Title
Interrogating marine virus-host interactions and elemental transfer with BONCAT and nanoSIMS-based methods

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Abstract
While the collective impact of marine viruses has become more apparent over the last decade, a deeper understanding of virus-host dynamics and the role of viruses in nutrient cycling would benefit from direct observations at the single-virus level. We describe two new complementary approaches - stable isotope probing coupled with nanoscale secondary ion mass spectrometry (nanoSIMS) and fluorescence-based biorthogonal non-canonical amino acid tagging (BONCAT) - for studying the activity and biogeochemical influence of marine viruses. These tools were developed and tested using several ecologically relevant model systems (\textit{Emiliania huxleyi}/EhV207, \textit{Synechococcus sp.} WH8101/Syn1, and \textit{Escherichia coli}/T7). By resolving carbon and nitrogen enrichment in viral particles, we demonstrate the power of

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nanoSIMS tracer experiments in obtaining quantitative estimates for the total number of viruses produced directly from a particular production pathway (by isotopically labeling host substrates). Additionally, we show through laboratory experiments and a pilot field study that BONCAT can be used to directly quantify viral production (via epifluorescence microscopy) with minor sample manipulation and no dependency on conversion factors. This technique can also be used to detect newly synthesized viral proteins. Together these tools will help fill critical gaps in our understanding of the biogeochemical impact of viruses in the ocean.

Introduction

Marine viruses impact food webs by shaping microbial community structure, serving as agents of gene transfer, and altering the direction and magnitude of carbon and nutrient flow (Fuhrman 1999, Wommack and Colwell 2000, Suttle 2007, Weitz and Wilhelm 2012). These microscopic predators infect the oceanic primary producers (e.g., *Prochlorococcus*, *Synechococcus*, Sullivan et al. 2003, and eukaryotic phytoplankton, Brussaard 2004a) and nitrogen fixers (e.g., *Trichodesmium*, Hewson et al. 2004), as well as globally dominant pelagic (SAR11, Zhao et al. 2013 and SAR116, Kang et al. 2013) and particle-associated heterotrophic bacteria (Bacteriodetes, Holmfeldt et al. 2013) and grazers (Massana et al. 2007).

Recent technological advances have changed our ability to study and understand the ecology of marine viruses (Brum and Sullivan 2015). Metagenomics has enabled scientists to have a better understanding of viral diversity and biogeography (Angly et al. 2006, Brum et al. 2015) as well as the predictive power to connect viruses with their hosts at larger community scales (Edwards et al. 2016, Roux et al. 2016). However, quantitative estimates of viral-induced mortality and the degree to which carbon and nutrients are cycled through virioplankton remain poorly constrained. Our current estimates of viral mortality are based primarily on indirect inferences and modeling (reviewed in Fuhrman 1999, Wommack and Colwell 2000, Suttle 2005, Breitbart et al. 2008) and are insufficient to capture the spatial and temporal variability needed to properly incorporate viruses into global ocean models (Weitz et al. 2014). Furthermore, the viral shunt paradigm — the idea that viral lysis will “keep carbon small” by reducing the flow of carbon moving upwards in the food web (Wilhelm and Suttle 1999, Suttle 2007, Weitz and Wilhelm 2012) — remains virtually unquantified due to technical challenges of making such
measurements in natural populations. Recent measurements hint that virus-infected cells may be preferentially grazed (Evans and Wilson 2008, Frada et al. 2014) and that viruses, better than prokaryotes and eukaryotes, predict 89% of variability in global ocean carbon flux from the surface waters to the deep sea — with cyanophages the best predictor among them (Guidi et al. 2016). As the decades-old viral shunt paradigm is being reevaluated with conceptual models (Weinbauer et al. 2004), and with new data suggesting that viruses can increase, rather than decrease carbon export in the oceans (Vardi et al. 2012, Guidi et al. 2016), independent quantitative methods to assess viral activity and their biogeochemical influence is critical.

The development and application of single-cell resolved isotopic (nano-scale secondary ion mass spectrometry or nanoSIMS) and fluorescence-based activity (biorthogonal non-canonical amino acid tagging or BONCAT) approaches in microbial ecology have enabled scientists to directly track cell-specific microbial activity and associated influence on biogeochemical cycling in an environmental context (e.g., Musat et al. 2008, Dattagupta et al. 2009, Sheik et al. 2013, 2014, Kopf et al. 2015, Terrado et al. 2017). While the 50 nm spatial resolution of the nanoSIMS is technically sufficient for resolving individual viruses, the combination of stable isotope probing and nanoSIMS has not yet been tested for the study of carbon and nitrogen transfer between environmental viruses and their host microorganisms. More recently, the BONCAT method (Dieterich et al. 2006) coupled with fluorophore addition via copper catalyzed ‘click chemistry’ has been used to visualize translationally active microorganisms in diverse environments (Hatzenpichler et al. 2014, 2016, Samo et al. 2014); however, the adaptation of BONCAT for fluorescently tracking virus-host interactions and rates of viral production has not been explored.

In this study, we tested the feasibility and constraints of using BONCAT and nanoSIMS to directly quantify new viral production and the transfer of carbon and nitrogen from host to virus during cell lysis. These methods were validated using 3 virus-host model systems including: *Emiliania huxleyi* strain CCMP374 and its ~200nm coccolithovirus EhV207, *Synechococcus sp.* strain WH8101 and its ~80nm virus Syn1, and *Escherichia coli* and its ~50 nm phage T7) with an eye towards laying a foundation for interrogating virus-host interactions within complex microbial communities. These specific model systems were selected because their infection dynamics are well characterized and, in the case of EhV207 and Syn1, they represent ecologically relevant marine virus-host pairs. The use of stable isotope probing coupled
to nanoSIMS analysis extends the ability to study the flow and fate of carbon and nitrogen through the smallest active members of marine food webs. Viral BONCAT shows promise as a fluorescence-based microscopy method for directly quantifying the proportion of newly synthesized lytic viruses (i.e. viral production), extending the utility of relatively rapid, viral-targeted DNA stains and epifluorescence microscopy assays (Hara et al., 1991; Nobel and Fuhrman 1998). Taken together, these new single viral particle resolved isotopic and fluorescence-based assays will help fill critical gaps in our understanding of the activity and biogeochemical impact of viruses in ocean ecosystems.

Materials and Methods

Viral-host cultures and isotope experimental incubations

*Emiliania huxleyi* (CCMP strain 374) and its lytic virus EhV 207 (~200 nm capsid) was used as a eukaryotic host-virus model system. *E. huxleyi* was obtained from the National Center for Marine Algae and Microbiota, and its virus was isolated in 2001 from the English Channel (Schroeder et al. 2002) and kindly provided by M. Allen (Plymouth Marine Lab). *E. huxleyi* was grown on an enriched artificial seawater media (ESAW without silicate; Harrison et al. 1980). Batch cultures were maintained at 18°C on a 14:10 light:dark cycle at ~150 µmol photons m\(^{-2}\) s\(^{-1}\). EhV207 was propagated by infecting cultures grown in f/2 minus silicon. Upon complete host lysis, viral lysates were filtered through a 0.45-µm syringe filter to remove cellular debris and stored at 4°C in the dark. Viral abundance was determined using SYBR Gold (Life Technologies) and flow cytometry (Brussaard 2004b).

*Synechococcus* sp. (strain WH 8101, Waterbury et al. 1986) with lytic virus Syn1 (~80 nm capsid) was used as a cyanobacterial host-virus model system. *Synechococcus* WH8101 culture was kindly provided by Dr. Gabrielle Rocap (University of Washington, School Oceanography). Prior to the experiments, the culture was acclimated to growth on 100% artificial seawater Amp1 media (Rippka 1988) using a 14:10 light:dark cycle. The Amp1 was prepared as indicated by Moore et al. (2007). Syn1 virus was propagated using unlabeled batch cultures of *Synechococcus* WH8101 grown on a 50/50 mix of Amp1 and SN (Waterbury and Wiley 1988) media without isotopically labeled carbon or nitrogen under low light conditions (20 µmol photons m\(^{-2}\) s\(^{-1}\)).

Single isotope-label virus enrichment experiments were done with hosts grown on either a) \(^{13}\text{C}\)-labeled media, b) \(^{15}\text{N}\)-labeled media, or c) an unlabeled control. *Synechococcus* sp. was
grown on Amp1 media with either a) 6 mM $^{13}$C-sodium bicarbonate (NaH$^{13}$CO$_3$; 50% label) and 400 µM unlabeled ammonia (NH$_4$) (Exp. 1), b) 400 µM $^{15}$NH$_4$ (100% label) and 6 mM unlabeled-NaHCO$_3$ (Exps. 2 and 3), or c) 400 µM unlabeled-NH$_4$ and 6 mM unlabeled-NaHCO$_3$. Unlabeled cultures were used to inoculate new labeled-cultures for virus enrichment experiments. These newly inoculated cultures were then passed two times into fresh isotopically-labeled media (splitting the culture 1:5 into new media after 7 and 8 days for Exps. 1 and 3, respectively). After the second transfer, exponentially growing host cells were sampled to provide the level of host labeling prior to infection. Host cells were fixed with glutaraldehyde (0.5% final concentration) at room temperature for 15 minutes, flash frozen and stored at -80°C. Syn1 was then added at a multiplicity of infection (MOI) of ~5 for 24 hours. After 24 hours, fresh media (with the appropriate label) was used to dilute the culture approximately 1:3. Cells were lysed to completion (approximately 7 days) at which point viruses were collected. Virus samples were filtered through a 0.2-µm filter, fixed with glutaraldehyde (0.5% final concentration) for 15 minutes at room temperature, flash frozen and stored at -80°C until further analysis.

An additional short-term isotope enrichment experiment was performed with *Synechococcus* sp. and Syn1 to determine if isotope transfer could be detected in the viral particles after pulsing a labeled substrate rather than growing the cultures for many generations on labeled substrates. Cultures grown in unlabeled Amp1 media were transferred to media with 400 µM $^{15}$NH$_4$ (100% label) at a 1:5 dilution and incubated for 24 hours. After 24 hours, host cells were sampled and viruses were added at an MOI of 1. The culture was lysed to completion (approximately 5 days) at which point viral samples were collected as described above.

