Light regulation of coccolithophore host–virus interactions

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Summary

- Viruses that infect photoautotrophs have a fundamental relationship with light, given the need for host resources.
- We investigated the role of light on Coccolithovirus (EhV) infection of the globally distributed coccolithophore, Emiliania huxleyi. Light was required for EhV adsorption, and viral production was highest when host cultures were maintained in continuous light or at irradiance levels of 150–300 μmol m⁻² s⁻¹. During the early stages of infection, photosynthetic electron transport remained high, while RuBisCO expression decreased concomitant with an induction of the pentose phosphate pathway, the primary source of de novo nucleotides. A mathematical model developed and fitted to the laboratory data supported the hypothesis that EhV replication was controlled by a trade-off between host nucleotide recycling and de novo synthesis, and that photoperiod and photon flux could toggle this switch.
- Laboratory results supported field observations that light was the most robust driver of EhV replication within E. huxleyi populations collected across a 2000 nautical mile transect in the North Atlantic.
- Collectively, these findings demonstrate that light can drive host–virus interactions through a mechanistic interplay between host metabolic processes, which serve to structure infection and phytoplankton mortality in the upper ocean.

Introduction

Virus infection is a primary mechanism of high lysis rates of phytoplankton populations (van Boekel et al., 1992; Bratbak et al., 1993; Brussaard et al., 1995; Valiela, 1995; Agusti et al., 1998). It is estimated that algal viruses turn over more than a quarter of the photosynthetically fixed carbon, fueling microbial foodwebs and short-circuiting carbon export to higher trophic levels and the deep sea (Fuhrman, 1999; Suttle, 2007) by releasing dissolved organic matter into the surrounding water. At the same time, viral-induced transparent exopolymer particle production (Joassin et al., 2011; Vardi et al., 2012) suggests infection may stimulate vertical sinking flux and enhance biological pump efficiency. Despite the impact on phytoplankton communities, viral-induced mortality is not routinely accounted for in models of ecosystem processes and carbon export, due to inadequate mechanistic and quantitative understanding of the environmental factors that regulate host–virus interactions.

Emiliania huxleyi and its associated Coccolithovirus (EhV) is a highly studied marine eukaryotic algal host–virus model system due to its ecological relevance and the availability of genetically diverse hosts and EhV strains in culture (Schroeder et al., 2002; Bidle & Vardi, 2011). Massive spring blooms of E. huxleyi in the North Atlantic (Townsend et al., 1994; Tyrell & Merico, 2004) are routinely terminated by the giant, lytic, double-stranded DNA containing EhVs (Bratbak et al., 1993; Schroeder et al., 2002; Lehahn et al., 2014; Laber et al., 2018; Sheyn et al., 2018), and studies have revealed that host–virus interactions are critically controlled through a lipid-based chemical arms race and subcellular regulation of autophagy and programmed cell death pathways (Bidle et al., 2007; Vardi et al., 2012; Rose et al., 2014; Rosenwasser et al., 2014; Schatz et al., 2014; Sheyn et al., 2016). These studies have collectively set the stage for detailed exploration into the environmental conditions that regulate key cellular pathways during infection.

As one of the most fundamental and readily measured resources in the ocean, light critically regulates E. huxleyi distribution, growth, and productivity, and in turn viral genome replication and virion production given the need for host resources (e.g. nucleotides, lipids, proteins, and energy). Decreased photosynthetic efficiency (Evans et al., 2006; Bidle et al., 2007) and increased nonphotochemical quenching (Llewellyn et al., 2007) during EhV infection suggest an uncoupling of photosynthetic electron flow and impaired photo-physiology (Evans et al., 2006), findings that are supported by transcriptomic analysis of photosynthetic-related genes during infection (Rosenwasser et al., 2014; Gilg et al., 2016). However, the
mechanistic connection between light, EhV infection, and host metabolism has yet to be fully characterized.

Here, we investigated the role of light in regulating host metabolism and infection dynamics and developed a mathematical model that explicitly resolves and quantifies the relationship between light and viral production. The cellular and biochemical basis of this interaction, combined with quantitative statistical analyses, helped to explain the vertical distribution of EhV abundance in natural *E. huxleyi* populations collected across a 2000 nautical mile transect in the North Atlantic. Our findings highlight the importance of light-driven host subcellular processes in mediating viral infection of *E. huxleyi* and vertically structuring these interactions in nature.

**Materials and Methods**

**Culture and virus maintenance**

*Emiliania huxleyi* strain 374, obtained from the National Center for Marine Algae and Microbiota, was batch grown in f/2 medium −Si at 18°C. Cultures were maintained under different photoperiods (14 h : 10 h, light : dark (LD) or continuous light) at an irradiance level of 150 μmol m$^{-2}$ s$^{-1}$, or on a 14 h : 10 h, LD cycle at different irradiance levels (25, 150, 300, 500, 1000 and 2000 μmol m$^{-2}$ s$^{-1}$) for >2 months before experiments and kept optically thin by frequent transfers (c. 2–4 d) to prevent cell lysis.光学 thinness was determined using a Coulter Multisizer II (Beckman Coulter, Brea, CA, USA) or an InFlux Model 209S Mariner flow cytometer (BD Biosciences, Franklin Lakes, NJ, USA). *Coccolithovirus strain EhV201* (hereafter referred to as EhV) was kindly provided by Dr Michael Allen (Plymouth Marine Laboratory). Viral lysates were obtained by infecting *E. huxleyi* at a virus to host ratio of c. 5–10, based on total virus abundance. Upon lysis, debris was removed by 0.45 μm pore-size polycarbonate filtration (Millipore, Burlington, MA, USA). Virus abundance was measured in 0.5% glutaraldehyde fixed samples using SYBR Gold (Molecular Probes, Eugene, OR, USA) and flow cytometry (Brussaard, 2004).

