Impaired viral infection and reduced mortality of diatoms in iron-limited oceanic regions

Chana F. Kranzler[®]¹[⊠], Mark A. Brzezinski², Natalie R. Cohen[®]^{3,4}, Robert H. Lampe[®]^{3,5}, Michael Maniscalco[®]², Claire P. Till[®]^{6,7}, James Mack¹, Jason R. Latham¹, Kenneth W. Bruland⁶, Benjamin S. Twining[®]⁸, Adrian Marchetti³ and Kimberlee Thamatrakoln[®]^{1⊠}

Diatom primary productivity is tightly coupled with carbon export through the ballasted nature of the silica-based cell wall, linking the oceanic silicon and carbon cycles. However, despite low productivity, iron (Fe)-limited regimes are considered 'hot spots' of diatom silica burial with enhanced carbon export efficiency, raising questions about the mechanisms driving the biogeochemistry of these regions. Marine viruses are classically recognized as catalysts of remineralization through host lysis, short-circuiting the trophic transfer of carbon and facilitating the retention of dissolved organic matter and associated elements in the surface ocean. Here we used metatranscriptomic analysis of diatoms and associated viruses, along with a suite of physiological and geochemical metrics, to study the interaction between diatoms and viruses in Fe-limited regimes of the northeast Pacific. We found low cell-associated diatom virus diversity and abundance in a chronically Fe-limited region of the subarctic northeast Pacific. In a coastal upwelling region of the California Current, transient iron limitation also substantially reduced viral replication. These observations were recapitulated in Fe-limited cultures of the bloom-forming, centric diatom, *Chaetoceros tenuissimus*, which exhibited delayed virus-mediated mortality in addition to reduced viral replication. We suggest Fe-limited diatoms escape viral lysis and subsequent remineralization in the surface ocean, providing an additional mechanism contributing to enhanced carbon export efficiency and silica burial in Fe-limited oceanic regimes.

iatoms contribute up to 40% of marine primary production¹ and have silicified cell walls that ballast substantial vertical flux of particulate organic matter out of the surface into the deep ocean². In high-nutrient, low-chlorophyll regions of the open ocean and along some coastal upwelling margins, iron (Fe) availability limits diatom production³⁻⁵ and may contribute to the decoupling of primary productivity and export efficiency in these waters⁶⁻⁸. Fe-limited waters are biogeochemically distinct, characterized by enhanced drawdown of silicic acid relative to nitrate by Fe-limited diatoms, which leads to high cellular silicon (Si):nitrogen (N) and Si:carbon (C)^{9,10}. These regions can also exhibit elevated diatom carbon export efficiency6-8 and markedly high burial of biogenic silica (bSiO₂) relative to organic carbon in underlying sediments¹¹. These 'hot spots' of silica burial contribute to widespread limitation of diatom silica production rates in the surface ocean, linking the Fe, Si and C cycles, and may impose a global constraint on the drawdown of atmospheric CO_2 and export production¹²⁻¹⁴. Together, these observations underscore the importance of elucidating the underlying chemical, physical and biological interactions that dictate the fate of diatom organic matter and associated elements in the ocean.

Viral infection of microbes is a major conduit for elemental cycling in the ocean, transforming the biogeochemical flux of nutrients and carbon, and toggling the balance between export and remineralization processes^{15,16}. By fuelling the microbial loop with dissolved organic matter released during host lysis, viruses circumvent the transfer of particulate organic matter to depth or higher trophic levels through the 'viral shunt'¹⁷. We recently reported accelerated virus-mediated mortality in Si-limited diatoms, demonstrating a role for nutrient availability in facilitating the viral shunt¹⁸. However viral infection can also 'shuttle' organic matter out of the surface ocean by stimulating processes that facilitate export such as particle aggregation^{19,20} and spore formation²¹, highlighting the dynamic, biogeochemical consequences of host–virus interactions. Here we explored diatom host–virus dynamics in Fe-limited regimes of the northeast Pacific Ocean.

Biogeochemically distinct phytoplankton assemblages

Our study encompassed five biogeochemically distinct sites in the iron mosaic of the California upwelling zone⁹ (CUZ) and along a natural Fe gradient (Line P) in the subarctic northeast Pacific Ocean (Fig. 1a). Iron supply to the CUZ (sites S1, S2 and S4), delivered via riverine inputs and sediment resuspension of the continental shelf, is constrained primarily by shelf width along this boundary, generating a mosaic of steep along-shore gradients in bioavailable Fe²². In contrast, Fe is largely supplied along Line P (sites P4 and P26 or Ocean Station Papa²³) through atmospheric deposition and horizontal advection with bioavailable Fe decreasing from near to offshore, resulting in a classical Fe-limited, high-nutrient, low-chlorophyll region (Fig. 1a). Among these sites, $bSiO_2$ concentrations ranged from 0.27 to 12.9 μ M and chlorophyll *a* (chl *a*) concentrations (>5 μ m) ranged from 0.08 to 12.3 μ gl⁻¹, with the

¹Department of Marine and Coastal Sciences, Rutgers University, New Brunswick, NJ, USA. ²Marine Science Institute and the Department of Ecology Evolution and Marine Biology, University of California, Santa Barbara, CA, USA. ³Department of Marine Sciences, University of North Carolina, Chapel Hill, NC, USA. ⁴Marine Chemistry & Geochemistry, Woods Hole Oceanographic Institution, Woods Hole, MA, USA. ⁵Integrative Oceanography Division, Scripps Institution of Oceanography, University of California, San Diego, La Jolla, CA, USA. ⁶Department of Ocean Sciences, University of California, Santa Cruz, CA, USA. ⁷Chemistry Department, Humboldt State University, Arcata, CA, USA. ⁸Bigelow Laboratory for Ocean Sciences, East Boothbay, Maine, ME, USA. ^{Sa}e-mail: chana.kranzler@marine.rutgers.edu; thamat@marine.rutgers.edu

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Fig. 1 | **Biogeochemical**, **physiological and molecular characteristics of initial phytoplankton communities in the northeast Pacific. a**, Mean surface nitrate concentration and sites (circles); symbol size depicts biogenic silica concentration. **b**,**c**, Relative transcript abundance of the major eukaryotic phytoplankton taxa (**b**) and diatom genera (**c**). **d**, Dissolved Fe concentrations. **e**, Si(OH)₄:NO₃ (mean \pm standard error; diamonds represent individual measurements, n=3). **f**, NO₃:Fe; dashed line denotes potential Fe limitation (12 µmol:nmol)²⁷. **g**, Si stress (V_{amb} : V_{enh}); ~1 indicates no stress. n.d. indicates no data. **h**, Photosynthetic efficiency (F_v/F_m). **i**, Ps-n ILL²⁴; values >0.5 and <-0.5 (dashed lines) indicate Fe-limited and Fe-replete growth, respectively.

lowest biomass by both metrics at S4 and P26, and the highest at S1 (Fig. 1a and Extended Data Table 1). Taxonomic distribution of mapped reads using metatranscriptomic assemblies confirmed diatoms were transcriptionally active at all sites, accounting for 4–57% of the total reads, with the highest proportion at S1 and the lowest at P26 (Fig. 1b and Supplementary Data 1). Within diatoms, *Chaetoceros, Pseudo-nitzschia* and *Thalassiosira* genera accounted for 25–78% of the transcripts at all sites, with notable representation of *Proboscia* at S2 (46%) and *Fragilariopsis* at P26 (19%; Fig. 1c and Supplementary Data 1).

