Seasonal dynamics of natural *Ostreococcus* viral infection at the single cell level using VirusFISH

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SIGNIFICANCE STATEMENT

One of the major challenges in viral ecology is the detection of viral-host interactions in complex communities, with the aim of assessing the impact of viruses in the populations of their host. Here we present the proof of concept that Virus fluorescence in situ hybridization (VirusFISH) can be used to visualize and monitor viral infection dynamics in natural occurring populations of Ostreococcus, a cosmopolitan picoeukaryote that form occasional blooms in nature. This technique can be easily implemented for other viral-host systems, allowing to visualize their interaction at the single-cell level.

ABSTRACT

Ostreococcus is a cosmopolitan marine genus of phytoplankton found in mesotrophic and oligotrophic waters, and the smallest free-living eukaryotes known to date, with a cell diameter close to 1 µm. Ostreococcus has been extensively studied as a model system to investigate viral–host dynamics in culture, yet the impact of viruses in naturally occurring populations is largely unknown. Here, we used Virus Fluorescence in situ Hybridization (VirusFISH) to visualize and quantify viral-host dynamics in natural populations of Ostreococcus during a seasonal cycle in the central Cantabrian Sea (Southern Bay of Biscay). Ostreococcus were predominantly found during summer and autumn at surface and 50 m depth, in coastal, mid-shelf and shelf waters, representing up to 21% of the picoeukaryotic communities. Viral infection was only detected in surface waters, and its impact was variable but highest from May to July and November to December, when up to half of the population was infected. Metatranscriptomic data available from the mid-shelf station unveiled that the Ostreococcus population was dominated by the species O.
*Lucimarina*. This work represents a proof of concept that the VirusFISH technique can be used to quantify the impact of viruses on targeted populations of key microbes from complex natural communities.

**KEYWORDS**

VirusFISH; natural communities; *Ostreococcus* spp.; viruses; infection dynamics.
INTRODUCTION

Quantifying the *in situ* impact of viruses on their hosts is one of the major challenges in marine viral ecology, and requires the detection of both specific viruses and their hosts within complex natural communities. Due to the absence of a universal phylogenetic marker for viruses, this has been achieved through PCR amplification of conserved marker genes within specific viral families (Chen and Suttle, 1995; Larsen *et al.*, 2008; Lehahn *et al.*, 2014; Gran-Stadniczeñko *et al.*, 2019), or through metagenomics (e.g. Mizuno *et al.*, 2013; Roux *et al.*, 2017). However, the identity of the viral host from metagenomic data can only be inferred based on the known host of cultured relatives, or by correlation with the presence of potential host sequences (Lima-Mendez *et al.*, 2015; Nishimura *et al.*, 2017). Other recently developed techniques to detect virus–host interactions in complex communities are digital droplet PCR (Lim *et al.*, 2017), and single-cell genomics (Roux *et al.*, 2014; Castillo *et al.*, 2019). Additionally, metatranscriptomics have also been used to follow some infection dynamics (Zeigler Allen *et al.*, 2017; Alonso-Sáez *et al.*, 2018). Yet, the impact of viruses on their host populations, as the proportion of infected cells, is not directly measured by any of those techniques, even though it is crucial information to understand virus–host population dynamics.

Promising alternatives to explore *in situ* viral-host interactions are phageFISH (Allers *et al.*, 2013), and its variants direct-geneFISH (Barrero-Canosa *et al.*, 2017), VirusFISH (Castillo *et al.*, 2020), or single-molecule FISH (Vincent *et al.*, 2021). Although their use in nature has been limited, phageFISH was recently used to pinpoint the host of an archaeal virus previously identified in an environmental metagenomic dataset (Hochstein *et al.*, 2016) and to demonstrate viral lysis of an autotrophic key player in the Earth’s crust (Rahlff *et al.*, 2020). Likewise, single-molecule FISH was recently used to quantify
active viral infection in an induced *Emiliania Huxleyi* bloom during a mesocosm experiment (Vincent *et al.*, 2021).

