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Viral lysing can alleviate microbial nutrient limitations and accumulate recalcitrant dissolved organic matter components in soil

Di Tong D^{1,2}, Youjing Wang D^{1,2}, Haodan Yu D^{1,2}, Haojie Shen D^{1,2}, Randy A. Dahlgren D³ and Jianming Xu D^{1,2}

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Viruses are critical for regulating microbial communities and biogeochemical processes affecting carbon/nutrient cycling. However, the role of soil phages in controlling microbial physiological traits and intrinsic dissolved organic matter (DOM) properties remains largely unknown. Herein, microcosm experiments with different soil phage concentrates (including no-added phages, inactive phages, and three dilutions of active phages) at two temperatures (15 °C and 25 °C) were conducted to disclose the nutrient and DOM dynamics associated with viral lysing. Results demonstrated three different phases of viral impacts on CO_2 emission at both temperatures, and phages played a role in maintaining Q_{10} within bounds. At both temperatures, microbial nutrient limitations (especially P limitation) were alleviated by viral lysing as determined by extracellular enzyme activity (decreased V_{angle} with active phages). Additionally, the re-utilization of lysate-derived DOM by surviving microbes stimulated an increase of microbial metabolic efficiency and recalcitrant DOM components (e.g., SUV254, SUV260 and HIX). This research provides direct experimental evidence that the "viral shuttle" exists in soils, whereby soil phages increase recalcitrant DOM components. Our findings advance the understanding of viral controls on soil biogeochemical processes, and provide a new perspective for assessing whether soil phages provide a net "carbon sink" vs. "carbon source" in soils.

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INTRODUCTION

Soil microorganisms are major agents of soil organic carbon (SOC) cycling, both as decomposers and contributors to the long-term persistence of SOC [1–3]. Annually, about 60 Pg SOC are mineralized to CO_2 by soil microorganisms and released into the atmosphere, thereby being a major regulator of global warming [4, 5]. To date, the role of abiotic factors in regulating microbes and thereby the SOC decomposition rate has been well established [6, 7]. However, the impact of microbial predators on microbial communities and their functions, through top-down control, remains elusive.

Viruses as the most abundant biological entity can dramatically influence microbial abundance and community structure through viral lysing [8–10]. In oceans, phages (viruses infecting prokaryotes) lyse 20–40% of prokaryotes each day, which significantly affects the marine carbon cycle where they account for an estimated 6–26% of marine carbon exchange [11, 12]. Although there is no direct evidence to what extent soil bacteriophages affect soil carbon cycling, the abundance of soil phages (roughly 10^7-10^{10} g⁻¹ soil) and the frequency of phage infection of bacterial cells are both higher than those found in aquatic environments [13–15]. The "viral shunt" mechanism refers to the cellular matter liberated by viral lysing that can be reutilized by surviving heterotrophic microbes. This further stimulates the recycling and mineralization of dissolved

organic matter (DOM) into CO₂, and hinders nutrient and energy flows to higher trophic levels [16, 17]. This process in the marine system can release 145 Gt carbon and boost the microbial utilization of DOM by up to 27% [18-20]. The viral shunt has been documented in the soil environment using a ¹³C labeling approach, whereas the quantity of carbon involved remains unexplored [21]. Another pathway by which phages influence carbon fluxes is termed the "viral shuttle". This mechanism describes the cell debris and transparent exopolymeric polysaccharides, as well as other lysis products, which are relatively recalcitrant to microbial decomposition, and therefore tend to aggregate and be retained in the soil environment for long time periods [22]. This concept is only discussed in a few review articles and is believed to act as a carbon sink in oceans [23]. To date, the viral shuttle has never been documented in soils, not to mention the possible mechanisms involved therein.

According to their life cycle, phages can be divided into virulent and temperate categories, whose infective properties are markedly different [24]. Virulent phages only act as predators controlling bacterial communities via lysing cells, based on the Kill-the-Winner model (KtW) [8]. In contrast, temperate phages can integrate their DNA or RNA into the host's chromosomes and replicate with their host, known as the lysogenic cycle [25]. When host cells cannot feed temperate phages for the replication of progeny, they switch

¹Institute of Soil and Water Resources and Environmental Science, College of Environmental and Resource Sciences, Zhejiang University, Hangzhou 310058, China. ²Zhejiang Provincial Key Laboratory of Agricultural Resources and Environment, Zhejiang University, Hangzhou 310058, China. ³Department of Land, Air and Water Resources, University of California, Davis, CA, USA. ^{Sem}enail: jmxu@zju.edu.cn