A suite of biogeochemical, physiological and molecular diagnostics were used to determine the initial nutrient regime at each site^{24–26} (Fig. 1d–i and Extended Data Table 1). In the CUZ, the site with the highest biomass, S1, was characterized by \geq 13.7 µM nitrate and silicic acid (Extended Data Table 1), the highest dissolved Fe (3.39 nM; Fig. 1d), Si(OH)₄:NO₃ > 1 (Fig. 1e) and a NO₃:Fe ratio below the threshold for anticipated Fe limitation²⁷ (Fig. 1f). All metrics suggest a nutrient-replete regime at S1. Consistent with these observations, kinetic limitation of Si uptake (that is, Si stress; $V_{ambient}$: V_{enh} ; Fig. 1g), photosynthetic efficiency (variable fluorescence [F_v]:maximum fluorescence [F_m]; F_v/F_m ; Fig. 1h) and the iron limitation index for *Pseudo-nitzschia*²⁴ (*Ps-n* ILI; Fig. 1i) did not show any physiological or molecular signatures of nutrient limitation at S1. Despite its geographic proximity to S1, S2 had substantially lower biomass and dissolved Fe (0.42 nM) and was depleted in silicic acid relative to nitrate (Fig. 1a,d–e; and Extended Data Table 1). Furthermore, the diatom assemblage exhibited a Si stress value of 0.51, indicating substantial substrate limitation of Si uptake (Fig. 1g). Although F_{ν}/F_m and Ps-n ILI at S2 were not consistent with Fe limitation at the time of sampling (Fig. 1h–i), elevated NO₃:Fe suggested the potential for impending Fe limitation with further nutrient consumption (Fig. 1f). Site S4 was characterized by high nitrate and silicic acid ($\geq 16.9 \,\mu$ M), 0.31 nM dissolved Fe and the lowest initial biomass (Fig. 1a,d and Extended Data Table 1). Similar to S2, F_{ν}/F_m and Ps-n ILI did not indicate Fe limitation at S4 (Fig. 1h,i), but elevated NO₃:Fe suggested the potential for iron limitation (Fig. 1f). No signs of Si limitation were apparent at S4 with Si(OH)₄:NO₃ \approx 1 and a Si stress value of 1.25 (Fig. 1e,g).

Along Line P, biomass (bSiO₂ and chl *a*) was higher nearshore at site P4, with silicic acid and nitrate $\leq 2.2 \,\mu$ M, 0.64 nM dissolved Fe (Fig. 1a,d and Extended Data Table 1) and depletion of nitrate relative to silicic acid (Si(OH)₄:NO₃>1; Fig. 1e), suggesting nitrate limitation at the time of sampling. The NO₃:Fe ratio, F_v/F_m and *Ps-n* ILI at P4 did not indicate iron limitation (Fig. 1f,h–i). In contrast, the assemblage at P26 displayed strong signs of Fe limitation including low biomass, silicic acid and nitrate both $\geq 11.7 \,\mu$ M, and 0.05 nM dissolved Fe (Fig. 1a,d and Extended Data Table 1). These observations were further supported by a greatly elevated NO₃:Fe ratio, impaired photosynthetic efficiency and a *Ps-n* ILI above the threshold for Fe-limited growth²⁴ (Fig. 1f,h–i), consistent with its classification as a chronically Fe-limited regime²⁸.

At each site, shipboard incubations were performed with added iron (+Fe) or desferrioxamine B (DFB; Extended Data Fig. 1), a strong Fe-binding ligand used to rapidly induce Fe limitation, alongside an unamended control (Ctrl). Together, these treatments enabled the characterization of the physiological and molecular response to fluctuating iron supply as previously reported²⁴⁻²⁶, and provided further support for the initial underlying nutrient regime at each site. At S1, there was no measurable response in biomass or $F_{\rm v}/F_{\rm m}$ to +Fe^{25,26} (Extended Data Fig. 1), further establishing its status as nutrient replete. The assemblage at S2 also did not respond to +Fe (Extended Data Fig. 1), but Si stress increased in severity by the first timepoint (t_{48}) , even after an initial spike of Si to this incubation (see Methods), dropping from 0.51 to 0.16 ± 0.03 (Extended Data Fig. 2) and further confirming Si limitation across all treatments. At S4, the assemblage displayed a positive growth response to +Fe relative to the Ctrl (Extended Data Fig. 1), suggesting the onset of iron limitation in this recently upwelled community^{25,26,29}. Along Line P, the P4 assemblage exhibited no growth response to +Fe; in contrast, both chl *a* concentration and F_v/F_m sharply increased at P26 in response to +Fe, supporting severe Fe limitation at this site. Together, these sites encompassed diverse nutrient regimes, ranging from nutrient replete (S1) to limited for silicon (S2), nitrate (P4) and iron (P26; Extended Data Table 1).

