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Quantifying interspecific coagulation efficiency of phytoplankton

Jørgen L. S. Hansen¹, Thomas Kiørboe².*

¹Marine Biological Laboratory, Strandpromenaden 5, DK-3000 Helsingør, Denmark
²Danish Institute for Fisheries Research, Charlottenlund Castle, DK-2920 Charlottenlund, Denmark

ABSTRACT: Non-sticky latex beads and sticky diatoms were used as models to describe mutual coagulation between sticky and non-sticky particles. In mixed suspensions of beads and Thalassiosira nordenskjoeldii, both types of particles coagulated into mixed aggregates at specific rates, from which the interspecific coagulation efficiency could be calculated. Stickiness between beads and T. nordenskjoeldii was 50% of that of T. nordenskjoeldii in monospecific suspensions, and this ratio remained constant throughout 12 experiments covering 1 order of magnitude variation in the stickiness level of T. nordenskjoeldii. Mutual coagulation between Skeletonema costatum and the non-sticky cells of Ditylum brightwellii also proceeded with half the efficiency of S. costatum alone. The latex beads were suitable to be used as ‘standard particles’ to quantify the ability of phytoplankton to prime aggregation of suspended particles.

KEY WORDS: Phytoplankton stickiness, Interspecific coagulation

INTRODUCTION

Aggregates consisting of live phytoplankton cells are frequently formed during diatom blooms. Sinking rates of these aggregates may exceed those of free, unaggregated cells by orders of magnitude, and the process of aggregation has been acknowledged as the key mechanism behind the massive sedimentation of phytoplankton often occurring subsequent to diatom blooms (Smetacek 1985, Passow 1991, Riebesell 1991, Kiørboe et al. 1994).

One important mechanism of aggregate formation is physical coagulation, which can be described as the physical collision of free, suspended cells and their subsequent adhesion into aggregates (Jackson 1990). The stickiness coefficient, α, expresses the probability of adhesion following the collision of 2 single cells, and models of coagulation processes have shown that this property of algae occasionally determines the dynamics of blooms: a bloom of sticky algae cannot exceed a certain density, the critical concentration, where coagulation balances algal growth (Jackson 1990).

Measurements of phytoplankton stickiness have been made on a number of diatoms grown in cultures (Kiørboe et al. 1990, Kiørboe & Hansen 1993, Drapeau et al. 1994) and in field and mesocosm experiments (Kiørboe et al. 1994, Dam & Drapeau 1995), and it has been shown that several diatom species are sticky. Field studies of diatom blooms have confirmed that phytoplankton aggregates indeed form by means of physical coagulation and that the rates and temporal pattern of their sedimentation can be predicted from coagulation theory (Kiørboe et al. 1994). In order to calculate aggregation rates, these studies have considered the phytoplankton community to be of uniform stickiness. However, phytoplankton communities consist of assemblages of species; aggregation occurs between species, and measurements of stickiness of cultured diatoms suggest variation between species which can differ by orders of magnitude (Kiørboe & Hansen 1993).

Hansen et al. (1995) modelled coagulation of a multispecies bloom by assuming that only certain species were sticky and that sticky species could cause non-sticky species to aggregate. The model showed that interspecific aggregation may cause a fast succession of species. It was assumed that interspecific stickiness

*Addressee for correspondence. E-mail: tk@dfi.min.dk

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equals the average of the intraspecific stickiness coefficients. However, the sticking efficiency between different species has never been measured, and it is not clear how stickiness coefficients measured in monospecific suspensions should be applied to multispecies diatom communities.

This study applies coagulation theory to heterogeneous particle assemblages and presents a technique for measuring stickiness between different populations of particles. In a mixture of non-sticky latex beads and cultured diatoms the aggregation rate of each type of particle is measured separately and the stickiness coefficient of interspecific collisions is calculated.

