Enhancement of viral production by addition of nitrogen or nitrogen plus carbon in subtropical surface waters of the South Pacific

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ABSTRACT: We tested the hypothesis that viral production is limited by nutrient availability in oligotrophic subtropical surface waters of the South Pacific. Nutrient (C, N, P) addition experiments were conducted at 2 stations (Stn SX18, 39° 60' S, 169° 60' W; Stn SX22, 19° 60' S, 169° 60' W) to examine the responses of viral production (3H-thymidine incorporation) and related variables in bottle-contained surface waters. At Stn SX18, viral production was enhanced in bottles amended with N and C with concomitant increases in bacterial production and growth, suggesting that both viral and bacterial production were co-limited by N and C. At Stn SX22, additions of N accelerated viral production with no consistent response of bacterial production, which might be an indication that prophage production was induced due to the alleviation in N limitation of host bacteria. In both experiments, nutrient-induced increases in viral production were not accompanied by increases in viral abundance, suggesting a possibility that viral production was balanced by loss. Our results suggest that viral production is limited by the availability of N or N+C in subtropical waters of the South Pacific, implying that viruses affect the mode of nutrient-induced changes in bacterial production and microbial trophic transfers in oligotrophic oceanic waters.

KEY WORDS: Viruses · Viral production · Nutrient limitation · Oligotrophic surface waters · South Pacific subtropical gyre

INTRODUCTION

Lytic viral production and associated destruction of host bacterial cells account for a large fraction (10 to 50%) of bacterial mortality in various aquatic environments, playing an important role in controls of bacterial communities, food webs, and biogeochemical cycling (Fuhrman 1999, Wilhelm & Suttle 1999, Weinbauer 2004). Previous studies have found that additions of nutrients stimulate lytic viral production in estuarine (Hewson et al. 2001), coastal (Tuomi et al. 1995, Wilson et al. 1998), neritic (Williamson & Paul 2004), and freshwater (Weinbauer et al. 2003, Lymer & Vrede 2006) environments, leading to the proposition that the enhancement of the production of host bacteria due to the alleviation of nutrient limitation results in increased viral production. Factors that account for this response in viral production may include increased encounter probability between host and viruses (density-dependent effect, Thingstad 2000), increased host activity (growth-dependent effect, Middelboe 2000), and induction of prophage production due to alterations in the metabolic status of the lysogenic host (lysogenic decision effect, Williamson et al. 2002). Enhancement of lytic viral production in response to nutrient additions has important implications for aquatic ecosystems, since it results in reduced transfer efficiency of bacterial production to higher trophic levels (Fuhrman 1999, Wilhelm & Suttle 1999). However, little is known about how viral populations respond to nutrient additions in oligotrophic open waters of subtropical gyres, the largest oceanic domain on earth, where bacterial growth is limited by carbon and nutrient availability.

Previous studies conducted in coastal and neritic waters have suggested that P availability limits lytic viral production; these studies include investigations conducted in the Gulf of Mexico (Williamson & Paul 2004) and the Raunefjorden (Tuomi et al. 1995, Wilson et al. 1998). These results might be explained by the high nucleic acid to protein ratio (hence high P:N) of viruses (Bratbak et al. 1993). However, C and N can also limit viral production depending on the environment (Tuomi & Kuuppo 1999, Williamson & Paul 2004). Stoichiometric constraints on viral production have yet to be fully clarified (Wikner et al. 1993). To our knowledge, no previous studies have examined the responses of viral production to P, N, and C additions for a complete set of single or combined additions of a suite of nutrients, making it difficult to assess which element limits (or co-limits) lytic viral production.

Here we tested the hypothesis that viral production is limited by nutrient availability in the subtropical gyre of the South Pacific. This region is characterized by low chlorophyll (chl) a concentrations and low primary productivity (Longhurst 1998), with few data available on processes mediated by bacteria and viruses. We conducted nutrient addition assays using surface waters and found that N or N+C additions enhance viral production with coupled or uncoupled responses of bacterial production and growth.