Due to the mounting evidence of the role that viruses may play in phytoplankton bloom termination, quite a lot of attention has been paid to viruses of bloom forming species, like the coccolithophore *Emiliania huxleyi* (Wilson *et al.*, 2002; Laber *et al.*, 2018; Sheyn *et al.*, 2018). Under high host cell abundances, such as those found in *E. huxleyi* blooms, the probability of encountering a virus with its host is high, which may result in a fast viral propagation through the host population (Suttle, 2000). By contrast, the impact of viruses on picoeukaryotic hosts that may form occasional blooms but are generally present at low abundances (Zingone, 1999; O’Kelly *et al.*, 2003; Countway and Caron, 2006) could be more challenging, and it has been little explored. A few studies have focused on the spatial and temporal dynamics of viruses targeting those occasional blooming phytoplankton species, particularly *Micromonas* (Cottrell and Suttle, 1991, 1995; Zingone, 1999; Baudoux *et al.*, 2015) but also *Ostreococcus* (Bellec *et al.*, 2010). Nevertheless, these studies used either plaque assays or most probable number approaches, and therefore they did not provide direct evidence of *in situ* interactions between viruses and hosts.

Recently, we implemented VirusFISH to detect, visualize and follow viral–host dynamics of *Ostreococcus tauri–Ostreococcus tauri virus 5* (OtV5) in culture (Castillo *et al.*, 2020). *Ostreococcus* (Mamiellaceae) is the world’s smallest free-living eukaryote known, with a cell diameter of ~1 µm (Courties *et al.*, 1994; Derelle *et al.*, 2006; Sanchez *et al.*, 2019). The genus comprises several species that can be ubiquitously found from the coast to the open ocean, and from mesotrophic to oligotrophic waters (Demir-Hilton *et al.*, 2011; Tragin and Vaulot, 2019). The number of sequenced *Ostreococcus* viruses is constantly increasing, and nowadays the complete genomes of many *Ostreococcus*
viruses are available (e.g. Derelle et al., 2008, 2015; Weynberg et al., 2011; Monier et al., 2017). Yet, nothing is known about the Ostreococcus virus–host relationships in situ, and how they change over a temporal scale. Our VirusFISH approach combines a Catalyzed Reporter Deposition Fluorescence in situ Hybridization detection of Ostreococcus species with the general OSTREO01 probe (Not et al., 2004) and viral probes originally designed for the detection of OtV5, but that target most Ostreococcus virus sequenced to date (Castillo et al., 2020).

Here we used VirusFISH to study the monthly dynamics of viral infection in natural populations of Ostreococcus over a full seasonal cycle. We demonstrate that VirusFISH is a powerful tool for assessing virus–host interactions in the environment, even when the hosts are present at low abundances.

MATERIALS AND METHODS

Environmental sampling

Samples were collected monthly along a coastal-shelf gradient in the Cantabrian Sea (Southern Bay of Biscay, near Xixón, Spain) at three stations: E1, coastal (30 m maximum depth (max depth); 43.58° N, 5.61° W), E2, mid-shelf (110 m max depth; 43.67° N, 5.58° W) and E3, shelf (160 m max depth; 43.78° N, 5.55° W) (Fig. 1), from January 2011 to December 2012, at the surface (5m) and at 50m depth (except for E1 where the maximum depth was 30 m). Temperature and salinity were measured by a SeaBird 25 CTD. Samples for chlorophyll a (Chl a) concentration were collected by sequentially filtering 200 mL seawater onto 20, 2 and 0.2 µm polycarbonate filters. Chl a concentration in the picoplankton size fraction (pChl, <2 µm) corresponded to the amount collected onto 0.2
µm filters after passing through 20 µm and 2 µm filters. Filters were kept frozen at −20°C and processed within two weeks, as explained in Calvo-Díaz and Morán (2006). Picoeukaryotic abundances were acquired by flow cytometry using 1.8 mL subsamples fixed with paraformaldehyde + glutaraldehyde (1% and 0.5% final concentration, respectively) as described in (Calvo-Díaz and Morán, 2006). A total of 120 samples for Chl a and picoeukaryotic abundances were analyzed.