**MATERIALS AND METHODS**

Batch cultures of the diatoms *Skeletonema costatum, Thalassiosira nordenskjoldii* and *Ditylum brightwellii* were cultivated in B1 culture medium (Hansen 1989) with silicate (25 mg l−1, or 75 mg l−1 for *D. brightwellii*), at 10°C and a light intensity of 60 μE m−2 s−1 in a 12 h light:12 h dark cycle. In all experiments we used cells in exponential growth. Culture samples for measurements of stickiness were ‘washed’ prior to the experiments in order to remove exopolymeric material from the suspension of cells which may interfere with stickiness measurements (Kierboe & Hansen 1993). This was achieved by 2 to 4 times of reverse filtration through a 6 μm plankton gauze followed by resuspension in fresh culture media, thereby reducing the volume fraction of old culture medium to less than 1 %. Sub-samples were stained with alcian blue to visualise any mucus particles (transparent exopolymeric particles, TEP) present in the samples as described by Kierboe & Hansen (1993).

Stickiness of the diatoms was quantified by the use of a couette device (van Duuren 1968) following the procedure described by Kierboe & Hansen (1993). Briefly, a couette consists of 2 cylinders, one inside the other. The outer cylinder rotates, and this generates a well-defined laminar shear in the fluid-filled annular gap between the 2 cylinders. The magnitude of the shear can be calculated from the rotation speed and the dimensions of the cylinders. The couettes were sampled at 15 to 20 min intervals and the particle concentrations monitored by an ELZONE 180 (Particle data, Inc.) particle counter equipped with a 120 or 240 orifice tube. The stickiness coefficient, α, can be calculated from the exponential decline in particle concentration (C) over time t (Kierboe et al. 1990)

\[
C_t = C_0 \exp \left( \frac{-7.824 \alpha \gamma}{\pi} t \right)
\]

where \( \phi \) is the volume fraction of particles and \( \gamma \) is the shear rate (s−1) in the couettes. The experiments were conducted in 2 to 4 couettes simultaneously at shear rates of 8 or 10 s−1 over 1 to 3 h periods. Individual aggregates were examined microscopically after the experiment.

Estimates of stickiness between different types of particles were obtained using the following technique: non-sticky latex beads were added to a suspension of diatoms and the rate at which the beads coagulated was compared with the coagulation rate of cells alone. Latex beads with an average size of 13.34 μm were used together with *Thalassiosira nordenskjoldii* (19 μm equivalent spherical diameter, ESD). Because the beads were smaller than the cells, it was possible to distinguish the 2 populations of particles on the electronic particle counter (Fig. 1). The decline in their concentrations was followed simultaneously. The beads were counted in a size window ranging from 6 to 15.6 μm, and the cells in a window ranging from the smallest cell size up to the maximum limit of the 240 μm orifice tube. The formation of a dimer composed of 1 cell and 1 bead causes 1 particle to ‘disappear’ from the lower window, while the formation of a dimer composed of 2 cells causes 1 particle less in the higher window. Thus, cell-bead and cell-cell coagulation was registered as a decline in particle concentration in the lower and the higher size windows, respectively.

First, an experiment was conducted with beads alone, which confirmed that these particles were non-sticky. Next, the effect of the presence of beads on the estimate of cell-cell stickiness was examined. In 1 experiment the coagulation of cells was measured in 2 simultaneously run couettes, 1 containing beads and diatoms and 1 only with diatoms. Two other test experiments were conducted: 4 couettes were run with varying concentrations of beads but with a constant algal concentration and a second series with varying algal concentrations and a constant concentration of beads.

![Fig. 1 Thalassiosira nordenskjoldii. Size distribution of particle volume of latex beads (open bars) and cells of T. nordenskjoldii (hatched bars)](image-url)
Following these experiments, another 12 experiments were conducted over a period of 3 mo with *T. nordenskiöldii* cultures of varying cell-cell stickiness. Based on the same principles, an experiment was conducted with colonies of *Skeletonema costatum* (16 μm ESD) together with *Ditylum brightwellii* (38 μm ESD). Three couettes were run simultaneously, 1 with *S. costatum* (4800 colonies ml⁻¹), 1 with *D. brightwellii* (800 cells ml⁻¹), and 1 containing the 2 species together (each at half the above concentrations). In the 2 couettes containing *S. costatum*, the concentration was monitored only in the size window covering *S. costatum* (9 to 30 μm) whereas coagulation of *D. brightwellii* cells alone was followed in the size window from 30 μm to the maximum limit of the 240 μm orifice tube.