_E. huxleyi_ CCMP 374 was grown in ESAW media with either a) 1.2 mM NaH$^{13}$CO$_3$ (100% label) and 553 µM unlabeled-nitrate (NO$_3$) (Exp. 1), b) 553 µM $^{15}$NO$_3$ (100% label) and 1.2 mM unlabeled-NaHCO$_3$ (Exp. 2), or c) 553 µM unlabeled-NO$_3$, and 1.2 mM unlabeled-NaHCO$_3$. _E. huxleyi_ cultures were grown on labeled media for a minimum of 10 generations prior to the start of virus enrichment experiments. At the start of each experiment, exponentially growing cultures were harvested by centrifugation (3,000 x g, 10 min) to provide the level of host labeling prior to infecting with EhV207 at an MOI of 5. EhV207 viruses were added directly to the culture with no additional transfers or dilutions. Upon host lysis (~ 3 d), viral lysates were collected by filtering through a 0.45-µm syringe filter, fixed in 0.5% glutaraldehyde
for 15 minutes at 4°C, then flash frozen in liquid nitrogen and stored at -80°C until further analysis. A dual-labeled viral enrichment experiment (Experiment 3) was set up similar to single-label experiments with the exception that *E. huxleyi* was grown in ESAW media containing both 1.2 mM NaH$^{13}$CO$_3$ (100% label) and 553 µM $^{15}$NO$_3$ (100% label) for ~6 months prior to the start of virus enrichment experiments.

Adsorption control experiments were performed with *E. huxleyi* to determine the influence of fixation on nitrogen adsorption to the outside of the cell. *E. huxleyi* was grown in unlabeled ESAW. Cells were harvested and fixed in glutaraldehyde (2% final concentration) at 4°C for 15 minutes. Cells were washed 3 times with unlabeled media and resuspended in either $^{15}$NO$_3$ or $^{15}$NH$_4$ labeled media and stored at 4°C for 2 days prior to being stored at -80°C.

**NanoSIMS sample preparation and microscopy**

Aliquots (10-20 µL) of fixed virus particles and host cells were spotted separately onto custom-cut silicon wafers (WSI08-1108001, Active Business Company GmbH). Prior to spotting the samples, fiduciary marks were added to the silicon wafers using a laser microdissection (Leica LMD7000) microscope. These marks were used as reference points to locate and precisely correlate fluorescence microscopy and nanoSIMS images. After viral particles and host cells were deposited on the surface of the silicon wafers, samples were rinsed with 0.02-µm filtered water and air-dried. Viruses and hosts were stained using SYBR gold (Life Technologies, 10X final concentration; 1:1000 dilution of 10,000X stock) for 15 minutes in the dark, washed with 0.02-µm filtered water, and air-dried. Wafers were subsequently imaged by fluorescence microscopy on the LMD microscope using a FITC filter set and a 60X (for hosts cells) or 150X (for viral particles) dry objective. *E. huxleyi*, *Synechococcus sp.*, and EhV were large enough for imaging with the dry objective, but the fluorescence was not bright enough to reliably image the smaller Syn1 viral particles. Instead, Syn1 was imaged using a 100X oil immersion objective on an upright epifluorescence microscope (Olympus BX51) with FITC filter set. Water soluble vectashield (Vector Laboratories) was used as anti-fade mounting solution and subsequently removed with 0.02-µm filtered water with water prior to nanoSIMS analysis. Gold nanoparticles (100 nm, OD 1, stabilized suspension in 0.1 mM PBS; Sigma-Aldrich) were tested as possible fiduciary markers with Syn1 (Exp. 3) to assist with locating pre-imaged viruses for nanoSIMS analysis. 1 µL of gold bead suspension was mixed with the sample.
directly on the wafer. Gold beads were mapped with both reflected light filter and using a 100X oil immersion objective in the same regions the viral particles were imaged.

NanoSIMS instrument conditions and analysis

All samples were analyzed with a CAMECA NanoSIMS 50L (CAMECA, Gennevilliers, France) housed in the Division of Geological and Planetary Sciences at the California Institute of Technology. LMD marked and fluorescence microscopy mapped sample regions were located using the NanoSIMS CCD camera. Viruses were analyzed using a ∼0.8-1 pA primary Cs⁺ beam current with a nominal spot size of ∼100 nm without pre-sputtering. Host cells were first pre-sputtered with 30-70 pA primary Cs⁺ beam current for 1 frame (approximately 11 minutes) and then analyzed using a 1 pA primary Cs⁺ beam current. Five masses were collected in parallel \( \left( ^{12}\text{C}^-, ^{13}\text{C}^-, ^{14}\text{N}^{12}\text{C}^-, ^{15}\text{N}^{12}\text{C}^-, ^{28}\text{Si}^- \right) \) using electron multipliers (EMs). During two runs, \(^{12}\text{C}^{13}\text{C}^-\) and \(^{12}\text{C}^2^-\) were collected instead of \(^{12}\text{C}^-\) and \(^{13}\text{C}^-\). \(^{197}\text{Au}^-\) was also collected when testing the use of gold beads with nanoSIMS. For all analyses, the beam was rastered over a square region (ranging from 3x3 µm to 10x10 µm for viruses and 10x10 µm to 20x20 µm for host cells) for approximately 11 min per analytical plane/frame. At least 15 and 10 frames were collected per analysis for virus and host, respectively. All ion images were recorded at 256 × 256 pixel resolution with a dwell time of 10 ms/pixel. Mass calibration was done every ∼30 min for all ions.

NanoSIMS data analysis

Raw data from ion images were processed using the open-source MATLAB plugin Look@NanoSIMS (LANS; Polerecky et al., 2012). Ion images from multiple frames were corrected for EM dead time and quasi-simultaneous arrival (QSA) effect and then aligned and accumulated. In most cases \(^{14}\text{N}^{12}\text{C}^-\) was used to align images. However, for the smallest viral particles (Syn1) aligning images across multiple planes was sometimes difficult due to low count rates and \(^{197}\text{Au}^-\) was used for aligning images whenever gold nanoparticles were included in the wafer preparation (Exp. 3). Regions of interest (ROIs) were drawn using either the \(^{14}\text{N}^{12}\text{C}^-\) or \(^{15}\text{N}^{12}\text{C}^-\) ion image to identify the outline of the viral particle or host cell. All ROIs represent individual viral particles or individual host cells.
Isotope calculations of viruses and their hosts were based on raw ion counts exported from LANS (Polerecky et al. 2012). All calculations and data processing was performed in R (R Development Core Team 2013) using the package lans2r. When the molecular ions $^{13}$C$^{12}$C and $^{12}$C$_2$ were collected, the $^{13}$C/$^{12}$C ratio was calculated as half of the $R_{\text{sample}}$ from the raw ion counts of $^{13}$C$^{12}$C and $^{12}$C$_2$. Counting error was propagated to the isotope ratio by standard error propagation based on a Poisson distribution (see Appendix S1 for details). Instrumental isotope fractionation of the nanoSIMS was determined and corrected for within each run by comparing the NanoSIMS acquired $^{13}$C/$^{12}$C and $^{15}$N/$^{14}$N ratios for all combined natural abundance (unlabeled) particles analyzed (virus and host) against the expected circum-natural abundance ratios ($^{13}$C/$^{12}$C = 0.011; $^{15}$N/$^{14}$N = 0.0036; Fig. S1, Appendix S1). Due to the high isotope enrichments associated with tracer experiments, it is more intuitive to compare fractional abundances rather than isotope ratios. Ion ratios were converted to fractional abundance using the relationship $F_{\text{sample}} = R_{\text{sample}}/(1+R_{\text{sample}})$, where $F$ is the fractional abundance and $R$ is the ratio of either $^{13}$C/$^{12}$C or $^{15}$N/$^{14}$N. Errors from counting statistics and instrumental mass fractionation were propagated for conversion to fractional abundance (see Appendix S1 for more details).

**Viral BONCAT - experimental incubations, microscopy, digital image analysis and data analysis**

Bioorthogonal non-canonical amino acid tagging (BONCAT) experiments were done with *Escherichia coli* K12 (kindly provided by D. Tirrell; Caltech) and T7 (kindly provided by R. Phillips; Caltech) as well as with *E. huxleyi* and EhV (described above). Incorporation of L-homopropargylglycine (HPG), a methionine analog, into new protein has been previously demonstrated for *E. coli* (Beatty et al., 2005; Wang et al., 2008), but not for *E. huxleyi*.

*E. coli* K12 was grown at 37°C in M9 minimal medium: 0.5 g NaCl, 2.0 g of glucose, 1.0 g NH$_4$Cl, 12.8 g Na$_2$HPO$_4 \times 7$ H$_2$O, 3.0 g KH$_2$PO$_4$, 492 mg MgSO$_4 \times 7$ H$_2$O, 11 mg CaCl$_2$, 100 mg thiamine per 1 L of deionized water. HPG (Click Chemistry Tools, L-homopropargylglycine resuspended in H$_2$O; 100 µM final concentration) was added to exponentially growing *E. coli* (OD600 = 0.3) and incubated for approximately one generation (OD600 = 0.6) prior to adding virus. A control treatment without HPG was run in parallel to the experimental treatment. Phage T7 was then added (MOI of 2) and the culture was lysed to completion. Upon host lysis, viral samples were filtered through a 0.2-µm filter to remove cell debris, fixed with paraformaldehyde (PFA; 2% final concentration) for 15 minutes at 4°C and
stored in the -80°C until further analysis. For proteomics, 0.2-µm filtrate was concentrated using 100 kDa Amicon ultra centrifugal filter (EMD Millipore). Approximately 4 mL of culture was concentrated into 250-500 µL and frozen at -80°C until extraction and digestion.