**VirusFISH: sample preparation, labeling and analysis**

Samples for FISH (4 mL) were collected at surface and at 50 m depths from the 3 different stations (except for E1, where the maximum depth was 30m) from January to December 2012, as described in (Arandia-Gorostidi et al., 2017). Samples were fixed with 3% freshly filtered formaldehyde and cells were collected onto 0.2 µm pore-size polycarbonate filters. Filters were kept at −80°C until their analysis. These samples were originally collected to perform FISH on bacterial groups which explains the low volume filtered. For the VirusFISH, cells and viruses were hybridized and analyzed as described in (Castillo et al., 2020). Briefly, samples were treated with alcohols to remove pigments, then cells were hybridized with the OSTREO01 probe for CARD-FISH, labeled with Alexa488, and after, viruses were hybridized with the 11 viral probes designed for *Ostreococcus* viruses labeled with Alexa594 (Castillo et al., 2020). Although the probes were originally designed for the *Ostreococcus tauri* virus 5 (OtV5), the genome region targeted by the probes is highly similar for most *Ostreococcus* virus sequenced to date (Fig. S1A, Table S1). Thus, our probes putatively target most *Ostreococcus* viruses. In order to further test this, we applied VirusFISH to a different *Ostreococcus*-virus system: *Ostreococcus mediterraneus* strain MA3 with the virus OmV2 (Yau et al., 2020), and compared the VirusFISH visualization to previous results obtained with the *O. tauri* -
OtV5 system (see supplementary methods and Fig. S1B,C). This confirmed that the
probes hybridize to other *Ostreococcus* viruses.

*Ostreococcus* cells were observed by epifluorescence microscopy under blue light
(475/30 nm excitation, 527/54 BP emission, and FT 495 beam splitter) and *Ostreococcus*
viruses under orange light (585/35 nm excitation, 615 LP emission, and FT 570 beam
splitter). All pictures were taken using the same intensities and exposure times (300 ms
for the blue light and 1 s for the orange light). For each sample, 4 random transects,
between 6 and 10 mm each, were performed to visualize and count infected and non-
infected *Ostreococcus*. The number of inspected *Ostreococcus* cells ranged from non-
detected to 20 cells per sample. A total of 57 samples were evaluated with VirusFISH.