Divergent signatures of infection across nutrient regimes

To explore the prevalence of viral infection across these ecophysiologically distinct diatom communities, we used metatranscriptomic analysis of size-fractionated biomass to simultaneously detect both host (diatom) and cell-associated viral transcripts, the latter used as a proxy for viral infection. Diatom virus contigs were identified using a homology-based query with the taxonomic gene markers replicase (Rep) and RNA-dependent RNA polymerase (RdRp) from known single-stranded (ss) DNA- and ssRNA-containing diatom viruses, respectively. No sequences with homology to ssDNA diatom viruses were recovered. In contrast, we identified 144 unique cell-associated ssRNA diatom virus-like contigs (Supplementary Data 3; Expect (E) value < 10⁻³). Phylogenetic placement within *Picornavirales* indicated that 79 of these contigs grouped among known diatom viruses (Extended Data Fig. 3) and were selected for further analysis. Selected contigs had a median length of 392 bp, but ranged from 124 to 8,999 bp, indicating the recovery of several near-complete viral genomes (~9kb in length; Extended Data Fig. 3). Viral contig reads were normalized to total diatom transcripts to enable comparison across samples with different diatom biomass¹⁸ and assess the contribution of individual and total virus contigs in the initial sample (t_0) and throughout each incubation. Although a subset of diatom virus contigs were shared across samples, hierarchical clustering of contig abundance revealed unique distribution patterns at each site (Fig. 2a). Diatom viruses were strikingly sparse in the Fe-limited P26 assemblage, with only three virus contigs identified, low viral diversity and low total cell-associated diatom virus abundance (Fig. 2). S1 and S2 had the highest number of distinct virus contigs (44 and 39 of the 79, respectively; Fig. 2a), with four dominant, highly abundant contigs at S2. Among all sites, S2 had ~1,000-fold higher total cell-associated diatom virus abundance (Fig. 2c), and was the only site in which a majority (56–75%) of viral contigs present at t_0 increased in abundance across all treatments (Extended Data Fig. 4). Furthermore, 13 additional virus contigs appeared at t_{48} , while only three virus contigs disappeared from at least one treatment (Extended Data Fig. 5). These data illustrate an actively progressing viral infection in the severely Si-stressed assemblage at S2, notably consistent with our previous findings that Si limitation facilitates virus infection in diatoms¹⁸.

The low diatom virus diversity and abundance at P26, taken together with diagnostic signatures of severe Fe limitation (Fig. 1 and Extended Data Fig. 1), raised the possibility of a relationship between Fe availability and viral infection in diatoms. The actively infected diatom community at S2 allowed us to test the hypothesis that Fe limitation impairs virus replication in the manipulative shipboard incubations. Differential expression analysis of diatom transcripts at t_{48} identified 105, 471 and 1,407 contigs in the DFB/Ctrl, Ctrl/+Fe and DFB/+Fe comparisons, respectively (Fig. 3 and Supplementary Data 4). No striking differences in iron homeostasis-related transcripts were observed between Ctrl and DFB treatments (Fig. 3a). However, when compared with +Fe, both Ctrl and DFB treatments exhibited clear overrepresentation of established Fe-limitation-responsive genes, including Fe-starvation-induced proteins (ISIPs) and plastocyanin (petE), as well as a suite of antioxidant genes such as peroxiredoxin (prx) and thioredoxin-dependant peroxidase (tpx; Fig. 3b,c). In addition, haem oxygenase (HMOX), widely recognized for its cytoprotective response to oxidative stress³⁰, with a potential role in repartitioning cellular Fe³¹, was among the most highly upregulated transcripts (798-fold, false discovery rate (FDR) = 1.00×10^{-9} for DFB/+Fe; 891-fold, FDR = 7.43×10^{-10} for Ctrl/+Fe), consistent with lab studies of Fe-limited diatoms^{32,33}. In the +Fe treatment, contigs involved in Fe-dependent photosynthetic electron transport and nitrogen assimilation were overrepresented, in line with previous studies^{25,34}, suggesting that the addition of Fe prevented the onset of Fe limitation (Fig. 3b,c). These transcriptional patterns demonstrate unequivocal Fe limitation in both Ctrl and DFB treatments at t_{48} of the S2 incubation, concurrent with Si limitation (Extended Data Fig. 2).

Interestingly, the molecular delineation between Fe-replete (+Fe) and Fe-limited (Ctrl and DFB) conditions at S2 could also be extended to cell-associated diatom virus contig abundance. Overall, the aforementioned increase in diatom virus contig abundance from t_0 to t_{48} was more pronounced in the +Fe treatment (median fold change of ~3 and ~2 in Fe-replete and Fe-limited treatments, respectively; Extended Data Fig. 5). In pairwise comparisons at t_{48} , 55% of viral contigs did not differ in abundance between Fe-limited treatments, compared with 30% of contigs between +Fe and Ctrl or DFB treatments (Fig. 3d-f). Of the 70% of viral contigs that exhibited a ≥2-fold change in the +Fe treatment, 86% and 78% increased in the +Fe/Ctrl and +Fe/DFB comparisons, respectively (Fig. 3e,f),



Fig. 2 | Metatranscriptomic analysis of cell-associated diatom viruses. a, Heatmap of diatom virus contig abundance across sites and incubations. Dendrograms reflect hierarchical clustering (rows, contigs; columns, samples coloured by treatment). White denotes contigs below detection. **b**,**c**, Shannon diversity index (*H*; **b**) and sum total of diatom virus contigs (**c**) at each site. Boxes depict median, upper and lower quartiles; whiskers denote values $1.5 \times$ the interquartile range. Symbols represents individual samples from Ctrl (yellow), DFB (blue) and +Fe (orange) treatments. Shape denotes timepoints (diamonds, t_0 ; circles, t_1 = 24-48 h; triangles, t_2 = 48-96 h). Letters denote statistically significant groups (ANOVA and Tukey's HSD post-hoc test; Supplementary Data 2).

suggesting Fe addition facilitated virus replication. Taken together, these data support the hypothesis that the onset of Fe limitation impairs viral replication in an actively infected diatom assemblage.

Impaired viral infection in an Fe-limited model diatom

To directly test whether Fe limitation and its associated subcellular response disrupts viral replication, we grew and infected laboratory cultures of the cosmopolitan, bloom-forming diatom, *Chaetoceros tenuissimus*, under Fe-limiting conditions. Fe limitation was diagnosed by a decrease in specific growth rates, chlorophyll autofluorescence and electron transport rates, and further confirmed by +Fe 'rescue' experiments (Extended Data Fig. 6). Fe-limited cells were also characterized by a 51.4% increase in cellular bSiO_2 (Extended Data Fig. 6), consistent with previous work documenting increased silica production under Fe limitation^{35,36}. In addition, Fe limitation also resulted in a 51.1% increase in bulk cellular antioxidant capacity (Extended Data Fig. 6), consistent with previous work demonstrating an enhanced ability of Fe-limited diatoms to combat oxidative stress^{32,33,37}.