For *E. huxleyi*, HPG (resuspended in DMSO; 100 µM final concentration) was added to an exponentially growing culture in unlabeled ESAW minus Si media and incubated under 14:10 LD dark cycle at 150 µmol photons m⁻² s⁻¹ for 24 hrs. A control treatment with an equivalent volume of DMSO was run in parallel to the experimental treatment. After 24 h, a subsample of host cells was harvested by centrifugation to examine the incorporation of HPG by the host. EhV207 (MOI of 5) and an additional pulse of HPG (100 µM final) were then added to the remaining culture. Upon host lysis, viral samples were filtered through a 0.45 µm filter, fixed with glutaraldehyde (0.5% final concentration) for 15 minutes at 4°C, flash frozen in liquid nitrogen and stored at -80°C until further analysis. For proteomics, 0.45-µm filtrate was concentrated using 100 kDA Amicon ultra centrifugal filter (EMD Millipore). Approximately 20 mL of culture was concentrated into 250-500 µL and frozen at -80°C until extraction and digestion.

An additional field experiment with HPG additions was performed with seawater collected from the Scripps Pier in La Jolla, California (32.87 N, 117.26 W). Surface waters were collected at 1700 h on 9 October 2016 and stored in a polycarbonate carboy at 10°C until the experiment was started at 0800 h on 10 October 2016. Seawater was placed into 250 mL incubation bottles. Treatment bottles (in replicate) were amended with 100 µM of HPG relative to negative controls (e.g., no HPG additions). Incubations were maintained at 18.7°C (October pier temperature average: https://tidesandcurrents.noaa.gov) under a 12h:12 h light:dark cycle at 100 µmol m⁻² s⁻¹ of photosynthetically active radiation (PAR). Samples for time points T₀, T₄₈, and T₇₂ h were taken to quantify virus concentration and incorporation of HPG. For this initial field test, fast growing populations with lytic cycles ranging from 12-48 hours were targeted. Therefore, sampling was delayed by 48 hours in order for host cells to grow in the presence of HPG and viral lysis to occur. Viral concentration was determined using traditional epifluorescence microscopy methods (Nobel and Fuhrman 1998; Patel et al., 2007). In short, glutaraldehyde fixed samples (1 mL) were gently filtered (0.5 mmHg) onto 0.02 µm Anodisc filters with HA backing filters, dried and stained for 15 min with SYBR Gold (0.25% final concentration). After staining, the filters were dried in the dark and mounted onto slides with
0.1% p-phenylenediamine anti-fade mounting solution. Samples for determining HPG incorporation into viruses were prepared differently in order to concentrate viruses from a larger volume of seawater and reduce background fluorescence for image analysis. Seawater samples (8 mL in duplicate) were concentrated using an Amicon Ultra Centrifugal filter (Millipore, 100K, UFC810096) to approximately 30 µL following the manufacturer’s recommendations (i.e., 4000 rcf for 15 minutes). Samples were fixed with glutaraldehyde (0.5% final concentration), flash frozen and stored at -80°C until further analysis.

Prior to the copper (I)-catalyzed cycloaddition, or click reaction, all samples (cultured and natural seawater) were spotted (2-5 µL) directly onto Teflon printed glass slides (Electron Microscopy Sciences, PTFE Printed Slides), air-dried, rinsed with 0.02-µm filtered water and air-dried again. Samples were put through an ethanol dehydration series (50:50, 80:20, 96:4 v/v EtOH:H2O) prior to incubation with freshly prepared click solution (4 µL of dye-premix [CuSO4, 0.1 mM; THPTA, 0.5 mM; TAMRA-azide fluorophore, 2 µM] was added to 245 µL of buffer solution [sodium ascorbate, 5 mM; aminoguanidine hydrochloride, 5 mM; 1X PBS]). Click solution (20 µL) was spotted onto the sample and incubated in a humid chamber (a sealed 50 mL falcon tube with water on tissue paper in the bottom) at room temperature in the dark for 30 minutes. Following incubation, the sample was washed with 1X PBS, water and 50:50 v/v EtOH:H2O. Samples were counterstained for 15 min with SYBR Gold (0.25% final concentration), washed with water, and air-dried.

Viral samples were analyzed with an Olympus BX51 upright epifluorescence microscope using a 100X objective (UPLFLN 100X 1.3 NA). Digital images were captured with a QIClick 12bit CCD digital camera using two filter sets: a FITC filter set (excitation 480/40 nm; emission 535/50 nm) and a Cy3 filter set (excitation 545/30 nm; emission 620/60 nm). Incorporation of HPG by viruses was quantified from these digital images by measuring the fluorescence signal of the azide-containing TAMRA fluorophore (i.e., click signal in Cy3) relative to the fluorescence signal from the SYBR Gold (i.e., DNA signal in FITC) within individual viral particles using a custom Matlab analysis pipeline (see Appendix S2 for more details and link to analysis code). In short, FITC and Cy3 images were corrected for uneven background illumination (based on a disk structuring element) and aligned with one another. Regions of interest (ROIs) were selected by applying a signal threshold with a size cutoff to the FITC image. In fluorescence microscopy, the resolution (the minimum distance required to resolve two particles from one another - \( R = \frac{\lambda}{2} \) /
[2xNA] = 0.184 μm - set by the objective numerical aperture [N; UPLFLN 100X 1.3N] and the wavelength of light used [λ; 480 nm for FITC filter]) does not limit the ability to accurately count small fluorescently labeled objects as long as they are separated from one another by more than the resolution distance. The centroid of each fluorescent ROI can therefore be used to locate the ROI with nanometer precision, far beyond the resolution limits of the microscope (Churchman et al., 2005, Yildiz and Selvin 2005, Waters 2009). Image analysis of non-clicked T7, Syn1 and EhV was used to confirm that individual viral particles (across a range of sizes; 50-200 nm) would be identified as ROIs based on their FITC signal from SYBR Gold DNA staining. Finally, green (FITC) and red (Cy3) fluorescence values were recorded for each ROI and data was normalized by image exposure time. Histograms of the red-to-green (R:G) ratios of individual viral particles (at least >1000 particles per treatment) were used to visualize differences in the viral populations from each experiment and/or treatment. A Wilcoxon Rank-Sum test was used to determine if the R:G ratio distributions between treatments with and without HPG were significantly different from one another. In order to quantify the number of newly synthesized viral particles and evaluate confidence intervals around our estimates, we coupled a rank-sum test with a bootstrap approach to determine the probability that a random sample drawn from the sample R:G distribution is greater than a random sample drawn from the negative control R:G distribution (see Appendix S2 for details and link to code). To assess under what conditions our theoretical approach accurately captures viral production (e.g., the percent of newly synthesized viral particles), we then tested our ability to determine the number of newly synthesized viruses from a simulated data set of T7 by this method. For the field data, viral production rates were calculated using the equations from Noble and Fuhrman (2000), but rather than track the disappearance of fluorescently labeled viruses, we essentially tracked the disappearance of unlabeled viruses as fluorescently labeled viruses were produced (see Appendix S2 for details). Viral turnover (days) was estimated by dividing the average number of viral particles by the virus production rate.

**Proteomics**

Protein was extracted from concentrated T7 and EhV in a 1:1 ratio with lysis buffer (200 mM Tris-HCl pH 8, 4% SDS) and a protease inhibitor (1 tablet for 10 mL of reaction) in a screw-cap tube. The sample was boiled for 15 minutes in water bath. Protein was quantified...
using Bradford Protein Assay (Bio-Rad Laboratories). A filter-aided sample preparation (FASP) procedure was used to clean (e.g., remove SDS) and digest proteins in preparation for proteomic analysis (Wiśniewski et al. 2009). Eluted peptides were lyophilized, resuspended in 100 μL of 0.2% formic acid, and desalted by HPLC with an Optimized Technologies C8 peptide Macrotrap (3x8mm; 200 μg maximum capacity). Desalted peptides were then lyophilized and stored at -20°C until mass spectrometry analysis.

Liquid chromatography-mass spectrometry was carried out on a Q-Exactive HF Orbitrap mass spectrometer (Thermo Fisher Scientific, Bremen, Germany) in the Proteome Exploration Laboratory at the Beckman Institute at Caltech. For the LC-MS/MS experiments, samples were loaded onto a custom-packed column (ReproSil-Pur 120 C18_AQ 1.9 μM; 50 μm internal diameter x 20 cm long). The column was enclosed in a column heater operating at 60°C. The peptides were separated with a 120 min elution gradient (0–40% solvent B) at a flow rate of 220 nL min⁻¹. The Orbitrap was operated in data-dependent acquisition mode to alternate automatically between a full scan (m/z 300–1700) and 12 subsequent HCD (higher energy collisional dissociation) MS/MS scans in the orbitrap.