**Identification of *Ostreococcus* spp. and *Ostreococcus* virus sequences in
metatranscriptomes**

Metatranscriptomic information from the continental shelf station (E2) during 2011 and
2012 was retrieved from (Alonso-Sáez *et al.*, 2020). Metatranscriptomic reads, previously
quality trimmed and cleaned of rRNA sequences, were screened for *Ostreococcus* spp.
(OS) and *Ostreococcus* virus (OV) sequences. First, a BLASTn database was constructed
of the four *Ostreococcus* species nuclear genomes (*O. tauri* RCC4221, *O. lucimarinus*
CCE9901, *Osterococcus* sp. RCC809 and *O. mediterraneus* RCC2590) and the 13
complete *Ostreococcus* spp. virus genomes sequenced to date. The Genbank accession
numbers of the genomes used were as follows. *O. tauri*: CAID0100001.2–
WMKK0100001.1–WMKK01000022.1, OtV1: FN386611.1, OtV2: FN600414.1,
OtV5: EU304328.2, OtV6: JN225873.1, OIV1: MK514405.1, OIV2: KP874736.1, OIV3:
HQ633060.1, OIV4: JF974316.1, OIV5: HQ632827.1, OIV6: HQ633059.1, OIV7: MK514406.1 and OmV1: KP874735.1 and OmV2 (MN688676). The *Ostreococcus* sp. RCC809 genome was obtained from the JGI Genome portal (https://genome.jgi.doe.gov/portal/ – accessed 28 February 2014). Second, the metatranscriptomic reads were queried against OS and OV genomes by BLASTn (BLAST 2.2.26+), accepting high scoring pairs with e-value <1e−5, identity >75% and query coverage >75%. This nucleotide identity cut-off was chosen as it corresponds to the average nucleotide identity between *Ostreococcus* spp. (*O. tauri* and *O. lucimarinus*), as well as between representatives of *Ostreococcus* virus clades (OtV5 and OtV6), and thereby avoids retrieving reads that originate from related Mamiellophyceae and prasinoviruses. Average nucleotide identities were calculated with the ANI server (http://enve-omics.ce.gatech.edu/ani). Third, metatranscriptomic reads matching OS and OV genomes from each sample were counted, assigned to the species corresponding to the top BLASTn hit. Finally, OS and OV read counts were expressed as counts per 100,000 reads to adjust for variation in per sample sequencing depth.

**Transcriptome coverage of *Ostreococcus* viruses**

To determine which regions of the viral genomes were expressed, metatranscriptomic reads were aligned to available genomes of viruses infecting *Ostreococcus* using BWA version 7.17 (Li and Durbin, 2009) with default parameters. The resulting alignments were visualized in IGV version 2.5.3 (Robinson *et al.*, 2011) as a Sashimi Plot. Read counts for the predicted coding sequences (CDS) were counted for each genome using the HTSeq version 0.9.1 (Anders *et al.*, 2015) package with the function htseq-count (default parameters except for -t CDS -i = “locus_tag”) taking the alignment files from
BWA and the predicted CDS from the published GenBank genome annotations as input files.

**Statistical analysis**

Correlation analyses were performed using Pearson correlation. All statistical analyses were accomplished with the JMP 9.0.1 (JMP®, Version 9.0.1. SAS Institute Inc., Cary, NC, 1989-2019.) or R 3.5.3 (R Development Core Team, 2016) software.

**RESULTS**

**Environmental setting**

In surface waters, the temperature ranged from ~12.3°C in winter to ~21.2°C in summer during 2011 and 2012 (Fig. S2). In contrast, salinity was rather constant throughout the year at an average of 35.7, with occasionally lower values in winter and autumn, and a marked decline in April 2012 at all the three stations (~35, Fig. S2). Chlorophyll a (Chl a) concentration at the surface in 2011 and 2012 peaked during spring and autumn reaching values of ~1 µg L⁻¹ at the three stations, and also in summer at station E1. At 50 m depth, there was much less variation than at the surface in temperature (range: 12–18.3 °C) and salinity (range: 35.6–35.8). In contrast, Chl a showed three peaks: in June at E2 in 2011, in May at E3 in 2012, and in late summer at both stations in 2012 (Fig. S2). Between June and November, a subsurface chlorophyll maximum (DCM) developed at around 40–50 m at the two shelf stations. As expected, nitrate (NO₃) and phosphate (PO₄) concentrations were in general lower at the surface than at the DCM for all stations and reached their maximum values during winter at both depths (Table S2).

Abundance of autotrophic picoeukaryotes (PE) was in general two-fold higher in surface waters than at 50 m depth (Fig. 2). At the surface, PE reached maximum abundances in
April and November for all three stations, with E1 also showing high values in summer, coincident with the peak in Chl \(a\). At 50 m depth, PE were almost absent during winter but from late spring to autumn their abundance ranged between 5,000 and 20,000 cells mL\(^{-1}\) (Fig. 2).

