Stuck in the mud: suspended sediments as a key issue for survival of chrysomonad flagellates

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ABSTRACT: The effect of suspended fine sediments on chrysomonad flagellates was investigated for ‘Spumella-like’ flagellates in laboratory studies and exemplarily for a flagellate community (with a focus on chrysomonads) originating from the oligomesotrophic Lake Mondsee, Austria, using different clay minerals and silicate beads. In the community experiment, the abundances of Spumella-like flagellates decreased significantly after introduction of suspended clays, but on the community level suspended clays did not negatively affect flagellate abundance. In order to understand this taxon-specific response, the influence of different clay characteristics, specifically of particle concentration and size, was investigated for Spumella-like flagellates in laboratory studies. We used ultra-microbacteria, i.e. typical bacterioplankton of Lake Mondsee, as food. For comparison we also investigated growth rates of the Spumella-like flagellates feeding on the large bacterium Listonella pelagia. These experiments confirmed that at bacterial abundances realized in oligotrophic and mesotrophic lakes, the flagellates are severely food limited. The presence of suspended sediments generally decreased the growth at any food concentration tested. This decrease was reflected in shifts of the growth kinetic parameters, i.e. in a positive correlation between the half-saturation constant and the threshold food concentration and a negative correlation between the maximal growth rate and the suspended sediment concentration. The clearance rates strongly decreased when small and large particles were present, but were only slightly affected for intermediate-sized particles. We assume that small particles in the size range of ingestible bacteria interfere with the feeding process and cause lower clearance rates, while intermediate-sized particles may serve as substrate for the attachment of flagellates and to subsequently optimize their clearance rates.

KEY WORDS: Turbidity · Growth rates · Microbial food web · Grazing · Chrysophytes/Chrysomonads · HNF · Protists · Turbulence

INTRODUCTION

Despite the general importance of suspended sediments in freshwater systems, specifically after precipitation or flood events (e.g. Serruya 1974, Threlkeld 1986, Lenzi & Marchi 2000), their interference with the feeding and growth of microbial components is severely understudied and largely unknown. In lakes, suspended sediment concentrations are typically in the range of several milligrams per liter, but may increase several orders of magnitude within hours following precipitation (Dokulil 1979, Knowlton & Jones 1995). In streams, suspended sediment concentrations are generally high, and, specifically, in many lowland streams average concentrations are between 100 and 20,000 mg l⁻¹ (Arruda et al. 1983, Eisma 1993, Brondson & Naden 2000, Orton & Kineke 2001). Despite the ubiquity of suspended sediments and the presumably great effect on microorganisms and particularly on protists, these interrelationships have not been thoroughly studied.

In the present study, we investigated the effect of suspended sediments on bacterivorous protists, for a common and generally abundant flagellate group, i.e. for colorless chrysophytes, often a dominant component of the microbial food web. Finlay & Esteban (1998) stated that heterotrophic heterokonts, and specifically chrysomonads, are ‘probably the most...
abundant of the heterotrophic flagellates in the plankton (e.g. *Paraphysomonas* and ‘*Spumella-like*’ flagellates) and the most important [...] grazers of bacteria-sized microorganisms’. Annually, on average, 20 to 50% of the pelagic heterotrophic nanoflagellate (HNF) biomass in freshwaters is formed by small heterokont taxa, mainly colorless chrysophytes (i.e. chrysomonads) and bicosoecids (Salbrechter & Arndt 1994, Carrias et al. 1998, Arndt et al. 2000).

Based on theoretical assumptions and the few available experimental studies (Englund et al. 1993, Jack & Gilbert 1993, Boenigk & Novarino 2004), suspended sediments are expected to affect pelagic protists in several ways. On the one hand, suspended sediments may decrease feeding efficiency and protist population growth, due to interference with the feeding process and/or mechanical damage to fragile protist cells especially in turbid environments (Boenigk & Novarino 2004). Interference with feeding can specifically be expected for interception-feeding pelagic flagellates (e.g. chrysomonads). On the other hand, suspended solids may serve as a substratum for the attachment of chrysomonad flagellates, thus indirectly triggering their feeding efficiency (Christensen-Dalsgaard & Fenchel 2003, Kierboe et al. 2004). These effects of suspended solids are further modified as dissolved substances attach easily to suspended sediment particles. Thus, the bioavailability of nutritive as well as harmful substances is modified (Boenigk et al. 2005a), which may not only affect the flagellates, but also their bacterial prey (Lind et al. 1997). Besides these effects on nutrient availability, sediment particles may provide an indirect grazing protection for bacteria, as bacterivorous predators confronted with high abundances of indigestible nanoparticles may fail to ingest bacteria at high rates (England et al. 1993, Jack & Gilbert 1993). The potential response of chrysomonad flagellates to suspended sediments is therefore expected to be complex. Our understanding of these interactions, however, is severely limited. The lack of experimental studies is probably due to the fact that suspended sediments have been largely disregarded as an important factor for heterotrophic microorganisms. The particle characteristics that are responsible for shifts in the protist population response are unknown, and the ecological significance of suspended particles for heterotrophic flagellate taxa is largely restricted to rough assumptions (Boenigk & Novarino 2004).

