Carbon dioxide: Signal for Excystment of *Naegleria gruberi*

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SUMMARY

Exposure of cysts of *Naegleria gruberi* to slightly increased environmental CO₂ caused excystment. Excystment was also induced by addition of proline or by an increase in the cyst population density, but both did so by increasing the amount of CO₂ produced by the bacteria (*Aerobacter aerogenes*) which contaminated the cyst suspensions. Molecular CO₂ would seem to be an excellent signal to induce excystment of a phagotrophic soil amoeba since the presence of CO₂ would indicate an environment favourable for growth of the amoebae. Once excystment is initiated, it can proceed to completion in atmospheric CO₂.

INTRODUCTION

An adaptive environmental signal for excystment should lead to emergence of the organism into an environment in which there would be a high probability of survival and growth. In the laboratory numerous stimuli can induce the excystment of protozoan cysts. These stimuli include shifts in temperature (Johnson & Evans, 1940) and pH value (Darby, 1929), the addition of water (Garnjobst, 1928) or of chemicals such as carbohydrates (Barker & Taylor, 1933), glycine (Beers, 1945), dicarboxylic acids and acetate (Haagen-Smit & Thimann, 1938), or viable bacteria (Crump, 1950). This diversity of stimuli might reflect the variety of organisms studied, each with its specific stimulus, a variety of stimuli for a given organism or laboratory artifacts. Certain observations are particularly suggestive. Colpoda excysts in response to all the chemicals listed above, to bacteria (Singh, 1941), and even to appropriate agitation (Barker & Taylor, 1933). One frequent stimulus, the presence of viable bacteria, causes numerous changes in the environment, only one of which may be the signal for excystment. In addition, viable bacteria, often of unknown type and unknown quantity, are usually present with the other agents studied, and possibly alter the added agent in an unknown way. The effect of bacteria on chemically induced excystment has usually been ignored or disposed of with a statement such as 'in all crucial experiments the cysts become active long before the bacteria could possibly produce an effect' (Beers, 1945).

*Naegleria gruberi* is a small phagotrophic soil amoeba which under appropriate conditions forms cysts or transforms into flagellates. Though the stimulus for naegleria excystment has not been studied, excystment has been described (Wilson, 1916; Rafalko, 1947; Schuster, 1968). One of the most dramatic cytological

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changes during excystment is the reappearance of contractile vacuoles, which are not present in the uninduced cyst. Active amoeboid motion precedes the emergence of the amoeba through one of several pores in the cyst wall. For our purposes a cyst is considered to be 'full' as long as the cytoplasm is entirely within the cyst wall, but once emergence begins and the cytoplasm is partially within and partially without, the cyst is 'transitional'. The emerging amoeba leaves behind an empty cyst wall. The excysted amoebae develop flagella, but this is without apparent effect on excystment.

This paper describes a study of three environmental variables—proline, CO₂, cyst population density—which induce the excystment of *Naegleria gruberi*, and of the role of bacteria in the process.

**METHODS**

*Growth and preparation of cysts.* *Naegleria gruberi* NB-1 and the details of procedures used for cultivating the amoebae and performing counts will be described by Fulton & Dingle (to be published). Amoebae were grown in association with *Aerobacter aerogenes* on autoclaved agar medium, NM, containing (g/1. distilled water): Difco Bacto-peptone, 2.0; glucose, 2.0; K₂HPO₄, 1.5; KH₂PO₄, 1.0; Difco Bacto-agar, 20. About 10⁶ cysts and 0.1 ml. of an overnight broth culture of *Aerobacter aerogenes* were spread over the surface of NM agar in a 100 mm. diam. Petri plate, and the plate incubated at 38-39°. Under these conditions the cysts excysted, and the resulting amoebae grew exponentially until they reached stationary phase at 2-8 x 10⁶ amoebae/plate. Some amoebae began to encyst after about 20 hr and all encysted within 48 hr. The plates of cysts were stored at 34° and used 4-6 days after plating.