**Dynamics of *Ostreococcus* and its viral infection during an annual cycle**

Using VirusFISH we tracked the abundance of *Ostreococcus* and virally infected *Ostreococcus* cells during 2012. *Ostreococcus* cells were counted as infected when the red fluorescence of the VirusFISH probe (see methods section) overlapped with the green signal of the *Ostreococcus* CARD-FISH probe (Fig. 3). The contribution of *Ostreococcus* to the picoeukaryotic assemblages over the seasonal cycle ranged from non-detectable to 20.8% in surface waters, averaging 2.6% (180 *Ostreococcus* mL\(^{-1}\)), and from non-detectable to 8.9% at 50 m depth, averaging 1.7% (184 *Ostreococcus* mL\(^{-1}\)) (Table S3). In surface waters at station E1, *Ostreococcus* abundances started to increase in late spring and reached the highest values in summer (1,226 *Ostreococcus* mL\(^{-1}\), Fig. 4). At the mid-shelf (E2) *Ostreococcus* cells displayed two relative maxima in July and November–December (508 – 361 *Ostreococcus* mL\(^{-1}\), Fig. 4), and at the shelf station (E3) we obtained similar results as in E2, with two relative maxima in July and November (578 – 342 *Ostreococcus* mL\(^{-1}\), Fig. 4). Remarkably, *Ostreococcus* cells could not be detected in August at the two stations more distant from shore (E2 and E3), whereas they showed maximal abundances at the coastal station E1 (Fig. 4A). At 50 m depth, *Ostreococcus* cells were also mainly found in summer and autumn, with the exception of October (Fig. 4B). Although *Ostreococcus* abundances reached higher values in surface waters than at 50 m depth, year-round average values were similar for both depths and among stations (i.e. E1 surface: 208.3±105 cells mL\(^{-1}\); E2 surface: 127.9±53.6 cells mL\(^{-1}\), E2 50 m:
151.1±38.3 cells mL\(^{-1}\); E3 surface: 121±54.1 cells mL\(^{-1}\), E3 50 m: 133.7±54.8 cells mL\(^{-1}\)).

In surface waters of E1, viral infection was observed in June, July, September, November and December, representing from 11 to 60% of the cells (22 – 78 infected *Ostreococcus* mL\(^{-1}\), Fig. 4). In E2, infected cells were visualized in late spring to early summer, representing from 7 to 50% of the cells (25 – 36 infected *Ostreococcus* mL\(^{-1}\), Fig. 4). In E3, we could only detect infected cells in November, which accounted for 25% of the *Ostreococcus* population (86 infected *Ostreococcus* mL\(^{-1}\), Fig. 4). Thus, the impact of viruses on *Ostreococcus* cells in surface waters along the coastal-shelf gradient analyzed here was variable, but infection took place mostly from May to June and from November to December (Fig. 4A, Table 1). Contrary to surface samples, at 50 m depth no infected cells could be detected at any time (Fig. 4B).

There was a significant positive relationship between the abundance of *Ostreococcus* and the abundance of autotrophic picoeukaryotes (Pearson correlation analysis. R=0.42, p-value=0.0016, n=54). The number of infected cells was also positively correlated with the abundance of autotrophic picoeukaryotes (Pearson correlation analysis. R=0.43, p-value=0.0013, n=54), but not with *Ostreococcus* abundance (Table S4).

**Detection of *Ostreococcus* and *Ostreococcus* viruses in metatranscriptomes**

Both *Ostreococcus* species (OS) and their viruses (OV) were detected in metatranscriptomic samples collected during 2011 and 2012 at the mid-shelf station (E2), except for May and July 2011, when OV were not detected, coincident with very low abundances of host transcripts (Fig. 5). The relative abundance of OS transcripts displayed a maximum in November, was second highest in April and remained low in the spring and summer months of May and July both in 2011 and 2012 (Fig. 5, upper panel).
The relative abundance of OV transcripts was more variable between sampling years. The highest values of OV transcripts were observed in April and November, being notably higher in 2011 than in 2012, and the lowest in May and July. The relative abundance of viral transcripts in relation to the abundance of host transcripts was higher in April 2011, pointing to a larger infection event at this sampling time (Fig. 5, lower panel).