When replete cultures were infected with the ssDNA virus, CtenDNAV, diatom host abundance decreased dramatically 5 days post infection (dpi) along with a marked increase in the proportion of dead cells (Fig. 4a). In contrast, when Fe-limited cultures were infected, the decline in host abundance was much more gradual and only apparent 8 dpi (Fig. 4a). A ~600-fold increase in extracellular virions was apparent in replete cultures 2 dpi ('pre-lysis'), compared with only a ~10-fold increase in Fe-limited cultures, indicating that early viral production was significantly impaired by Fe limitation in C. tenuissimus (Fig. 4b). In addition, the final viral titre ('post-lysis'), collected after host lysis, and burst size (the number of viruses produced per host cell) was significantly attenuated in Fe-limited cells compared with replete cells (Fig. 4b,c). Delayed host mortality was also observed in Fe-limited C. tenuissimus infected with a ssRNA virus, CtenRNAV (Extended Data Fig. 7), demonstrating a conserved response to these genetically distinct viruses. These data demonstrate impaired viral replication, reduced extracellular virus production and delayed host mortality under Fe limitation, and are consistent with previous work demonstrating decreased viral production in Fe-limited cultures of the green alga, Micromonas, and the prymnesiophyte, Phaeocystis³⁸.

Our observations of upregulated antioxidant genes in Fe-limited diatom communities (Fig. 3) and elevated cellular antioxidant

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Fig. 3 | **Diatom molecular response and cell-associated diatom virus abundance during Fe limitation at S2. a-c**, Differential expression of diatom contigs as a function of mean contig abundance between DFB/Ctrl (**a**), Ctrl/+Fe (**b**) and DFB/+Fe (**c**). Filled symbols are statistically significant (FDR < 0.05). Colours denote genes involved in Fe limitation (blue), antioxidant activity (purple), photosynthesis (red) and nitrogen assimilation (orange/yellow). See Methods for abbreviations. **d-f**, Pairwise comparisons of cell-associated diatom virus contig abundance (mean \pm standard error, n = 3) between DFB/Ctrl (**d**) +Fe/Ctrl (**e**) and +Fe/DFB (**f**) at t_{48} . Insets show highly abundant contigs. Colour denotes \geq 2-fold increase (red) or decrease (blue). Unity slopes (dashed lines) indicate no difference.



Fig. 4 | Diatom host-virus dynamics in Fe-limited *C. tenuissimus.* **a**, Host abundance and percent dead cells (symbol size) in replete (orange) and Fe-limited (blue), uninfected (open) and infected (filled) cultures with CtenDNAV (biological triplicates with LOESS regression). **b**, Virus abundance in replete (orange) and Fe-limited (blue) cultures at t_0 (initial), 2 dpi (pre-lysis) and post-lysis (6–8 days for replete; 8–12 days for Fe-limited). ****P* < 0.001, two-way ANOVA, Tukey's HSD post-hoc test. **c**, Viral burst size in infected replete and Fe-limited cultures. Boxes depict median (line), upper and lower quartiles; whiskers denote values 1.5× the interquartile range. ****P* < 0.001, unpaired two-tailed *t*-test.

capacity in Fe-limited *C. tenuissimus* (Extended Data Fig. 6) suggest the mechanism underlying this response may lie at the intersection of Fe homeostasis and reactive oxygen species (ROS) metabolism. Both Fe-dependent^{38–40} and ROS-dependant^{41–43} impacts on infection dynamics have emerged as focal points in diverse host–virus systems, ranging from bacteria to humans. While the production of ROS is well-established as an early host defence response to pathogen

invasion⁴¹, recent work suggests that ROS production can also be co-opted by viruses to support viral replication and trigger cell death^{42,43}. Intriguingly, *HMOX*, a significantly upregulated transcript in the Fe-limited treatments at S2 (Fig. 3), is also associated with substantial antiviral activity in diverse viral lineages through broad cytoprotective mechanisms and the inhibition of programmed cell death pathways⁴⁴. In picornaviruses specifically, *HMOX* reduces viral replication through the attenuation of virus-induced ROS production⁴⁴. In nutrient-replete *C. tenuissimus*, we observed a two- to threefold increase in ROS production during viral infection, concomitant with a decrease in host abundance (Extended Data Fig. 8). We also found that priming the host antioxidant system with a sub-lethal dose of hydrogen peroxide (H_2O_2), a well-described ROS, resulted in delayed infection dynamics (Extended Data Fig. 8), similar to those observed during Fe limitation. We posit that efficient ROS scavenging due to elevated cellular antioxidant capacity contributes to reduced viral production and delayed mortality in Fe-limited diatoms.

Biogeochemical implications

In the vast Fe-limited regions of the global ocean, diatom silica production^{9,10}, species composition⁴⁵, grazing⁴⁶ and reduced microbial mediated remineralization⁶ have been invoked as mechanistic explanations for enhanced silica burial¹¹ and efficient diatom carbon export^{6,8}. Viruses, the most abundant biological entity in the ocean, are regarded as widespread mediators of remineralization and nutrient recycling. However, the discovery that environmental conditions such as temperature, salinity⁴⁷ or light⁴⁸ influence host-virus dynamics and the emerging concept of a 'viral shuttle' are transforming our view of viruses within a global biogeochemical framework⁴⁹. Reduced viral replication and delayed mortality in Fe-limited diatoms, together with our previous findings that silicon limitation facilitates infection and accelerates mortality¹⁸, highlight the fundamental imprint that nutrient availability has on host-virus interactions. Notably, our observations encompass both open ocean and coastal assemblages, experiencing chronic or transient Fe limitation, and allude to a shared cellular response that impedes infection even in regions with co-occurring Si limitation⁵⁰, establishing an 'ecophysiological hierarchy' with the impact of Fe superseding that of Si.

We suggest these dynamic, nutrient-driven host-virus interactions shape the fate of diatom carbon, silica and associated elements. Whereas diatoms infected in Si-limited regimes would enter the 'viral shunt', diatoms in Fe- or Fe/Si co-limited regimes would be retained in the particulate fraction and poised to enter the 'viral shuttle'. Impaired infection in Fe-limited diatoms, concomitant with increased mineral ballast, would further promote export, curbing the supply of carbon, nutrients and trace metals that would otherwise be released through lysis in the surface ocean.

