Optimization of extraction and estimation of viruses in silty freshwater sediments

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ABSTRACT: The present study focused on the optimization of procedures for the extraction of viruses from silty freshwater sediments for subsequent enumeration. Viral abundance in 2 different shallow backwater systems of the River Danube (Austria) ranged from $1.45 \times 10^9$ to $9.58 \times 10^9$ particles ml$^{-1}$ wet sediment. The highest virus yields from the bulk of the sediments were obtained by 1 min sonication ($3 \times 20$ s intervals, with 10 s interruptions). An increase in sonication time of up to 5 min decreased viral counts by an average of 15%. Since dissolved DNA within sediment samples could bind to the nucleic acid stain and thereby inflate viral estimates, sediment samples are often treated with DNase before the staining procedure. Moreover, they are usually centrifuged and diluted to a high extent in order to avoid interference of particulate material with virus counting. Centrifugation led to a reduction of viral numbers by 2 to 36% compared to untreated samples and did not reduce the background fluorescence; thus counting of viruses was not facilitated. Diluting 2000× with Milli-Q water always provided an average of 19% lower viral numbers than diluting 4000×. Treatment with DNase had no significant effect on virus counting, with viral numbers in untreated samples being on average 96% of those in DNase-containing samples. Additionally, 2 different nucleic acid stains were compared—viruses stained with SYBR Gold fluoresced brighter than those stained with SYBR Green I and fluorescence lasted longer, while background fluorescence was reduced sufficiently, thus facilitating virus counting. Viral numbers using SYBR Gold were on average twice of those obtained with SYBR Green I. The mean efficiency of virus extraction was 88.8% using the protocol outlined in this paper, and was thus slightly higher than that obtained in previous sediment investigations.

KEY WORDS: Virus extraction · Virus counting · Epifluorescence microscopy · Silty freshwater sediment

INTRODUCTION

In freshwater and marine environments, viruses are the most numerous component of plankton, with abundances ranging from $10^4$ to $10^8$ viruses ml$^{-1}$ (Wommack & Colwell 2000). Through infection and lysis, viruses exert a direct effect on the mortality of heterotrophic bacterioplankton, cyanobacteria and phytoplankton (Fuhrman 1999, Wilhelm & Suttle 1999, Wommack & Colwell 2000). They also play a role in biogeochemical cycling and control of host-species diversity (Fuhrman 1999, Wilhelm & Suttle 1999). The study of viruses in aquatic systems requires a reliable method for estimating the levels of these biological entities. The available methods for the determination of viral abundance in aquatic systems include counting by transmission electron microscopy (TEM) (Bergh et al. 1989, Bersheim et al. 1990), by flow cytometry (Chen et al. 2001, Brussaard 2004) and epifluorescence microscopy (EFM) (Hara et al. 1991, Hennes & Suttle 1995, Xenopoulos & Bird 1997, Noble & Fuhrman 1998). The latter technique allows rapid enumeration, avoiding the use of expensive equipment necessary for estimation by TEM or flow cytometry.

Viral abundance in sediment systems is 10- to 1000-fold higher than in the overlying water column (e.g. Maranger & Bird 1996, Steward et al. 1996, Drake et al. 1998, Hewson et al. 2001, Lawrence et al. 2002, Fischer et al. 2003, Hewson & Fuhrman 2003). Studies using EFM to estimate viruses in sediments applied a variety of protocols, without testing whether the procedures carried out before enumeration (e.g. virus dislodgement from particles, centrifugation, dilution) affect viral counts (Danovaro & Serresi 2000, Ricciardi-Rigault et al. 2000, Danovaro et al. 2001, 2002, Fischer et al. 2003, 2004, Hewson & Fuhrman 2003, Hewson et al. 2003, Middelboe et al. 2003, Mei & Danovaro 2004). A single study aimed at designing a protocol for determination of benthic viral numbers was done only on marine sediments (Danovaro et al. 2001); freshwater systems were completely neglected. An added complication in the enumeration of naturally occurring benthic viral populations is the uncertainty as to whether differences in extractability of viruses among different sediment types exist. Hence, it would be useful to test whether the protocol designed for virus enumeration in marine sediments is adaptable to freshwater sediments, and to which extent improvements of the methodology are required.