Thermo raw files were processed and searched with MaxQuant (v. 1.5.3.30) (Cox and Mann 2008, Cox et al. 2011). For E. coli / T7 samples, all UniProt E. coli entries (4311 sequences) and all UniProt T7 entries (57 sequences) were searched along with a contaminant database containing proteins such as keratin and trypsin (247 sequences). For E. huxleyi / EhV207 samples, all UniProt E. huxleyi entries (35700 sequences) and EhV207 entries (470 sequences) were searched along with the same contaminant database as used for the E. coli search. Trypsin was specified as the digestion enzyme with up to two missed cleavages allowed. Protein N-terminal acetylation (+42.0106), Met oxidation (+15.9949), Met->Aha conversion (-4.9863), and Met->Hpg conversion (-21.9877) were specified as variable modifications. Carbamidomethylation of cysteine (+57.0215) was specified as a fixed modification. Match between runs, LFQ, and iBAQ were enabled. Precursor ion tolerance was 4.5 ppm after recalibration and fragment ion tolerance was 20 ppm. Additionally, a decoy database was constructed to determine the false discovery rate by reversing the target database. Using the decoy database, the protein, peptide, and modified site level false discovery rates were set to be less than 1%.
Viral BONCAT sensitivity testing

The ability to detect newly synthesized viral particles via Viral BONCAT depends on the number of HPG substitutions per viral particle and the influence of an HPG substitution on the measured R:G ratio. By modeling variations in these parameters (based on knowledge from T7), we explored the sensitivity of Viral BONCAT to detect newly synthesized viral particles under different scenarios of viral activity (i.e., the fraction of the viral pool that is actively turning over or % active). The model is briefly described below, but additional details and links to download the code can be found in Appendix S2. The number of detected substitutions ($n_{\text{subs}}$) and the impact of an HPG substitution on the R:G ratio ($d_{\text{RG}}$) were estimated from T7 data (12 and 0.0491, respectively). However, depending on where the substitution sites are located and the size of the viral proteome, these parameters may vary; therefore, we considered a range of values for each parameter. We simulated the production of new viruses by adding a random number of methionine substitutions (modeled with a binomial distribution each increasing the R:G fluorescence ratio by a specific amount) to a subset (x percent) of the negative control R:G ratios with a different number of particles in play. Using 1000 particles drawn from the T7 negative control distribution, we explored how the following conditions impacted the probability distributions of HPG-labeled particles: $n_{\text{subs}} = 1, 5, 12$; $d_{\text{RG}} = 0.01, 0.03, 0.05$; % active = 5, 25, 50, 75, 100. In an effort to model the variability that individual substitutions may have on the actual changes in R:G, we also explored how percent variations in $d_{\text{RG}}$ (± % of the average $d_{\text{RG}}$) for individual substitutions influenced the shape of the probability distributions for HPG-labeled viruses.

Results

Quantification of stable isotopes in viral particles using stable isotope probing and nanoSIMS

In both the cyanobacterial and eukaryotic algal-virus model systems, we were able to visualize enrichment in carbon-13 and nitrogen-15 in individual viral particles on the nanoSIMS and trace the transfer of $^{13}$C (bicarbonate) and $^{15}$N (nitrate or ammonium) incorporated by the photosynthetic hosts to their viruses after host lysis (Fig. 1). Both sets of virus-host pairs showed enrichment in stable isotope amended treatments relative to unlabeled controls (Figs. 2, 3 and 4). As a result of slight differences in the length of time grown in the presence of the isotopically
labeled substrate (see Methods for details), Synechococcus sp. and E. huxleyi C and N enrichment varied across different experiments. However, independent of the absolute level of enrichment detected, the transfer of $^{13}$C and $^{15}$N-labeled biomolecules from host to newly synthesized viral particles was reproducible (Figs. 2, 3 and 4) in both single and dual isotope ($^{13}$C and $^{15}$N) labeling experiments.

For both model systems the isotopic enrichment in the viral particles was similar to that of the host whenever the host culture was fully acclimated to isotopically labeled nutrients (Figs. 2 and 4). Synechococcus sp. cells and Syn1 particles (~80 nm capsid) recovered from stable isotope probing experiments with $^{13}$C-bicarbonate, had an average fractional abundance ($^{13}$C/$^{13}$C+$^{12}$C) of 21± 1.0 at % (atom percent) and 15 ± 4.7 at% $^{13}$C, respectively, approximately 15 times greater than natural abundance (~1.1 at% $^{13}$C ; Fig. 2, Exp.1). After amendment with ammonium ($^{15}$N-labeled), measured populations of Synechococcus sp. and Syn1 had an average fractional abundance of 63 ± 1.7 at% and 75 ± 7.1 at% $^{15}$N (Exp. 2) and 82 ± 2.0 at% and 69 ± 8.5 at% $^{15}$N (Exp. 3), respectively, approximately 200 times over the natural abundance value of ~0.36 at% $^{15}$N (Fig. 2). NanoSIMS analysis of Synechococcus sp. and Syn1 from unlabeled control experiments demonstrated these particles had fractional abundances close to natural abundance (Synechococcus sp. was 1.11 ± 0.03 at% $^{13}$C and 0.365 ± 0.02 at% $^{15}$N; Syn1 was 1.19 ± 0.07 at% $^{13}$C and 0.37 ± 0.19 at% $^{15}$N for $^{13}$C and $^{15}$N, respectively).

After incubation with $^{13}$C-bicarbonate, E. huxleyi and its virus EhV207 (200 nm capsid) had an average fractional abundance of 30 ± 1.3 at% and 30 ± 4.1 at% $^{13}$C, respectively, approximately 30 times over natural abundance (Fig. 3, Exp.1) and after incubation with $^{15}$N-nitrate, the eukaryotic host and recovered EhV207 virus had an average fractional abundance of 81 ± 2.0 at% and 72 ± 2.7 at% $^{15}$N respectively, approximately 200 times over natural abundance (Fig. 3, Exp. 2). After incubation with both $^{13}$C-bicarbonate and $^{15}$N-nitrate, E. huxleyi and EhV207 had an average $^{13}$C fractional abundance of 15 ± 1.0 at% and 15 ± 2.9 at% $^{13}$C, respectively, and an average $^{15}$N fractional abundance of 95 ± 1.2 at% and 85 ± 2.5 at% $^{15}$N, respectively (Fig. 3, Exp. 3). E. huxleyi and EhV in unlabeled controls remained at natural abundance (E. huxleyi fractional abundances were 1.1 ± 0.07 at% $^{13}$C and 0.37 ± 0.007 at% $^{15}$N; EhV207 fractional abundances were 1.0 ± 0.27 at% $^{13}$C and 0.37 ± 0.07 at% $^{15}$N).

Short-term (24-hour) stable isotope probing experiments conducted with Synechococcus sp. and its virus Syn1 (Fig. 4) are more analogous to the type of experiment possible in the field.
Even with this relatively short-term incubation, detection of the transfer of $^{13}$C and $^{15}$N-labeled macromolecules between host bacteria and virus was possible by nanoSIMS. Notably, there was an offset in $^{15}$N-enrichment between paired host and virus (with greater enrichment in the virus), with Synechococcus sp. and Syn1 containing an average fractional abundance of $19 \pm 2.8$ at% and $64 \pm 4.9$ at% $^{15}$N, respectively. This offset was not observed in the acclimated Synechococcus cultures grown for multiple generations in the presence of the isotope label (Fig. 2).

The potential for abiotic adsorption of the $^{15}$N-labeled nitrate and ammonium to the cell surface was experimentally tested in formaldehyde killed controls and determined to be minimal, with no significant contribution to the experimental enrichment signal (Fig. S2). These findings are consistent with the results from SIMS $^{15}$NH$_4$ adsorption control experiments reported in Orphan et al. (2009) and Dekas et al. (2009).

Viral particles had significantly lower ion counts (within an analytical frame as well as across analytical frames) relative to the host cells (Fig. 1, Fig. S3). The larger eukaryotic viral particles (~200 nm, EhV207) generated average 582 and 583 $^{12}$C$^-\ and\ ^{14}$N$^{12}$C$^-\$ counts per particle per frame while the smaller viral particles (~70 nm, Syn1) yielded an average of 438 and 209 $^{12}$C$^-\ and\ ^{14}$N$^{12}$C$^-\$ counts per particle per frame. Host cyanobacteria and algal cells had much higher average C and N counts per cell per frame ($E.\ huxleyi = 8.7x10^5\ ^{12}$C$-,\ 13.5x10^5\ ^{14}$N$^{12}$C$-$; Synechococcus sp. = $8.8x10^4\ ^{12}$C$-,\ 2.3x10^5\ ^{14}$N$^{12}$C$-$), consistent with increased biovolume. As a result of the low ion yield for the submicron viral particles, the statistical error for our virus stable isotope ratios was much larger than for host cells (See Appendix S1 for details on error propagation). Merging data across multiple analytical frames increased the total counts to $10^4$ and $10^3$ for EhV207 and Syn1, respectively, reducing the error around each estimate (Fig. S3, Appendix S1). Overall, our data indicate that the propagated relative error (1$\sigma$) from counting statistics on the $^{13}$C and $^{15}$N fractional abundance estimate of an individual viral particle ranges between 1-3 atom % (Figs. 2, 3, and 4). The total diversity (population standard deviation) in the fractional abundance isotopic composition of the viral particles in each tracer acclimated culture system ranged from 2.5-3.7% for EhV207 (n=214 particles measured) and 4.9-9.7% for Syn1 (n=260 particles measured). See Appendix S4 for additional details on methodological considerations for viral isotope measurements.
**Viral-BONCAT: detection of HPG incorporation in cultured virus-host models using epifluorescence microscopy**

BONCAT coupled with fluorophore addition via copper (I)-catalyzed azide-alkyne cycloaddition (i.e. click chemistry) was used successfully to demonstrate the transfer of HPG, a methionine analog, from newly synthesized proteins in cultured host cells to their viruses during lytic infection. Initial BONCAT experiments were conducted with co-cultures of *E. huxleyi* and its virus EhV207 as well as with *E. coli* and an infecting T7 phage. Incorporation of HPG by host cells (*E. huxleyi* and *E. coli*) was confirmed by epifluorescence microscopy using standard BONCAT protocols (Hatzenpichler et al., 2014; Figs. 5F and 6F). Consistent with previous reports, HPG addition (100 µM final concentration) did not significantly alter host growth relative to incubations without HPG (Fig.S4; Dieterich et al. 2006, Bagert et al. 2014, Hatzenpichler et al. 2014, 2016). After lysis of HPG-labeled host lysis, individual viral particles were analyzed for incorporation of HPG using our modified BONCAT protocol and visualized by epifluorescence microscopy. The fluorescence intensity and number of viral particles were quantified from digital images by measuring the fluorescence signal of the azide-containing TAMRA fluorophore (R = click signal in Cy3) relative to the fluorescence signal from the SYBR Gold (G = DNA signal in FITC) within individual viral particles. Fluorescence image analysis of T7 (50 nm), Syn1 (80nm) and EhV (200 nm) confirmed that individual viral particles across a range of sizes could be identified in digital images based on their FITC signal from SYBR Gold stain (Fig. S5). These cultured viruses were also used to determine an appropriate size cutoff that could be applied during digital image analysis. The minimum size for all viruses considered was constrained to 1 pixel (0.1159 µm) determined by CCD camera resolution (QIClick 12bit) and we conservatively chose an upper size limit of 0.45-µm (approximately 4 pixels; Fig. S5) to capture viral particles that were < 200 nm. While these sizes are not reflective of actual capsid size, our analyses show that the application of these cutoffs effectively captured the particle size range of interest (Fig. S5).