Online content

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Methods

Cruise sampling and measurements. Incubation experiments were conducted on two separate cruises-one within the CUZ, 3-26 July 2014, onboard the RV Melville (MV1405) and another along Line P in the subarctic northeast Pacific Ocean, 7-23 June 2015, onboard the Canadian Coast Guard Ship John P. Tully (Line P cruise 2015-09). Details on incubation setup are reported elsewhere^{25,26}. Briefly, surface water collected using trace-metal-clean techniques was distributed into acid-cleaned 10 l cubitainers. Treatments, in triplicate, included an unamended control (Ctrl), a 200 nM addition of DFB to bind and reduce bioavailable iron, and the addition of 5 nM FeCl₃ (+Fe). At site S2, 15 µM of Si(OH)4 was added to all treatments to support growth of diatoms due to the initially low Si(OH)₄ concentration (<4.7 µM). At site P4, 10 µM of NO₃⁻ was added to all treatments to support growth due to low initial NO3⁻ concentrations (<1.5 µM). Cubitainers were incubated on deck with flowing surface seawater for temperature control and screened to approximately 30% surface irradiance and sampled at dawn. Each site was sampled at the start of the incubation (initial, t_0) and at two timepoints (t_1 and t_2) throughout each experiment (24 and 48 h at S1; 48 and 72 h at S2; 48 h and 72 h at S4; 24 h and 48 h at P4; 48 and 96 h at P26). Methods for analysing chl a, bSiO₂, photosynthetic efficiency, and dissolved nutrients and iron are described elsewhere²⁵. Mean surface nitrate concentrations shown in Fig. 1a were obtained from the World Ocean Database⁵¹.

Silica production rates were measured using the radioisotope ³²Si as described in ref. ⁵²; 320 ml samples received 260 Bq ³²Si(OH)₄ (specific activity >40 kBq (µg Si)⁻¹) followed by incubation in deckboard flow-through incubators screened with neutral density filters to 55% of surface irradiance. After 4–7h, samples were filtered onto 1.2 µm polycarbonate filters (Millipore) and quantified as described in ref. ⁵³. Gross production rates (ρ in µM Si d⁻¹) were normalized to bSiO₂ concentration to determine specific rates⁵⁴ (V_b d⁻¹). Si stress was assessed using paired samples that measured silica production rates at the ambient silicic acid concentration (V_{amb}) and after an 18 µM addition of non-radioactive sodium silicate (V_{enb}). Under the assumption that the 18 µM Si(OH)₄ addition saturates Si(OH)₄ uptake (that is, approximates the maximum uptake rate, V_{max}), the ratio $V_{amb}V_{enh}$ represents the fraction of maximum uptake rate supported by the ambient [Si(OH)₄]¹⁵, with values of ~1 indicative of Si-replete populations and values approaching zero indicative of severe Si stress. Values between 0.92 and 1.0 are considered within the error of detection⁵⁶.

Metatranscriptomic analysis. Biomass for metatranscriptomic analysis was collected by filtration onto 0.8 µm Pall Supor filters (142 mm) via peristaltic pumping, immediately flash frozen in liquid N_2 and stored at $-80\,^{\rm o}\mathrm{C}$ until extraction onshore. Biomass samples were collected at the start of the incubation (initial, t₀) and at 24h (S1), 48h (S2), 48h and 72h (S4), 24h and 48h (P4), and 96 h (P26). The initial RNA extraction, sequencing, assemblies and annotations are described elsewhere25,26. Briefly, RNA was extracted using the ToTALLY RNA Total RNA Isolation Kit (Ambion Thermo Fisher Scientific). Metatranscriptomic library preparation was performed with the Illumina TruSeq Stranded mRNA Library Preparation Kit and HiSeq v4 reagents with poly-A selection of mRNA. Samples were barcoded and sequenced on an Illumina HiSeq 2000. Biological triplicates were sequenced for each treatment, except for samples with low yields (for example, t₀ and Line P samples), in which triplicates were pooled into one sample. Merged pairs and non-overlapping paired-end reads were used to assemble contigs using ABySS v1.5.2 with varied k-mer sizes⁵⁷. Subassemblies from each site were merged with Trans-ABySS v1.5.358 and redundant contig removal verified with GenomeTools v1.5.159. Only contigs with 100% identity were merged. In cases where a shorter contig was contained within a longer contig and 100% identical, the longer contig was retained. Contigs shorter than 120 bp were discarded. Functional annotation of eukaryotic transcripts was assigned by best homology (lowest E value) with BLASTX v2.2.31 and the Kyoto Encyclopedia of Genes and Genomes (release 75)60. Taxonomic classification and additional functional annotation of hypothetical proteins were assigned using MarineRefII, a custom reference database maintained by the Moran Lab at the University of Georgia that includes transcriptomes from the Marine Microbial Eukaryote Transcriptome Sequencing Project⁶¹. Read counts were estimated from the merged assembly containing contigs from all sites and incubations using the quasi-mapping method implemented in Salmon v0.7362. No pre-filtering of reads was performed-contigs with normalized counts >0 were considered to be present. In our analysis of relative transcript abundance, major eukaryotic phytoplankton taxa and diatom genera that were <5% of the total mapped reads were classified as 'other'.

Downstream metatranscriptomic analyses were performed in the open source program R version 3.5.1 Feather Spray⁶³. *Ps-n* ILI values were calculated as described in ref. ²⁴. For sites S1 and P26, the index was calculated from unamended deckboard incubations, due to low initial abundance of *Pseudo-nitzschia* transcripts. For analysis of diatom transcript abundance, EdgeR v3.12.0⁶⁴ was used to calculate the mean counts per million (CPM) and fold change of diatom transcripts in pairwise comparisons between treatments, using the exactTest with edgeR's estimate of tagwise dispersion across all samples. *P* values were adjusted for multiple testing with the FDR-controlling procedure⁶⁵ and FDR=0.05 was used as the cutoff for differentially expressed genes. Log fold change (M) versus mean abundance (A) plots were produced using the R package ggplot2⁶⁶. Abbreviations

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for analysed contigs are as follows: iron starvation induced protein 1 (*ISIP1*), phytotransferrin (*pTF*, formerly *ISIP2a*), iron starvation induced protein 2B (*ISIP2b*), iron starvation induced protein 3 (*ISIP3*), flavodoxin (*fldA*), plastocyanin (*petE*), ascorbate peroxidase (*APX*), cytochrome c peroxidase (*CcP*), glutaredoxin (*grx*), haloperoxidase (*HPO*), haem oxygenase (*HMOX*), peroxiredoxin (*prx*), thioredoxin-dependent peroxiredoxin (*tpx*), thioredoxin (*trx*), ferredoxin (*petF*), ferredoxin-ADP + reductase (*petH*), cytochrome c_6 (*petJ*), photosystem II oxygen-evolving enhancer (*psbO*), cytochrome b_6f (*petC*), nitrate transporter (*NRT*), nitrate reductase (*NIR*), nitrite reductase (*NIR*), glutamate synthase (*gltS*), urea transporter (*URT*) and ammonium transporter (*AMT*).