**Viral infection**

Exponentially growing *E. huxleyi* cultures were infected with EhV (virus-to-host ratio: 5), c. 1–2 h into the light phase, unless otherwise stated. In comparative experiments (e.g. LD vs continuous light or LD vs dark), infections were performed concurrently using the same stock of freshly propagated viral lysate. For dark experiments, EhV was added at the onset of the light period to allow adsorption and internalization. At the end of the light phase, a subset of the cultures was wrapped in foil and kept dark for the remainder of the experiment. Control cultures continued on the normal LD cycle. For 3-(3,4-dichlorophenyl)-1,1-dimethylurea (DCMU) experiments, 5 μM was added to control and infected cultures 24 h postinfection (hpi). Burst size was calculated by dividing the total number of viruses produced by the number of host cells lost (i.e. maximum host abundance minus final host abundance). Cell biomass was collected for enzyme activity measurements and immunoblots by filtration onto 47 mm diameter, 1.2 μm pore-size polycarbonate filters, snap frozen in liquid nitrogen, and stored at −80°C.

**Photosynthetic measurements**

Electron transfer rates and the maximum photosynthetic efficiency of photosystem II (PSII), expressed as $F_{v}/F_{m}$, were measured using a custom-built fast fluorescence induction and relaxation system (Gorbunov & Falkowski, 2005). The minimum ($F_{o}$) and maximum ($F_{m}$) fluorescence yields, $F_{v}/F_{m}$ ($F_{m}−F_{o}/F_{m}$ for dark-adapted cells and $ΔF′/F′_{m}$ under ambient light), and $σ_{PSII}$ (the functional absorption cross-section of PSII) were calculated from the analysis of fluorescence induction at the microsecond time scale. The photosynthesis vs irradiance curves were reconstructed from measurements of electron transport rates ($P = E × σ_{PSII} × ΔF′/F′_{m}$) as a function of ambient irradiance $E$ (μmol m$^{-2}$ s$^{-1}$).

**Viral adsorption**

Free, extracellular virus abundance was used as a proxy for adsorption, where the disappearance of viruses from the medium is as an indication of viral adsorption onto host cells. Nonspecific adsorption was assessed by adding viruses to cell-free media. A drop in free virus was detected within 5 min, but returned to initial concentrations after 15 min, suggestive of nonspecific adsorption, presumably to the wall of the flask, and subsequent release of viruses. Therefore, measurements earlier than 30 min were not used.

**Pentose phosphate pathway activity**

NADPH production by the two rate-limiting enzymes of the pentose phosphate pathway (PPP), glucose-6-phosphate dehydrogenase and 6-phosphogluconate dehydrogenase, was measured via a modified protocol (Tian et al., 1994; Šindelář et al., 1999). Filters were thawed and cells were scraped in TEM buffer (20 mM tri(hydroxymethyl)aminomethane hydrochloride (Tris-HCl), 1 mM EDTA, 2.5 mM magnesium chloride, pH 7) with a fresh razor blade, transferred to tubes, and sonicated three times on ice in 20 s intervals (Misonix; Microson, Farmingdale, NY, USA; power setting of 2–3). After centrifugation (4°C, 10 min, 20 817 g), the supernatant was assayed in 96-well microtitre plates in a total volume of 300 μl containing 100 μl of cell extract, 150 μl NADP+ (Sigma-Aldrich), 3 mM glucose 6-phosphate (Santa Cruz Biotechnology, Dallas, TX, USA), 3 mM 6-phosphogluconate (Sigma-Aldrich), and TM buffer (100 mM Tris-HCl, 12.5 mM magnesium chloride, pH 8). Negative controls were cell extracts and TM buffer, without substrate or NADP+. Enzyme activity was determined from the slope of the reaction, measured every 30 s at 340 nm (SpectraMax M3, Molecular Devices, San Jose, CA, USA) between 10 and 30 min, and an NADPH standard curve (Caymen Chemical, Ann Arbor, MI, USA). Protein concentrations were determined using a BCA Protein Assay Kit (Thermo Fisher Scientific, Waltham, MA, USA).
Quantitative immunoblot analysis

Protein extraction and quantitative immunoblots were performed as described (Brown et al., 2008; Thamatrakoln et al., 2013). Immunoblots were probed with primary antibodies (Agrisera, Vännäs, Sweden) against PsbD (a proxy for PSII) and RbcL (a proxy for RuBisCO) at a dilution of 1 : 40 000, followed by a horseradish-peroxidase-conjugated secondary antibody at 1 : 10 000 dilution. Chemiluminescence detection was performed using ECL Select (GE Healthcare, Chicago, IL, USA) and imaged using a ChemiDoc XRS+ CCD imager (Bio-Rad, Hercules, CA, USA). Adjusted volume values were obtained, and standard curves of serially diluted recombinant proteins were used to estimate the amount of protein as described (Brown et al., 2008; Thamatrakoln et al., 2013).

Nonlinear model of host–virus nucleotide dynamics

A mathematical model was developed to characterize and disentangle the effect of host growth, host nucleotide recycling and de novo nucleotide synthesis on host–virus population dynamics. By extension to Wikner et al. (1993), the model resolves time variation in the concentration of nucleotides (nucleotides ml⁻¹) within uninfected hosts (S), infected hosts (I), internal viruses produced by host nucleotide recycling (P), internal viruses produced by de novo nucleotide synthesis (D), and free (extracellular) viruses (V). Direct measures of the individual nucleotide pools were not made given methodological limitations. Rather, the model was constrained with total cellular nucleotides (S + I + P) and free viral nucleotides (V), which were converted from host and virus abundance respectively and known genome size (Supporting Information Table S1). We hypothesized that host growth, nucleotide recycling and de novo synthesis each have a distinct and quantifiable influence on the dynamics of total cellular and free viral nucleotides and that these distinct signatures would lead to constraint on the light dependence of nucleotide synthesis rates. Parameter definitions and units are in Table S1.

\[
\frac{dV}{dI} = -v_p \phi SV + v_p \delta (P + D) \tag{Eqn 5}
\]

In Eqn 1, hosts synthesize nucleotides at rate \( \mu_h \). The linear interaction term \( \phi SV \), controls the total flux of nucleotides from susceptible hosts and free, extracellular viruses to newly synthesized, intracellular viruses. The parameter \( v_p \) represents the portion of this flux that originates from the virus, and \( 1 - v_p \) represents the portion that originates from the host, with

\[
v_p = \frac{V_{ind}}{V_{ind} + S_{ind}} \tag{Eqn 6}
\]

where \( V_{ind} \) and \( S_{ind} \) are the nucleotide contents of an individual virus and host respectively (Table S1).