We focused our investigations on chrysomonad flagellates for 2 reasons: (1) colorless chrysophytes are among the most dominant bacterivores in many aquatic habitats (see above) and (2) despite the positive effect of suspended sediments on chrysomonad flagellates, as reported by Boenigk & Novarino (2004), these flagellates are theoretically expected to be negatively affected by suspended sediments, thus demanding further in-depth investigations. We hypothesized that suspended sediments negatively affect chrysomonad flagellates and that the strength of this effect depends on particle concentration and size characteristics. In order to test these hypotheses, we investigated the effect of suspended particles on pelagic chrysophytes using a natural community taken from Lake Mondsee, Austria, and a set of specific laboratory experiments testing the effect of suspended sediments on the growth of ‘*Spumella-like*’ flagellates.

**MATERIALS AND METHODS**

**Origin of strains, culture conditions and origin of clays.** Single-cell, colorless chrysophytes are common and abundant in the plankton of many lakes and streams (see ‘Introduction’). Experiments were conducted using ‘*Spumella-like*’ flagellates (*Spumella* sp., Chrysophyceae, Chromulinaceae) as model organisms. The flagellate strain JBC07 (61.1 ± 24.8 µm³) was isolated from the eutrophic shallow lake Tai Hu, China, and the flagellate strain JBM10 (69.3 ± 17.8 µm³) was isolated from a pond located in Mondsee, Austria (Boenigk et al. 2005b). Both flagellate strains are affiliated with the C3 cluster of the Chrysophyceae and are common freshwater flagellates (Boenigk et al. 2005b). They are able to predate on ultramicrobacteria (Boenigk et al. 2005c). The axenic protist strains were cultured in NSY basal medium (Hahn et al. 2003) on a diet of the heat-killed bacterial strain *Listonella pelagia* CB5 and were transferred weekly to fresh medium. Cultures were kept at 16°C in permanent light (8.9 µE m⁻² s⁻¹).

The large bacterial strain *Listonella pelagia* CB5 (0.38 ± 0.2 µm³) and the ultramicrobacterial strain MWH-MoNR1 (0.04 ± 0.017 µm³; closest known relative *Clavibacter michiganensis* [Microbacteriaceae]) were used as living food bacteria in the experiments (Hahn & Höfle 1998, Hahn 2003). The bacterial strains were cultured in NSY basal medium supplemented with equal amounts (1 g each) of nutrient broth, soytoine and yeast extract (Hahn et al. 2003). Before the experiments were started, bacteria were repeatedly centrifuged for 15 min at 5000 × g and washed with NSY basal media to remove traces of DOM. Microscopic analysis showed that the flagellate strain MWH-MoNR1, which was used in the suspended sediment experiments, does not aggregate or attach to the suspended particles.

All flagellate and bacterial strains originate from oligomesotrophic lakes and represent common freshwater organisms.
Two natural clays and artificial silicate particles were used in the experiments: a kaolinite-dominated clay (gray clay: 84% kaolinite, 3% other clay minerals, zeta potential — 47 mV) and a kaolinite/montmorillonite clay (yellow clay: 40% montmorillonite, ~60% kaolinite; zeta potential — 26 mV), both originating from a clay pit in the Eifel, Germany (Fig. 1); Kaolinite, i.e. the end-product of weathering of silicates, and montmorillonite are among the dominant clay minerals in most soils of the temperate region. Therefore, these minerals are expected to dominate the clay and silt fraction of suspended sediments in surface waters after precipitation and were thus selected for our experiments.

