrane structures within the VFs by TEM, during the time points analyzed, but it is possible that the VFs are heterogeneous in their composition throughout the infection, even within a single cell, and that membrane structures may be transiently present. Therefore, differences between the observations of our study and that of others could be attributed to differences in the infection time points analyzed.

Given that the VFs were not bound by a membrane, we hypothesized that they could be formed by liquid-liquid phase separation (LLPS). This phenomenon is potentially a means of cytoplasmic compartmentalization without the need for membrane-bound organelles (21), and is known to be involved in the replication of other viruses, for example vesicular stomatitis virus, rotavirus, rabies virus, influenza A viruses, and measles virus (18, 22–25). Structures that form through LLPS typically show fusion and fission, rapid recovery of fluorescence after photobleaching, and are dissolved by 1,6-hexanediol (26, 27). We have previously demonstrated that IBDV VFs move rapidly in the cytoplasm and coalesce together (17). Here, we extend these observations by demonstrating that VF FRAP was rapid, and the VFs were dissolved in the presence of 4% 1,6-hexanediol. Taken together, these data are consistent with the Birnaviridae VFs being formed through LLPS.

FIG 6 PAs of virions and electron-dense regions of the cytoplasm were observed in infected cells by TEM. DF-1 cells were either transfected with GFP1-10 and infected 24hpt with IBDV strain PBG98-VP1-GFP11 virus (A), or DF-1 cells were infected with the ARV strain S1133 (B), or the recombinant wt IBDV strain PBG98 at an MOI of 1. At 18 (A), 20 (B), or 10 (C) hpi, cells were fixed, prepared, and imaged by TEM.
Next, we transfected cells with individual IBDV proteins tagged to GFP and observed that GFP-VP3 formed VFLs, but that other IBDV proteins did not. It is possible that the presence of the GFP tag may have interfered with the ability of the other proteins to form VFLs, and so we cannot conclude from our study that untagged versions of these proteins do not form these structures. However, we are confident the VFL structures we observed following GFP-VP3 expression are not an artifact of the GFP tag, as our data are consistent with recent studies that have expressed untagged VP3 alone in cells and found it forms distinct cytoplasmic puncta (16). Here, we extend these observations by demonstrating that the GFP-VP3 VFLs show some FRAP, recovering to 46% of the original intensity over the course of the experiment, compared to GFP-VP4 which showed no FRAP. This indicates that VP3 spontaneously forms cytoplasmic VFLs with some liquid-like properties, however, optimal LLPS may require the full complement of proteins and RNA to be in the VF.

We demonstrated that the VFs show a high level of colocalization with VP3 (Manders’ coefficient 0.9), but a lower colocalization with the VP2 capsid protein (MC 0.6). This observation is consistent with Dalton & Rodriguez (2014), who observed a lower colocalization of VP2 with VP1, VP3 or dsRNA at early time points postinfection (13). For this to be the case, the site of virus replication (the VFs) must be distinct from the PAs of virions that we and others have observed by TEM (28). To reconcile this, we conducted CLEM, and demonstrated that the VFs were discrete electron dense regions of the cytoplasm, separate from the PAs. However, key questions remain, including whether any cellular proteins or mRNAs are recruited to the VF, and what processes

**FIG 7** VFs correlated with discrete electron-dense regions of the cytoplasm that were distinct from the PAs. DF-1 cells were seeded onto gridded coverslips, transfected with GFP1-10, and infected with the PBG98-VP1-GFP11 virus 24hpt at an MOI of 1. Eighteen hpi, cells were stained with Hoechst 33342, and cells on selected grid squares were imaged live by confocal microscopy. Scale bar, 20 μm (A). An image of the boxed region is shown enlarged (B). The same cells were then processed and imaged by TEM. The same cell from figure A and B is shown in (C). The confocal image was then overlaid on top of the TEM image and the two correlated (D). An image of the boxed region is shown enlarged (E).
occur there. For example, it remains unknown whether translation occurs in the VF, or whether components of the innate immune pathway are sequestered in the VF, as they are in the inclusion bodies of respiratory syncytial virus (29). Moreover, whether there is an ultrastructure or compartmentalization within the VF remains unknown. While the TEM images show a homogeneous electron dense region, it is possible that proteins are not evenly distributed within the VF. Imaging using superresolution microscopy would help to shed light on this. Finally, the dynamics of virion assembly and PA formation remains poorly understood. We did not observe VFs that contained visible virus particles by TEM, unlike avian reovirus. One hypothesis for this is that individual virus particles formed in the VF may be rapidly repelled into the cytoplasm due to a charge incompatibility. When multiple virions are expelled from a VF, they may accumulate together and form a lattice due to electrostatic interactions between the capsids. Alternatively, VP2 may accumulate in a VF until it reaches a critical concentration and then spontaneously forms many capsids, forming a PA. In this way, a PA may be a remnant of a VF. Birnaviridae are thought to undergo random packaging rather than selective packaging (30), and while both these models would be consistent with random packaging, much work is left to do to determine the process of virion assembly.