Diatom virus contigs were identified using a homology-based query (TBLASTN, E value < 10⁻³) with RdRp and Rep genes from known diatom-infecting ssRNA (family Marnaviridae) and ssDNA viruses (family Bacilladnaviridae), respectively. Given the poly-A tailed genomes of ssRNA diatom viruses, poly-A selection during metatranscriptomic library preparation would capture ssRNA viral genomes as well as actively replicating viruses. Diatom virus RdRp read counts were normalized to the length of each contig and sequencing depth yielding viral contigs per million (C_v) , analogous to transcript per million normalization67 used for comparing relative gene expression between samples. To assess the degree of infection across samples with different diatom biomass loads, C, was normalized to total diatom transcript per million¹⁸, to yield normalized cell-associated diatom virus contig abundance (nC_v) , similar to group normalization methods used in previous studies34,68,69. The use of total diatom transcript abundance as a proxy for diatom biomass was previously validated as an indicator of diatom biomass using microscope counts and bulk measurements²⁶. Heatmaps and dendrograms using Euclidean distance and hierarchical clustering were produced with the R package gplots v3.0.1.270. Shannon diversity indices for diatom virus diversity were calculated using the R package vegan⁷¹

For phylogenetic analysis of viral contigs, reference *Picornavirales* RdRp amino acid sequences were obtained from the National Center for Biotechnology Information, trimmed and aligned using ClustalW⁷² in Mega 7⁷³. A maximum likelihood tree was constructed from the reference alignment using RAxML v8.2.10 with the PROTGAMMAWAGF substitution model and 100 bootstrap replicates⁷⁴. Putative diatom virus contigs (*E* value < 10^{-3}) were aligned to the reference alignment using hmmalign and the hidden Markov model (HMM) profile of the reference alignment (built using hmmbuild in HMMER; hmmer.org). Tree files were packaged using taxtastic v0.8.3 and environmental contigs were placed on the reference tree using pplacer v1.1.alpha19⁷⁵. The tree was exported and the posterior probability was calculated using guppy v1.1⁷⁵.

Culturing conditions and infection experiments. *Chaetoceros tenuissimus* Meunier, isolated in the coastal waters of Japan⁷⁶, and its associated viruses were kindly provided by Y. Tomaru (National Research Institute of Fisheries and Environment of Inland Sea, Japan). Host cultures were maintained in modified SWM-3 media⁷⁷ with 0.2 mM silicate, 2 mM nitrate and 2 nM Na₂SeO₃⁷⁶, at 15 °C on a 12:12 light:dark cycle at ~120 µmol photons m⁻² s⁻¹. Culture work was performed in sterile polystyrene filter-capped culture flasks or in acid-washed polycarbonate culture flasks to minimize iron contamination. Cell abundance was measured by flow cytometry (BD Accuri C6; 488 nm excitation, >670 nm emission) using chlorophyll fluorescence and forward scatter.

For Fe-limitation experiments, exponentially growing cultures in replete media were centrifuged at $4,000 \times g$ for 10 min and washed and resuspended in either SWM-3 without added Fe or in an artificial seawater media^(36,7) (ESAW) without added iron and buffered with $100 \,\mu$ M EDTA⁸⁰. Once an Fe-limited phenotype was confirmed (~3–5 days), cultures were subdivided and one set was infected with CtenDNAV⁸¹ or CtenRNAV⁷⁶ at a multiplicity of infection of 10. Data shown are representative of five independent experiments (n=15). For experiments with hydrogen peroxide (H₂O₂), a sub-lethal concentration was determined using a dose curve of 0–150 μ M H₂O₂ (Sigma; 30% v/v) on exponentially growing cultures (-4.5 × 10⁵ cells ml⁻¹). Cell abundance and photosynthetic efficiency were monitored and 100 μ M H₂O₂ was selected for downstream infection experiments s24h after exposure to 100 μ M H₂O₂. All statistical analyses of data collected from laboratory cultures were performed in R version 3.5.1 Feather Spray⁶³.

Cellular biogenic silica and particulate organic carbon. Dissolved silicon and $bSiO_2$ in laboratory cultures were measured using standard protocols^{82,83}. Samples for particulate organic carbon measurements were collected on pre-combusted GF/F filters, wrapped in pre-combusted foil and stored at $-20^{\circ}C$ until processing. Filters were thawed and dried at 60°C and then placed in a glass desiccator with concentrated hydrochloric acid for 24h to dissolve particulate inorganic carbon. Samples were analysed on a CNS elemental analyser (Carlo Erba NA 1500).

Biophysical measurements. Maximum photochemical quantum yield of photosystem II (photosynthetic efficiency; F_v/F_m) was measured using a custom-built fluorescence induction and relaxation system⁴⁴, providing measurements of F_o (minimum) and F_m (maximum) fluorescence yields. Maximum efficiency of photosystem II was calculated as the ratio between variable fluorescence (F_v) and F_m , where $F_v/F_m = (F_m - F_o)/F_m$. Prior to measurements,

cultures were acclimated in low light for 15 min. Maximal rates of photosynthetic electron transport (P_{max}) were retrieved by fitting photosynthesis versus irradiance curves with the exponential rise function, $P = P_{max}(1 - \exp(E/E_k))$, where *E* is the irradiance and E_k is the light saturation parameter.

Cellular antioxidant capacity. Fe-limited and replete cultures were collected by filtration on 1.2 µm polycarbonate membrane filters and resuspended in buffer (5 mM potassium phosphate, 0.9% sodium chloride, pH = 7.4). Cells lysates were generated through three flash freeze–sonication cycles (30 sec sonication on ice). Samples were then centrifuged at $10,000 \times g$ at 4°C for 15 min to collect cell debris and supernatants were transferred to fresh tubes and stored at -80 °C until further analysis. The total antioxidant capacity in cell extracts was determined using the antioxidant assay kit (Cayman Chemical), according to the manufacturer's instructions. Absorbance at 750 nm was measured in a SpectraMax M3 microplate reader and the cellular antioxidant capacity is expressed as Trolox equivalents (mM per cell).

Diagnostic staining and flow cytometry. The relative proportion of live/ dead cells in cultures was determined by staining with SYTOX Green (Thermo Fisher Scientific) at a final concentration of 1 μ M for 15 min in the dark at room temperature and subsequent analysis using flow cytometry (BD Accuri C6; 488 nm excitation, 533/30 emission). Unstained samples were used to set a threshold for background green fluorescence. The relative number of dead cells in a population (percent positively stained) was determined as those that exceeded the background fluorescence of the unstained sample. Intracellular ROS were determined by staining with 2',7'-dichlorodihydrofluorescein diacetate (CM–H₂DCFDA; Thermo Fisher Scientific) at a final concentration of 5 μ M for 1 h in the dark at room temperature and subsequent analysis using flow cytometry (BD Accuri C6; 488 nm excitation, 533/30 emission). Intracellular ROS fluorescence was calculated by subtracting the mean background cellular fluorescence of unstained populations (relative fluorescence units per cell) from the mean fluorescence of positively stained populations.