In addition, artificial silicate particles (sicastar, micromode: 0.1 µm diameter, Art.-No. 43-00-102; 0.4 µm diameter, Art.-No. 43-00-402; 0.8 µm diameter, Art.-No. 43-00-802, zeta potential — 61.5 mV; 1.5 µm diameter, Art.-No. 43-00-153; 3 µm diameter, Art.-No. 43-00-303; 15 µm diameter, Art.-No. 43-00-154) were used as standardized basis particles. Silicate beads possess a silanol surface and therefore show a strong negative surface charge, as is also the case in natural clays. In contrast to natural particles, the surface layer of these particles and their chemical composition is defined.

Influence of suspended particles on the growth of a freshwater flagellate community. The sample, containing a natural assemblage of flagellates, was collected from the surface water of Lake Mondsee. The water was gently filtered through a 11 µm polycarbonate filter (Millipore) to exclude larger predators such as planktonic ciliates and metazoans.

The sample was divided into 4 subsamples of 180 ml each. Three of the 4 subsamples were supplemented with artificial particles, e.g. gray clay, yellow clay and 0.8 µm beads, to a final concentration of 10 mg l⁻¹ (corresponds to approximately 4.2 × 10⁶, 2.9 × 10⁵ and 1.49 × 10⁷ particles ml⁻¹ for the gray clay, the yellow clay and the 0.8 µm beads, respectively). The concentration of 10 mg l⁻¹ is well within densities reported in natural systems, and the particle size range is typical for lakes and lowland streams (Dokulil 1979, Ritchie et al. 1986). The samples were again subdivided into 3 replicates of 60 ml each. Experiments were run in 100 ml Schott flasks and incubated on an overhead shaker (Stuart STR4) at 10 rotations min⁻¹, 15°C and permanent light (8.9 µE m⁻² s⁻¹). Single-cell colorless chrysophytes (Spumella-like), pigmented chrysophytes (Ochromonas/Chromulina) and total number of heterotrophic flagellates were counted daily in live subsamples for 7 d using a Zeiss Axiovert 135 at 200 × magnification under phase contrast. In total, at least 80 flagellate cells were counted per sample.

Growth of Spumella-like flagellates feeding on different food sources. Growth rates of Spumella-like flagellates feeding on live prey, e.g. the large bacterium Listonella pelagia CB5 and the small bacterium MWH-MoNR1, were determined. Growth of the flagellate strain JBC07 was tested at different bacterial abundances between 0.1 and 200 × 10⁶ bacteria ml⁻¹ and for a no-food control. Flagellates were adapted to the respective food conditions 24 h before initiation of the experiment. All experiments were run in NSY inorganic basal medium at 16°C in the dark. Initial experimental volume was 50 ml in 100 ml SCHOTT flasks. Start abundance of flagellates was adjusted to 1–2 × 10⁴ flagellates ml⁻¹. The first subsamples were taken after 3 h, and subsamples of 3 ml were taken every 3 h for a period of 15 h. All subsamples were fixed with formaldehyde (2% f.c.), stored at 4°C and further processed within 2 d. Preliminary experiments proved that DAPI occasionally did not stain the flagellate cells, even when using a high concentration of stain and long incubation times (data not shown). Therefore, a staining protocol using a mixture of SYBR Green I (Molecular Probes) and DAPI was
applied. Briefly, 1 ml of the subsample was stained with SYBR Green I (10 000-fold final dilution of the stock solution) and DAPI (20 µg ml–1 f.c.) for 30 min, filtered onto black 0.2 µm Nucleopore filters (Millipore) backed by 0.45 µm polycarbonate filters (Millipore) and stored at –20°C until inspection. Flagellates were counted under an epifluorescence microscope using UV and blue excitation. At least 80 individuals were counted per sample. Growth rates were calculated from the exponential growth phase.

Growth rates for strain JBM10 were taken from Boenigk et al. (2006). These growth rates were obtained in a similar way as for the flagellate strain JBC07.

Growth rates of flagellates feeding on the small and large bacterium were manually fitted to the Michaelis-Menten equation using non-linear regression, such that:

\[
\mu = \mu_{\text{max}} \times \frac{c - i}{K_m + c - i}
\]

where \(\mu_{\text{max}}\) is the maximal growth rate (d–1), \(c\) is the bacterial abundance (bacteria ml–1), \(i\) is the bacterial threshold food concentration for growth (bacteria ml–1), and \(K_m\) is the half-saturation constant (bacteria ml–1).