Our study is not without limitations. First, the GFP signal was only visible from 8 hpi using our equipment. Second, it has been shown that rotavirus VFs become less liquid over time (18). We only conducted FRAP and 1,6-hexanediol treatment at 16 hpi, and we did not compare different time points. However, that FRAP was rapid and VFs dissolved by hexanediol at this time point suggests that the VFs do not lose fluidity over time in the same way as rotavirus VFs. Third, we conducted our studies within DF-cells which are immortalized fibroblast cells, whereas the tropism of IBDV in vivo is in B cells. It is challenging to image B cells, but owing to their more spherical nature, large nucleus, and their nonadherence it is possible that the IBDV VFs within the cytoplasm have different dynamics. Finally, our TEM and CLEM experiments were only conducted at later time points in infection, and it would be beneficial to conduct a time course, including earlier time points.

In summary, Birnaviridae VFs are sites of de novo RNA synthesis, are not bound by a membrane, show properties consistent with liquid-liquid phase separation, and are distinct from the PAs observed by TEM. This new information enhances our fundamental knowledge of the replication of a family of viruses responsible for disease of economic importance to poultry and aquaculture, that could be used to develop strategies to better control disease or optimize their therapeutic application in the future.

MATERIALS AND METHODS

Cell-lines and antibodies. DF-1 cells (chicken embryonic fibroblast cells, ATCC number CRL-12203) were maintained in Dulbecco’s modified Eagle’s medium (DMEM) (Sigma-Aldrich, Merck, UK), supplemented with 10% heat-inactivated fetal bovine serum (hiFBS) (Gibco, Thermo Fisher Scientific, UK). The primary antibodies used in this study were raised against tubulin (SantaCruz Biotechnology, UK), PDI (Enzo Life Science, Farmingdale, NY), VP4 (a kind gift from Jóse Castón [CNB-CSIC, Spain]), VP2, and VP3 (31). Primary antibodies were diluted 1:100 and secondary antibodies conjugated to Alexa fluor 488-or 568 (Invitrogen, Thermo Fisher Scientific, UK) were diluted 1:500 in 4% bovine serum albumin (BSA) (Sigma-Aldrich, Merck, UK).

Viruses. The cell-culture adapted Avian Reovirus (ARV) strain, S1133, was obtained from Charles River (Wilmington, MA, USA), and a molecular clone of the cell-culture adapted IBDV strain, PBG98, was rescued as previously described (17). Three alanine residues were added to the N terminus of the GFP11 tag as a linker and a stop codon was added to the C terminus to make the amino acid sequence: AAARDHMVLHEYVNAAGIT-Stop. The nucleotide sequence encoding this was added to IBDV segment B, which encodes VP1, prior to the 3’ noncoding region, and the recombinant virus, PBG98-VP1-GFP11, was rescued as previously described (17). Stocks of the viruses were grown in DF-1 cells and titrated in 96-well plates (Falcon, Corning, UK) seeded with DF-1 cells at a density of 4 × 10⁴ cells per well. Wells were infected in quadruplicate with a 10-fold dilution series of supernatant. After 5 days, wells were inspected for signs of cytopathic effect (CPE) and the titer of virus determined as per the method by Reed and Muench (32) and expressed as Tissue Culture Infective Dose-50 (TCID₅₀).

Immunofluorescence microscopy. DF-1 cells were seeded onto coverslips (TAAB, UK) in 24-well plates (Falcon, Corning, UK) at a density of 1.6 × 10⁵ per well and transfected with GFP1-10 using lipofection 2000 (Thermo Fisher Scientific, UK), 20 4 h prior to infection with the PBG98-VP1-GFP11 virus. At the relevant time points, cells were fixed with a 4% paraformaldehyde solution (Sigma-Aldrich, Merck, UK) and then spontaneously forms many capsids, forming a PA. In this way, a PA may be a remnant of a VF.
Expression of IBDV GFP-fusion proteins in DF-1 cells. Plasmids encoding GFP fused to individual viral proteins (VP1, VP2, VP3, VP4 and VP5) were synthesized by Geneart (Thermo Fisher Scientific, UK). The GFP tag was fused to the N terminus of VP1,3,4 and 5, and to the C terminus of VP2 as the N-terminal tag did not yield a signal with this protein. The inserts were cut out of the Geneart plasmids using restriction enzymes and ligated into the multiple-cloning site of the pSF-CAG-KAN expression vector (Sigma-Aldrich, UK). DF-1 cells at 70% confluence were transfected in 24 well plates with 0.5 μg of each plasmid using Lipofectamine 2000, according to the manufacturer’s instructions. Twenty-four hours posttransfection, cells were fixed with 4% paraformaldehyde solution, permeabilized with 0.1% triton-X, and stained with DAPI. Cells were washed in water, mounted in Vectashield and imaged with a Leica TCS SP5 confocal microscope.