In Eqn 2, infected hosts synthesize nucleotides at rate \( \mu_i \), and nucleotides are either converted to viral progeny at rate \( v_p \mu_v \) or lysed at rate \( v_p \). The parameter \( v_p \) is included to mimic a delay in internal viral production and host lysis due to internal assembly of viral progeny. This delay parameterizes the latent period (i.e. the time during which the virus is replicating prior to host lysis). Unlike previous models that mimic the latent period with a time lag (e.g. Wang, 2006), \( v_p \) forces a delay due explicitly to the internal depletion of host resources by the virus. Specifically, \( v_p \) forces the rate of lysis to increase as host resources are converted to intracellular viruses:

\[
v_p = \frac{P + D}{I + P + D} \tag{Eqn 7}
\]

In Eqn 7, as host nucleotides \( I \) are converted to internal viral nucleotides \( P + D \), \( v_p \) increases from very low values and then saturates close to one, which in turn causes the rate of lysis \( v_p \delta \) in Eqn 2 to increase, and then saturate at a fixed rate.

Nucleotides required for viral genome replication can be obtained in two ways, either through recycling of existing host nucleotides or de novo nucleotide synthesis. Recycling, by definition, leads to a depletion of host nucleotides, whereas de novo synthesis does not. Eqn 3 accounts for the accumulation of internal viruses by nucleotide recycling at rate \( v_p \mu_v \), and the loss of internal viruses due to lysis at rate \( v_p \delta \). Eqn 4 accounts for the production of internal viruses by de novo synthesis at rate \( v_p \mu_v \), as well as accumulation of internal viruses due to adsorption, and losses due to host lysis at rate \( v_p \delta \). Having the rate of nucleotide recycling and de novo nucleotide synthesis dependent on \( v_p \) and accounts for the time it takes for a virus to co-opt the host biosynthetic machinery. Note that the sum of Eqns 3 and 4 represents a single mass balance for the total number of internal nucleotides, \( P + D \). We separate \( P \) and \( D \) because it is only appropriate to count nucleotides generated via recycling \( P \), with host nucleotides inferred with measured cell abundance and known nucleotide content (i.e. genome size; Table S1). Nucleotides generated via de novo synthesis are assumed to be ‘invisible’ until host lysis.
Eqn 5 assumes the free virus nucleotide concentration $V$ is a balance between production of free viruses by host lysis at rate $v_l\delta$, and uptake of viruses due to adsorption to host cells at rate $v_r\phi S$. For all experiments, model solutions were obtained by numerically integrating Eqns 1–5 using initial conditions that matched the experimental observations.

Parameter constraint with the Metropolis algorithm

Our goal was to quantify the effect of host growth, nucleotide recycling and de novo nucleotide synthesis on total cellular $(S+I+P)$ and free viral $(V)$ nucleotide dynamics. We used the Metropolis algorithm (Metropolis et al., 1953) to test formally whether rates of nucleotide synthesis could be uniquely constrained with knowledge of total cellular and free-virus dynamics. We leveraged this constraint to explore the light dependence of viral replication. The Metropolis algorithm is a Monte Carlo type simulation that quantifies parameter uncertainty using a random walk to explore parameter combinations that explain observed dynamics. When the observations are unable to constrain model parameters, the random walk does not converge, and the algorithm predicts ever wider parameter distributions. When the observations are sufficient to constrain model parameters, the algorithm converges on unique parameter distributions.

The five model parameters controlling infection dynamics are virus adsorption $\phi$, host growth rate $\mu_h$, host nucleotide recycling rate $\mu_r$, de novo nucleotide synthesis rate $\mu'_v$, and lysis rate $\delta$. The adsorption parameter was fitted using data from adsorption experiments. The remaining parameters $\theta = \{\mu_h, \mu_r, \mu'_v, \delta\}$ were fitted using population-level observations, labeled here $D(t_i)$, of host-virus dynamics. The Metropolis algorithm starts with an initial parameter set $\theta_{current}$ manually determined so that the model solutions are reasonably close to the observations. A new parameter set, $\theta_{proposed}$, is then drawn from appropriate distributions. We used lognormal distributions with means matching the old distribution, and standard deviations chosen manually to ensure average acceptance ratios within the range 23–44% to promote efficient convergence (Gelman et al., 1996). The new parameters were chosen from lognormal distributions to prevent negative values. Solutions $y(t_i|\theta_{proposed})$ to the model with the new parameter set are then calculated. For both the old and the new models, the sum-of-squared difference between the model and the observations is divided by the measurement uncertainty of the observations, assumed here to be the average variance of each triplicate $\sigma^2$.

$$
\chi^2_{current} = \sum_i \frac{D(t_i) - y(t_i|\theta_{current})}{2\sigma^2}
$$

Eqn 8

$$
\chi^2_{proposed} = \sum_i \frac{D(t_i) - y(t_i|\theta_{proposed})}{2\sigma^2}
$$

Eqn 9

A likelihood function is defined in the following way:

$$
P(D|\theta) = \exp(-\chi^2)
$$

Eqn 10

A likelihood ratio is then calculated by taking the ratio of the new to the old likelihood function:

$$
\frac{P(D|\theta_{proposed})}{P(D|\theta_{current})} = \exp(-\chi^2_{proposed} + \chi^2_{current})
$$

Eqn 11

If the likelihood ratio is larger than a random number $r$ between 0 and 1, the new parameter set becomes $\theta_{current}$, and the process is repeated. If the likelihood ratio is less than $r$, the proposed set of parameters is disregarded and the process repeated with the old parameter set. Whenever a new parameter set is adopted, those parameters are recorded. Over time, enough parameters are stored to estimate parameter distributions that minimize error between the model and the observations. The distributions represent the full range of parameters that give suitable fits with respect to uncertainty in the observations.