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1248

to the lytic cycle [26]. Different from KtW, lysogenic phages can help their host survive in harsh environments by integrating auxiliary metabolic genes (AMGs) into the host's genome, referred to as the Piggyback-the-Winner model (PtW) [10, 27, 28]. Additionally, temperature can alter interactions between phages and hosts, by altering the viral latent period, viral and host production rates, burst size, and so on [29, 30]. Moreover, different microbial species have different affinities for various DOM chemical components [31–34]. Based on the impact of diverse viral life models on bacterial communities and different microbial affinities for DOM components, we posit that the presence of viruses will alter the fractionation of DOM chemical components, thereby affecting the biogeochemical cycling of carbon.

Recent studies demonstrated that soil nutrient limitations are a crucial factor controlling microbial metabolism and carbon cycling in terrestrial ecosystems, especially phosphorus limitation [35]. This leads to the question: Can soil nutrient limitation be alleviated by release of intracellular materials by viral lysing? Moreover, the role of temperature in regulating the viral lysis rate and infection process maybe a critical factor in a rapidly warming world [29, 30, 36]. To address these knowledge gaps, we hypothesize that microbial nutrient limitation will be alleviated by active phages and the microbial metabolic efficiency will be increased due to lysate DOM, for which microbes have a strong affinity. Further, we posit that these processes will vary as a function of temperature.

There are few studies examining the impact of soil viruses on soil carbon and nitrogen cycling [37–39], let alone on soil carbon dynamics from the perspective of microbial trait-based strategies and DOM components. To test our hypotheses and disentangle the impacts of soil phages on SOM dynamics, soil bacteria and phages were reinoculated in sterilized soil and incubated at two different temperatures (15 °C and 25 °C). The objectives of this study were to (1) explore how soil phages influence microbial respiration; (2) disclose the impact of phages on the bacterial community and trait-based strategies; and (3) reveal if soil phages promote the accumulation of recalcitrant DOM components, consistent with the viral shuttle mechanism.

MATERIALS AND METHODS

Soil sampling

Soil was collected from a natural forest in Jiangxi Ecological Science and Technology Park of Soil and Water Conservation (115°42′08″-115°43′06″E, 29°16′37″-29°17′40″N). This is the center of the red soil region of China, located in De'an County, Jiangxi Province; additional details of the study area can be found in elsewhere [40]. Five $2 \times 2 m$ subplots were established at each corner and the center of three randomly established large plots (20 × 20 m), and the 15 subplot composite samples (0–20 cm) were combined to form a single equal-weight composite representative of the study area. The sample was transported to the laboratory on ice. Plant residues and coarse fragments were removed from the soil sample by passing through a 2-mm sieve, and the <2-mm soil fraction divided into two parts: (1) air-dried for physicochemical analysis and (2) stored fresh at 4°C for extraction of soil bacteria and phages.

Preparation of phage and bacterial suspensions from soil

Soil phage and bacterial suspensions were extracted from 1.2 kg of soil suspended in 1.2 L saline magnesium buffer (SM; per L containing 5.8 g NaCl, 2 g MgSQ₄:7H₂O, 50 mL 1 M Tris-HCl (pH 7.5)). The suspension was vortexed for 3 min at maximum intensity, shaken for 2 h at 250 rpm and then centrifuged at 4 °C for 20 min $(2500 \times g)$ [41, 42]. The supernatant was passed through a 2-µm cellulose acetate filter, before being filtered and concentrated using tangential flow filtration (TFF) with two different poresize filters (0.22 µm and 100 kDa) [17, 37]. The retentates obtained (>0.22 µm) were used as bacterial suspensions, and the concentrated filtrates were used as phage suspensions (100 kDa-0.22 µm) and no-phage suspensions (<100 kDa) [38]. The final volumes of retentates and filtrates were 500 mL, 80 mL and 620 mL, respectively. In order to simulate the in-situ environment, we added 1.2 ml of phage suspensions and 3.6 ml bacterial suspensions to each serum bottle containing 35 sterilized

air-dried soil, which approximated real-world phage and bacteria concentrations. We then adjusted the soil moisture to 40% of field capacity with 4.9 ml of no-phage water (filtered through 100 kDa).