**Effect of suspended particles on Spumella-like flagellates.** The presence of suspended sediments in these experiments demanded permanent (over head) shaking to keep particles in suspension. Shaking avoided sedimentation of the suspended particles, and microscopic analysis provided evidence that the particles were suspended homogenously and did not aggregate in the suspension during the experiment (data not shown). Preliminary experiments further showed that the flagellates needed a longer time to acclimatize to these conditions. Therefore, flagellates were acclimatized to the experimental conditions (i.e. food concentration and turbulence [shaking at 10 rotations min–1]) for 3 d before the experiments were started. The ultramicrobacterium MWH-MoNR1 was used as live prey. The food concentration was checked daily using epifluorescence microscopy and adjusted by either adding food or diluting the medium. Experiments were carried out in 100 ml flasks mounted to an overhead rotator (Stuart STR4). Generally, abundances of flagellates at the beginning of the experiments were approximately 5000 ml–1, except at low food concentrations when flagellate abundances were adjusted to approximately 1000 ml–1.

**First set of experiments — influence of particle concentration on growth:** To assess the growth rates at different particle loads, flagellates were acclimatized to food concentrations of 0.1, 0.2, 1, 3, 5, 10, 20 and 50 × 10^6 bacteria ml–1. When the experiment was started, silicate beads with 0.8 µm diameter were added at 5 different concentrations (0, 1, 5, 20 and 100 mg l–1). All experiments were run in 3 replicates. Final sample volume at the start of the experiments was 30 ml, and subsamples of 3 ml were taken every 6 (JBC07) and 4 h (JBM10) for a period of 24 h for analysis. Subsamples were fixed, and 1 ml of each subsample was stained and inspected using epifluorescence microscopy as described above (see ‘Growth of Spumella-like flagellates feeding on different food sources’ section, above). All experiments were run in 3 replicates.

Growth rates of flagellates were calculated from the exponential growth phase. Growth rates from each of the treatments containing different concentrations of suspended particles were fitted to the Michaelis-Menten equation (see ‘Growth of Spumella-like flagellates feeding on different food sources’ section, above) using non-linear regression.

**Second set of experiments — influence of particle size on the feeding process:** In another set of experiments, the effect of particle size on the flagellate strain JBC07 was investigated. We determined clearance rates as we expected that the interference with feeding and the attachment to particles, consequently increasing filtration efficiency, are the major components affecting flagellate–particle interaction. In these experiments the food concentration was adjusted to 3 × 10^6 bacteria ml–1. Particles of different sizes (0.1, 0.4, 0.8, 1.5, 3 and 15 µm) were added at a final particle load of 10 mg l–1 (corresponds to approximately 7.6 × 10^9, 1.2 × 10^9, 1.49 × 10^9, 2.26 × 10^9, 2.8 × 10^9 and 2.3 × 10^9 particles ml–1 for the 0.1, 0.4, 0.8, 1.5, 3 and 15 µm beads, respectively) and at a final particle abundance of 1.49 × 10^7 particles ml–1 (corresponds to approximately 0.02, 1.25, 10, 66, 527 and 66 000 mg l–1 for the 0.1, 0.4, 0.8, 1.5, 3 and 15 µm beads, respectively). Final sample volume at the start of the experiments was 25 ml, and subsamples of 3 ml were taken every 2 h for a period of 24 h. Subsamples were fixed, and 1 ml of each subsample was stained to check bacterial abundances using epifluorescence microscopy as described above (see ‘Influence of suspended particles on the growth of a freshwater flagellate community’ section, above). At least 400 bacterial cells were counted per sample. All experiments were run in 3 replicates.

Clearance rates (h–1) of the flagellate strain JBC07 feeding on the ultramicrobacterium MWH-MoNR1 were calculated according to the following formula:

\[
F = \frac{g \times 10^6}{N_{\text{prey}}} \div 24
\]

where \(g\) is the grazing rate of Spumella (\(g = \mu_{\text{prey experimental}} - \mu_{\text{prey control}}\)) and \(N_{\text{prey}}\) is the mean bacterial abundance during the period of incubation.
**Statistical analysis.** All statistical tests, specifically the ANOVA and t-tests for comparison of growth rates and clearance rates, were performed using the software package SigmaStat 2.03.

**RESULTS**

**Incubation experiment**

Bacterial abundances did not significantly vary during the period of incubation in any of the treatments (ANOVA, p >> 0.05) and were 5.02 ± 0.57 and 4.69 ± 0.15 × 10⁶ bacteria ml⁻¹ (control treatment), 4.69 ± 0.48 and 4.37 ± 0.15 × 10⁶ bacteria ml⁻¹ (0.8 µm beads), 3.14 ± 0.26 and 3.14 ± 0.37 × 10⁶ bacteria ml⁻¹ (gray clay) and 4.03 ± 0.89 and 3.66 ± 0.29 × 10⁶ bacteria ml⁻¹ (yellow clay) at the beginning and the end of the experiment, respectively.