By repeating the fitting exercise for both the light and the dark treatments, estimates of parameter distributions for $\mu_h$, $\mu_r$, $\mu'_v$, and $\delta$ were generated. All simulations were repeated numerous times from different initial parameter guesses to ensure consistent convergence. Parameters were assumed to be constrained by the data if the variance in the posterior distribution did not increase with respect to the number of iterations of the optimization algorithm. In some cases, this led to poor constraint; for example, in the continuous light treatments. By looking for differences in the fitted parameter distributions between different light treatments, we were able to discern whether key processes, namely de novo synthesis or recycling of host nucleotides, were dependent on light intensity.

Field sampling and statistical analysis

Emiliania huxleyi populations (and associated EhVs) were collected during the North Atlantic Virus Infection of Coccolithophores Expedition (http://www.bco-dmo.org/project/2136) using Niskin bottles on a 24-position rosette equipped with an SBE conductivity–temperature–depth profiler (Sea-Bird Scientific, Bellevue, WA, USA). Cell biomass (and associated viruses) was collected from c. 3–51 of 200 µm mesh-filtered seawater onto large (142 mm diameter), 0.8 µm pore-size polycarbonate filters (Millipore) to minimize clogging. Filters were immediately submerged in 10 ml extraction buffer (100 mM Tris-HCl pH 8, 250 mM EDTA pH 8, 100 mM sodium chloride, and 1% sodium dodecyl sulfate) and homogenized at maximum speed on a Vortex Genie (Mo Bio, Carlsbad, CA, USA) for 10 min in the presence of 2 ml molecular-grade zirconium beads (equal amount of 100 µm and 400 µm diameter beads; OPS Diagnostics, Lebanon, NJ, USA). Filters were subjected to three freeze–thaw cycles via submerging in liquid nitrogen and thawing at 50°C, homogenized for 5 min after each cycle and stored at −80°C. Upon processing, filters were thawed and 5 ml was incubated for 1 h at 50°C with 100 µg Proteinase K, followed by phenol : chloroform : isoamyl alcohol, 25 : 24 : 1 v/v/v extraction. Nucleic acids were precipitated with two volumes of 100% ethanol and 0.2 M sodium chloride. Following centrifugation (20 min, 10 000 g), the DNA
pellet was washed with 20 ml of 70% ethanol, centrifuged again (10 min, 10,000 g) and then resuspended in 1× TE buffer (pH 8). Trace impurities that could inhibit enzymatic amplification reactions were removed using the PowerClean® DNA Clean-Up Kit (Mo Bio). Quantitative PCR using SYBR® Green was used to quantify *E. huxleyi* (via cytochrome oxidase I) and EhV (via major capsid protein) (Coonen, 2011). Tenfold serially diluted genomic DNA (extracted from flow cytometry quantified *E. huxleyi* (RCC1216) and EhV (EhV86) was used to calibrate the cytochrome c oxidase I- and major capsid protein-specific quantitative PCR. Data were analyzed by linear regression in R using the lm() function and model fit verified with the plot(model) function. All data were log10-transformed for consistency.

## Results

### EhV infection, the PPP, and photosynthesis

Genes associated with *de novo* nucleotide synthesis, namely those involved in the PPP, were found to be upregulated during infection of *E. huxleyi* (Rosenwasser *et al*., 2014). To determine whether these observed increases in transcript abundance translated into higher rates of nucleotide synthesis, we measured the enzymatic activity of key enzymes of the PPP during the early stages (<24 hpi) of EhV infection. EhV infection arrested host growth within 24 hpi, concomitant with an increase in the production of extracellular viruses (Fig. S1). PPP activity in uninfected controls remained steady at ∼9 nM NADPH min⁻¹ μg protein⁻¹ (Table 1). In contrast, PPP activity increased in infected cultures by c. 1.6-fold between 4 and 14 hpi, reaching maximum levels that were >2.5-fold higher than uninfected controls by 24 hpi. Given that the PPP shares enzymes with the Calvin cycle, upregulation of the former should result in the downregulation of the latter. Measured protein expression levels of RuBisCO (Fig. 1a), the rate-limiting enzyme of the Calvin cycle, showed that the induced PPP activity was coincident with a 50% decrease in the biochemical potential of the Calvin cycle. In contrast, PsbD (a proxy for PSII) protein expression was similar between control and infected cells (Fig. 1a), suggesting photosynthetic electron transport was not inhibited in infected *E. huxleyi*.

No significant difference was observed in the electron transport rate between control and infected cells 5 hpi (data not shown) or 24 hpi (Fig. 1b). We further tested the role of PSII in viral production by inhibiting PSII activity with DCMU (Fig. 1c). There was a c. 15% drop in cell abundance in infected cultures that was not observed in infected cultures without DCMU (Fig. 1c, inset). Associated viral abundance was reduced by c. 50% in infected cultures treated with DCMU (Fig. 1d), which was not due to lower host growth, as the decrease in host abundance in the presence of DCMU was less (c. 30%) than the accompanying decrease in viral production. The infection was carried out for an additional 48 h to assess whether viral replication in DCMU-treated cultures was delayed, but cultures failed to produce the equivalent number of viruses as untreated, infected cultures, even at 96 hpi.

### Light critically regulates infection

Light heavily impacted virus adsorption to *E. huxleyi* cells. There was little adsorption when EhVs were added to *E. huxleyi* at the onset of the dark period (see the Materials and Methods section). Only upon transition to the light 10 h later was there notable adsorption, observed as a 78% drop in free EhV abundance within 2 h (Fig. 2a). By contrast, when EhV was added at the onset of the light period, a near immediate drop of 44% in free EhV abundance was observed (Fig. S2a), consistent with previous studies (Mackinder *et al*., 2009).