Experimental design

Experiments were conducted in 250-mL serum bottles with 35 g sterilized air-dried soil (50 kGy γ -radiation over 24 h). Microcosm treatments consisted of (1) bacterial suspensions with no added phage (SM); (2) bacterial suspensions with autoclaved dead phage (DV) [38]; and (3) bacterial suspensions with different phage dilutions (no dilution (V), twofold dilution (2⁻¹ V) and tenfold dilution (10⁻¹ V)). All treatments were incubated at two different temperatures (15 °C and 25 °C) with four replicates. The total number of microcosms were 5 soil phage states (no phage, inactive phage, and active phage with 0-, 2- and tenfold dilution) × 2 temperatures × 3 sampling times (days 7, 28 and 98) × 4 replicates = 120 total microcosms.

Soil properties and gas emission analysis

A pH meter was applied to measure soil pH in a 1:2.5 soil: distilled water ratio after a 30 min equilibration. Soil organic carbon (SOC) and soil total nitrogen (TN) were quantified by an elemental analyzer. Available P (Bray-P) was extracted with 0.03 M NH₄F/0.025 M HCl and quantified using the molybdenum blue spectroscopy method [43, 44]. Microbial biomass carbon (MBC), nitrogen (MBN) and phosphorus (MBP) were measured by CHCl₃ fumigation-extraction [45, 46]. Soil total P (TP) was measured using a digestion method [47, 48]. Dissolved organic matter (DOM) as characterized by organic carbon (DOC) and nitrogen (DON) was extracted with Milli-O water at a ratio of 1:10 (w/v) and determined using a Multi C/N TOC analyzer. Extracted DOM samples were kept in the dark at 4 °C for further characterization. Soil NH4⁺ and NO3⁻ were extracted by 1 M KCl from fresh soil, and determined using a flow injection analyzer [49]. The measurement of extracellular enzyme activity, putative microbial physiological traits and calculation of microbial metabolic efficiency are described in Supplementary Information.

Soil CO₂ fluxes from the incubation studies were determined at 2, 5, 7, 14, 17, 21, 28, 35, 42, 49, 56, 63, 70, 77, 84, 91 and 98 days using gas chromatography (GC-200 Plus Shimadzu, Japan) as described by [50–52]. The respiration rate (R) of CO₂ (mg CO₂ kg soil⁻¹ d⁻¹) was calculated as [52]:

$$R = \rho \times \frac{V}{M} \times \frac{dc}{dt} \times \frac{273}{T}$$
(1)

where ρ , V, M, dc/dt and T denote the density of CO₂ at standard temperature and pressure (1.98 g/L), volume of incubation jar (L), mass of soil (g), slope of the linear regression for gas concentration gradient through time and incubation temperature (K), respectively. Cumulative CO₂ emission was calculated as the sum of the area bounded by the rate using the trapezoidal rule.

DOM spectral analysis

DOM concentration of incubated soil was expressed as the DOC concentration determined by a total organic carbon (TOC) analyzer [53]. The UV-Vis spectra of DOM were assessed for a series of indices as proxies for chemical characteristics of the DOM (e.g., SUV254, SUV260 and spectral slope ratio (S_R) representing the aromaticity, hydrophobicity and molecular weight, respectively), via an ultraviolet-visible spectrophotometer [54–56]. SUV254 and SUV260 were calculated as the ratio of UV absorption at 254 and 260 nm to the DOC concentration [57, 58].

After diluting DOC concentrations to 8 mg C L⁻¹, excitation-emission matrix (EEM) spectra of DOM were measured by fluorescence spectrometry, with a scan rate of 2000 nm min⁻¹ [59]. The excitation wavelength range was 200–500 nm (5-nm sampling interval), whereas the emission wavelength range was 250–600 nm (2-nm sampling interval). EEM fluorescence spectra were visualized in MATLAB 2012b with DOM Fluor toolbox [56, 58]. The humification index (HIX) was calculated for an excitation wavelength of 254 nm, by dividing the peak area of the emission wavelengths between 435 and 480 nm by the sum of the areas between 435–480 nm and 300–345 nm [60, 61].