In the control treatment the flagellate community showed low, but significant growth (μ = 0.05 ± 0.02 d⁻¹; t-test; p = 0.021). In the presence of suspended sediments, community growth was significantly higher for silicate beads and the kaolinite-dominated clay (gray clay) (μ = 0.14 ± 0.05, p = 0.017 and μ = 0.13 ± 0.004, p = 0.019, respectively), but not for the kaolinite/montmorillonite (yellow clay) (μ = 0.06 ± 0.02, p = 0.938). In contrast to this community response, the abundance of solitary chrysophytes decreased in all treatments (ANOVA; Tukey test; p = 0.001, p = 0.023 and p = 0.014 for beads, kaolinite-dominated clay and kaolinite/montmorillonite, respectively; Fig. 2). For *Ochromonas*/Chromulina, suspended sediments did not significantly alter the mortality rate (ANOVA; Tukey test; p >> 0.05 for all clay types).

**Growth in the absence of suspended sediments**

Growth rates of the flagellate strain JBC07 as a function of food concentration corresponded to saturation kinetics (Michaelis-Menten equation). Maximum growth rate when fed with the large bacterium CB5 (μmax) was 2.8 d⁻¹, the half-saturation constant (Km) was 1.2 × 10⁶ bacteria ml⁻¹ and the threshold food concentration (i) for positive growth was 0.3 × 10⁶ bacteria ml⁻¹. When the ultramicrobacterial strain MWH-MoNR1 was used as bacterial food, i was 6.5 × 10⁶ bacteria ml⁻¹, Km was 22 × 10⁶ bacteria ml⁻¹ and μmax was 2.3 d⁻¹ (Fig. 3). Growth rates of the flagellate strain JBM10 growing on the bacteria CB5 and MWH-MoNR1 are provided in Boenigk et al. (2006).

**Growth in the presence of different concentrations of suspended sediments**

Growth rates in the presence of suspended sediments still corresponded to saturation kinetics (Michaelis-Menten equation). The growth kinetic parameters depended on the concentration of suspended sediment (S, mg l⁻¹), i.e. μmax decreased, whereas Km and i increased (Figs. 4 & 5). i increased linearly with S: i = 0.23 × S + 2.7617 (r² = 0.99) and i = 0.229 × S + 1.201 (r² = 0.99) for *Spumella* strains JBC07 and JBM10, respectively. Similarly, Km increased linearly with S: Km = 0.194 × S + 8.01 (r² = 0.98) and Km = 0.179 × S + 5.596 (r² = 0.99) for *Spumella* strains JBC07 and JBM10, respectively. μmax of the flagellates decreased exponentially with increasing S: μmax = 1.90 × e⁻⁰.⁰₃₇₇ × S (r² = 0.99) and μmax = 1.36 × e⁻⁰.₁₂₈₄ × S (r² = 0.96) for *Spumella* strains JBC07 and JBM10, respectively (Fig. 5). Based on the coefficient of determination (adjusted r²), the models explained 90.4 and 85.4% of the observed variation for the growth rates of JBC07 and JBM10, respectively (Fig. 4), whereas food concentration alone explained only 70.9 and 39.6% for JBC07 and JBM10, respectively.

**Effect of particle size on the clearance rate**

We suspected that the predominant negative effect of suspended particles is largely due to interference with feeding, and further that this interference is correlated to particle size. We therefore investigated the effect of particle size on the clearance rate.

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Fig. 2. Growth rates of a flagellate community originating from Lake Mondsee and specifically of solitary chrysophytes, i.e. ‘*Spumella*-like’ flagellates and *Ochromonas*/Chromulina, in the presence of different suspended particles. *: growth rates that are significantly different (ANOVA; Tukey test) from the respective growth rate in the control treatment.
Fig. 3. *Spumella* sp. Growth rate as a function of (A) bacterial abundance and (B) bacterial biovolume for the flagellate strains JBC07 and JBM10 based on non-linear regression models (for explanation see ‘Materials and methods’): \( \mu = 2.8 \times (c - 0.3 \times 10^6)/(1.2 \times 10^6 + c - 0.3 \times 10^6) \) and \( \mu = 2.3 \times (c - 6.5 \times 10^6)/(22 \times 10^6 + c - 6.5 \times 10^6) \) for the flagellate strain JBC07 feeding on the large and the small bacterium, respectively. Growth rates of the flagellate strain JBM10 are provided in Boenigk et al. (2006). Gray area indicates bacterial abundances and biovolumes usually realized in oligotrophic to mesotrophic environments.