EhV production, defined as the number of EhVs measured upon lysis, was also light dependent. Comparative infection dynamics in the dark were performed after EhV was allowed time to adsorb to host cells in the light. Although uninfected *E. huxleyi* cultures did not grow in the dark (Fig. 2b), cell lysis and declines in host abundance were observed for both infected cultures regardless of light or dark treatment (Fig. 2b). However, the 71% lower virus production in the infected, dark treatment (Fig. S2b) resulted in a 3.5-fold lower burst size (i.e. the number of viruses produced per *E. huxleyi* cell) compared with infected cultures grown on an LD cycle (Fig. 2c).

Under continuous light, *E. huxleyi* growth and Fv/Fm were similar to cultures grown on a LD cycle (Figs 2d, S2c), indicating cells were not physiologically stressed, consistent with previous observations (Nielsen, 1997). However, infected cultures under continuous light produced c. 50% more viruses (Fig. S2d), resulting in a 1.5-fold higher burst size (Fig. 2e). Infected cultures in continuous light also had nearly twofold higher PPP activity than LD infected cultures (average over the entire experiment was 4.7 ± 1.4 nM⁻¹ and 2.5 ± 0.6 nM⁻¹ NADPH min⁻¹ μg protein⁻¹ respectively), with significant increases observed as early as 6 hpi under continuous light compared with 20 hpi under LD conditions (Fig. 2f,g).

We also tested how photon flux impacted host–virus interactions. The maximum steady-state growth rate μmax at irradiance

| Table 1 NADPH production rate with time, hours postinfection (hpi), in control and infected *Emiliania huxleyi* cultures. |
|---------------------------------------------------------------|
| **NADPH production rate (nM min⁻¹ μg protein⁻¹)**             |
|                  | 4 hpi | 14 hpi | 19 hpi | 24 hpi | 28 hpi |
|------------------|-------|--------|--------|--------|--------|
| Control          | 10.9 ± 0.9 | 9.1 ± 0.3 | 8.4 ± 0.9 | 9.2 ± 0.4 | 7.2 ± 1.4 |
| Infected         | 10.6 ± 0.2 | 14.0 ± 1.9* | 18.4 ± 2.3* | 26.1 ± 1.3** | 23.6 ± 1.5** |

Mean ± SD for biological triplicates is shown; t-test: *P < 0.01; **P < 0.001.
levels of 150, 300, and 500 $\mu$mol m$^{-2}$ s$^{-1}$ was similar ($t$-test $P$-value > 0.05; in the range 0.89–0.96 ± 0.10–0.14 d$^{-1}$), whereas $\mu_{\text{max}}$ was 0.45 ± 0.09 d$^{-1}$ and 0.74 ± 0.10 d$^{-1}$ at irradiance levels of 25 $\mu$mol m$^{-2}$ s$^{-1}$ and 200 $\mu$mol m$^{-2}$ s$^{-1}$ respectively (Fig. 3a). At all irradiance levels except 2000 $\mu$mol m$^{-2}$ s$^{-1}$, $F_0/F_m$ was similar, suggesting growth at an irradiance of 25 $\mu$mol m$^{-2}$ s$^{-1}$ was limited by light rather than impaired photosynthetic efficiency (Fig. 3a). The lower $F_0/F_m$ at an irradiance of 2000 $\mu$mol m$^{-2}$ s$^{-1}$ suggests the decreased growth may have been due to impaired photophysiology. When infected at an irradiance of 25, 150 or 300 $\mu$mol m$^{-2}$ s$^{-1}$, host lysis occurred c. 4 d postinfection (dpi; Fig. 3b). By contrast, lysis occurred within 2–3 dpi at an irradiance > 500 $\mu$mol m$^{-2}$ s$^{-1}$ (Fig. 3b). EhV production was maximal at intermediate irradiance (150 and 300 $\mu$mol m$^{-2}$ s$^{-1}$, Fig. 3c) and was approx. five-fold lower at irradiances of 25 and 500 $\mu$mol m$^{-2}$ s$^{-1}$, and 16-fold lower at irradiances of 1000 and 2000 $\mu$mol m$^{-2}$ s$^{-1}$ (Fig. 3c). When expressed as a burst size, the intermediate irradiance levels produced the highest number of viruses per host (Fig. 3d). This relationship was robust, being observed across a large number ($n = 5$–25) of discrete samples (Fig. S3).

Photoperiod and irradiance serve as a switch between de novo and recycled nucleotide synthesis

Given the increased PPP activity concomitant with decreased RubisCO expression, we hypothesized enhanced viral production in the light was due to a concerted strategy to increase nucleotide synthesis and viral genome replication, while maintaining host integrity. To test this, we developed a mathematical model to quantify the dependence of free virus population dynamics on the rate of de novo nucleotide synthesis (Eqns 1–5; Fig. S4), accounting for the potentially interacting and compensating effects of host growth and recycling of host material. Our premise was that host growth $\mu_h$, nucleotide recycling $\mu_r$ and de novo nucleotide synthesis $\mu_v$ each have distinct and constrainable effects on population dynamics. We leveraged the dependence of host–virus population dynamics on these internal processes to infer their light dependence.

In Fig. 4 we show the contrasting effect within the model of nucleotide recycling $\mu_r$ and de novo nucleotide synthesis $\mu_v$ on population dynamics. Increasing the rate of nucleotide recycling leads to an earlier decline of the host population (Fig. 4a) and...
lower viral production due to the premature lysis of the host (Fig. 4b). Increased *de novo* synthesis also leads to early host decline (Fig. 4c) but results in enhanced viral production (Fig. 4d). The early decline of the host with increased *de novo* nucleotide synthesis is a consequence of our assumption that the rate of lysis is influenced positively by the accumulation of...
internal viral nucleotides (Eqns 3–5, 7). However, the concomitant increase in the rate of de novo nucleotide synthesis allows increased viral production. The contrasting dynamics in Fig. 4(a, b) and Fig. 4(c, d) respectively provide a distinct signature that allows the model to make predictions about nucleotide recycling $\mu_v$ and de novo synthesis $\mu_{\text{n}}$, based on host and virus population dynamics.