The red-to-green (BONCAT-cy3/DNA-Sybr Gold) fluorescence ratios (R:G) of all viral particles were pooled within treatments to compare the distribution of R:G ratios across treatments (Figs. 5 and 6). For both the EhV and T7 populations, a higher R:G ratio was observed in viral particles produced from host cultures incubated with HPG relative to the negative controls (host incubation without HPG) (mean (µ); $\mu_{HPG} = 0.31$, $\mu_{NEG} = 0.08$ and $\mu_{HPG} =$
Wilcoxon Rank-Sum test was used to confirm the negative control and positive treatment distributions were significantly different from one another (p-values < 0.05). Using probability statistics coupled to bootstrapping, we estimated the percentage of newly synthesized particles. For T7, 99.2% of viral particles were designated as HPG-labeled (e.g., newly synthesized) with a lower and upper 95% confidence interval of 98.9% and 99.4%, respectively. For the eukaryotic virus EhV, 94.5% of viral particles were designated as HPG-labeled (e.g., newly synthesized) with a lower and upper estimate of 91.2% and 97.2%, respectively.

**Detection of HPG incorporation into newly synthesized viral proteins via proteomics**

To independently confirm the successful transfer of host-associated HPG-labeled proteins or peptides into infecting viral particles, we conducted proteomic experiments on BONCAT-labeled *E. coli* T7 and *E. huxleyi* EhV207 and ran a targeted screen for HPG-labeled viral proteins. Proteomic analysis of the concentrated viral lysates confirmed the incorporation of HPG into select viral proteins. Here positive identification of peptide-spectrum matches was determined where the methionine (Met) in the peptide was replaced with an HPG, which results in a -21.9877 Da mass shift.

In the *E. coli* and T7 BONCAT experiment, 8 different T7 proteins associated with viral particle structure (e.g., procapsid and capsid formation as well as packaging the DNA into the capsid) were found to contain 12 HPG substitutions (Table 2). For the *E. huxleyi* and EhV 207 BONCAT experiment, 5 different EhV 207 proteins, including the major capsid protein and several putative uncharacterized proteins, were observed to contain 6 HPG substitutions. The annotated spectra for the HPG-labeled proteins are available in Appendix S3. The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE (Vizcaíno et al. 2016) partner repository with the dataset identifier PXD006580.

**Sensitivity of fluorescence detection for Viral BONCAT**

In BONCAT, the fluorescence intensity should be proportional to the total number of HPG substitutions for methionine within the viral proteome after the incubation. To assess the sensitivity of the BONCAT fluorescence assay to resolve newly synthesized viral particles, we ran an *in silico* analysis examining the influence of the number of Met substitutions and the R:G
ratio shift per substitution on the resultant R:G distributions (Fig S6, S7). By combining the T7 proteomics data with the R:G ratio distributions from the ~50 nm T7 particles after BONCAT labeling, we derived estimates for the number of detected substitutions (n\textsubscript{subs}) and the impact of an HPG substitution on the R:G ratio (d\textsubscript{RG}). 12 HPG substitutions were detected in the T7 proteome with a corresponding estimated mean shift in R:G ratio of d\textsubscript{RG}=0.049 per substitution (based on the mean difference in R:G between HPG labeled sample and negative control).

Assuming substitutions are binomially distributed, the number of average substitutions affected the shape of the expected R:G histogram, with an increasing number of substitutions leading to a wider R:G spread (Fig. S6). We also observed a shift in the HPG-positive distribution by accounting for different levels of variability around the d\textsubscript{RG} (Fig. S7). A higher level of variation (i.e., 60\%) resulted in a distribution with a more skewed tail.

To test our ability to accurately measure viral production, we modeled our T7 BONCAT data assuming a specific level of activity (e.g., the percent of the viral population actively turning over) across a wide range of the aforementioned parameters, and then determined if it was possible to back calculate the starting activity levels using the rank-sum based activity statistic (see Appendix S2 for more details) without prior knowledge of the input parameters (Fig. 7). This exercise revealed that it is possible to accurately estimate the number of newly synthesized viral particles using Viral-BONCAT in situations where >25\% of the viral population is actively turning over and there is >1 methionine substitution per viral particle with a resulting shift in R:G per substitution (d\textsubscript{RG}) of >0.03 (Fig. 7, S6, S7). Below this level of substitution and population turnover, we were unable to accurately calculate viral activity, as the small shift in the R:G per substitution resulted in greater error around the production measurement (Fig. 7).

Accurate estimates of production can be derived with a d\textsubscript{RG} of 0.01, as long as there are a greater number of substitutions per viral particle (i.e., n\textsubscript{subs} ≥ 5), or the fraction of the population turning over is higher (i.e., % active ≥ 25\%). This distinction between ‘background noise’ (e.g., BONCAT negative distributions) and the distribution associated with BONCAT positive particles becomes even more pronounced with higher d\textsubscript{RG}s (on the order of the 0.049 observed for T7; Fig. S6, S7). Based on our observations from treatment replicates, we also determined how systematic error in the observed R:G ratios due to systemic shift in signal intensity of the measured green fluorescence in either the samples themselves, or in the positive control, could limit the ability to measure viral production using BONCAT (Fig. S8). While the
limitations of a low number of substitutions and/or low R:G shifts per substitution remain the same, taking this systematic error into consideration increases the confidence intervals for the activity estimates.

**Estimating viral production from BONCAT**

Incubations using coastal seawater amended with HPG (100 µM) were conducted to assess the utility of Viral-BONCAT for estimating viral production in the field. In these experiments, the starting virus concentration was $2.63 \times 10^7 \pm 2.21 \times 10^6$ particles mL$^{-1}$. After 48 hours and 72 hours, virus concentrations were $2.72 \times 10^7 \pm 5.55 \times 10^6$ and $2.30 \times 10^7 \pm 5.08 \times 10^6$ particles mL$^{-1}$ (averages of replicates), respectively. Histograms of R:G ratios revealed that the population of viruses produced in the HPG treatments had a higher mean R:G ratio relative to the negative controls after 48 and 72 hours ($\mu_{\text{HPG}} = 0.2, \mu_{\text{NEG}} = 0.1$; Fig. 8, Table 3). A Wilcoxon rank-sum test was used to confirm the negative control and positive HPG treatment distributions were significantly different from one another (p-values < 0.05). We observed greater overlap in the R:G ratios between the negative controls and the HPG treatments from the field experiments compared to the Viral-BONCAT with our cultured model organisms. There was also a significant difference in R:G ratio distributions in HPG treatments between 48 hours and 72 hours (p-value < 0.05). Using probability statistics coupled to bootstrapping, we estimated the number of newly synthesized particles at each time point (combining both replicates). After 48 hours, 85.8% of viral particles were designated as HPG-labeled (e.g., newly synthesized) with a lower and upper 95% confidence interval of 78.7% and 91.1%, respectively. After 72 hours, 78.5% of viral particles were designated as HPG-labeled (e.g., newly synthesized) with a lower and upper 95% confidence interval of 68.2% and 86.7%, respectively. Virus production rates (calculated from the 48-hour data only) were estimated to be $2.1-3.0 \times 10^7$ particles mL$^{-1}$ day$^{-1}$ translating to a viral turnover of 0.8-1.2 days (Table 4).

**Discussion**

Viruses are integral to microbial food webs and play an important role in the fate and cycling of carbon and nutrients in the ocean. While our understanding of the impact of viruses in ocean ecosystems has increased through the use of metagenomics, model systems, and microscopy, we currently lack methods to directly assess the rates of new virus production and
the transfer of carbon and nitrogen derived from active microbial host cells. This study introduces two complementary microanalytical imaging methods using nanoSIMS and BONCAT that address this knowledge gap, enabling the direct quantification of newly synthesized viral particles and information about host-derived carbon and nitrogen sources within environmental samples. The ability to fluorescently tag newly synthesized host and viral proteins by BONCAT directly in environmental samples represents an easy to implement and versatile method for quantifying new viral production by epifluorescence microscopy. Likewise, the analyses of Carbon-13 and/or Nitrogen-15 enrichment in viral particles using nanoSIMS provides fundamental compositional data about newly produced viruses, including specific information regarding the source and transfer of recently assimilated carbon and nitrogen by host cells during viral infection and lysis. Here we discuss the development and optimization of Viral-BONCAT and nanoSIMS and demonstrate their potential for marine microbial ecology research using three diverse bacterial and algal host-virus model systems, representing a range in viral particle sizes and metabolic capabilities. We further demonstrate the use of BONCAT in natural field populations for assessing new virus production in a pilot study using surface water collected off of the coast of Southern California.

**Tracking elemental exchanges between viruses and their hosts via NanoSIMS**

Over the past 15 years, the use of secondary ion mass spectrometry (SIMS and nanoSIMS) for measuring the stable isotope composition of microbial cells from environmental samples has become a valuable tool for assessing the ecophysiology and specific roles of microorganisms in biogeochemical cycles (e.g., Orphan et al. 2001, Cliff et al. 2002, Musat et al. 2008, Dekas et al. 2009, Woebken et al. 2012, Eichner et al. 2017). To date, nanoSIMS-facilitated research in microbial ecology has focused on the analysis of bacteria, archaea and microeukaryotes. However, the use of high-resolution secondary ion imaging for analyzing the isotope composition of environmentally relevant nanometer scale viruses has not been developed. Given that individual viral particles are near the 50 nm resolution limit of the nanoSIMS and the isotopic analysis of these nanometer-sized particles introduces a number of analytical challenges, we initially assessed whether the small size and C and N content within individual viruses could be detected by nanoscale secondary ion imaging with cesium. Additionally, we assessed whether the secondary ion yield from viruses was sufficient for
detecting stable isotope $^{13}$C and $^{15}$N enrichment after growth of the host cells with $^{13}$C and $^{15}$N-labeled substrates.