## RESULTS

Staining with alcian blue did not reveal significant amounts of TEP in any of the cultures, and TEP was totally absent in the suspensions of washed cells. The aggregates formed by *Thalassiosira nordenskiöldii* in the couette device were dense globules of cells, and they did not absorb stain when exposed to Alcian blue. Thus, there was no indication that TEP had any influence on the coagulation, which agrees with previous studies of *Skeletonema costatum* (Kiarboe & Hansen 1993). In monospecific suspensions, vegetative cells of *Ditylum brightwellii* were consistently non-sticky whereas the stickiness coefficient of *T. nordenskiöldii* ranged between 0.1 and 0.7 and *S. costatum* between 0.1 and 0.6. The control experiment showed that the latex beads themselves were non-sticky.

In mixtures with sticky cells of *Thalassiosira nordenskiöldii*, the latex beads coagulated into aggregates, which appeared as a mosaic of beads and cells in the microscope. Due to aggregation formation, the numbers of free cells and latex beads declined exponentially, but at different rates depending on treatment. Parallel experiments run with the same algal concentration but with and without latex beads and with beads in different concentrations showed similar rates of exponential declines of cells (Fig. 2A). The slopes of the individual regressions did not differ significantly (analysis of variance; 0.30 > p > 0.10) and beads, thus, did not affect measurements of cell-cell stickiness coefficients. In these experiments the coagulation rates of beads were independent of their initial concentration, as indicated by similar slopes (0.30 > p > 0.10; Fig. 2B). In the experiment in which the concentration of cells was varied, coagulation rates of beads followed the coagulation rates of cells (Fig. 2C, D).

The coagulation rate of beads versus coagulation rate of cells in the same couettes revealed a coagulation rate between latex beads and cells of 54% of that of cells alone. This relative coagulation efficiency remained constant over a variation in coagulation rates spanning 2 orders of magnitude (Fig. 3A) and a variation in stickiness of *Thalassiosira nordenskiöldii* of 1 order of magnitude (Fig. 3B). The plots in Fig. 3A and B are equivalent, except that the latter compensates for variation due to variation in particle concentration and shear and reports values as averages (± SD) of replicate couettes.

The experiment with *Skeletonema costatum* and *Ditylum brightwellii* revealed a stickiness coefficient of *S. costatum* of 0.54 while *D. brightwellii* did not coagulate in a monospecific suspension. However, *D. brightwellii* appeared in aggregates in the couette containing both species. In the presence of *D. brightwellii*, the rate of decline of particle concentration in the *S.
costatum size window was only slightly less than that in the pure suspension of S. costatum (Fig. 4), even though the concentration of S. costatum was decreased by a factor of 2. As a consequence, the 'apparent' stickiness coefficient of S. costatum, calculated from Eq. (1) and considering only the volume concentration of S. costatum, was higher, i.e. 0.94.

**DISCUSSION**

The experiments with the latex beads demonstrate, for the first time, that coagulating diatoms indeed have a sticky surface, which also can prime aggregation of non-sticky particles, and that diatoms within aggregates are held together by adhesion of frustules. The fact that the cell-bead sticking efficiency strongly relates to the observed cell-cell stickiness supports the conceptual idea of stickiness as a quantifiable property.

It is possible to calculate the sticking efficiency between the Thalassiosira nordenskjoeldii cells and the latex beads. In a monodisperse suspension of particles of diameter d, the change in concentration of free particles is (McCave 1984):

\[
\frac{dC}{dt} = -1.3\alpha d^2 C^2 - \beta \alpha C^2
\]

which is equivalent to Eq. (1) since \(\pi d^3/6C = \varphi \beta = 1.3\gamma d^2\) is the collision kernel for shear coagulation. This equation describes the formation of a dimer as a loss of 1 particle from the suspended population. In order to describe the dynamics of a suspension consisting of 2 populations of particles, a and b, having diameters \(d_a\) and \(d_b\), and occurring in concentrations \(C_a\) and \(C_b\), Eq. (2) expands to

\[
\frac{dC_a}{dt} = \frac{dC_a}{dt} + \frac{dC_b}{dt} =
\]

\[
-\alpha_{aa}\beta_{aa}C_a^2 - 2\alpha_{ab}\beta_{ab}E_{ab}C_aC_b - \alpha_{bb}\beta_{bb}C_b^2
\]