To assess the light dependence of these rates, we first assessed whether the model with these assumptions was consistent with the observed host–virus population dynamics. We then asked, using a formal fitting procedure with explicit quantification of parameter uncertainty (Metropolis et al., 1953), whether there were clear differences in rates of recycling and de novo nucleotide synthesis between light treatments.

The model successfully captured host–virus nucleotide dynamics in cultures infected in the dark, on an LD cycle, and under continuous light (Fig. S5). For the dark and LD treatments, the Metropolis algorithm converged on unique parameter distributions (Fig. 5), demonstrating that the empirical measurements were sufficient to constrain model parameters in these conditions. The continuous light treatment had too few measurements to constrain parameter values (Fig. S5e,f).

The model output suggests that host nucleotide recycling and de novo nucleotide synthesis were both strongly dependent on light, and the contribution of nucleotides to viral genome replication by each pathway could be toggled by external light conditions. In dark treatments, the model predicted rapid nucleotide recycling, accompanied by relatively slow de novo synthesis (Fig. 5a,b). This prediction is due to rapid demise of host populations in the dark and low viral production (Figs 2b, 4a,b). Conversely, LD treatments showed slower recycling and more rapid de novo synthesis (Fig 5a,b), arising from delayed host lysis and a large burst of free-virions toward the end of the infection (Figs 2, 4c,d).
Fig. 4 Sensitivity analysis of infection dynamics to the rate of host nucleotide recycling $\mu_i$ and de novo synthesis $\mu'_i$ during the time-course of infection. When a higher rate of nucleotide recycling is assumed, (a) host abundance declines more rapidly and (b) free extracellular virus abundance decreases. When a high rate of de novo synthesis is assumed, (c) host abundance declines more rapidly due to faster internal progression of infection and (d) viral abundance increases. The distinct signature of these parameters on host–virus population dynamics allows their light dependence to be quantified.

Fig. 5 Rates of viral-mediated recycling of host nucleotides and de novo nucleotide synthesis are sensitive to irradiance level. Parameter estimates for the rates of host nucleotide recycling or de novo synthesis, obtained using the Metropolis algorithm to fit Eqns 1–5 to Emiliania huxleyi–Coccolithovirus nucleotide dynamics shown in Supporting Information Fig. S4. (a, b) Maximal recycling $\mu_i$ and de novo synthesis $\mu'_i$ in dark and light : dark (LD) treatments. Box edges are lower and upper quartiles, and whiskers are 5th and 95th quartiles of the posterior parameter distributions. (c, d) Estimates of $\mu_i$ and $\mu'_i$ for irradiance level experiments. Error bars are lower and upper quartiles either side of the medians.
Contrasting parameter predictions during LD vs dark treatments were insensitive to a wide range of virus adsorption rates \( \phi \), suggesting predictions of \textit{de novo} synthesis and recycling rates are robust to any uncertainty in our predictions of \( \phi \) (Fig. S6).

Repeating the fitting procedure for experiments at different light intensities revealed that the rate of nucleotide recycling and \textit{de novo} synthesis were also strongly dependent on irradiance level. Nucleotide recycling was lowest at intermediate irradiance (150–500 \( \mu \)mol m\(^{-2}\) s\(^{-1}\)) and highest at low and high irradiances (25, 1000–2000 \( \mu \)mol m\(^{-2}\) s\(^{-1}\); Fig. 5c). \textit{De novo} synthesis was opposite, being the highest at intermediate irradiance and lowest at low and high irradiance (Fig. 5d). In some cases, predictions of recycling and \textit{de novo} synthesis rates were poorly constrained by the experimental data (Fig. 5c,d), but the clear difference across light intensities in spite of this large error supports the hypothesis that key infection parameters are light dependent. Taken together, these model predictions demonstrate that slow recycling and rapid \textit{de novo} synthesis combine to impact EhV production and that light can lead to a systematic shift between nucleotide recycling and \textit{de novo} synthesis.

Light structures infection of natural populations

The termination of dense coastal and open ocean North Atlantic blooms of \textit{E. huxleyi}, which can span \( 10^5 \) km\(^2\) (Holligan \textit{et al.}, 1993; Tyrell & Merico, 2004) and last over a period of days to weeks, is commonly attributed to EhV infection (Brathak \textit{et al.}, 1993; Vardi \textit{et al.}, 2012; Lehahn \textit{et al.}, 2014). We used the 2012 North Atlantic Virus Infection of Coccolithophores Expedition as a platform to extend our laboratory-based mechanistic findings and test the hypothesis that irradiance structures the infection of natural populations of \textit{E. huxleyi}. We previously demonstrated these populations were at various stages of infection (Laber \textit{et al.}, 2018) using a suite of structurally distinct glycosphingo- and betaine-like lipids that represent functional biomarkers of the \textit{E. huxleyi}–EhV infection process (Vardi \textit{et al.}, 2009, 2012; Fulton \textit{et al.}, 2014; Hunter \textit{et al.}, 2015). Using targeted, quantitative PCR we quantified the intracellular abundance of \textit{E. huxleyi}-associated EhVs (expressed as a virus-to-microbe ratio, VMR) at three to five depths (97 discrete samples) in the euphotic zone at 21 stations across the c. 2000 nautical mile cruise track (Figs 6, S7). At 18 of the 21 stations, there was a subsurface VMR maximum that corresponded to c. 1–10% of the surface irradiance \( I_0 \). The relationship between VMR and percentage surface \( I_0 \) at these stations was similar to that between burst size and irradiance level seen in laboratory cultures (Fig. 3d), in that there was an optimal percentage surface \( I_0 \) that supported the highest VMR; above and below that optimum, VMR decreased. Taking the entire dataset together, there was a significant positive relationship between VMR and depth (\( P = 6.19 \times 10^{-4} \), slope 0.02, \( R^2 = 0.12 \); linear regression with log-transformed VMR and depth), despite the inherent variability in pooling across stations. These observations corroborate active, subsurface infection of these populations (Laber \textit{et al.},
Table 2  Statistical analysis of North Atlantic environmental samples collected during the North Atlantic Viral Infection of Coccolithophores Expedition.