Using two well-defined model photosynthetic eukaryotic (E. huxleyi) and cyanobacterial (Synechococcus sp. WH8101) hosts and their viruses (EhV207, ~200 nm capsid diameter; Syn1, ~80 nm capsid diameter), we demonstrated the ability to not only resolve viral particles using nanoscale secondary ion imaging, but also successfully measure $^{13}$C and $^{15}$N enrichment in individual viral particles after host lysis. Initial tests were based on host cells grown for multiple generations in the presence of $^{13}$C-bicarbonate and a nitrogen source (either $^{15}$N-ammonium for Synechococcus sp. WH8101 or $^{15}$N-nitrate for E. huxleyi) followed by infection and viral particle recovery after lysis. NanoSIMS analysis of EhV207 and Syn1 produced a detectable and stable ion yield and sufficient $^{12}$C-, $^{13}$C-, $^{12}$C$^{14}$N- and $^{12}$C$^{15}$N- ions for stable isotope ratio measurements from individual viral particles (Figs. 2, 4). By increasing the number of cycles and length of analysis time for each frame (i.e., dwell time), combined with the use of isotopically enriched host cells, we obtained sufficient precision to reliably differentiate $^{13}$C and $^{15}$N-enriched viral particles above the negative controls and background. Merging the data across individual frames increased the total ion counts for individual viral particles and reduced the error around each estimate (Fig. S3, Appendix S1). Alternative analytical approaches to increase the secondary ion yield by increasing the intensity of the Cs+ primary ion beam current resulted in rapid sputtering and loss of these submicron-sized particles (see Appendix S1 for additional details).

The degree of variability and statistical uncertainty in the $^{13}$C and $^{15}$N nanoSIMS measurements was related to particle size, with greater variability in the measured $^{13}$C and $^{15}$N enrichment for the 80 and 200 nm viral particles (both within and across viral ROIs) relative to their larger host cells. Additionally, there was greater variability in the measured $^{13}$C and $^{15}$N for the smaller Syn1 viruses relative to EhV207 viruses (Figs. 2, 3, 4). These observations are consistent with our expectations based on the counting statistics. Incorporating this variability as propagated error (1σ) around each estimate translates to approximately 1-3 atom % uncertainty in the fractional C and N abundance measurements for individual viral particles. While nanoSIMS analysis can likely be extended to analyze viruses smaller than Syn1, the error around individual particle measurements will increase as the total ion counts per virus target decreases. Despite the inherent uncertainty in the nanoSIMS stable isotope analysis, this method is still able
to reliably differentiate $^{13}$C and $^{15}$N enriched viral particles after labeling of host cells from control populations of unlabeled viruses and, as discussed below, is also capable of resolving the proportion of enriched viruses from the total viral population from short term stable isotope probing experiments.

Overall, these culture-based experiments demonstrate that it is possible to measure $^{15}$N or $^{13}$C enrichment in viral particles above natural abundance values. Due to the low C and N ion yield per particle, however, also it is important to note that the uncertainty surrounding measurements of individual viral particles by nanoSIMS is large and complicates interpretation of the specific level of carbon or nitrogen enrichment. The absolute level of enrichment for an individual nanometer scale viral particle is confounded by high error from counting statistics and may also be influenced by inherent variability in the host $^{13}$C or $^{15}$N enrichment associated with growth rate differences and/or temporal lags between $^{13}$C/$^{15}$N-substrate uptake and viral lysis events. Therefore, the power of the nanoSIMS stable isotope tracer experiments is in obtaining quantitative estimates for the proportion of viruses that have been produced directly or indirectly from primary productivity (i.e., from infection of active autotrophic host cells) or from host organisms metabolizing a specific nutrient (e.g., nitrogen) or carbon source out of the total viral population recovered from the environment, rather than providing a direct comparison of the magnitude of enrichment levels between host microorganisms and virus.

The initial nanoSIMS lab experiments using fully $^{13}$C and $^{15}$N labeled host cells were valuable for our initial development and testing of this method. However, in order to track viral-host dynamics in naturally occurring communities using stable isotope probing, pre-enrichment of environmental host cells over many generations is not feasible. As such, we also tested the sensitivity of nanoSIMS to detect $^{15}$N transfer between the *Synechococcus* sp. host and Syn1 after short (24-hour) incubation times with $^{13}$C-bicarbonate and $^{15}$N-ammonium. These experiments verified the ability to resolve $^{13}$C and $^{15}$N enrichment within the 80 nm Syn1 particles after a short-term incubation experiment. Notably, the level of $^{15}$N-enrichment of Syn1 (64 at%) was offset from values measured in the host *Synechococcus* (19 at%), a phenomenon that was not observed in the acclimated cultures grown for multiple generations in the presence of the isotope label (Fig. 2). The reason for this offset is not understood at this time. However one possible explanation is that the host microorganisms still retained a significant fraction of “older” unlabeled biomass resulting in partial enrichment of the host after the 24 hour
incubation, whereas the viral particles produced after infection at the 24-hour mark are comprised heavily of newly synthesized macromolecules synthesized during the incubation, enriched in the isotopically labeled substrate. Alternatively, this difference might also be explained by the offset in timing between the measurement of the host cells (subsample collected just prior to infection) and viruses recovered after host lysis. It is possible that over the 3-day infection period, the *Synechococcus* cells continued to assimilate additional $^{15}$N prior to cell lysis, resulting in additional $^{15}$N enrichment of the host. This temporal offset between host cells and virus is unavoidable in field measurements, as host cells can no longer be measured after lytic infection.

Despite the caveats associated with the stable isotope labeling approach, the results from our experiments demonstrate the ability to combine stable isotope probing with nanoSIMS for quantifying the fraction of $^{13}$C or $^{15}$N enriched viruses produced after infecting metabolically active hosts following assimilation of a specific isotopically labeled substrate. Coupled with $^{13}$C-bicarbonate or different forms of $^{15}$N-labeled nitrogen not only allows for the quantification of newly produced viral particles, but also offers a mechanism for directly linking viruses with the cycling of specific carbon and nitrogen substrates in the environment. This includes assessing the proportion of newly fixed carbon in autotrophic hosts that is shunted into the free viral fraction over time. The differentiation of isotopically-enriched viruses from unlabeled viral particles during short term SIP experiments indicates that tracking the dynamics of substrate assimilation by environmental host microorganisms and associated viral production should be possible in the field.

*Estimating viral production using Viral BONCAT*

Accurate estimates of viral production in the ocean are critical for determining the influence of viral lysis on phytoplankton and bacterial mortality. A variety of methods have been employed to estimate viral production; these include, but are not limited to the incorporation of radiolabeled substrates into viral DNA (e.g., $^{3}$H-thymidine or $^{32}$P; Steward et al. 1992a,b, Fuhrman and Nobel 1995), TEM-based observations of visibly infected cells (e.g., Proctor et al. 1993, Steward et al. 1996), quantifying viral decay rates (e.g., Heldal and Bratbak 1991, Bratbak et al. 1992), using fluorescently labeled viruses as tracers (e.g., Noble and Fuhrman 2000, Ohno

The viral-BONCAT and nanoSIMS methods developed in this work complement these earlier approaches while also providing additional information about viral production in ocean systems, either through linking viruses to specific host metabolisms using stable isotope labeling or by concomitant protein tagging of active viruses and their corresponding host cells. The aforementioned stable isotope analysis of single viral particles by nanoSIMS provides quantitative, compositional data about the proportion of newly synthesized viruses and specific elemental exchanges between hosts, viruses, and the dissolved organic carbon/nitrogen pools in the ocean. However, due to the specialized instrumentation and detailed sample preparation, this method is inherently lower throughput and time consuming with large sample sets (Table 1). Fluorescence based methods such as the general protein-targeted Viral-BONCAT are comparatively inexpensive and easy to employ with minimal sample manipulation and experimental set up. Nucleic-acid based fluorescence staining of viruses is widely used in microbial ecology studies to visualize and count populations (e.g., Nobel and Fuhrman 1998, Allers et al. 2013, Sherr et al. 1987, Hennes et al. 1995, Zhang et al. 2010, Ohno et al. 2012). By targeting proteins rather than nucleic acids, Viral-BONCAT potentially targets a broader range of viruses, such as single stranded RNA/ DNA viruses, which are currently missed with SYBR staining methods, but may represent a large proportion of the viral assemblage in the surface ocean (Steward et al. 2013, Roux et al. 2016b). As such, assuming that microbial hosts in the system under investigation actively incorporate HPG and transfer the modified methionine and/or proteins to viruses with enough HPG to generate a fluorescence signal above background, this approach should serve as a powerful complement to established methods of virus quantification and production measurements. See Appendix S4 for additional details on methodological considerations for measuring viral activity via Viral-BONCAT.

The Viral-BONCAT method was successfully developed and optimized using two cultured viral-host systems: the heterotrophic, bacterium *E. coli* and its 50 nm phage T7; and the photosynthetic eukaryote *E. huxleyi* and its ~200 nm virus EhV207. Similar to that of bacteria and archaea (Hatzenpichler et al. 2014, Samo et al. 2014), viral methionine content (at least for T7, Syn1 and EhV) is about 3% of the total amino acid pool (Table S1). Incorporation of host-derived HPG-labeled amino acids (or newly synthesized proteins) into viruses was
distinguishable from BONCAT negative viral controls based on the R:G fluorescence ratio (Figs. 5, 6). Our hypothesis, based on lytic culture dynamics, assumed that 100% of the viruses would become HPG labeled if their host was HPG-labeled. A small fraction of unlabeled viruses were initially added to each culture as infection inoculum, accounting for only ~1% of the total virus population after host lysis. The final estimates for the percent of newly synthesized viruses in the T7 (98-99%) and EhV (87-94%) Viral-BONCAT experiments are consistent with this prediction. Interestingly, about 5-10% of the EhV viruses recovered at the end of the experiment remained unlabeled based on R:G ratios. Further work is needed to confirm the difference between *E. huxleyi* and *E. coli* viral production. Given the more nuanced, ‘animal-like’ infection phase dynamics of the *E. huxleyi*-EhV system (Mackinder et al. 2009, Grimsley et al. 2012), it is conceivable the BONCAT-negative particles may have been associated with an early ‘chronic’ phase of infection with viruses released by budding prior to active assimilation of HPG by hosts. This type of budding behavior is not observed in the cyanobacterial virus model system.