**MATERIALS AND METHODS**

**Study site, sample collection, and experimental setup.** Sample waters were collected at 2 stations deployed in the South Pacific (Stn SX18: 39° 60’S, 169° 60’W; Stn SX22: 19° 60’S, 169° 60’W) during a cruise by RV ‘Hakuho-Maru’ (KH 04-5) between January and February 2005. The sampling stations are located in the South Pacific Subtropical Gyre (SPSG) Province, although Stn SX18 is close to the convergence zone (Longhurst 1998). Surface water samples were collected with a clean bucket, filtered through 150 µm nylon mesh to eliminate large plankton, and contained in 20 l polyethylene tanks. One liter each of the filtrate was dispensed into polycarbonate bottles (1 l capacity, Nalgene). Either P (NaH2PO4), N (NH4Cl), or C (glucose) was added alone or in combination to sample waters in order to obtain a complete set of sole or combined additions of elements (P, N, C, P+N, P+N+C, N+C, P+N+C, and a non-addition control) with a fully replicated (3 bottles for each treatment) factorial design. The final concentration of each compound added was 1 µM. The bottles were incubated for 24 h at in situ temperature in the dark. The bucket, tanks, nylon mesh, and bottles used for the experiments were rinsed before use with 10% hydrochloric acid followed by vigorous rinsing with Milli-Q water. During sample collection and handling, gloves were worn and care was taken to minimize contamination. However, a clean bench and metal-free reagents to minimize trace metal contamination were not used in this study. Potential effects of trace metals (e.g. iron) on viral production and bacterial activities (Church et al. 2000) were outside the scope of this study.

**Abundances of bacteria and viruses.** For the determination of bacteria and viral abundances, subsamples were withdrawn from original sample waters and from incubation bottles at the end of incubation. Subsamples were preserved with 0.2 µm filtered paraformaldehyde (final concentration 2%), stored in a refrigerator, and used for the preparation of slides according to Noble (2001). Slides were prepared within 12 h after sampling. Triplicate slides were prepared for each bottle. Two ml of samples were filtered through a 0.02 µm Anodisc filter (Whatman), followed by rinsing with 0.02 µm filtered Milli-Q water. Filters were placed on 100 µl of SYBR Green I solution (Molecular Probes, final dilution 2.5 × 10^{-3}), and stained for 15 min in the dark. After staining, excess dye was removed with Kimwipes (Kimberly-Clark). The filters were then mounted with an anti-fade mounting solution (4:1 mixture of AF-1, Citifluor, and VectaShield, Vector Labs) to be stored frozen until later analysis. Bacteria and viruses were counted using an epifluorescence microscope (Olympus BX61; filter U-MWIB2) equipped with a 100 W Hg lamp (HSH-1030L). At least 300 cells of bacteria and 200 virus-like particles, distinguished on the basis of shape and brightness (Noble & Fuhrman 1998), were counted for each slide. The coefficient of variation (CV) for triplicate slides was 1 to 10% and 1 to 17% for bacterial abundance and viral abundance, respectively.