| Parameter                  | Units (log_{10}) | $P$-value | Slope | $R^2$ | $n$ | Outliers? |
|----------------------------|------------------|-----------|-------|-------|-----|-----------|
| Physical                   |                  |           |       |       |     |           |
| Depth                      | m                | $6.19 \times 10^{-4}$ | 0.02  | 0.12  | 97  | No        |
| Light                      | Surface irradiance (%) | $5.42 \times 10^{-6}$ | -0.44 | 0.15  | 97  | No        |
| Salinity                   | PSU              | > 0.1     | -29.01| 0.03  | 97  | Yes       |
| Mixed-layer depth          | m                | 0.24      | 0.82  | 0.015 | 97  | No        |
| Temperature                | °C               | 0.91      | 0.75  | 0.01  | 97  | No        |
| Chemical                   |                  |           |       |       |     |           |
| Nitrate and nitrite        | μM               | 0.29      | -0.42 | 0.012 | 94  | Yes       |
| Ammonium                   | μM               | 0.57      | 0.19  | 0.004 | 97  | No        |
| Phosphate                  | μM               | 0.30      | 0.22  | 0.014 | 81  | Yes       |
| Dissolved O$_2$            | ml l$^{-1}$      | 0.02      | -5.73 | 0.06  | 97  | Yes       |

All stations from the cruise were pooled within each analysis to probe the prediction from laboratory studies that the intracellular virus-to-microbe ratio (VMR) would vary with depth. Subsequent analyses examined which component of depth (physical or chemical) drives this relationship. All data were analyzed by linear regression against the VMR, with both predictors and VMR log-transformed; parameter, units, significance ($P$-value), slope, goodness of fit ($R^2$), and sample size ($n$) of regression lines, and the presence of outlying values are shown. Bold $P$-values are those with significance < 0.001.

2018; Sheyn et al., 2018), with the degree of infection correlating with depth. We then analyzed physical (light, temperature, salinity, mixed layer depth) and chemical (phosphate, nitrate and nitrite, ammonium, dissolved oxygen) aspects of this depth distribution to determine which parameters were the most robust predictors of the VMR–depth relationship. Light was the only significant ($P = 3.14 \times 10^{-4}$) and robust ($R^2 = 0.13$) predictor of VMR (Table 2; Fig. S8) across depth, whereas the other parameters were either not significant ($P > 0.05$) or only weakly ($R^2$ ranged from 0.04 to 0.06, often prone to the impact of outliers) related to VMR.

Discussion

Elucidating the cellular processes that fundamentally regulate EhV infection is essential for developing a mechanistic framework for interpreting and modeling algal host–virus interactions. Given their inability to replicate without a host, it follows that obtaining nucleotides, either through host genome recycling or de novo synthesis, represents the first and foremost challenge to viral production. As the primary mechanism for de novo nucleotide synthesis, an increased PPP activity during infection has been documented in a wide range of systems, including animals, plants, and cyanobacteria (Bissell et al., 1973; Šindelář et al., 1999; Lachaise et al., 2001; Thompson et al., 2011). Widespread enrichment of genes associated with the PPP has also been observed in marine metaviromes (Enav et al., 2014).

Here, we show that EhV infection increases host PPP activity, consistent with increased PPP-associated gene expression (Rosenwasser et al., 2014). We suggest this induced activity allows EhV to meet the requirement for genome replication that cannot be supported by host nucleotide recycling alone. With a genome size of c. 400,000 bp (Nissimov et al., 2012) and burst sizes ranging from 500 to 1000 virions per host (this study; Castberg et al., 2002), EhV201 would conservatively require $2 \times 10^8$ nucleotides for replication. Even if all $1.42 \times 10^9$ nucleotides of the E. huxleyi genome (Read et al., 2013) were recycled, EhV would still require an additional $6 \times 10^7$ nucleotides to achieve a mature burst size. This disparity becomes even more pronounced when the discrepancy between the guanine–cytosine content of E. huxleyi (66%) and EhV (40%) is taken into account. Although the plastid genome could also provide a source of recycled nucleotides, our observations of high PsbD expression (a gene encoded by the plastid genome) in infected cultures argues against plastid genome recycling.

The importance of de novo nucleotide synthesis is further supported by reduced RubisCO protein levels and maintenance of electron transport rates and PSII activity. By diverting photosynthetic energy toward the PPP, viruses may enhance nucleotide synthesis while preserving energy that would otherwise be spent fixing unnecessary carbon. While functionally equivalent to similar observations in cyanophage (Lindell et al., 2004; Thompson et al., 2011), the mechanism by which EhV maintains host photosynthesis while decreasing RubisCO is unclear.

Light-dependent adsorption and entry of EhV to host cells is conceptually similar to cyanophage (Cséke & Farkas, 1979; Kao et al., 2005; Jia et al., 2010), and we hypothesize this dependence is related to the role lipid rafts play in facilitating EhV entry (Rose et al., 2014). Lipid rafts, membrane microdomains that sense extracellular stimuli and activate various signaling cascades through protein–protein interactions, have been implicated as entry sites of pathogens and viruses in other systems (Neilan et al., 1999; Chazal & Gerlier, 2003; Hajishengallis & Lambris, 2011). Proteomic analysis of purified lipid rafts from infected E. huxleyi cells suggests EhVs utilize a similar entry mechanism (Rose et al., 2014). Intriguingly, E. huxleyi lipids rafts were also found enriched in light-harvesting and Chl$a/b$ binding proteins (Rose et al., 2014). Although a role for lipid rafts in light capture is unknown, we posit that light regulates lipid raft production and thereby functionally limits entry of EhV during periods of darkness.