Within the frame of a long-term study on the carbon flow between the compartments of the microbial loop in the oxygenated layer of silty sediments of backwater systems (Fischer et al. 2003, 2004, Wielschtnig et al. 2003a,b), it was of primordial importance to consider the viral loop. Protocols on virus extraction and estimation in silty freshwater sediments are so far lacking, yet silty sediments are a frequent feature of most aquatic habitats with moderate or little water disturbance such as deep lakes, sheltered lakes, alluvial forests and oxbow lakes (Fenchel et al. 1998, Wetzel 2001). The present study was executed to optimize the EFM enumeration of viruses in silty freshwater sediments. We focused our attention on (1) virus dislodgement from particles (the first step in any analysis of benthic viruses is effective extraction from the sediment) (2) effect of centrifugation of samples (to remove coarse sediment particles and facilitate virus counting), (3) extent of dilution of sediment samples necessary to avoid interference of particulate material with virus counting, (4) the possible falsification of virus counts caused by extracellular DNA, and (5) the comparison of the commonly used nucleic acid stain SYBR Green I with SYBR Gold.

MATERIALS AND METHODS

Study site and sample collection. Samples were taken from the Kühwörther Wasser and the Alte Donau, 2 shallow (mean water depth 1 and 3.5 m, respectively) backwater systems of the River Danube, Austria. The investigated sites were characterized by muddy sediments consisting of 2 to 6% medium to coarse sand (≥250 µm), 6 to 14% fine sand (125 to <250 µm), 25 to 33% very fine sand (63 to <125 µm), 42 to 63% silt (2 to <63 µm), and 5% clay (<2 µm). Sediment cores were collected between August 2000 and May 2001 by hand-coring using Plexiglas tubes (inner diameter 5.8 cm, penetration depth 15 to 20 cm). The top 0.5 cm of the sediment of 5 cores was pooled by gentle mixing with a stirring magnet. Sediment parameters (bulk density, water content, organic matter content) were determined as described by Kirschner & Velimirov (1999). Porosity was calculated from the water content and bulk density of the sediment. Water content varied from 73 to 86%, porosity from 0.77 to 0.87, and organic matter content from 16.7 to 21.8 mg ml⁻¹ fresh weight. Since the sampling sites were similar as to granulometry, bulk density, porosity, water content and organic matter content, the results have been combined.

To establish a protocol for virus counting in silty freshwater sediments, aliquots (1 g) of the sediment slurry were diluted with 30 ml Milli-Q water, and fixed with electron microscopy-grade glutaraldehyde (3% final concentration). Samples were kept at 4°C in the dark until processing, which was always performed within 24 h. Since storage of aldehyde-fixed samples at 4°C reduces viral numbers (Xenopoulos & Bird 1997, Marie et al. 1999, Danovaro et al. 2001), even within the initial 24 h period of fixation (Brussaard 2004, Wen et al. 2004), the viral counts in the present study may be underestimates. All samples that are compared were processed simultaneously. They were treated with 0.02 µm-filtered sodium-tetrapyrophosphate for at least 20 min. A low final concentration of 5 mM was used in order to minimize the extraction of humic acids (Maranger & Bird 1996). Moreover, the use of high pyrophosphate concentrations makes the optical field opalescent under the EFM, thus complicating virus counting (Danovaro et al. 2001).