Regarding the phylogenetic affiliation of the OS and OV transcripts, we found that the Ostreococcus assemblage maintained the same rank species abundance profile in all samples, with O. lucimarinus as the most transcriptionally active species (51–91% of Ostreococcus reads), followed by O. tauri (6–47% of reads), while Ostreococcus sp. RCC809 and O. mediterraneus were minor contributors (both 1–4% of reads). This pattern was also reflected in the OV transcript pool, with O. lucimarinus virus transcripts dominating, followed by O. tauri viruses. O. mediterraneus viruses represented a minor fraction of the transcripts, whereas the only known virus infecting Ostreococcus sp. RCC809, OtV2 (Weynberg et al., 2011), was not detected (Fig. 5).

When the percentage of infected cells detected by VirusFISH was compared to the ratio of OV/OS transcripts we obtained consistent results, with samples where the number of infected cells was higher having higher OV/OS ratios (Fig. S3).

**Transcriptome coverage of Ostreococcus viruses**

To determine which genomic regions of the Ostreococcus viruses were being transcribed in the samples, all metatranscriptomic reads were aligned to the model Ostreococcus virus strain, OtV5, which is the virus that has received the most extensive molecular characterization (Derelle et al., 2008, 2017; Yau et al., 2016), and other Ostreococcus virus. O. lucimarinus viruses showed approximately twice the percentage of CDS transcribed compared to O. tauri infecting viruses (~20% compared to ~9%, Table S5).
The pattern of transcription between *Ostreococcus* viruses was comparable (Fig. S4), and despite the low read coverage (~100 reads, Table S5) due to the low sequencing depth of the metatranscriptomes, transcripts were distributed along the viral genome length. This indicates that the entire viral genomes were transcribed *in situ*. Furthermore, genes likely involved in transcription, DNA replication and capsid assembly were expressed, suggesting that *Ostreococcus* viruses were captured during active infection of their host cells. The most highly expressed gene was the major capsid protein, further suggesting the viruses were sampled during lytic replication.

**DISCUSSION**

Our results showed generally low abundances of *Ostreococcus* over the seasonal cycle, although this genus occasionally represented up to ~20% of total picoeukaryotic cells (Table S3). This is in agreement with previous results showing that *Ostreococcus* in coastal and shelf sea waters present <5·10³ cells mL⁻¹ at the surface and DCM (Zhu *et al.*, 2005; Countway and Caron, 2006; Cardol *et al.*, 2008), unlike in lagoons such as the Thau Lagoon (NW Mediterranean, France), where *O. tauri* can dominate the phytoplankton assemblage based on flow cytometric signatures (Chrétiennot-Dinet *et al.*, 1995; Vaquer *et al.*, 1996). Despite *Ostreococcus* can produce sporadic blooms, increasing two orders of magnitude over its basal concentration and accounting for up to 70% of the total picoeukaryotic community (O’Kelly *et al.*, 2003; Countway and Caron, 2006), we did not observe any of these blooms over our seasonal cycle. With a few exceptions, the highest contribution of *Ostreococcus* to the picoeukaryotic assemblage occurred in summer (Table S3). This might indicate that this tiny picoeukaryote is better adapted than other members of the picoeukaryotic assemblage to grow under low inorganic nutrient conditions that characterize the period from April-May to October in these coastal waters.
Indeed, this period is usually characterized by maxima in picophytoplankton biomass (Calvo-Díaz and Morán, 2006; Calvo-Díaz et al., 2008).