Viral replication and production were also impacted by light, with fewer viruses produced when infected cultures were maintained in the dark. As obligate photoautotrophs, prolonged
darkness represents a significant physiological stress, and the lack of viral production may simply be due to host senescence. However, if cell mortality was significant, we would expect a dramatic decrease in host abundance. Rather, we observed no net change in cell abundance, suggesting either equal growth and mortality or that cells were neither dividing nor dying. By contrast, when host cultures were maintained under continuous light, viral production was enhanced, further supporting the hypothesis that light is required for maximum viral production. In addition, elevated basal PPP activity in continuous light cultures compared with LD cultures suggests increased EhV production is supported by the availability of additional nucleotides through enhanced \textit{de novo} synthesis.

Despite evidence that algal viruses have the potential to be a primary control on natural communities (Bratbak \textit{et al.}, 1993; Schroeder \textit{et al.}, 2003; Vardi \textit{et al.}, 2012; Laber \textit{et al.}, 2018; Sheyn \textit{et al.}, 2018), their impacts are not quantitatively assessed in ecosystem models resolving coupled interactions between phytoplankton, grazers, and bacteria. This is in part due to the lack of empirical data on key parameters, such as encounter and adsorption with hosts ($\phi$), replication rates (e.g. $\mu_a$, $\mu_v$) and host lysis ($\delta$) (Record \textit{et al.}, 2016). Our empirically based mathematical model of EhV production hypothesizes that viral production is influenced by the rate of nucleotide recycling and \textit{de novo} synthesis. In the dark, when host metabolism is suppressed, viral production is diminished due to a combination of rapid nucleotide recycling and diminished \textit{de novo} synthesis. When hosts grow rapidly at intermediate irradiance, reduced host nucleotide recycling can support enhanced viral production because host integrity and growth are preserved, leading to a larger pool of available host nucleotides for viral genome replication. At high light, viral production is diminished due to a combination of rapid nucleotide recycling and diminished rates of \textit{de novo} synthesis. Together, these insights form a base with which to explore ecosystem effects of host–virus metabolism in contrasting aquatic light environments.

Our laboratory- and modeling-based data suggest EhV adsorption, and subsequent conversion and/or synthesis of nucleotides, is enhanced in the light and that maximum viral replication occurred at mid-range irradiance levels. At high light, viral production may be inhibited by high reactive oxygen species production (Sheyn \textit{et al.}, 2016), whereas at low light there may simply not be enough \textit{de novo} nucleotide synthesis to support high viral production. Alternatively, the shorter latent period (i.e. the time until host lysis) at the higher irradiance levels (>500 $\mu$mol m$^{-2}$ s$^{-1}$) may itself represent an advantageous strategy whereby the virus induces host lysis to ensure virion release. This has been observed in bacteriophage, whereby the optimal lysis time is dependent on host abundance and physiology (Wang, 2006). We also cannot rule out a direct effect of light on the viruses. Although ultraviolet radiation is the major cause of viral decay in the ocean (Suttle & Chen, 1992; Noble & Fuhrman, 1997), photosynthetically active radiation has also been documented to facilitate decay (Traving \textit{et al.}, 2014; Wei \textit{et al.}, 2018). Exploring the effect of photosynthetically active radiation on viral abundance, infectivity, and adsorption needs to be further investigated to better understand the mechanism by which light impacts the latent period and viral production. Regardless, our data suggest virus activity in the upper ocean may be vertically structured by light. Within stations, the highest intracellular EhV abundance measured in natural populations occurred at depths corresponding to c. 1–10% of the surface irradiance ($I_0$), near the base of the euphotic zone, with lower abundances above and below this level. This was not due to the number of available hosts, given that VMR is normalized to host abundance; and importantly, these data are consistent with our laboratory findings that EhV replication is maximal between irradiances of 150 and 300 $\mu$mol m$^{-2}$ s$^{-1}$, levels that would correspond to c. 7–15% surface $I_0$ on a sunny day (c. 2000 $\mu$mol m$^{-2}$ s$^{-1}$). When the entire dataset is pooled (97 discrete samples across 2000 nautical miles) and tested against a variety of environmental parameters, light was the only one found to have a significant relationship with VMR, with increasing light leading to lower VMR. The somewhat low, but significant, $R^2$ value (0.13) may be partially attributed to sampled populations being at different stages of infection (Laber \textit{et al.}, 2018), but nonetheless suggests a relationship between light and infection. Although we lose the resolution to disentangle the relationship seen at individual stations when we pool all of the samples, taken together with our laboratory data, these findings demonstrate that light can serve as a driving mechanism by which EhV infection can be vertically structured in natural populations.

Our laboratory-, field-, and modeling-based work form a mechanistic and empirical foundation on which to constrain key parameters regulating algal host–virus interactions. Our finding that light regulates critical aspects of EhV replication and production opens the door for mechanistic models of host–virus impacts on ecosystem function across a gradient of light environments. This, in turn, provides a novel and rigorous means to disentangle competing controls on phytoplankton populations, and the impact of viral infection on carbon cycling and nutrient dynamics.

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Author contributions

KT and KDB conceived the study. KT designed the experiments. KT, LH, CM and JRL performed the experiments, DT and MJF performed the modeling. MJLC contributed the
quantitative PCR data from the North Atlantic field samples, FN performed the flow cytometry, BK analyzed the field data and contributed statistical analysis. KT, DT and KDB wrote the paper with contributions from BK, MJLC and MJF.

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Supporting Information

Additional Supporting Information may be found online in the Supporting Information section at the end of the article:

**Fig. S1** Time course of host and viral abundance during a typical infection experiment.

**Fig. S2** Additional host and EhV abundance data for dark and continuous light infection experiments.

**Fig. S3** Additional EhV abundance and burst size data for infections at different irradiance levels.

**Fig. S4** Fitted model simulations to adsorption experiments.

**Fig. S5** Mathematical model capturing host–virus population dynamics in different light conditions.

**Fig. S6** Sensitivity of key fitted parameters to the rate of virus adsorption ($\phi$).

**Fig. S7** Depth profile of *Emiliania huxleyi* and EhV abundance.

**Fig. S8** Linear regression analysis of VMR and physical and chemical parameters measured across the North Atlantic.

**Table S1** Model parameter definitions and units.

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