**Bacterial production.** Bacterial production was determined from the incorporation rate of thymidine (TdR) using a centrifuge method (Kirchman 2001). Triplicate subsamples (1.5 ml, contained in screw-capped centrifuge tubes, SSI) and 1 trichloroacetic acid (TCA)-killed control were amended with [methyl-3H] TdR (89 Ci mmol⁻¹, Amersham, TRK686, final concentration 10 nM) to be incubated for 12 to 17 h at in situ temperature in the dark. Extraction by precipitation with 5% cold TCA was followed by cold ethanol rinsing using a temperature controlled desktop centrifuge (18000 × g at 4°C for 10 min for each run; Eppendorf, 5417R). The extracts were then completely dried and mixed with scintillation cocktail (1 ml, Ultima Gold, Packard Instruments) for the radioassay using a Wallac 1400 scintillation counter with cor-
Viral production. Viral production was determined by the TdR method with enzyme digestions according to Noble & Steward (2001) with modifications. Triplicate subsamples (7 ml each) were contained in polypropylene tubes (14 ml capacity, BD Falcon), amended with [methyl-3H] TdR (89 Ci mmol^{-1}, Amersham, TRK686, final concentration 10 nM), and incubated for 24 h at in situ temperature in the dark. The incubation was terminated by filtering the samples through 0.2 µm syringe filters (Acrodisc, Pall). Triplicate filtrates (1.5 ml each) were contained in screw-capped microcentrifuge tubes (2 ml capacity, SSI) to be incubated (room temperature for 1 h) with a mixture of nucleases [1 U DNase I [Sigma, D5025], 1 U RNase A [Sigma, R4875], and 5 U Micrococcal nuclease [Worthington, NFCP] µl^{-1}]. After incubation, samples were heated (100°C, 1 min) to denature the enzymes, followed by cooling for 10 min on ice. In order to hydrolyze viral capsids and bacteria-derived protein, which might be non-specifically radiolabeled because of TdR catabolism (Brittain & Karl 1990), we treated the samples with Proteinase K (100 µg ml^{-1} final concentration, Sigma, P2308, 37°C for 1 h). After heating (100°C for 1 min) and cooling (on ice for 10 min), each sample was spiked with 40 µl of a carrier solution (50 µg ml^{-1} final concentration each of DNA [Sigma, D4522], RNA [Sigma, D4522], and BSA [Sigma, B4287]) and 80 µl of ice-cold 100% TCA. Precipitates were collected by centrifugation (18000 × g at 4°C for 10 min; Eppendorf, 5417R), resuspended in ice-cold 5% TCA, and extracted again by centrifugation. The precipitates were then hydrolyzed with 50 µl of 5% TCA at 90°C for 30 min. After cooling, 1 ml of scintillation cocktail (Ultima Gold, Packard Instruments) was added to each tube for the radioassay (see above). TCA-killed controls were prepared for each treatment. However, because of high variability in radioactivity of TCA-killed controls, a fixed value (40 dpm) was used as a blank for all treatments. Ratios of radioactivity of sample to blank generally exceeded 7. The CVs of triplicate measurements were 2 to 49%.

Abundance of heterotrophic nanoflagellates. Abundances of heterotrophic nanoflagellates (HNF) in original sample waters were counted according to Sherr & Sherr (1983). Subsamples (50 ml), fixed with glutaraldehyde (final concentration 2%), were double-stained with 4’,6-diamidino-2-phenylindole (DAPI) and fluorescein isothiocyanate (FITC). Cells were collected on 0.8 µm black Nucleopore filters (Whatman) and counted under an epifluorescence microscope (Olympus BX61; filters used were U-MWU2 and U-MWIB2 for DAPI and FITC, respectively).

RESULTS

Environmental and microbial variables

At the time of our sampling, surface water temperature and salinity at Stn SX18 were 17.2°C and 35.0 PSU, respectively, whereas corresponding values at Stn SX22 were 28.6°C and 35.1 PSU (Table 1). The low concentrations of chl \( a \) (0.10 µg l^{-1}, and 0.09 µg l^{-1} for Stns SX18 and SX22, respectively) and nitrate (<0.01 µmol l^{-1} for both stations) in surface waters that we collected (Table 1) are typical for the SPSG Province (Longhurst 1998).

Bacterial production rates (TdR incorporation rates) were 9.6 ± 0.6 and 13.0 ± 0.9 pmol TdR l^{-1} d^{-1} at Stns SX18 and SX22, respectively. These values are close to the range of the TdR incorporation rates determined in other subtropical regions (e.g. Hoppe et al. 2006). Viral production determined from TdR incorporation varied in the range of 0.2 ± 0.1 pmol TdR l^{-1} d^{-1} (Stn SX18) and 12.0 ± 1.8 pmol TdR l^{-1} d^{-1} (Stn SX22); these values fall within a range of viral incorporation rates of TdR reported in other marine systems, including coastal and high latitude oceanic regions (Steward et al. 1996, Helton et al. 2005). Other chemical and biotic variables are summarized in Table 1.