Virus abundance. Virus abundance was measured in lysates generated by filtering infected cultures through a 0.22 µm pore size filter to remove cellular debris. Lysates were stored at 4 °C and processed within one month of collection. The abundance of infectious virus particles (infectious units ml⁻¹) was measured using the most probable number assay^{18,85} (MPN) and calculated using the EPA-MPN calculator⁸⁶. Extracellular virus abundance data were collected from 2–6 independent experiments. Burst size was calculated by dividing the final virus abundance by the decrease in host abundance.

Data availability

All cruise-related data are publicly available at the Biological & Chemical Oceanography Data Management Office (CUZ: project number 559966, https:// www.bco-dmo.org/deployment/559966; Line P: http://www.waterproperties.ca/ linep/2015-009/index.php). Metatranscriptome sequencing data are available in the NCBI sequence read archive (SRA) under the BioProject accession numbers PRJNA320398 and PRJNA388329. Assembled contigs, read counts and functional annotation of contigs are available at http://marchettilab.web.unc.edu/data. All data generated or analysed during this study are included in this published article and its Supplementary Information files. Source data are provided with this paper.

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Author contributions

C.F.K. and K.T. conceived the project, designed the laboratory experiments and wrote the manuscript. C.F.K. performed metatranscriptome and statistical analyses. C.F.K. conducted the laboratory culture-based experiments. J.M. and J.R.L. provided technical support on laboratory experiments. M.M. assisted with RdRp phylogenetic analyses. K.T., M.A.B., N.R.C., M.M., C.P.T., A.M., B.S.T. and K.W.B. participated, collected and analysed samples on the CUZ cruise; B.S.T. and N.R.C. participated, collected and analysed samples on the Line P cruise. A.M., N.R.C. and R.H.L. extracted the RNA and generated the metatranscriptome data. All authors provided comments on the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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Correspondence and requests for materials should be addressed to C.F.K. or K.T.

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Extended Data Fig. 1 | See next page for caption.

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Extended Data Fig. 1 Bulk parameters measured during incubations at each site. a, Size-fractionated chlorophyll a (>5 µm; µg L⁻¹), **b**, biogenic silica (µM), and **c**, maximum photochemical quantum yield of photosystem II (photosynthetic efficiency; F,/F_m) throughout each incubation. Treatments included an initial sample (t_o; open), and unamended Control (Ctrl; yellow), DFB (blue) and +Fe (orange). **d**, Fold-change in chlorophyll a (open bars) and biogenic silica (filled bars) between +Fe and Ctrl treatments at each timepoint. Note that at sites S2 and P4, 15 µM of Si(OH)₄ and 10 µM of NO₃⁻ were added, respectively, to all treatments due to low initial concentrations (Si(OH)₄ at S2 < 4.7 µM; NO₃⁻ at P4 < 1.5 µM). Mean and standard error of triplicate incubations are shown along with individual replicates (diamonds). Statistical significance of the community response to iron addition is depicted (panels **a-c**) for each time point with ****P* < 0.001, ***P* < 0.05 by analysis of variance (ANOVA) followed by Tukey's HSD post hoc test. Additional statistical analysis is available in Supplementary Data 2.



Extended Data Fig. 2 | Silicon stress during incubation experiments at each site. Kinetic limitation of Si uptake, or Si stress (V_{amb} : V_{enh}), at each site for initial samples (t_0 ; open diamonds) and unamended Control (Ctrl; yellow), DFB (blue) and +Fe (orange) treatments throughout each incubation experiment. Each point represents a distinct treatment (color) and time point (symbol) within each incubation (diamonds, t_0 ; circles, t_1 , 24–48 h; triangles, t_2 , 48–96 h). See methods and Supplementary Fig. 1 for individual time points. Values approaching zero are indicative of severe Si stress and values ~1 are indicative of silicon replete populations. The boxes depict the median (horizontal line) and upper and lower quartiles of the data with whiskers encompassing data points within 1.5× of the interquartile range. ***P < 0.001 by Kruskal-Wallis with Dunn's multiple comparison test. No Si stress data was collected during the P4 and P26 incubations, or at t_2 in the S1 and S2 incubations for Ctrl and +Fe treatments.



Extended Data Fig. 3 | Phylogenetic analysis and size distribution of diatom virus-like contigs. a, Maximum likelihood phylogenetic tree of RNA-dependent RNA polymerase (RdRp) amino acid sequences within the order *Picornavirales* and placement of homologous contigs identified in metatranscriptomes. Bootstrap values >50 are shown (100 replicates). Triangles denote sequences that fall within the family *Marnaviridae* with blue triangles denoting the genus *Marnavirus*, which is distinctly comprised of dinoflagellate viruses and red triangles identifying putative diatom viruses that were selected for downstream analysis. **b**, Size distribution (bp) of diatom virus-like contigs. The abbreviations, names and NCBI database accession numbers of the amino acid sequences used to construct the reference alignment are: AglaRNAV, *Asterionellopsis glacialis* RNA virus, BAP16719; CsfrRNAV, *Chaetoceros socialis radians* RNA virus 1, YP_002647032; RsRNAV, *Rhizosolenia setigera* RNA virus 01, YP_006732323; CtenRNAV01, *C. tenuissimus* RNA virus 01, YP_009505620; CtenRNAVII, *C. tenuissimus* RNA virus type-II, BAP99818; CspRNAV2, *Chaetoceros* species RNA virus 02, BAK40203; HaRNAV, *Heterosigma akashiwo* RNA virus, AAP97137; CPSMV, cowpea severe mosaic virus, NP_619518; BPMV, Bean pod mottle virus, NP_612349; PYFV, Parsnip yellow fleck virus, BAA03151; RTSV, Rice tungro spherical virus, NP_042507;SBV, Sacbrood virus, AIZ75645; PV, Human poliovirus 1, CAA24461; AIV, Aichi virus 1, ADN52312; BQCV, Black queen cell virus, NP_620564; TrV, Triatoma virus, NP_620562; CPV, Cricket paralysis virus, NP_647481; DCV, *Drosophila* C virus, NP_044945; TSV, Taura syndrome virus, NP_149057.