High-throughput sequencing and analysis of bacterial community

Soil DNA was extracted using the MoBio PowerSoil DNA Isolation Kit (MP Biomedicals, USA) following manufacturer's instructions. The primers 515 F/907 R were used to amplify the V4-V5 region of the 16 S rRNA gene

and sequenced on an HiSeq PE250 sequencing platform (Illumina, USA) [31, 62]. Bacteria reads were in the range of 48,477–89,042 per sample, and the screened sequences were clustered into operational taxonomic units (OTUs) using UPARSE, with a similarity cutoff of 97% [48, 49]. For each representative sequence, annotated taxonomic information was determined using the SILVA (version 138) database [63]. The obtained sequences were used to assess bacterial α-diversity (i.e., Shannon index) and principal coordinates analysis (PCoA) was applied to represent the shift in the bacterial community based on a Bray–Curtis dissimilarity matrix, with the vegan package (version 2.5–7) [32].

Statistical analyses

Redundancy analysis (RDA) was applied to reveal relationships between bacterial communities and various soil/microbial parameters using the vegan package in R 3.6.3. Spearman heatmap plots and other plots were visualized by Origin. One-way ANOVAs with subsequent least-significantdifference (LSD) tests were used to determine differences among measured parameters (e.g., V_{angle} , V_{length} , CO_2 emission, CUE, SUV254, HIX SUV260). The main statistically significant factors for predicting CO₂, qCO₂ and CUE were identified by random forest analysis (R package: rfPermute), whereby a higher percentage of mean squared error (%IncMSE) indicates a more important factor [64, 65]. A correlation network between bacterial taxa and measured parameters was used to identify soil taxa strongly interacting with Vanale and recalcitrant DOM (e.g., SUV254, SUV260 and HIX). Spearman correlations were considered as strong when r > 0.5 and significant when p < 0.05; correlation matrices were visualized with Gephi [66, 67]. SPSS Version 24 (SPSS, Chicago, IL, USA) and R version 3.6.3 were used for statistical analysis with a significant level of p < 0.05, unless otherwise stated.

RESULTS

Microbial respiration dynamics at different temperatures

All five soil virus treatments (i.e., SM, DV, V, 2^{-1} V and 10^{-1} V) displayed three phases of CO₂ released during soil incubation (Fig. 1). In the first phase (first two weeks after microbe/virus inoculation), the CO₂ efflux rate at both 15 °C and 25 °C was lower in active viral groups, compared to the SM and DV groups. During the second and third phases, the trend for CO₂ efflux rates varied between temperatures. At 25 °C, the second phase (days 14 to 42) CO₂ efflux rate with active viruses was higher than that in SM and DV groups, although the CO₂ emission rate was different among active viral dilutions. During the third phase (days 42-98), the CO₂ efflux rate in the SM and DV groups again exceeded those of the active groups (Fig. 1A). In contrast to the 25 °C treatments, the phase 2 (days 15 to 35) CO_2 efflux rate at 15 °C for the 10^{-1} V group with active viruses was higher than the SM and DV groups. For phase 3 at 15 °C, the CO₂ efflux rate in all active viral groups was significantly higher than in the SM and DV groups, which persisted for the duration of the experiment (Fig. 1B).

Total cumulative CO₂ fluxes during the incubation are shown in Fig. S1. To assess the mechanisms associated with the three phases of CO₂ effluxes, three reference time points were selected to characterize each phase: Days 7, 28 and 98. Cumulative CO₂ emissions with active viruses at 25 °C were significantly lower than the SM and DV groups (p < 0.05) in the first phase (Fig. S2a). During the second phase, cumulative CO₂ efflux in active viral groups increased and exceeded those of the SM and DV groups. Cumulative CO₂ production in active viral groups followed the order: $25-2^{-1}V > 25-V > 25-10^{-1}V$ (Fig. S2c). By the end of the incubation, cumulative CO₂ production changed to the order: 25- $DV > 25-10^{-1} V > 25-SM \approx 25-V \approx 25-2^{-1} V$ (Fig. S2e). In contrast, cumulative CO₂ production with active viruses at 15 °C showed the order of $15-10^{-1}$ V > $15-2^{-1}$ V > 15-V throughout the entire incubation period (Fig. S2b, d, f). Compared to the 15-SM and 15-DV groups, cumulative CO₂ production in active viral groups was lower at the early stage. For the entire incubation period, cumulative CO₂ production followed: $15-10^{-1}$ V > $15-2^{-1}$ V > $15-2^$ V > 15-DV > 15-SM (Fig. S2f).