The experimental BONCAT data from cultured virus-host systems (Figs. 5, 6) and predictions from the sensitivity model (Figs. 7, S7, S8) identified three main factors that influence the ability to quantify viral production using this method. These include: the number of methionine substitutions ($n_{\text{subs}}$) per virus, the magnitude of the shift in R:G ratio per substitution ($d_{\text{RG}}$), and the percentage of the viral assemblage actively turning over (% active). These parameters were modeled as discrete values to provide benchmarks for our sensitivity analysis and data interpretation. Further tests of Viral-BONCAT with other virus-host model systems will enable better characterization of the variability in the number of substitutions and the impact of each substitution ($n_{\text{subs}}$) on the R:G ratio ($d_{\text{RG}}$).

As a pilot study, we demonstrated the use of Viral-BONCAT with coastal seawater to detect BONCAT positive viral-like particles within a mixed community and estimate viral production rates in the field (Fig. 8). Unlike our model culture studies, the methionine substitution rate and the $d_{\text{RG}}$ is expected to vary among the diverse viral assemblages in environmental samples. Assuming all newly synthesized viral particles become HPG-labeled after lysing actively growing host cells, the relative proportion of HPG labeled vs. unlabeled viral particles provides a direct quantification of the number of newly synthesized viral particles produced over a given time period. Furthermore, if the decay rate is similar between HPG-labeled and unlabeled viral particles, comparing the number of HPG-labeled viruses relative to
total viruses across two time points enables an estimation of net viral production rates (e.g., production-decay). Using probability statistics (as was done with the culture model systems), we were able to extract the number of fluorescently labeled viruses that were significantly greater than the unlabeled control and calculate viral production rates within the microbial community in surface waters collected off the Scripps Pier off of La Jolla, CA. However, viral production rate estimates are difficult to compare because they inherently depend on the concentration of viruses present in the sample; therefore, viral turnover is more amenable to comparisons across studies as it focuses on the relative proportion of the total viral assemblage, rather than an absolute concentration. The estimated viral turnover from this experiment was 1.5 days, consistent with previous measurements in near shore southern California regions (Steward et al. 1992b, Nobel and Fuhrman 2000).

This preliminary field study demonstrates the suitability of Viral-BONCAT for estimating viral production in a mixed or natural community. Focused, comparative studies are still needed to determine the best time-scales over which to sample and the frequency at which HPG should be added to the incubations. Shorter incubation times could provide hourly production rates, but the experimental timescales need to allow for adequate turnover of the viral population (>25%) and long enough to ensure newly synthesized viral particles have sufficient HPG incorporation (>1 methionine substitution). The concentration of bioavailable environmental methionine relative to the amount of added HPG will also be important to consider as this ratio has been shown to influence HPG uptake (Samo et al. 2014). In environments where cells are very active, continuous HPG additions may be needed to obtain sufficient labeling of the host populations (and viral particles) over time. This could be one plausible explanation for why we observed a reduction in the percentage of HPG-labeled viral particles between 48 and 72 hours. Another possibility is that the host cells, from which those viruses were produced, did not take up HPG; HPG incorporation by the host is a prerequisite for Viral-BONCAT. Notably, active HPG assimilation has been demonstrated in diverse archaea, bacteria, and microeukaryotes and in environmental samples (Hatzenpichler et al., 2015, 2016, Samo et al. 2014; Babin et al. 2016, this study). Likewise, even microorganisms classified as obligate photo- or chemolithoautotrophs have been shown to actively assimilate amino acids from the environment (e.g. Ouverney and Fuhrman 2000). Additional studies to test the applicability of this method in different environments as well as parallel methodological
comparisons of virus production estimates using more standard approaches (e.g., virus reduction) are also needed. Overall, this technique advances our ability to assess the activity of viruses in the environment and offers an independent and direct approach to quantify viral production with minor sample manipulation and no dependency on conversion factors.

Future directions and implications for understanding viral-host interactions in the ocean

Our current understanding of the biogeochemical impact of marine viruses is limited by our ability to measure exchanges of nutrients and carbon between viruses and their environment. Direct efforts to quantify the influence of the ‘viral shunt’ are rare, with most studies indirectly estimating viral turnover rates and elemental release. The lack of straightforward and reliable approaches for estimating viral production and rates of viral mortality on phytoplankton and bacteria in the ocean remains one of the biggest obstacles for incorporating viral-mediated processes into global models of nutrient and energy cycling. Furthermore, tracing the fate of newly fixed carbon and nitrogen into different components of microbial food webs (e.g., phytoplankton, grazers, viruses and dissolved organic matter) is of great interest for understanding the direction, rates and magnitude of energy flow in marine ecosystems. The combination of measuring the fate of pathway specific stable isotope labels via nanoSIMS and general viral production by fluorescence via Viral BONCAT now enables the direct quantification of specific sources of C and N moving through the viral pool. The use of Viral-BONCAT provides a relatively inexpensive and rapid assessment of the fraction of free viroplankton that were recently involved lysing an active host cell. Coupled with incubations amended with different sources of $^{13}$C-labeled carbon or $^{15}$N-labeled nitrogen provides a mechanism for directly quantifying the proportion of active viroplankton that are involved in mediating the turnover of newly fixed carbon and nitrogen (via photosynthesis, chemosynthesis, diazotrophy, ammonium, or nitrate assimilation) relative to viruses involved in the lysis of heterotrophs over many different temporal and spatial scales. Future work will expand the use of stable isotope tracers beyond those used in this study (bicarbonate, nitrate, ammonium) to directly quantify the proportion of active viroplankton involved in mediating the turnover of carbon, nitrogen and potentially sulfur substrates in the surface oceans, mesopelagic or even more complex sediments and soils. Additional avenues could include creating isotopically labeled viral particles and trace the movement of enriched C and N from viruses into...
phytoplankton, grazers, heterotrophic bacteria, and dissolved organic matter. Coupling these activity- and isotope-based techniques with other powerful microbial ecology methods (e.g., flow cytometry cell sorting, genomics, proteomics) opens up exciting possibilities for viral ecology studies (Brussaard et al. 2016).

Another potential application of Viral-BONCAT is to identify specific viral proteins produced during the lytic infection. Combining BONCAT with proteomics (e.g., Babin et al. 2016, Bagert et al. 2016) provides information about newly synthesized viral proteins and potentially changes in protein expression over time in tandem with taxonomic information (e.g., capsid protein sequences) in the context of their natural environment. HPG substitution into specific proteins was confirmed by mass spectrometry from experiments using BONCAT-positive T7 and EhV viruses, with many of the detected HPG-labeled proteins involved in procapsid and capsid formation and DNA packaging (Table 2). Moniruzzaman et al. (2017) recently demonstrated the utility of the Major Capsid Protein (MCP) as a marker for infection dynamics and diversity in natural populations. Therefore, tracking changes in HPG-labeled viral proteins over time in other cultures and extending this to natural environments (e.g., tracking HGP incorporation into MCP) provides us an exciting opportunity to characterize infection dynamics at the protein level.

Proteomics coupled with BONCAT can also be used to better characterize viral diversity and biogeography, an important and growing area of exploration in the field of marine virology (Breitbart et al. 2002, 2004, Hurwitz et al. 2015, Brum et al. 2015, Roux et al. 2016a). One major advantage of protein targeted Viral BONCAT is that, unlike currently available fluorescent nucleic acid stains that are limited to double stranded DNA viruses, BONCAT has the potential to also label single-stranded DNA and RNA viruses (Steward et al. 2013, Roux et al. 2016b). Successful sorting of HPG-labeled viruses (via fluorescence signal) prior to metagenomic or metatranscriptomic analysis would allow for a comparison of active virus populations (e.g., FCM sorted viral genomes) among the total viral pool (e.g., bulk viral metagenomes) and help to highlight factors that influence active viral communities over space and time and govern enhance viral success in the ocean (Angly et al. 2006, Breitbart et al. 2012).

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References


Orphan VJ, Turk KA, Green AM, House CH (2009) Patterns of 15N assimilation and growth of


Table Legends

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**Table 2.** Proteomics identification of HPG-labeled T7 and EhV proteins from Viral BONCAT experiments.

**Table 3.** Percentage of BONCAT positive viral-like particles and corresponding R:G ratios from HPG incubation experiments with coastal seawater. Percent labeled is given as the average ± 95% confidence interval.

**Table 4.** Comparison of Viral BONCAT estimate of virus turnover in coastal seawater with published estimates of turnover and viral production rates using different methods.

Figure Legends

**Figure 1.** NanoSIMS ion images of cultured hosts and viruses. Example ion images (\(^{14}\)N\(^{12}\)C-) of *Emiliania huxleyi* 374 (A), *Synechococcus* WH8101 (B), EhV 207 (C) and Syn1 (D). Color scale represents total ion counts. Large particles in D are gold beads.

**Figure 2.** Compiled nanoSIMS data from single-label experiments with cyanobacterial-virus model system. Whisker plots showing fractional abundance of *Synechococcus sp.* WH8101 and virus Syn1 after incubation with \(^{13}\)C-bicarbonate (Exp. 1), \(^{15}\)N-ammonia (Exps. 2, 3) and control incubations (right panel). Fractional abundance (y-axis) reveals enrichment above background for carbon (top row, natural = 1.09 at%) and nitrogen (bottom row, natural = 0.36 at%).