Comparison of SYBR Green I and SYBR Gold viral counts. The original SYBR Green I and SYBR Gold stock solutions (10 000× concentrate in DMSO, Molecular Probes) were diluted with autoclaved Milli-Q water to final concentrations of 1 and 0.25%. Sediment
samples were sonicated for 1 min (3 x 20 s, with 10 s interruptions) at 70 W using a Branson Sonifier 450 (Branson Ultrasonics Corporation), and diluted (4000 ×) with Milli-Q water. Aliquots (1 ml) were then filtered through 0.02 µm pore-size Al₂O₃ Anodisc membrane filters backed by a 0.2 µm pore-size cellulose nitrate filter at approximately 20 kPa vacuum. The Anodisc membranes were dried and laid sample-side up on a drop of the staining solution for 15 min in the dark (Noble & Fuhrman 1998, Chen et al. 2001, Noble 2001). The filters were then dried and mounted on a glass slide with a drop of Citifluor (Glycerol/PBS solution-AF1; Agar Scientific) containing 0.1% p-phenylene-diamine (made freshly from a frozen 10% aqueous stock; Sigma Aldrich Chemie). All preparations were done under subdued light. Filters were examined at a magnification of 1250× with a Leica DMRB microscope (Leica) equipped with an HBO 50 W mercury lamp (excitation wavelength 450 to 490 nm, cut-off filter 515 nm). Total counts of 30 to 50 randomly selected fields usually exceeded 200 viruses per subsample. The inventory of viruses from bulk sediment included particle-adsorbed viruses as well as viruses in pore water. Viral numbers are expressed as ‘particles ml⁻¹ wet sediment’.

**Virus dislodgement.** Sediment samples were sonicated for 0, 1, 2, 3, 4, and 5 min at 70 W. To prevent overheating, sonication was at intervals of 20 s with 10 s interruptions. Additionally, after samples were treated 3 x for 20 s, sonication was interrupted for 5 min, during which time the samples were kept on ice. After further dilution, viral numbers were estimated as described in the foregoing subsection.

**Centrifugation time.** After sonication, samples were centrifuged (Sigma Laboratory Centrifuge 3K30, fixed-angle rotor No. 12156, Sigma Laborzentrifugen) under different conditions: at 1000 × g for 5 and 3 min; at 500 and 200 × g for 5 min; at 400 and 800 × g for 1 min (Gerba et al. 1977, LaBelle & Gerba 1982, Tsai et al. 1983, Chalapati Rao et al. 1984, Hu 1998, Danovaro et al. 2001, 2002). The supernatant was further diluted, and viruses in the supernatant were counted.

**Dilution of sediment samples.** Sediment samples were sonicated and subsequently diluted 2000 × or 4000 × with Milli-Q water, before virus counting.

**DNase-test.** Aliquots (1 g) of the sediment slurry were diluted, treated with pyrophosphate and sonicated; 250 Kunitz units of DNase I from bovine pankreas (Sigma) were added to 1 ml aliquots and incubated for 30 min at room temperature (Suttle 1993). Additional aliquots (1 ml) without DNase were incubated under the same conditions and served as controls. After fixation with glutaraldehyde, sonication and further dilution, the viruses in samples containing DNase and in untreated samples were counted. To confirm the DNase-digested DNA in water from the Kühwörther Wasser, its activity was assayed following the protocol of Drake et al. (1998): 250 Kunitz units of DNase were added to 40 µg ml⁻¹ DNA (Type I, ‘highly polymerized’ from calf thymus, Sigma) in water samples. The temporal change in absorbance at 260 nm within 10 min, monitored using a Hitachi U-2000 spectrophotometer (Hitachi), was 0.002 min⁻¹ (data not shown).

**Virus extraction efficiency.** A known amount of viruses concentrated from the water column of the study sites by ultrafiltration (Noble & Fuhrman 1999) was added to samples (1 g) of the sediment slurry (= virus-amended samples), and shaken at 250 rpm for 30 min at room temperature to allow adsorption of viruses to sediment particles (Gerba et al. 1977, LaBelle & Gerba 1979, Tsai et al. 1983). After fixation with glutaraldehyde, treatment with pyrophosphate, sonication and further dilution, viruses were counted by EFM. Extraction efficiency was determined by dividing virus counts in virus-amended sediment by the sum of counts in unamended sediment samples plus counts of the virus ultrafiltration-concentrate (= control), and multiplying this ratio by 100.

**Statistical analysis.** For the computation of correlations between data, we used Spearman’s rho correlation. For analysis of variance, the nonparametric Mann-Whitney U-test was applied according to Zar (1974). In all statistical analyses, probabilities of ≤0.05 and ≤0.01 were considered significant and highly significant, respectively. We used SPSS version 10.0.8 (SPSS) software.