*Ostreococcus* viral infection dynamics was variable throughout the year, with the percentage of infected cells ranging from non-detected to 60%. This supports the view that viruses may have an impact in controlling the abundance of *Ostreococcus* cells, as it has been suggested in other field studies where infected *Ostreococcus* cells were visualized with TEM (O’Kelly et al., 2003). Most *Ostreococcus* viruses isolated to date have fast infection cycles in culture, with clearance of the culture observed usually within two days (Derelle et al., 2008; Zimmerman et al., 2019; Castillo et al., 2020). However, coexistence between the host and the virus for longer periods of time has also been observed (Thomas et al., 2011; Yau et al., 2020). In the bloom-forming algae *Emiliania huxleyi* it was recently shown that virulent virus may prevalently display temperate infection dynamics in nature, switching only to a lethal infection when the physiology of the cells become compromised due to high cell densities (Knowles et al., 2020). Nonetheless, this may be different in species that show predominantly low abundances, like *Ostreococcus*. Moreover, our monthly sampling frequency was likely not enough to detect episodes of boom and bust in the *Ostreococcus* populations or to quantify the role of viruses in controlling their dynamics. Similarly, Johannessen et al. (2017) reported that Haptophyte and virus community composition and diversity varied substantially during an annual cycle without any clear pattern of covariance. In a study conducted over three years, both *Micromonas pusilla* and its viruses were shown to fluctuate widely on smaller time scales (i.e., weekly sampling, Zingone et al. 1999). Indeed, a tipping point during infection dynamics after which the infection rapidly propagates has been reported in both lab experiments and in natural populations (Zimmerman et al., 2019; Castillo et al., 2020; Vincent et al., 2021), as well as strong diel cycle infection dynamics (Aylward et al.,
Altogether, these observations suggest that high frequency samplings (i.e. over hourly to daily scales) should be carried out to fully apprehend virus–eukaryote interactions in nature. In any case, our work is the first approximation that directly assessed the impact of viruses on a picoeukaryotic population under non-bloom conditions in nature. The fact that *Ostreococcus* were found in very different abundance levels across the spatial and temporal gradient studied here was important to test the performance of VirusFISH on this model microorganism.

The use of a general *Ostreococcus* CARD-FISH probe does not allow to distinguish between species. However, metatranscriptomic data from surface waters unveiled that the dominating species was *O. lucimarinus*. A previous study has shown that this species inhabits waters from the surface to the DCM (Rodríguez *et al.*, 2005) and it is the most widely distributed, whereas *O. tauri* and *O. mediterraneus* are mostly restricted to the surface layer of coastal waters and lagoons (Rodríguez *et al.*, 2005; Tragin and Vaulot, 2019). Thus, it is likely that most *Ostreococcus* cells found in our samples belonged to *O. lucimarinus*.

The metatranscriptomic data also indicated that *O. lucimarinus* coexisted with several viruses infecting this species. *Ostreococcus* viral transcriptional activity was higher in 2011 than in 2012, when we did the VirusFISH analyses. However, even in 2011 their transcriptional activity was low relative to that of the hosts suggesting that the impact of viruses on the *Ostreococcus* populations was only moderate (Fig. 5). High *Ostreococcus* viral transcriptional activity relative to that of their putative hosts has been shown based on metatranscriptomics in the Baltic Sea (Zeigler Allen *et al.*, 2017). Therefore, we may have missed large infection events due to our monthly sampling frequency. A recent transcriptomic study on an infection of Prasinovirus upon *Ostreococcus* has shown that the viral attack occurs mostly at night (Derelle *et al.*, 2017),
which may also explain the low viral transcriptional activity detected in our samples, that
were taken around noon. Finally, other factors that may contribute to this relatively low
viral activity are the coexistence of distinct transcriptional states during infection
dynamics (Vincent et al., 2021), and the co-occurrence of susceptible and resistant host
phenotypes (Yau et al., 2020) to the array of Ostreococcus viruses present at each
sampling time-point (Fig. 5). A combination of metatranscriptomics with VirusFISH
analyses performed with higher sampling frequency should help gain a clearer insight
into the virus–host dynamics of natural populations of Ostreococcus.