<table>
<thead>
<tr>
<th>Stn</th>
<th>Temp. (°C)</th>
<th>Salinity (PSU)</th>
<th>PO(_4)^{3-} (µM)</th>
<th>NO(_3) (µM)</th>
<th>Chl a (µg l^{-1})</th>
<th>Bacterial abundance (×10^6 cells l^{-1})</th>
<th>Bacterial production (pmol TdR l^{-1} d^{-1})</th>
<th>Viral abundance (×10^9 viruses l^{-1})</th>
<th>Viral production (pmol TdR l^{-1} d^{-1})</th>
<th>HNF abundance (cells l^{-1})</th>
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<tr>
<td>SX18</td>
<td>17.2</td>
<td>35.0</td>
<td>0.18</td>
<td>&lt;0.01</td>
<td>0.10</td>
<td>2.6 ± 0.2</td>
<td>9.6 ± 0.6</td>
<td>5.0 ± 0.4</td>
<td>0.2 ± 0.1</td>
<td>500 ± 170</td>
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<td>(39° 60' S, 169° 60' W; 22 Jan 2005)</td>
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<tr>
<td>SX22</td>
<td>28.6</td>
<td>35.1</td>
<td>0.09</td>
<td>&lt;0.01</td>
<td>0.09</td>
<td>1.8 ± 0.6</td>
<td>13.0 ± 0.9</td>
<td>11.0 ± 0.7</td>
<td>12.0 ± 1.8</td>
<td>120 ± 25</td>
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<td>(19° 60’ S, 169° 60’ W; 31 Jan 2005)</td>
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Responses of bacteria and viruses to nutrient additions

In order to examine the responses of bacteria and viruses to nutrient additions, we compared 3 variables among treatments. (1) TdR incorporation rates of bacteria and viruses determined at the end of the incubation for 24 h; these values are regarded as indices of bacterial production and viral production and are denoted hereafter as BP_{24} and VP_{24}, respectively. (2) Increases in the abundances of bacteria and viruses during the incubation were determined, i.e. (abundance at the end of the incubation for 24 h) – (abundance at the beginning of the incubation); these values are regarded as indices of numerical responses of bacteria and viruses and are denoted hereafter as ΔN_b and ΔN_v, respectively. (3) Cell-specific incorporation rates of TdR by bacteria were determined at the end of the incubation for 24 h; these values are regarded as indices of bacterial growth rates and are denoted hereafter as BG_{24}.

At Stn SX18, BP_{24} differed significantly among treatments (ANOVA with a Bonferroni correction, p < 0.001), with much higher (11- to 19-fold) values in the N+C and P+N+C treatments (BP_{24}, 160 pmol TdR l^{-1} d^{-1}) relative to those in other treatments, including the non-addition control (9 to 14 pmol TdR l^{-1} d^{-1}; Fig. 1A). ΔN_b varied in the range of −0.2 to 0.6 × 10^8 cells l^{-1}, which did not differ significantly (p > 0.05) among treatments (Fig. 1B). BG_{24} differed significantly (p < 0.001) among treatments, with much greater (12- to 22-fold) values for the N+C and P+N+C treatments (62 to 63 × 10^{-20} mol TdR cell^{-1} d^{-1}) relative to other treatments (3 to 5 × 10^{-20} mol TdR cell^{-1} d^{-1}; Fig. 1C). For VP_{24}, the equal-variance criterion was not met for a multiple comparison because of high variability associated with the estimate of mean VP_{24} in the P+N treatment; the coefficient of variation (CV) of VP_{24} for the P+N treatment was 79%, which was much greater than CVs of VP_{24} for other treatments (11 to 24%). Therefore, in the following comparison of VP_{24}, we excluded the estimate for the P+N treatment. The result indicates that VP_{24} varied significantly (p < 0.001) among treatments, with much greater (4- to 8-fold) values for the N+C and P+N+C treatments (15 to 19 pmol TdR l^{-1} d^{-1}) relative to other treatments (2.6 to 3.5 pmol TdR l^{-1} d^{-1}; Fig. 1D). ΔN_v varied in the range of −1.5 to −0.5 × 10^8 viruses l^{-1}, with no significant difference among treatments (p > 0.05; Fig. 1E).