**RESULTS AND DISCUSSION**

**Viral abundance**

Following our protocol (see Fig. 6), the viral numbers in the sediment systems investigated varied from 1.45 × 10⁹ to 9.58 × 10⁹ particles ml⁻¹ wet sediment, and were thus in the same order of magnitude as those previously reported for sediments of aquatic environments, including marine (e.g. Danovaro & Serresi 2000, Hewson et al. 2001, Danovaro et al. 2002) and freshwater (e.g. Maranger & Bird 1996, Ricciardi-Rigault et al. 2000, Fischer et al. 2003, 2004) sediments.

**Comparison of SYBR Green I and SYBR Gold viral counts**

SYBR Gold has so far been used to stain viruses in samples from the water column (McDaniel et al. 2001, Yager et al. 2001, Williamson et al. 2002, Anesio et al.
2004, Breitbart et al. 2004, Brussard 2004, Lisle & Priscu 2004, Wen et al. 2004) and sediment (Fischer et al. 2003, 2004) of aquatic systems, in soil samples (Williamson et al. 2003) and human feces (Breitbart et al. 2003). Direct comparison of staining viruses with SYBR Gold and the commonly used dye SYBR Green I has been made in 4 studies: (1) Chen et al. (2001) reported that even when no antifading-solution was used, the fluorescence of SYBR Gold-stained viruses in marine water samples was stable for more than 2 min under EFM, while the SYBR Green I signal faded within 30 s. Using flow cytometry to enumerate Cyanophage P49, the mean fluorescence per P49 virus determined with SYBR Gold was about 2 times higher than that with SYBR Green I. However, Chen et al. (2001) did not directly compare viral counts with SYBR Gold and SYBR Green I by EFM in their investigation. 

(2) This also applies to the study of Brussaard (2004), who used flow cytometry to examine a variety of different pelagic viruses, including representatives of phytoplankton viruses, cyanobacteriophages, coliphages, marine bacteriophages and natural mixed marine virus communities stained with SYBR Green I and SYBR Gold. The use of SYBR Gold resulted in slightly lower green fluorescence signals of the stained viruses. No significant effect was recorded for the total virus counts when all tested viruses were taken into account, but for some of the individual viruses and a natural sample, staining with SYBR Gold resulted in reduced total counts compared to staining with SYBR Green I. 

(3) Using EFM, Wen et al. (2004) found that staining with SYBR Green I and SYBR Gold yielded apparently identical numbers of pelagic viruses. (4) The only attempt to compare the ability of different nucleic acid dyes to stain viruses in sediment samples was by Hewson & Fuhrman (2003), who counted marine benthic viruses by EFM using SYBR Green I, SYBR Green II and SYBR Gold. They made only a brief comment that the emission color of SYBR Green II and SYBR Gold was close to that of autofluorescent humic materials in

Fig. 1. SYBR Gold (0.25% final concentration)-stained sample from silty freshwater sediment filtered onto a 0.02 µm pore-size membrane. The larger bright yellow-green particles are considered to be bacteria (B), the smaller particles viruses (V). Bacteria and viruses are easily distinguishable from humic substances (HU). (Leica DMRB microscope and Kappa CF 15/4 MCC camera)
sediiments, and could not be used easily for benthic virus counts. Unfortunately, Hewson & Fuhrman (2003) did not detail the results of virus estimates with the different stains. For routine virus counting in marine sediments, they used SYBR Green I.

In the present study, viruses in sediment samples stained with SYBR Gold yielded a bright and stable yellow-green fluorescence under the EFM (Fig. 1). The brightness of the SYBR Gold signal varied, likely due to different viral genome sizes among viruses (Chen et al. 2001). Viral particles could be easily distinguished from bacteria by their relative size and shape. In comparison with SYBR Green I, the fluorescence of SYBR Gold-stained viruses was not only brighter, but also lasted longer, while background fluorescence was reduced sufficiently, thus facilitating virus counting.