The temperature sensitivity of CO_2 emissions (Q_{10}) was calculated, to investigate how soil phages influence Q_{10} . The Q_{10}



Fig. 1 Dynamics of CO₂ emission rate (mean ± std dev) with added viruses incubated over 98 days. 25 °C (A) and 15 °C (B). SM, no added viruses; DV, inactive viruses; V, active viruses with no dilution; 2^{-1} V, active viruses with twofold dilution; 10^{-1} V, active viruses with tenfold dilution; 25 and 15 signify different temperatures (25 °C and 15 °C) during the incubation.

values of active phage groups were significantly lower than those in SM and DV groups, and were maintained within the bounds of 1–3 after the cultivation was stable (Fig. S3).

Soil microbial metabolic efficiency and resource acquisition traits

To illustrate the influence of soil viruses on microbial metabolic efficiency and resource acquisition traits, qCO_2 and CUE were used to quantify the metabolic efficiency, and resource acquisition traits were assessed by V_{length} and V_{angle} . For microbial metabolic efficiency, qCO_2 showed no observable differences among different groups or temperatures (Fig. S4), whereas CUE significantly changed after the addition of active viruses (Fig. 2). At 25 °C, the CUE of groups inoculated with active viruses was higher than the SM and DV groups and decreased with increasing dilution (Fig. 2A). At 15 °C, only the CUE of the 15-V group was higher than the 15-SM group, even though the trend of CUE with active viruses declined with dilution. The addition of inactive viruses seemed to increase the CUE relative to the no virus addition group (SM) at 15 °C (Fig. 2B).

Microbial acquisition strategies can be elucidated from V_{length} and V_{angler} where the former is used to represent the relative C vs. nutrient (N and P)-acquiring enzyme activities and the latter



Fig. 2 Carbon use efficiency (CUE) (mean ± std dev). 25 °C (A) and 15 °C (B). SM, no added viruses; DV, inactive viruses; V, active viruses with no dilution; 2^{-1} V, active viruses with twofold dilution; 10^{-1} V, active viruses with tenfold dilution; 25 and 15 signify different temperatures (25 °C and 15 °C) during the incubation. Different lowercase letters represent significant differences at p < 0.05.

expresses the relative P vs. N-acquiring enzyme activities. All groups were deemed plentiful with respect to organic carbon (V_{length} was less than 1), albeit a higher V_{length} occurred for active viral groups (Fig. S5). Adding active viruses into the soil seemed to alleviate nutrient constraints, especially for phosphorus, based on the decrease in V_{angle} (Fig. 3). The effect of alleviated nutrient constraints depended on temperature, with the lowest V_{angle} occurring for the tenfold dilution group at 25 °C, whereas the opposite was found at 15 °C.

Accumulation of recalcitrant DOM components

The DOC concentration across the different groups showed no significant differences. In contrast, the optical properties of DOM were diverse among treatments, as measured by 3D-EEM fluorescence spectroscopy and UV–vis spectroscopy (Figs S6, S7). After addition of soil viruses at 25°C, SUV254 and HIX showed a considerable increase of 37.9–98.1% and 48.3–105.3%, respectively (Fig. 4A, C). This indicates a higher degree of humification and aromaticity, characteristics that make DOM more resistant to microbial decomposition. At 15 °C, only the 15-V group with the highest viral inoculation produced higher SUV254 (p < 0.05), as compared to the 15-SM and 15-DV groups (Fig. 4B). Inoculating soil viruses increased HIX at 15 °C, except for the 15–10⁻¹ V group,



Fig. 3 Microbial resource acquisition strategies. 25 °C (**A**) and 15 °C (**B**). V_{angle} quantifies the relative P vs. N-acquiring enzyme activities; higher V_{angle} represents relatively higher P vs. N acquisition strategies. SM, no added viruses; DV, inactive viruses; V, active viruses with no dilution; 2^{-1} V, active viruses with twofold dilution; 10^{-1} V, active viruses with tenfold dilution; 25 and 15 signify different temperatures (25 °C and 15 °C) during the incubation. Different lowercase letters represent significant differences at p < 0.05.

which indicates that the DOM aromaticity increased by 22.5% in the 15-V group and 44.2% in the $15-2^{-1}$ V group (Fig. 4D). SUV260, as a proxy for DOM hydrophobicity, showed changes in treatment groups similar to SUV254 (Fig. S7).