The range of BP_{24} values observed in the Stn SX22 experiment (200 to 610 pmol TdR l^{-1} d^{-1}) was higher than that in the Stn SX18 experiment (9 to 160 pmol TdR l^{-1} d^{-1}), which might be related to higher water temperature at Stn SX22 (29°C) than at Stn SX18 (17°C). At Stn SX22, BP_{24} differed among treatments

![Fig. 1](image-url)
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(p < 0.001), with either lower (P+C treatment, 200 pmol TdR l–1 d–1) or higher (P+N treatment, 610 pmol TdR l–1 d–1) values relative to the control (440 pmol TdR l–1 d–1) and other treatments (400 to 520 pmol TdR l–1 d–1; Fig. 2A). $\Delta N_b$ varied in the range of 1.4 to 4.9 × 10^8 cells l–1; $\Delta N_b$ in the P+N+C treatment was significantly (p < 0.05) higher than corresponding values relative to the control (Fig. 2B). $BG_{24}$ differed significantly (p < 0.05) among treatments, with higher values in the N treatment (160 × 10^{-20} mol TdR cell–1 d–1) relative to the control (100 × 10^{-20} mol TdR cell–1 d–1; Fig. 2C). $VP_{24}$ varied in the range of 8.3 to 34.0 pmol TdR l–1 d–1 and differed among treatments (p < 0.001). $VP_{24}$ in the N, P+N, N+C, and P+N+C treatments were 2- to 3-fold greater than those in the control and other treatments, which varied in the range of 8.3 to 14.0 TdR l–1 d–1 (Fig. 2D). $\Delta N_v$ was in the range of −3.3 to 0.4 × 10^8 viruses l–1, which did not differ among treatments (p > 0.05; Fig. 2E).

**DISCUSSION**

Our study is the first to demonstrate viral responses to nutrient additions in oligotrophic (chl a < 0.1 µg l–1) oceanic environments. The enhancement of viral production ($VP_{24}$) in the N+C and P+N+C treatments of the SX18 experiment suggests that viral production was co-limited by N and C. This response of viral production agreed well with that of bacterial production ($BP_{24}$) and growth ($BG_{24}$) following nutrient additions, indicating that viral production was enhanced because of increasing bacterial growth. Because the numerical response of bacteria ($\Delta Nb$) to the enrichment was minimal (probably because bacterial growth was balanced by the loss due to HNF bacterivory; Strom 2000), it is unlikely that the enhancement of viral production was due to the increasing encounter probability between host and viruses. The above results are consistent with the hypothesis that viral production depends on the growth of the host (Moebus 1996a,b, Middelboe 2000), which in turn is limited by nutrients in marine environments (Bratbak et al. 1993, Tuomi et al. 1995, Hewson et al. 2001, Williamson & Paul 2004, Bongiorni et al. 2005).