For dilution of the stain stock solutions, we used the standard protocol of Noble & Fuhrman (1998) and Noble (2001), who recommended increasing the concentration of SYBR Green I usually used (2.5 × 10⁻³ dilution of the stock solution, i.e. final concentration of 0.25%) for samples of specific environments, i.e. when the viruses did not stain brightly. We diluted both the stock solution of SYBR Green I and SYBR Gold to a final concentration of 1 and 0.25%, respectively, and compared the resultant viral counts (Fig. 2). SYBR Green I-counts at different dye concentrations did not differ significantly (p > 0.90, n = 9), and the precision of the counts was similar, with an average coefficient of variation (CV) for triplicate samples of 30 ± 12% versus 26 ± 10%. In contrast, counts for different dye concentra-
tions of SYBR Gold differed significantly (p < 0.05, n = 9). Viral numbers estimated after staining with the 1% SYBR Gold-solution were on average only 56 ± 5% of those obtained with the 0.25% solution. The results of the 2 concentrations were correlated (p < 0.05, n = 9). At first sight, it seems paradoxical that a higher concentration of the SYBR Gold dye yielded lower viral counts; because of the brighter fluorescence signal of the viruses, one would expect more accurate counts. However, a 1% final concentration of SYBR Gold also led to very brightly stained bacterial cells, and the resultant fluorescent halos around these cells as well as the higher background fluorescence obtained with samples stained with 1% SYBR Gold probably masked the fluorescence of the stained viruses, resulting (at least partly) in lower counts. Brussard (2004) also reported reduced viral abundance at high dye concentrations (0.005 to 0.01% dilution of commercial stock solution of SYBR Green I for counting viruses by flow cytometry).

Direct comparison of SYBR Green I-virus counts with SYBR Gold-counts revealed that the results of the 2 methods were correlated (p < 0.05, n = 9). However, staining with SYBR Gold yielded significantly higher viral numbers (p < 0.05, n = 9), implying that staining viruses with SYBR Green I led to underestimation of benthic viral abundance. Virus estimates using SYBR Gold at 1 and 0.25% final concentrations were, respectively, 1.5 ± 0.20 and 2.7 ± 0.45 times higher than those obtained with SYBR Green I. Also, SYBR Gold-counts were more precise, with an average CV of 16 ± 3% for the 1% solution and 8 ± 1% for the 0.25% solution, respectively, compared to SYBR Green I CVs of 30 ± 12 and 26 ± 10%.

As mentioned at the beginning of this subsection, Wen et al. (2004) found no difference in pelagic viral counts using SYBR Green I and SYBR Gold. However, they used a very high concentration of both dyes (4% final concentration of original stock solution), and the present investigation has shown that the difference between viral counts using SYBR Green I and SYBR Gold is lower when the dyes are diluted to a final concentration of only 1% compared to 0.25%.

Based on our results, we recommend the use of SYBR Gold nucleic acid stain diluted to a final concentration of 0.25% for routine virus estimation in sediments. Because of the high amount of detritus and the presence of sediment particles, virus counting in sediment samples is more difficult than counting of pelagic viruses. The
stability and brightness of SYBR Gold fluorescence combined with the reduced background fluorescence represents a distinct advantage for the enumeration of benthic viruses by EFM. Moreover, SYBR Gold is a sensitive fluorescence stain for detecting double- and single-stranded DNA and RNA, whereas SYBR Green I is a fluorochrome mainly useful for detecting double-stranded DNA. The manufacturer (Molecular Probes) claims that it can also be used to detect single-stranded DNA and RNA; but its sensitivity is lower than for double-stranded DNA. This may have partly contributed to the difference between the SYBR Green I- and SYBR Gold counts. Another advantage of SYBR Gold is that the price is only half that of SYBR Green I.

Virus dislodgement

The use of pyrophosphate as an eluent to dislodge viruses from particles is well established (e.g. Lawrence et al. 2002, Hewson & Fuhrman 2003, Hewson et al. 2003), and in some studies sediment samples have been additionally sonicated (e.g. Maranger & Bird 1996, Danovaro & Serresi 2000, Danovaro et al. 2001, 2002, Mei & Danovaro 2004). Middelboe et al. (2003) reported that the addition of sodium pyrophosphate significantly increased dislodgement of viruses from marine sediment relative to untreated samples, and that the application of sonication treatment further increased the release of viruses. The efficiency of their combined pyrophosphate–sonication treatment was 65 to 78% of the total extractable viruses.