**Figure 3.** Compiled nanoSIMS data from single and dual label experiments with eukaryotic algae-virus model system. Whisker plots showing fractional abundance data from single-label (Exp. 1,2) and dual-label (Exp. 3) incubations with coccolithophore *E. huxleyi* and its virus EhV 207 after incubation with \(^{15}\)N-nitrate and/or \(^{13}\)C-bicarbonate as well as control incubations (right panel).
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**Figure 4.** Compiled nanoSIMS data from short-term experiments with cyanobacterial-virus model system. Whisker plots showing fractional abundance of *Synechococcus sp.* WH8101 and virus Syn1 after incubation with $^{15}$N-ammonia (left panel) for 24 hours. An unlabeled control incubation (right panel) was run in parallel. Fractional abundance (y-axis) reveals enrichment above background for nitrogen (natural = 0.36 at%).

**Figure 5.** Visualization of BONCAT labeled *E. huxleyi* virus (EhV 207) and fluorescence analysis. Examples of images of EhV particles from experiments of *E. huxleyi* grown with (A, B) and without (C, D) HPG. The click reaction was performed with a TAMRA-Azide dye (detected with a Cy3 filter cube) and particles were counter stained with SYBR gold (detected with a FITC filter cube). Histogram (E) the red to green fluorescence ratio (R:G) of viral particles from all particles analyzed across 16 images (viral particles $n_{HPG} = 3713$, $n_{CONTROL} = 6522$). Red bars are from particles in the HPG labeled treatment, and blue bars are from viral particles in the no HPG control. F) Visualization of BONCAT labeled *E. huxleyi* compared to no HPG control. The click reaction was performed with CR-110-Azide dye (detected with FITC filter cube) and cells were counter stained with DAPI (detected with a DAPI filter cube). Note – images in figure have been background corrected.

**Figure 6.** Visualization of BONCAT labeled *E. coli* virus (T7) and fluorescence analysis. Examples of images of T7 particles from experiments of *E. coli* grown with (A,B) and without (C,D) HPG. Click reaction was performed with a TAMRA-Azide dye (detected with a Cy3 filter cube) and particles were counter stained with SYBR gold (detected with a FITC filter cube). Histogram (E) of the red to green fluorescence ratio (R:G) of viral particles from all particles analyzed across 10 images (viral particles $n_{HPG} = 31512$, $n_{CONTROL} = 20858$). Red bars are from particles in the HPG labeled treatment, and blue bars are from viral particles in the no HPG control. F) Visualization of BONCAT labeled *E. coli* compared to no hPG control. The click reaction was performed with CR-110-Azide dye (detected with FITC filter cube) and cells were
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**Figure 7.** Accuracy and precision of viral production estimates using Viral BONCAT. Results from modeling T7 data with a specific level of activity (e.g., the percent of the population actively turning over) and varying \( n_{\text{sub}} \), \( d_{\text{RG}} \) and different levels of variability around the \( d_{\text{RG}} \) to determine the ability to back calculate the original activity using the rank-sum bootstrap approach.

**Figure 8.** Visualization of BONCAT labeled viruses and fluorescence analysis from field tests. Fluorescence images of viral particles from field experiment in which seawater was amended with HPG for 48 hours (example images shown in A, B) and 72 hours (images not shown) relative to the negative control (example images shown C, D). Click reaction was performed with a TAMRA-Azide dye (detected with a Cy3 filter cube) and particles were counter stained with SYBR gold (detected with a FITC filter cube). Histograms of red to green fluorescence ratio of viral particles incubated with HPG after 48 hours (E) and 72 hours (F). The color of the bar indicates treatment. See Table 2 for details on percentage of viral particles labeled and number of particles analyzed. Note – images in figure have been background corrected.
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Table 1. Comparison of BONCAT and nanoSIMS approaches for studying viral interactions

<table>
<thead>
<tr>
<th>Benefit</th>
<th>BONCAT</th>
<th>nanoSIMS</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>assessment of production</td>
<td>assessment of C and N transfer from host to virus</td>
</tr>
<tr>
<td>Quantitative</td>
<td>semi-quantitative</td>
<td>quantitative</td>
</tr>
<tr>
<td>Throughput</td>
<td>high</td>
<td>low</td>
</tr>
<tr>
<td>Cost</td>
<td>inexpensive</td>
<td>expensive</td>
</tr>
<tr>
<td>Limitations</td>
<td>requires host to assimilate HPG</td>
<td>lower size limit (20-50 nm)</td>
</tr>
</tbody>
</table>

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Table 2. Proteomics identification of HPG-labeled T7 and EhV proteins from Viral BONCAT experiments.

<table>
<thead>
<tr>
<th>Virus</th>
<th>Protein Name*</th>
<th>Protein</th>
<th>HPG-Substitution Positions</th>
</tr>
</thead>
<tbody>
<tr>
<td>T7</td>
<td>Major capsid protein 10A; Minor capsid protein 10B</td>
<td>P19726;P19727</td>
<td>1;4;10;293</td>
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<tr>
<td>T7</td>
<td>DNA ligase</td>
<td>P00969</td>
<td>288</td>
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<tr>
<td>T7</td>
<td>Capsid assembly scaffolding protein (Gp9)</td>
<td>P03716</td>
<td>279</td>
</tr>
<tr>
<td>T7</td>
<td>Terminase, small subunit gp18</td>
<td>P03693</td>
<td>22</td>
</tr>
<tr>
<td>T7</td>
<td>Single-stranded DNA-binding protein gp2.5</td>
<td>P03696</td>
<td>137</td>
</tr>
<tr>
<td>T7</td>
<td>Peptidoglycan hydrolase gp16</td>
<td>P03726</td>
<td>150</td>
</tr>
<tr>
<td>T7</td>
<td>Portal protein</td>
<td>P03728</td>
<td>118;179</td>
</tr>
<tr>
<td>T7</td>
<td>Protein 6.5</td>
<td>P03800</td>
<td>1</td>
</tr>
<tr>
<td>EhV</td>
<td>Major capsid protein</td>
<td>G4YBV8</td>
<td>23</td>
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<td>EhV</td>
<td>Putative uncharacterized protein</td>
<td>G4YBJ4</td>
<td>97;105</td>
</tr>
<tr>
<td>EhV</td>
<td>Putative uncharacterized protein</td>
<td>G4YBX3</td>
<td>97</td>
</tr>
<tr>
<td>EhV</td>
<td>Putative uncharacterized protein</td>
<td>G4YCD5</td>
<td>91</td>
</tr>
<tr>
<td>EhV</td>
<td>Putative uncharacterized protein</td>
<td>G4YCE6</td>
<td>39</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Treatment</th>
<th>% HPG-Labeled</th>
<th>Viral Particles Analyzed</th>
<th>Mean R:G</th>
</tr>
</thead>
<tbody>
<tr>
<td>Positive HPG 48-hrs A</td>
<td>86.4 ± 0.7</td>
<td>32404</td>
<td>0.21</td>
</tr>
<tr>
<td>Positive HPG 48-hrs B</td>
<td>85.8 ± 0.7</td>
<td>27928</td>
<td>0.20</td>
</tr>
<tr>
<td>Positive HPG 72-hrs A</td>
<td>72.0 ± 0.9</td>
<td>26477</td>
<td>0.16</td>
</tr>
<tr>
<td>Positive HPG 72-hrs B</td>
<td>85.4 ± 0.8</td>
<td>31334</td>
<td>0.19</td>
</tr>
<tr>
<td>Negative Control</td>
<td>n/a</td>
<td>19884</td>
<td>0.11</td>
</tr>
<tr>
<td>Negative Control</td>
<td>n/a</td>
<td>19097</td>
<td>0.10</td>
</tr>
</tbody>
</table>
**Table 4.** Comparison of Viral BONCAT estimate of virus turnover in coastal seawater with published estimates of turnover and viral production rates using different methods.

<table>
<thead>
<tr>
<th>Viral Production Rates (viruses mL⁻¹ day⁻¹)</th>
<th>Viral turnover (days)</th>
<th>Geographical Location</th>
<th>Method</th>
<th>Citation</th>
</tr>
</thead>
<tbody>
<tr>
<td>16.4 x 10⁶</td>
<td>1.5</td>
<td>Southern California, nearshore</td>
<td>BONCAT</td>
<td>this study</td>
</tr>
<tr>
<td>12-230 x 10⁹</td>
<td>0.5-3.9</td>
<td>Southern California Bight, nearshore</td>
<td>Incorporation of radiolabeled substrate</td>
<td>Steward et al. 1992b</td>
</tr>
<tr>
<td>0 - 2.8 x 10⁹</td>
<td>8.9-30</td>
<td>Southern California Bight, offshore</td>
<td>Incorporation of radiolabeled substrate</td>
<td>Steward et al. 1992b</td>
</tr>
<tr>
<td>2.8 - 28 x 10⁶</td>
<td>1-2</td>
<td>Southern California region</td>
<td>Fluorescently labeled viruses as tracers</td>
<td>Nobel and Fuhrman 2000</td>
</tr>
<tr>
<td>5.3 - 43 x10⁶</td>
<td>5.1-27.8</td>
<td>Discovery Passage and Straight of Georgia (British Columbia, Canada)</td>
<td>Dilution technique</td>
<td>Wilhem et al. 2002</td>
</tr>
<tr>
<td>4.8 - 5.9 x 10⁷</td>
<td>n/a</td>
<td>Chesapeake and Delaware Bays</td>
<td>Tangential Flow diafiltration dilution method</td>
<td>Winget et al. 2005</td>
</tr>
<tr>
<td>0.08 - 1.7 x 10⁶</td>
<td>n/a</td>
<td>North Water</td>
<td>Frequency of visibly infected cells</td>
<td>Middleboe et al. 2002</td>
</tr>
<tr>
<td>0.68 - 15.6 x 10⁸</td>
<td>n/a</td>
<td>Coastal Bay, Norway</td>
<td>Viral decay rates</td>
<td>Heldal and Bratbak 1991</td>
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</tbody>
</table>