Live cell imaging. DF-1 cells were seeded into a Chambered number 1.0 Borosilicate Coverglass slide (Nunc, Lab-Tek, Sigma-Aldrich, Merck, UK) at a density of 8 × 10^4 per well and cultured in 1 mL of DMEM supplemented with 10% hiFBS. Twenty-four hours after seeding, cells were transfected with GFP-1-10 and then infected with the PBG98-VP1-GFP11 virus 24 hpt at an MOI of 1. Thirteen hours postsinfection (hpi), cells were treated with 10 μg/mL of actinomycin D to prevent cellular transcription for 4 h, after which 1 mM EU was added to the cells that became incorporated into newly synthesized RNA molecules. 1 h after EU treatment, cells were fixed, permeabilized with 0.1% triton-X, and stained with a Click-IT RNA imaging kit (Thermo Fisher Scientific) to visualize the EU, according to the manufacturer’s instructions. Twenty-four hours posttransfection, cells were fixed with 4% paraformaldehyde solution, permeabilized with 0.1% triton-X, and stained with DAPI. Cells were washed in water, mounted in Vectashield and imaged with a Leica TCS SP5 confocal microscope.

Visualization of 5-ethynyl uridine (EU) signal. DF-1 cells were transfected with GF1-10 and infected with the PBG98-VP1-GFP11 virus 24 h post-transfection (hpt) at an MOI of 1. Thirty-three hours postsinfection (hpi), cells were treated with 10 μg/mL of actinomycin D to prevent cellular transcription for 4 h, after which 1 mM EU was added to the cells that became incorporated into newly synthesized RNA molecules. 1 h after EU treatment, cells were fixed, permeabilized with 0.1% triton-X, and stained with a Click-IT RNA imaging kit (Thermo Fisher Scientific) to visualize the EU, according to the manufacturer’s instructions. Twenty-four hours posttransfection, cells were fixed with 4% paraformaldehyde solution, permeabilized with 0.1% triton-X, and stained with DAPI. Cells were washed in water, mounted in Vectashield and imaged with a Leica TCS SP5 confocal microscope.
Corelightive light and electron microscopy (CLEM). DF-1 cells were seeded onto gridded glass bottom dishes (iMaStTek) at a density of 8 × 10^4 per well and cultured in 1 mL of DMEM supplemented with 10% hiFBS. Twenty four hours after seeding, cells were transfected with GFPI-10 and then infected with the PBCG98-VPI-GFP1 virus 24 hpt. Eighteen hpi, cells were stained with Hoechst 33342 (Thermo Fisher Scientific) and selected grid squares were imaged live in L-15 media with a Leica TCS SP8 confocal microscope in a climate-controlled chamber. The samples were fixed in phosphate buffered 2% glutaraldehyde (Agar Scientific) for 1 h before being fixed for a further hour in aqueous 1% osmium tetroxide (Agar Scientific). The samples were dehydrated in an ethanol series; 70% for 30 min, 90% for 15 min and 100% three times for 10 min each. After infiltration of 100% epoxy resin (Agar Scientific) for 2 h, the samples were embedded and polymerized overnight at 60°C. The glass coverslips were removed with liquid nitrogen and the appropriate grid squares located. 80 nm thick sections were cut, collected onto copper grids (Agar Scientific) and grid stained using Ledom EM AC20. The specific cells imaged in the confocal were located and imaged at 100 kV in a FEI Tecnai 12 TEM with a TVIPS F214 digital camera.

SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

SUPPLEMENTAL FILE 1, MOV file, 2.3 MB.
SUPPLEMENTAL FILE 2, MOV file, 1 MB.
SUPPLEMENTAL FILE 3, MOV file, 9.6 MB.
SUPPLEMENTAL FILE 4, PDF file, 0.2 MB.

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