Despite the fact that investigations on viruses in sediments are becoming more frequent, to our knowledge only a single study has been conducted to determine the optimal sonication time: Danovaro et al. (2001) sonicated muddy and sandy marine sediments for 0, 1, 3, 8 and 15 min. The highest viral counts were obtained after 3 min sonication, and these values were significantly higher than values obtained without sonication. A further increase in sonication time decreased viral counts. However, virus sorption usually increases with increasing cation concentration in solution, particularly in the presence of divalent cations (Schijven & Hassanzadeh 2000, Jin & Flury 2002). Therefore, the protocol design for the extraction of benthic viruses from marine sediments may not be applicable to freshwater sediments. The present study thus aimed at detecting the optimal sonication time for the extraction of viruses from silty freshwater sediments. Highest viral counts were obtained after 1 min sonication, being on average 19 ± 8% higher than without sonication (Fig. 3). This difference was statistically significant (p < 0.03, n = 6). Viral counts after 2 and 3 min sonication were on average 94 ± 10% and 89 ± 9% of counts obtained after sonication for 1 min (p > 0.20, n = 6). Further increases in sonication time further decreased virus estimates, and sonication for 4 min or longer reduced virus counts significantly (p < 0.05, n = 6), with an average of 79 ± 11% of viral numbers detected after 1 min sonication. The CV was always low, ranging between 3 and 14%. From the trend observed during the experiments, we believed that higher viral counts after sonication >5 min would be very improbable; therefore we aborted the experiments, and a sonication time of 1 min was used in all further experiments.

When using sonication to dislodge viruses from sediment particles, it must be taken into account that this procedure may cause disruption of virus-infected bacterial cells, releasing mature viral particles, and thus falsifying the number of free viruses originally present in the sample. Assuming a production of $22.2 \times 10^7$ viruses ml$^{-1}$ h$^{-1}$ (maximum value for viral production measured previously in one of the backwater systems investigated; Fischer et al. 2003), we calculated that even if all viruses theoretically produced within 1 h were to be set free from infected cells during the sonication step, viral abundance would on average be overestimated by a factor of only 1.029 (range 1.021 to 1.038). Disruption of virus-infected bacteria
did thus not drastically falsify the numbers of free viruses counted in the present study.

**Centrifugation time**

The presence of sediment particles on the Anodisc filters complicates virus counting. Therefore, sediment samples have usually been centrifuged prior to virus estimation by EFM (e.g. Drake et al. 1998, Danovaro et al. 2001, 2002, Hewson et al. 2001, 2003, Lawrence et al. 2002, Hewson & Fuhrman 2003, Middelboe et al. 2003), under the assumption that this process does not remove viruses, and hence does not falsify viral counts. However, none of these studies tested whether this assumption is valid. Our experiments revealed that centrifugation of sediment samples led to an average of 15 ± 14% lower viral counts compared to untreated samples (Fig. 4). The difference in viral abundance was significant (p < 0.05, n = 3) in 2 cases (1000 × g for 30 min: 36% lower counts; and 200 × g for 5 min: 32% lower counts). Only centrifugation of samples at 800 × g for 1 min, as carried out by Danovaro et al. (2001, 2002), gave values similar to those of untreated samples. An average of 98% of viruses counted in untreated samples were present after centrifugation. However, in subsequent experiments we did not centrifuge samples prior to virus counting because centrifugation at 800 × g for 1 min did not reduce the background fluorescence, and thus virus counting was not facilitated. Moreover, counts of untreated samples were slightly more precise than counts of centrifuged samples, as indicated by a CV of 11% (average CV of all untreated samples of the different centrifugation experiments) versus 14% for centrifuged samples. The opposite trend would be expected since untreated samples should be more heterogenous (i.e. contain more particles) than centrifuged samples. We have no explanation for the higher precision of counts in untreated samples.

**Dilution of sediment samples**

To avoid interference of particulate material with virus counting, and to obtain an optimal number of viruses per microscopic field (15 to 30), it is necessary to dilute sediment samples to a high extent. We diluted samples by 2000× and 4000×, and compared viral counts. The 2000× dilution always produced significantly lower viral numbers (p < 0.05, n = 6) than the 4000× dilution (data not shown). Virus estimates after 2000× dilution (4.1 × 10⁹ to 5.3 × 10⁹ particles ml⁻¹) were on average only 81 ± 13% of those for 4000× diluted samples (5.0 × 10⁹ to 7.4 × 10⁹ particles ml⁻¹). The precision of the 4000× dilution (CV 12%) was less than that of the 2000× dilution (CV 6%). However, since naturally occurring viruses were being extracted, the best extraction was considered to be achieved by the method giving the highest yield of viruses. In subsequent experiments, we thus diluted sediment samples 4000×.