Linking microbial properties to microbial metabolic activity and recalcitrant DOM components

A correlation analysis was conducted to assess whether the decreased V_{angle} (index of alleviated nutrient constraints by viral lysis) was related to microbial metabolic activity and DOM optical properties. All of the measured indexes (e.g., CO_2 emission, SUV254, SUV260, HIX) showed significant negative correlations with V_{angle} (Fig. S8). These associations could indicate that the nutrients released by viral lysis increased metabolic activity (shown as CO_2 emission) and enriched recalcitrant DOM components (represented by SUV254, SUV260 and HIX). Further, random forest analysis indicated that both V_{angle} and the DOM indexes (e.g., HIX, C1, C2, C3) could explain the differences in CO_2 emission, q CO_2 and CUE (Fig. 5).



Fig. 4 Changes of recalcitrant DOM components (mean \pm std dev) after incubation. SUV254 (A, B) represents the aromaticity degree of DOM, and HIX (C, D) indicates the humification degree of DOM. SM, no added viruses; DV, inactive viruses; V, active viruses with no dilution; 2^{-1} V, active viruses with twofold dilution; 10^{-1} V, active viruses with twofold dilution; 25 and 15 signify different temperatures (25 °C and 15 °C) during the incubation. Different lowercase letters represent significant differences at p < 0.05.

Bacterial communities were significantly altered by soil phages regardless of temperature (Adonis, p < 0.05; Fig. S9). Redundancy analysis was applied to explore the potential effects of edaphic properties (filtered by a variance inflation factor (VIF < 10)) on bacterial community composition (Fig. S10). Results showed that the selected factors explained 55.0% of the variance in the bacterial community. Most factors were related to DOM properties and indexes related to bacterial physiological traits (e.g., HIX, SUV260, C3, DOC, V_{angle} and V_{length}). Bacterial taxa associated with V_{angle} and the optical properties of DOM (including SUV254, SUV260 and HIX) were identified based on Spearman correlations (r > 0.5, p < 0.05; Fig. 6). The correlation network between bacterial taxa and various factors suggested that nearly all bacterial species were negatively associated with $V_{\mbox{\scriptsize angle}}$ whereas they were positively correlated with HIX, SUV254 and SUV260 (Table S1).

DISCUSSION

The three phases of microbial respiration affected by phages Three phases of microbial respiration were observed in both temperature regimes (Fig. 1). We attribute these phase dynamics to a tradeoff between the rate of bacterial mortality caused by viruses and the utilization of nutrients acquired by survivors, which is consistent with previous studies [17]. However, the phases differed with temperature. Compared with control groups (SM and DV) at 25 °C, CO₂ released from the soil with active phage additions was initially lower, and then increased before finally decreasing again (Fig. 1A). This CO₂-release pathway can be explained as follow: (1) Active phages lysed bacterial cells resulting in a decreased bacterial population and a subsequent reduction in CO₂ emission. (2) Released intracellular organic matter from viral lysing was reutilized by the surviving microbes, which stimulated bacterial activity and increased microbial respiration. (3) Finally, continued viral lysing resulted in a lower microbial abundance and decreased CO₂ fluxes. In the last phase, the number of survivors were too few to support higher CO₂ fluxes relative to the control groups [39]. Furthermore, at 15 °C, phases 2 and 3 contrasted with the phenomena at 25 °C (Fig. 1B). In phase 2 at 15 °C, only microbial respiration from the lowest concentration of active phages $(15-10^{-1} \text{ V})$ was higher than that in control groups. During phase 3, CO₂ emissions from all active phage groups were higher throughout the remainder of the incubation. These results from both temperature regimes are in



Fig. 5 Mean predictor importance (percent increased mean square error, MSE) for microbial metabolic indexes, besed on Random Forest analysis. The metabolic quotient (qCO₂), (**A**), carbon use efficiency (CUE), (**B**) and CO₂ emission (**C**). NO₃⁻, soil nitrate-N; NH₄ + , soil ammonium-N; AP available phosphorus, TP total phosphorus, SOC soil organic carbon, TN total nitrogen, DOC dissolved organic carbon, DON dissolved organic nitrogen, N/P the ratio of total nitrogen and total phosphorus, C/P the ratio of soil organic carbon and total phosphorus, C/N the ratio of soil organic carbon and total nitrogen, BG β -1,4-glucosidase, CBH cellobiohydrolase, NAG N-acetyl- β -D-glucosaminidase, LAP leucine aminopeptidase, AP/MBC means normalizing acid phosphatase activities to per MBC. Enzyme/MBC represents specific enzyme activities calculated by normalizing activities to units/mg MBC; Ecoenzymatic vector V_{length} (V_{length}, relative C:nutrient acquiring ratio); HIX, humification index of DOM; C1 (humic-like), C2 (quinone-like), C3 (protein-like), 3D-fluorescence components of DOM; MD, represents the relative molecular mass of DOM; SUV254, represents the aromatic degree of DOM; SUV260, RIC microbial biomass carbon, MBN microbial biomass nitrogen, MBP microbial biomass phosphorus, RICHNESS richness of bacterial community, SHANNON alpha diversity of bacterial community, PCoA1 PCoA2 are the two-dimensional PCoA ordinations of taxonomic composition for the bacterial community. **p* < 0.05; ***p* < 0.01.

full agreement with the viral shunt mechanism as interpreted by previous studies [17, 22].