In contrast, at Stn SX22, we found that viral production ($VP_{24}$) was enhanced in treatments with N additions (N, P+N, N+C, and P+N+C), whereas bacterial production ($BP_{24}$) and abundance ($\Delta N_b$) did not display such a trend, except that $BP_{24}$ in the P+N treatment, $\Delta N_b$ in the P+N+C treatment, and $BG_{24}$ in the N treatment were significantly higher than the corresponding values in the other treatments. This uncoupling between bacterial and viral responses to nutrient additions at Stn SX22 may indicate that the induction of prophage production occurred among lysogenic populations of bacteria in response to the alteration of the metabolic status of host cells. In the bacteriophage $\lambda$, it has been suggested that a ‘lysogenic decision’ occurs toward the mode of lytic phase in response to changes in growth conditions from poor to rich media (Oppenheim et al. 2005). Consistent with this notion, William-
son et al. (2002) found that prophage induction was occasionally stimulated by P additions in Tampa Bay, Florida, USA. This led us to hypothesize that the enhanced viral production at Stn SX22 was due to the switching of lysogenic to lytic mode of infection of viruses due to metabolic alterations of bacteria, explaining the attenuation in responses of bacterial production to N enrichment. This hypothesis is consistent with the proposition that a lysogenic mode of viral infection is common in oligotrophic oceans (Jiang & Paul 1998). However, further studies are clearly needed to examine the relative importance of lytic infection and lysogeny in oligotrophic waters of the South Pacific.

Some methodological problems should be commented on. We determined viral production using a TdR approach (Noble & Steward 2001), which has advantages in sensitivity and simplicity over other techniques such as the dilution and the fluorescently labeled virus methods (Noble & Steward 2001, Helton et al. 2005). The TdR method is particularly suitable for nutrient addition studies in which a large number of samples (n = 24 for each experiment in the present study) must be handled promptly. However, conversion factors that relate TdR incorporation rate to viral production have been poorly constrained, especially in oligotrophic environments (Helton et al. 2005). Although conversion factors may vary depending on viral genome size, G+C content, and isotope dilution factors, little is known about variations of these variables in oceanic environments. Given that viruses can use host nucleic acids and that RNA-containing viruses would not be labeled, viral production determined by the TdR method would be too low (Helton et al. 2005). Conversely, the TdR method might overestimate viral production because of the inclusion of non-specifically labeled components in the fraction of viral DNA. It has been suggested that TdR catabolism can cause non-specific labeling of bacterial components (e.g. proteins) (Brittain & Karl 1990), which potentially results in the production of 3H-labeled colloidal debris in the <0.2 µm fraction. To minimize this effect, we treated the samples with proteases (see ‘Materials and methods’). However, proteins and DNA coated by liposome-like particles might not be digested by enzymes (Nagata & Kirchman 1997). Because of the above reasons, the data on viral production obtained using the TdR method in oligotrophic systems should be interpreted cautiously. However, biases associated with the above factors are probably not systematic with regard to the treatments used in this study (i.e. nutrient additions). Thus, although the accuracy of the estimates of viral production determined by the TdR method remains to be ascertained by future studies, we consider that our conclusions concerning patterns in responses (i.e. relative values) of viral production to nutrient additions are valid.

Our results suggesting that viral production is limited by either N (Stn SX22 experiment) or N+C (Stn SX18 experiment) but not by P (both Stn SX22 and SX18 experiments) contrast the results obtained in previous enrichment studies, which have generally suggested that viral production is limited by P in coastal and neritic waters (Bratbak et al. 1993, Tuomi et al. 1995, Williamson & Paul 2004). Bratbak et al. (1993) suggested that viral production may be more sensitive to P than N limitation because the nucleic acid-to-protein ratios (hence P:N ratios) of viruses are high. However, it has been demonstrated that viruses use host nucleotides for the synthesis of viral DNA (Wikner et al. 1993). This mechanism could facilitate effective scavenging of P resources from the host, alleviating P limitation of viral production. Therefore, it is probably not surprising that viral production is limited by the availability of N rather than P to be used for the synthesis of capsid protein. However, the mode of stoichiometric regulations of virus–host systems under nutrient-limited conditions is poorly understood (Wikner et al. 1993). It remains to be seen if N limitation of viral production is widespread in oligotrophic open waters of the Pacific where N is chronically depleted.