**DNase test**

It is conceivable that dissolved DNA within sediment samples could bind to SYBR Gold and thereby inflate viral estimates. In order to eliminate uncertainties in virus counting due to extracellular DNA interference, we tested the effect of DNase treatment on silty freshwater sediment samples. Virus estimates for untreated samples were 96 ± 4% of those for samples to which DNase had been added (Fig. 5). The results for the 2 protocols were positively correlated (p < 0.01, n = 12). Since the difference was not statistically significant (p > 0.60, n = 12), we omitted the addition of DNase in further experiments. Drake et al. (1998), who enumerated viruses in the pore water of marine sediments, obtained similar results. In contrast, Danovaro et al. (2001) detected significantly higher viral counts after DNase treatment of marine sandy sediments, and suggested that virus counting was facilitated in these samples through the reduction in fluorescence noise caused by extracellular DNA bound to the fluorochrome. Nevertheless, DNase should be used with caution, since it has repeatedly been reported to affect viral particles, consequently falsifying viral counts: (1) Using TEM,
Viral counts of virus-amended samples (i.e. sediment samples to which a known amount of viruses concentrated from the water column had been added and then extracted following the protocol of the present study) did not differ significantly from the control (i.e. the sum of counts in untreated sediment samples plus counts of the virus concentrate) (p > 0.10, n = 6); mean efficiency of virus extraction was 88.8 ± 11.8% (range 67.5 to 101.6%). In comparison, the treatment of sediment samples with pyrophosphate and sonication in other studies led to a virus extraction efficiency of 60 to 78%, which was tested by washing the sediment samples 2 or 3 times with Milli-Q water and virus-free seawater, respectively, after initial elution (Danovaro et al. 2001, Middelboe et al. 2003). A virus extraction efficiency of 85% in the initial elution step was achieved by treating sediment samples with an elution buffer consisting of pyrophosphate, EDTA and formaldehyde alone (without sonication), followed by 3 successive washes with the elution buffer (Hewson & Fuhrman 2003). However, the effect of sonication was not tested.

Extraction efficiency in the present study was higher than that in previous investigations, but nevertheless 11.2% of the added viruses were not extractable, probably because they were too firmly attached to sediment particles. Assuming that the extraction efficiency of the added and subsequently adsorbed viruses is a realistic reflection of the adsorption behavior of viruses in natural sediment samples, we therefore underestimated the benthic viral population by some 11%.

Thus, our protocol allows assessment of the fraction of ‘non-extractable’ viruses, whereas that used by other authors (washing the sediment samples several times, see above) does not provide any information on the virus fraction that may still remain resistant to the extraction procedure even after several washing steps.

A schematic overview of the protocol we recommend for counting viruses in silty freshwater sediments is given in Fig. 6.

Virus extraction efficiency

Viral counts of virus-amended samples (i.e. sediment samples to which a known amount of viruses concentrated from the water column had been added and then extracted following the protocol of the present study) did not differ significantly from the control (i.e. the sum of counts in untreated sediment samples plus counts of the virus concentrate) (p > 0.10, n = 6); mean efficiency of virus extraction was 88.8 ± 11.8% (range 67.5 to 101.6%). In comparison, the treatment of sediment samples with pyrophosphate and sonication in other studies led to a virus extraction efficiency of 60 to 78%, which was tested by washing the sediment samples 2 or 3 times with Milli-Q water and virus-free seawater, respectively, after initial elution (Danovaro et al. 2001, Middelboe et al. 2003). A virus extraction efficiency of 85% in the initial elution step was achieved by treating sediment samples with an elution buffer consisting of pyrophosphate, EDTA and formaldehyde alone (without sonication), followed by 3 successive washes with the elution buffer (Hewson & Fuhrman 2003). However, the effect of sonication was not tested.

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