Eq. (3) expresses the loss of particles due to both intraspecific and interspecific coagulation of the populations. In this case, \(C_a\) and \(C_b\) describe the total concentration of particles measured in the \(z\) size windows on the particle counter. The kinetic for interspecific coagulation is \(\beta_{ab} = 1.3\gamma (0.5d_a^2 + 0.5d_b^2)\), and \(E_{ab}\) is the contact efficiency between particles of a different size. According to Hill (1992), \(E_{ab} = 7.5p^2/(1 + 2p)^2\), where \(p = d_a/d_b\) when \(d_a < d_b\). In the present study we used \(E_{ab} = 9p^2/(1 + 2p)^2\) because the contact efficiency is included in the measurements of stickiness in monospecific suspensions of algae and \(E_{aa} = E_{bb} = 1\) (Hansen et al. 1995).

Since the latex beads (particle a) alone do not coagulate (\(\alpha_{aa} = 0\)), and because interspecific coagulation between algae and latex beads does not affect the concentration of particles larger than or equal to the size of cells (particle b), the decline in each of the particle size windows as monitored on the particle counter is as follows:

\[
\frac{dC_a}{dt} = -2\alpha_{aa}\beta_{aa}E_{ab}C_aC_b
\]

\[
\frac{dC_b}{dt} = -\alpha_{bb}\beta_{bb}C_b^2
\]

The ratio of the specific aggregation rates (or ratios of exponential declines) then equals
The observed ratio between the specific aggregation rates of beads and cells was 0.54 (Fig. 3). Inserting the diameters of the latex beads (13.34 μm) and cells (19 μm ESD) in Eq. (5) results in a ratio of \(\frac{a_{ab}}{a_{bb}} = 0.57\). Thus, the interspecific stickiness coefficient between beads and cells equals the average of the intraspecific values.

This may also be true for mutual coagulation of sticky and non-sticky diatoms. Following the basic equations above, it is possible to calculate the interspecific stickiness between the sticky colonies of *Skeletonea costatum* (species a) and the non-sticky *Ditylum brightwelli* (species b) from:

\[
\frac{dc_a}{dt}/c_a = -a_{aa}b_{ab}c_a - 2a_{ab}b_{ab}c_a c_b
\]

The rate of exponential decline of *S. costatum* in mixture with *D. brightwelli*, \(\frac{dc_a}{dt}/c_a\), was \(9.21 \times 10^{-5}\) s\(^{-1}\) (Fig. 4). Inserting sizes and initial concentrations of the 2 species, the shear rate (10 s\(^{-1}\)), and the stickiness coefficient of *S. costatum* \(a_{aa} = 0.54\) into Eq. (6) results in \(a_{ab} = 0.24\) or \(a_{ab} = 0.44 = 50\%\). Again, the interspecific stickiness equals the average stickiness of the 2 groups of particles. Although this last experiment needs to be repeated with different species, the result suggests that aggregate formation in multi-species blooms may be primed by a few sticky species. The results also verify the assumptions made by Hansen et al. (1995), who used the average stickiness of 2 species to calculate interspecific coagulation. Their models showed that mutual coagulation and sedimentation caused a fast succession within the *diatom* community. Thus, an understanding of how mutual coagulation works provides insight into the mechanisms driving species succession during diatom blooms and makes laboratory measurements of stickiness in monospecific suspension more useful for modelling multi-species blooms.

The non-spherical shape of most diatom species makes calculations of interspecific collision rates difficult and consequently measured stickiness values really express the combined effect of the hydrodynamical behaviour of the cells and their stickiness. Taking this problem into consideration, the spherical and non-sticky latex beads provide a suitable 'standard particle' for measuring the ability of the cells to prime coagulation of other particles in general.

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**LITERATURE CITED**


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