In both the Stn SX18 and SX22 experiments, despite significant effects of nutrient additions to viral production, viral abundance (ΔNv) showed no consistent trends, being close to 0 or negative in most cases. One hypothesis to explain this observation is that the loss of viruses occurred during the incubation at rates comparable to or exceeding those of viral production. To examine the extent and variations of viral loss, the rate of the loss of viruses (L, viruses l–1 d–1) was calculated as follows:

$$L = \rho - \left( N_{24} - N_0 \right)$$

where ρ is the production rate of viral particles (viruses l–1 d–1) averaged for samples collected at the beginning and at the end of the incubation, and N24 and N0 are viral abundances at the end and at the beginning of incubation, respectively. To convert the incorporation rate of TdR to ρ, we used a conversion factor of 6 × 1020 viruses mol-TdR–1 (Steward et al. 1992). Results indicate that substantial losses occurred in both Stn SX18 (1 to 6 × 109 viruses l–1 d–1) and SX22 (7 to 14 × 109 viruses l–1 d–1) experiments. By using all data collected from the 2 experiments, we found that viral loss rates are positively correlated with bacterial production, which varied by 70-fold (9 to 610 pmol TdR l–1 d–1) across experiments and among treatments (Fig. 3); 89% of variations in viral loss rates were accounted for by bacterial production. Note that the above calculation is based on an assumption that viral production is
related to the $^3\text{H}$-TdR incorporation rate with a fixed conversion factor. However, in their study on $^{32}\text{P}$ tracer fluxes in a virus-bacteria model, Wikner et al. (1993) suggested that the isotope dilution decreases (hence viruses with higher specific activities are produced) with increasing specific growth rate of bacteria. If this model-based prediction generally applies to viral-host communities in oligotrophic open waters, the variation in the isotope dilution might explain, to a certain extent, the pattern depicted in Fig. 3. This possibility should be examined in future studies.

Several factors could contribute to the loss of viruses in marine waters. These factors include (1) enzymatic degradation and adsorption to colloidal substances (Noble & Fuhrman 1997), (2) grazing by HNF (Gonzalez & Suttle 1993), (3) scavenging by settling particles (Proctor & Fuhrman 1991), and (4) solar irradiance (Suttle & Chen 1992), although the latter 2 factors could be discounted in our incubation experiments. Our results showing that viral loss rates increase with increasing bacterial production suggest that enzymes and colloidal particles are more abundantly produced when bacterial production is high. Alternatively, high bacterial production may result in high grazing activities by flagellates, which in turn can consume viruses (Gonzalez & Suttle 1993). Regardless of the exact mechanisms, our results suggest that viral loss might be closely coupled with bacterially mediated processes. We also point out that effects of nutrient additions to viruses could be largely underestimated if only viral abundance is examined as a response variable.

To conclude, our data support the hypothesis that nutrients limit viral production in oligotrophic surface waters of the South Pacific, with element(s) of deficiency being N or N+C. Enhancement of viral production was coupled with bacterial growth rather than abundance, although our results also suggest that N addition might induce prophage production with minimal responses in bacterial production. Several studies have found that bacterial production is enhanced by additions of NH$_4$ and glucose (but usually not by the addition of NH$_4$ alone) in oceanic regions (Kirchman et al. 1990, Cherrier et al. 1996, Kirchman & Rich 1997, Church et al. 2000), but these studies did not examine responses of viral production. Generally, it is predicted that gross growth efficiency (GGE) of bacteria increases with increasing availability of N (Goldman et al. 1987), leading to the increase in efficiencies of the transfer of carbon from dissolved organic matter (DOM) to higher trophic levels via a DOM-bacteria pathway. However, if the response of bacteria to N (or N+C) addition is generally accompanied by viral responses, as we have suggested here, the enhancement by nutrients of bacterial GGE can be largely compensated by the enhancement of the ‘viral shunt’ (Fuhrman 1999, Wilhelm & Suttle 1999), a possibility that should be examined in future studies.

Despite increasing viral production due to nutrient additions, we failed to detect responses in the abundance of viruses, which appears to be explained by close coupling of viral loss and bacterial production. Further studies are required to examine in greater detail the mechanisms underlying coupling and uncoupling of bacterial production, viral production, and viral loss in marine systems.

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