It is important to note that it is possible that the VirusFISH probes used were not able to
detect the full diversity of virus infecting the natural populations of Ostreococcus. This
could be the reason for the lack of detection of infection at 50 m depth. However,
according to Allers et al. (2013), a single probe is enough to visually detect one virus,
with the detection efficiency increasing with the number of viral probes used, and most
of the Ostreococcus virus sequenced to date are highly similar in at least 8 of the probes
used (Table S1). The fact that the VirusFISH results were consistent with the
metatranscriptomic data suggests that a broad range of Ostreococcus virus can indeed be
hybridized by the probes, further supporting the use of these probes to monitor natural
infection dynamics of this important picoeukaryote. VirusFISH can detect both early and
late infection stages, as in our experiments with cultures we were able to quantify infected
Ostreococcus cells when viral production was still negligible (Castillo et al., 2020). Thus,
we believe that the reason we did not detect any infected cells at 50 m is because the level
of infection was likely lower than in the surface, and the volume of sample filtered was
probably too small for the low abundance of Ostreococcus cells. In fact, this may also be
the reason why we did not detect infection at some of the surface samples over the
seasonal cycle.
In conclusion, we show that VirusFISH has strong potential to follow the dynamics of hosts and their infecting viruses in nature. It requires the previous knowledge of the viral genome, and preferably also the host genome to design the adequate probes (i.e. probes that do not target regions of the host genome that are similar to the virus), as well as the viral DNA material to use it as template to synthetize the probes. Having both elements, this approach can be easily implemented with any genome-sequenced virus–host system available in culture. Furthermore, VirusFISH can also be used to unveil unknown eukaryotic hosts of abundant viruses detected in metaviromes, by using that environmental DNA to synthetize the viral probes. Hence, VirusFISH opens avenues in viral ecology to tackle the role of viruses in controlling the abundance of key players in marine microbial communities, allowing to visually quantify the impact on specific host populations.

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R Development Core Team (2016) R: a language and environment for statistical computing.


FIGURES LEGEND

Figure 1. Location of the sampling stations. Abbreviations: E1: coastal station; E2: mid-shelf station; E3: shelf station. Samples were taken at 0 and 50 m depth, except for E1 where the maximum depth was 20 m.

Figure 2. Small autotrophic picoeukaryote abundances for coastal (E1), mid-shelf (E2) and shelf (E3) waters during a two-year period (2011 and 2012). A. Surface and B. 50 m depth. Note the difference in the y-axis between figures A and B. Note: for July and December 2012 data were not available at 50 m depth samples.

Figure 3. Micrographs of Ostreococcus cells in natural samples from the Cantabrian Sea. Upper and lower panel: infected Ostreococcus cells (arrow with round head), in which the red signal of the VirusFISH labeled viruses can be easily seen. Middle panel: a healthy non-infected Ostreococcus cell (asterisk) and a lysed Ostreococcus cell showing the viral cloud released from the cell with almost no cytoplasm (arrow with sharp head).

Figure 4. VirusFISH results for Ostreococcus cells abundance and infection by Ostreococcus viruses in 2012. A. Surface, B. 50m depth, in coastal (E1), mid-shelf (E2) and shelf (E3) waters. Note: April data was not available for surface samples.

Figure 5. Relative abundances of Ostreococcus spp. (OS) transcripts (upper plot) and Ostreococcus viruses (OV) transcripts (lower plot) detected in metatranscriptomes from the surface waters at the mid-shelf station (E2). Note the difference in y-axis between the graphs. Nov.: November.
Table 1. Impact of viruses on *Ostreococcus* populations in the coastal (E1), mid-shelf (E2) and shelf (E3) stations. See Figure 1 for the location of the stations. ND: Non-detected; ‘─’: No data available. Color intensity indicates increasing percentage of infected cells.

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