Fig. 4. *Spumella* sp. Effect of suspended clay and food concentration on the growth of bacterivorous flagellates. (A,C) Growth rates as functions of bacterial densities in the presence of 5 concentrations (0, 1, 5, 20 and 100 mg l\(^{-1}\)) of suspended silicate particles (0.8 µm). Dotted lines indicate the respective numerical response curves following Michaelis-Menten kinetics, allowing for an effect of suspended sediment concentration on the parameters \( i, K_m \) and \( \mu_{max} \) (see ‘Materials and methods’ for explanation). (B,D) Fit of observed growth rates to model assumptions. Allowing for effects of both food concentration and suspended sediment concentration on growth rates, the models explained 90.4 and 85.4% of observed variation (see ‘Results’). For the strain JBM10 the data points corresponding to 100 mg suspended sediment l\(^{-1}\) were partly excluded as maximal growth rate is near 0 and, consequently, Michaelis-Menten kinetics become an inappropriate approximation.
In the absence of suspended sediments, the clearance rate for the flagellate strain JBC07 feeding on the bacterial strain MWH-MoNR1 was 5.46 ± 0.14 nl flagellate⁻¹ h⁻¹. When pooling all data, the presence of suspended sediments affected flagellate clearance rates negatively (t-test; p = 0.002). We tested the effect of particle size both for a fixed particle concentration of 10 mg l⁻¹ and for a fixed particle abundance of 1.49 × 10⁷ particles ml⁻¹. Regarding the fixed particle concentration of 10 mg l⁻¹, a hump-shaped relationship with an optimum and decreasing values at lower and higher particle sizes (second-order polynomial regression model) was the best model to explain the change of clearance rates (r² = 0.74; p ≤ 0.001). Optimum particle size, however, was smaller, i.e. the smallest effect of suspended sediment particles was found for the 1.5 µm beads (Fig. 6).

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Similarly, when the different-sized sediment particles were added at a fixed abundance of 1.49 × 10⁷ particles ml⁻¹, a hump-shaped relationship with an optimum and decreasing values at lower and higher particle sizes (second-order polynomial regression model) again was the best model to explain the change of clearance rates (r² = 0.74; p ≤ 0.001). Optimum particle size, however, was smaller, i.e. the smallest effect of suspended sediment particles was found for the 1.5 µm beads (Fig. 6).

**DISCUSSION**

**Growth kinetic parameters are correlated to suspended sediment concentration**

The growth rates of the flagellate strain JBC07 feeding on both bacteria fit with literature data (e.g. Jürgens 1994, Rothhaupt 1996a), and were generally sim-
ilar to those of the flagellate strain JBM10 (Boenigk et al. 2006). However, in the second set of experiments, i.e. when the effect of suspended sediments was tested, threshold and $K_m$ values were somewhat lower. This may be due to the presence of traces of DOM in the latter experiments, i.e. either a direct stimulation of the flagellates or an indirect stimulation via bacterial growth. However, as bacterial abundances remained constant in our experiments (within the counting error) and the concentration of DOM was low, the responsible factors for the observed shifts in threshold and $K_m$ values remained vague. The fact that bacterial abundances did not significantly change in our experiments corresponded to our expectations, as growth rates of ultramicrobacteria are generally low, even under optimal conditions (Hahn 2003).

Direct interception implies that the predator propels water past itself and catches particles with which it collides (Fenchel 1987). Theoretical considerations suggest that suspended particles should affect the feeding efficiency of interception-feeding nanoflagellates, e.g. as individual particle handling is time consuming, a high density of non-food particles in the water is expected to strongly decrease the uptake of nutritive food and, therefore, negatively affect ingestion efficiency and subsequent growth of the flagellates (Boenigk & Arndt 2002). Corresponding to these theoretical considerations, suspended sediments decreased the growth rates of the investigated flagellates in our experiments. The growth rates reflected literature data (Jürgens 1994, Rothhaupt 1996b), but the respective growth kinetic parameters ($K_m$, $i$ and $\mu$) were correlated to the concentration of suspended sediments. Corresponding to the optimal foraging theory, the time periods for 1 instance of ingestion may be divided into the time spent ‘searching’ and the time spent ‘handling’. For a low frequency of contacts (i.e. low particle abundance), the time spent ‘handling’ becomes negligible within the time budget of the flagellate and thus flagellate feeding and growth should be approximately related linearly to particle concentration. For a high frequency of contacts (i.e. high particle abundance), the time spent ‘handling’ becomes increasingly important and the increase in the ingestion and growth rates with increasing particle abundance should therefore increasingly deviate from a linear correlation, but better fit saturation kinetics (Boenigk & Arndt 2000). This may explain the correlation between parameters of the numerical response and particle abundance in our experiment. Threshold value and $K_m$ (i.e. parameters mainly characterizing the course of the numerical response curve at low to intermediate food concentrations) were approximately linearly correlated with particle abundance, whereas $\mu_{\text{max}}$ (i.e. a parameter mainly characterizing the course of the numerical response curve at high food concentrations) was exponentially correlated with particle concentration.