The phase dynamics of microbial respiration are differentially affected by the temperature of the soil environment. The highest CO₂ emission for all active phage groups occurred in phase 2 at 25 °C, whereas the highest CO₂ emissions were somewhat delayed at 15 °C. This might results from the warmer temperature advancing the onset of the viral shunt and/or shortening the viral latent period [17]. Additionally, the difference in microbial respiration rates at different temperatures during phase 3 may be partially due to increased bacterial activity at 25 °C, which increased CO₂ emission in control groups and was not evident at 15 °C. This demonstrates that microbial activity increased at the higher temperature and without the influence of predators, thereby accelerating respiration rates [68-70]. Moreover, temperature could change interactions between phages and their hosts, such as altering the viral latent period, viral and host production rates, and burst size [29, 30].

Apart from temperature, cumulative CO_2 fluxes were significantly associated with the concentration of phages. Inoculating higher concentrations of active phages resulted in a slight increase or no obvious change in CO_2 fluxes, whereas adding a lower concentration of phages stimulated CO_2 emission (Fig. S2e, S2f). This infers that high inoculation rates caused increased microbial mortality, which subsequently decreased SOC mineralization, even though a more nutritionally-enriched environment is presumed relative to the low inoculation rate [39]. In addition, the Q_{10} values for active phage treatments appeared to play a role in maintaining Q_{10} values within the bounds of 1–3, which is in accordance with the global average Q_{10} value of 2.56 ± 0.86 [71]. However, the Q_{10} of SM and DV groups were significantly higher than those in active phages groups after the cultivation was stable, with Q_{10} values exceeding 3 at Day 98 (Fig. S3). These changes of Q_{10} among treatment groups suggest that soil phages could decrease the temperature sensitivity of CO_2 to warming, which could attenuate SOC mineralization affects due to global warming. However, the specific mechanisms warrant future investigation.

Lysate DOM alleviates microbial nutrient limitations

Previous studies revealed that viral lysate-derived DOM is relatively more bioavailable for microbial utilization, and can serve as an important substrate for microbial metabolism, particularly in harsh environments experiencing nutrient deprivation [72–75]. These studies found approximately 75% of lysate DOM was reused by surviving microbes, with 50–80% identified as bioavailable compounds, such as amino acids and carbohydrates [73, 74, 76]. Hence, the components of bioavailable DOM would have a higher affinity by microbes [34, 77]. This is consistent with our findings that microbial CUE increased in active phage groups (Fig. 2).

Recent studies report that soil microorganisms are often subject to P and N limitations at the global scale [78–80]. In pure culture systems, the release of phosphorus from host cells by viral lysing has been demonstrated, which suggests that viral lysates could be an important source of inorganic nutrients and markedly promote P regeneration [72]. Thus, to test our hypothesis that microbial nutrient limitation will be alleviated with active phages in the soil environment, the microbial physiological traits were assessed based on extracellular enzyme activity. The results supported our hypothesis by indicating that nutrient constraints (especially phosphorus limitation) were alleviated by lysate DOM, as demonstrated by a decrease in V_{angle} (Fig. 3).

Several previous studies found that microbial P limitation can have prominent impacts on soil carbon cycling [35, 81, 82]. For

1252



1253

Fig. 6 Correlation networks between measured variables (e.g., V_{angler} , SUV254, SUV260 and HIX) and associated bacterial taxa. 25 °C (A) and 15 °C (B). Different color of nodes represents different species (colored at the phylum level), and measured variables (labeled with characters). Red lines indicate positive interactions and green lines indicate negative interactions, at p < 0.05 and r > 0.5. The thickness of the lines indicates the strength of the correlations.

example, P limitation can stimulate more CO_2 released from soil owing to more enzymes being secreted by microbes in an attempt to obtain additional P from SOM [83–86]. Conversely, P limitation may constrain microbial activity, thereby increasing soil C stocks due to reduced SOC mineralization [87, 88]. The results of our study support the former premise, that alleviating phosphorus limitation by increasing lysate DOM increases microbial CUE. To our knowledge, this is the first report assessing nutrient restriction of microorganisms from the perspective of soil virus interactions with the microbial community.