Extended Data Fig. 4 | Temporal changes in cell-associated diatom viruses throughout each incubation. Log_2 fold change in diatom virus contig abundance (nC_v) at t_1 (circles, 24–48 h) or t_2 (triangles, when present, 48–72 h) compared to t_0 for Ctrl (yellow), DFB (blue) and +Fe (orange) treatments (n=3), *P < 0.05 by a one-way analysis of variance (ANOVA) followed by a Tukey HSD post hoc test. The boxes depict the median (horizontal line) and upper and lower quartiles of the data with whiskers encompassing data points within 1.5x of the interquartile range. See methods and Supplementary Fig. 1 for individual time points (symbols) in each incubation.



Extended Data Fig. 5 | Temporal changes in cell-associated diatom viruses throughout the S2 incubation. a, Log_2 fold change in diatom virus contig abundance (nC_v) between t_{48} and t_0 for Ctrl (yellow), DFB (blue) and +Fe (orange) treatments (n=3), *P < 0.05 by a one-way analysis of variance (ANOVA) followed by a Tukey HSD post hoc test. The boxes depict the median (horizontal line) and upper and lower quartiles of the data with whiskers encompassing data points within 1.5x of the interquartile range. **b**, Heatmap of mean abundance for each diatom virus contig (nC_v) identified in the initial (t_0) sample and at t_{48} across the treatments for both high (top panel) and low abundance (bottom panel) contigs. Contigs that were below detection are depicted in white. **c**, Abundance (nC_v) of each cell-associated diatom virus contig in Ctrl (left panel), DFB (middle panel) and +Fe (right panel) treatments at t_{48} vs t_0 . Mean \pm standard error is shown (n=3). Dotted lines denote unity-slope lines, indicating no difference between timepoints. Insets depict 'high' abundance contigs. In panels a and c, open and closed symbols denote 'high' and 'low' abundance diatom virus contigs, respectively.



Extended Data Fig. 6 | Diagnosing iron limitation in *Chaetoceros tenuissimus.* **a**, Specific growth rates (day^{-1}) , **b**, mean chlorophyll autofluorescence (RFU cell⁻¹) and **c**, electron transport rates $(e^- s^{-1} PSII^{-1})$ of C. *tenuissimus* cultures in replete (orange symbols) and Fe-limited (blue symbols) growth media. **d**, Cell abundance and **e**, maximum photochemical quantum yield of photosystem II (photosynthetic efficiency; F,/F_m) during Fe-limitation (blue) and after Fe addition (orange) in an Fe 'rescue' experiment. Each independent biological replicate (n=3) is shown by individual symbols with lines of best fit depicting a LOESS regression. **f**, cellular biogenic silica (µmol bSiO₂ cell⁻¹) and **g**, cellular antioxidant capacity (mM Trolox eq cell⁻¹) of replete and iron-limited *C. tenuissimus*. Statistical significance was determined using an unpaired, two-sided t-test. Different symbols denote biologically independent replicates across six independent experiments.



Extended Data Fig. 7 | The impact of Fe limitation on diatom host-virus dynamics in laboratory cultures of Chaetoceros tenuissimus infected with

CtenRNAV. Host abundance in replete (orange circles) and Fe-limited (blue triangles), uninfected (open symbols) and infected (closed symbols) cultures with the single stranded (ss) RNA-containing virus, CtenRNAV. Individual symbols represent independent biological replicates (n=3) with lines of best fit depicting a LOESS regression.



Extended Data Fig. 8 | See next page for caption.

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Extended Data Fig. 8 | **A** role for oxidative stress and reactive oxygen species during infection of *Chaetoceros tenuissimus*. **a**, Host abundance and **b**, intracellular levels of reactive oxygen species (ROS; assessed by the diagnostic, fluorescent stain H_2DCFDA in replete, uninfected (open, orange symbols/bars) and infected (closed, orange symbols/bars) *C. tenuissimus* cultures with CtenDNAV. Mean \pm standard error is shown (n = 3). **c**, Host abundance and **d**, maximum photochemical quantum yield of photosystem II (photosynthetic efficiency; F_v/F_m) following addition of different doses of hydrogen peroxide (H_2O_2 , 0-150 µM). Mean \pm standard error is shown for biological duplicates. Due to the observed decrease in cell abundance and photosynthetic efficiency, followed by physiological recovery, 100 µM H_2O_2 was chosen for downstream infection experiments. Time course of **e**, host abundance and **f**, photosynthetic efficiency during viral infection with CtenDNAV of untreated control cultures (orange circles) and cultures pre-exposed to a sub-lethal dose of H_2O_2 (100 µM; blue squares) for uninfected (open symbols) and infected (closed symbols) cultures (n=3). Data are representative of three independent experiments.

Extended Data Table 1 | Initial bulk parameters and underlying nutrient regime at each site

Site ID	Region	Location	Chl a	bSiO2	NO3	Si(OH)4	Fe	Temp	Salinity	Nutrient Regime
S1	CUZ	Coastal; 38°39.3N, 123°39.87W	12.3 ± 1.63	12.9	13.7 ± 1.39	20.8 ± 2.53	3.39	10.7	33.8	Replete
S2	CUZ	Coastal; 38°15.31 N, 123°57.98W	0.99 ± 0.07	2.96	10.3	4.69 ± 0.02	0.42	12.5	33.6	Si-limited
S4	CUZ	Coastal; 42°40N, 125°W	0.30 ± 0.01	0.27	16.9 ± 0.22	19.3 ± 0.11	0.31	10.9	33.4	Replete → Fe limited
P4	NE Pacific	Coastal; 48°39N, 126°40W	0.54 ± 0.34	1.4 ± 0.26	1.49	2.18	0.64	12.4	31.9	N-limited
P26	NE Pacific	Oceanic; 50°N, 145°W	0.08 ± 0.02	0.44 ± 0.09	11.7	15.6	0.05	10.5	32.5	Fe-limited

Additional chemical and biological parameters measured at sites in the California Upwelling Zone (CUZ) and along the Line P transect in the subarctic northeast Pacific (NE Pacific). Site designation, region and location are included with initial measurements of chlorophyll *a* (Chl *a*; μ g L⁻¹), biogenic Si (bSiO₂; μ M), dissolved nitrate (NO₃ and NO₂; μ M), dissolved silicic acid [Si(OH),; μ M], dissolved iron (Fe; nM), temperature (Temp; °C) and salinity (PSU). Nutrient regime summarizes the underlying nutrient condition characterized for each location. When available mean and standard deviations are shown from replicate samples (n=3).