The negative effect of suspended particles on the growth of the *Spumella*-like flagellates can most probably be generalized, as the strength of the effect of suspended particles on the growth rates was similar for both investigated flagellate strains.

In contrast to our findings, we found in a former study (Boenigk & Novarino 2004), a positive stimulation of the maximal growth rate of interception-feeding flagellates in the presence of clay particles. In that study, flagellates were directly transferred to the experiment from the stock culture. Further, experiments were run at satiating food concentrations using larger bacteria. Thus, residual growth and shifts in the feeding behavior at satiating food conditions may have overlaid the actual response to the experimental treatment. The results of that study must therefore be considered carefully. The discrepancy between these studies again highlights the importance of natural food concentrations and food particle size for estimating the effect of abiotic parameters on the growth of *Spumella*-like flagellates (cf. Boenigk et al. 2006).

**Strength of disturbance depends on particle size: mechanisms involved in the flagellate–particle interaction**

Interference with feeding, on the one hand, and attachment to particles to subsequently increase filtration efficiency, on the other hand, are the major components considered to affect the flagellate–particle interaction (Jack & Gilbert 1993, Christensen-Dalsgaard & Fenchel 2003, Boenigk & Novarino 2004). If this assumption is true, one would expect a strong effect of particle size in this interaction. In fact, we found that the particle size of suspended sediments is an important factor influencing the clearance rate of the *Spumella*-like flagellates. The clearance rates strongly decreased when small and large particles were present, but were only slightly affected by intermediate-sized particles of around 3 µm in diameter. When small sediment particles in the range of ingestible bacteria were present, clearance rates were low, indicating that these particles interfere with the feeding process (see above). For intermediate-sized particles (around 1.5 to 3 µm), the negative effect of the suspended particles on the feeding process became negligible.

We suspect that the ingestion of intermediate-sized particles, and therefore the interference with the feeding process, becomes increasingly counterbalanced by increasing filtration efficiency due to attachment.
Christensen-Dalsgaard & Fenchel (2003) reported that the highest feeding flow produced by the flagellate was observed for cells attached to large particles that were still small enough for the cells to be able to pull them. This was judged to be a result of wall effects caused by the proximity of the surface of the particle. While Christensen-Dalsgaard & Fenchel (2003) reported the best particle size to attach was 25 µm for *Paraphysomonas vestita*, optimal particle size for the *Spumella* strain used in this study was <15 µm. The reduction of the ingestion efficiency is assumed to depend on the cell size of the flagellate, the distance of the center of force of the flagellum to the cell body and the length of the attachment thread. The small cell size of our flagellate strain JBC07 and also the very short or even absent attachment thread may consequently explain the difference of the optimal particle size between *Spumella* (present study) and *P. vestita* (Christensen-Dalsgaard & Fenchel 2003). When the number of particles was held constant (instead of the total particle weight), these size effects were overlaid by concentration-dependent effects. In this case optimal particle size was even around 1.5 µm. We assume that mechanical damage of the flagellate cells became of quantitative importance for larger particles.

**Significance of suspended sediments for chrysomonad flagellate field populations**

Colorless chrysophytes are an often dominating component of the microbial food web (Salbrechter & Arndt 1994, Carrias et al. 1998, Finlay & Esteban 1998, Arndt et al. 2000). Suspended clays and silts are also common in many lakes worldwide (Dokulil 1979, Gliwicz 1986, Threlkeld 1986, Hart 1988, Knowlton & Jones 1995). Low annual mean concentrations of <10 mg l⁻¹ may increase to >200 mg l⁻¹ upon precipitation (Dokulil 1979, Knowlton & Jones 1995). Despite the ubiquity of suspended sediments, however, their effect on this common and abundant flagellate group has hardly been studied (England et al. 1993, Jack & Gilbert 1993, Boenigk & Novarino 2004). Further, despite the vast literature on suspended sediment concentrations, the biologically relevant particle characteristics, i.e. particle size distribution and particle abundance, are rarely mentioned.