Viruses contribute to accumulation of recalcitrant DOM in the soil environment

In our study, the alleviation of microbial P limitation appeared to promote SOC sequestration through accumulation of recalcitrant DOM (as indicated by higher SUV254, SUV260 and HIX; Fig. 4 and

S7). We found that V_{angle} showed a significant, negative correlation with several indexes of recalcitrant DOM. Moreover, the coupling of correlation and random forest analyses highlighted these DOM parameters as effective predictors of microbial metabolic efficiency and activity (Fig. 5 and S8). Transformation of microbial carbon into recalcitrant DOM components could represent an important soil carbon sink via a biological pump mechanism, with the concomitant re-utilization of the bioavailable intracellular organic matter increasing microbial activity, which is known as the viral shuttle mechanism [22, 23, 89, 90].

CDOM and FDOM are important fractions of environmental DOM (accounting for 20–70% of the overall DOM) that have been utilized to assess the quality and transformations of DOM [55, 57, 58]. Various studies investigated the relationships between viruses and DOM in aquatic environments and found that high viral lytic activity resulted in the production and accumulation of

FDOM, especially humic-like FDOM [73, 91–94]. These results are in agreement with our study that found a higher degree of humification and aromaticity in the accumulated DOM generated upon addition of active phages (Fig. 4, S6 and S7). Additionally, protists, phytoplankton and heterotrophic microbes are crucial factors regulating DOM dynamics [95–97].

Based on bacterial community composition and the correlation network between bacterial taxa and soil indexes, our results showed that phages have an appreciable impact on the bacterial community, and interactions between lysate DOM and bacteria play a strong role in determining the accumulation of recalcitrant DOM in soils (Fig. 6). Previous studies in oceans indicated a large accumulation of viral lysate DOM that is not readily utilized by microbes and therefore buildups in the recalcitrant DOM pool [23, 55, 60, 77, 98]. A possible explanation is that viral lysis suppresses the abundant microbial taxa and gives rare taxa time and space to propagate, which increases the utilization of bioavailable DOM and alters the microbial community [75, 99]. Additionally, viruses could change the microbial pathway for uptake of DOM compounds by introducing their genetic information into the host and reprogramming their metabolic pathways [16, 23, 100]. Our findings highlight that viral lysatederived DOM is not only metabolized and respired as CO₂, but other fractions are transformed into recalcitrant DOM that accumulates in the soil environment [22, 89]. This is the first study to provide direct evidence for the existence of the viral shuttle mechanism in soils.

CONCLUSION

This study contributes a more comprehensive understanding of viral effects on microbial respiration and DOM alteration in the soil environment. Microbial respiration displayed three distinct phases over the 98-day incubation period that were highly dependent on temperature and phage concentration. We ascribe these findings to the tradeoff between the rate of bacterial mortality caused by viruses and utilization of the nutrients released by the surviving microbial community. The viral lysatederived DOM can alleviate soil nutrient limitations, especially P limitation, as reflected by a lower V_{angle} . Furthermore, our results highlight that the bioavailable fraction of the lysatederived DOM can be re-utilized by the surviving microbes, whereas the recalcitrant DOM fraction (e.g., SUV254, SUV260 and HIX) can contribute to carbon sequestration in soils. This is the first study to demonstrate that both the viral shunt and viral shuttle mechanisms are active in the soil environment. This research provides new insights to guide future investigations that incorporate the role of viruses and the viral shuttle mechanism into nutrient cycling in soil environments.

DATA AVAILABILITY

The original sequence data were deposited into the National Center for Biotechnology Information Sequence Read Archive database under project ID PRJNA974558. Experimental data have been included as a supplement to this publication.

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AUTHOR CONTRIBUTIONS

Design of experiment: JX; soil sampling and microcosm incubation: DT, YW and HY; measurement of soil properties: DT and HS; statistical analysis: DT; visualization: DT; virting—original draft: DT and JX; writing—review and editing: JX, RA. Dahlgren and DT.

COMPETING INTERESTS

The authors declare no competing interests.

ADDITIONAL INFORMATION

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Correspondence and requests for materials should be addressed to Jianming Xu.

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1256