We could demonstrate that dominant protist taxa such as the chrysophytes are significantly affected by suspended sediments. Even slight shifts in the abiotic factors may make the difference between survival and death of the population at the food concentrations common in oligotrophic and mesotrophic lakes (i.e. 0.4 to 4 × 10⁵ µm³ ml⁻¹). As suggested by Boenigk et al. (2006), flagellates in oligotrophic and mesotrophic lakes are severely food limited and the bacterial biovolume should allow for low growth rates only. Thus, even slight shifts towards adverse abiotic conditions must be expected to lead to the mortality of at least some flagellate taxa. Accordingly, in our experiments, suspended clay concentrations of only 5 to 10 mg l⁻¹ made a difference between net growth and net mortality at bacterial biovolumes realized in the field.

It should be noted that field populations of bacteria are more variable in size than those in our experiment, and thus a bacterial abundance of 2 to 4 × 10⁶ bacteria ml⁻¹ in the field corresponds roughly to a biovolume of 1 to 4 × 10⁵ µm³ ml⁻¹. In our experiments, we used 1 ultramicrobacterial strain, which is representative for freshwater bacterioplankton, but, of course, much less variable in size. In consequence, the abundance allowing for a net growth of the flagellates was around 5 to 10 × 10⁵ bacteria ml⁻¹ in our experiments. Still, the available bacterial biovolume (i.e. around 1 to 4 × 10⁵ µm³ ml⁻¹) corresponded to that of bacterial field populations in oligotrophic and mesotrophic lakes.

In contrast to taxon-specific investigations, the effects of suspended sediments may not be striking on the community level (HNF or protists) and, thus, may lead to a false impression, i.e. that suspended particles do not matter. Accordingly, we found no negative effect of suspended clay on the community level in our study, but a strong effect on specific taxa. As we only investigated 1 model community, generalizations on the community level are, so far, difficult. The significance of particular particle characteristics such as surface load, form and ion exchange capacity is not well understood either (but see Boenigk et al. 2005a). The effect of suspended particles may differ markedly depending on the types of particles introduced and on the taxonomic composition of the flagellate community. Further studies on different flagellate communities and using different types of suspended sediments are therefore desirable. We suspect, however, that particle abundance and particle size distribution are mainly responsible for the specific particle–protist interaction and that purely pelagic flagellates may generally be negatively affected, whereas flagellate taxa that are commonly associated with sediments (i.e. benthic flagellates, soil flagellates and flagellates attached to substrate flocs [e.g. bodonids as observed in our experiment; data not shown]) may even be stimulated. Still, these interactions may be further modified by bacterial aggregation or biofilm formation, i.e. factors that were excluded from our experiment by choosing the model food bacterium.

Shifts in the concentration of suspended sediments in the field are also usually linked to shifts in turbulence and DOM supply. The latter factors have both been shown to affect flagellate growth directly and
indirectly via stimulation of bacterial growth (e.g. Peters et al. 2002, Dolan et al. 2003, Havskum et al. 2003). It may, therefore, be difficult to separate effects of suspended sediments from those of DOM supply and turbulence in the field, specifically as different taxa may respond differently to these factors. However, our results provide evidence of the significance of suspended sediments for the growth of bacterivorous protists, specifically of colorless chrysomonads.

In conclusion, suspended sediments are a crucial, but so far underestimated, factor for bacterivorous protists. Our results imply a strong taxon-specific response to suspended sediment load already at low suspended sediment concentrations. The ecological response of different bacterivorous protist taxa to suspended sediment load is consequently a key issue towards a better understanding of microbial dynamics, specifically in small or turbulent water bodies.

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LITERATURE CITED
Boenigk J, Wiedlroither A, Pfandl K (2005a) Heavy metal toxicity and bioavailability of dissolved nutrients to a bacterivorous flagellate are linked to suspended particle physical properties. Aquat Toxicol 71:249–259

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Rothhaupt KO (1996a) Laboratory experiments with a mixotrophic chrysophyte and obligately phagotrophic and phototrophic competitors. Ecology 77:716–724
Rothhaupt KO (1996b) Utilization of substitutable carbon and phosphorus sources by the mixotrophic chrysophyte Ochromonas sp. Ecology 77:706–715

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