Cysts were prepared for excystment experiments by harvesting each plate into 10 ml. demineralized water and centrifuging the suspension at 10° for 105 sec. at 500 g in a swinging bucket rotor. The supernatant fluid was discarded and the pellet washed twice more in demineralized water. The final pellet was resuspended under conditions suitable to the experiment. The cyst concentration was determined by a Coulter counting apparatus (Coulter Electronics, Hialeah, Florida, U.S.A.) and the bacteria by colony count.

*Excystment on agar.* Agar excystment experiments were done on autoclaved agar media containing 2% (w/v) Bacto agar and supplements as indicated for each experiment. A sample of a washed counted cyst suspension was spread over the surface of a plate, and the plate placed, agar upward, without its cover, in a desiccator of about 2 l. capacity which contained 50 ml. of a 10% (w/v) NaCl solution to maintain constant humidity. The gas phases were adjusted (see below), and the desiccators incubated at 34°. The beginning of incubation was the zero time for each experiment.

To evaluate excystment on agar, the organisms were examined *in situ*. The plates were removed from the desiccators and a circular coverslip 18 mm. in diameter was placed directly on the agar, with care taken to allow a film of liquid to flow between the agar and the coverslip. The organisms were examined with phase contrast optics at a magnification of ×400. When the plates were to be re-incubated they were returned to the desiccators and the gas phases readjusted. For each time point a fresh area of a plate was examined.
**Results**

**Inducers of Naegleria excystment**

*Proline.* In the course of other work, proline was observed to induce the excystment of *Naegleria gruberi*. Liquid suspensions of cysts excysted when L-proline was added at concentrations greater than $10^{-5} \text{M}$; the rate of excystment was similar and maximal at $10^{-4}$, $10^{-3}$, and $10^{-2} \text{M}$. The excysted amoebae immediately transformed into flagellates, and to avoid this we attempted to obtain excystment on agar plates where transformation does not occur. Contrary to the effect of proline in liquid, cysts placed on proline agar did not excyst, regardless of the concentration of proline. Extending the time of incubation, washing the agar, or varying the number of cysts plated did not result in excystment. The disparity between the results in liquid and on agar was resolved by a chance observation. On proline agar plates, uncovered cysts remained dormant while those under a coverslip excysted after an hour (Fig. 1). It seemed likely that the coverslip
created an environment stimulating excystment by interfering with gas exchange. One possibility was an increase in CO$_2$ concentration under the coverslip.

**Carbon dioxide.** Proline agar plates spread with cysts were incubated in atmospheres of different CO$_2$ concentrations (Table 1). Increased pCO$_2$ induced excystment, and the amount of excystment was a function of the pCO$_2$. Furthermore, when the pCO$_2$ was increased, excystment occurred independently of the presence of proline in the agar.

In liquid, where measurement of degree of excystment is much more precise, either proline or CO$_2$ induced excystment (Fig. 2). Under similar conditions of temperature and aqueous environment and with saturating concentrations of

![Figure 1](image1.png)

**Fig. 1.** The coverslip experiment. An agar plate buffered at pH 7.4 with 0.02 M-tris and containing 10$^{-8}$ M-L-proline was spread with 2 x 10$^6$ cysts and incubated at 34°C under standard conditions. A sample of the initial cyst suspension was fixed and counted. At hourly intervals a coverslip was applied to a new area of the plate and excystment evaluated in situ; at the same time the areas under the coverslips applied at one (A) and at 2 hr (B) were re-examined.

![Figure 2](image2.png)

**Fig. 2.** Induction of excystment in liquid by proline and by CO$_2$. Three flasks were prepared each containing 5 ml. of phosphate buffer, pH 6.8, with 8 x 10$^6$ cysts per ml. One flask also contained 10$^{-8}$ M-proline, and another flask was capped and its atmosphere adjusted to 0.95 % pCO$_2$. The flasks were shaken at 25°C with 100 strokes per minute. At the end of the experiment the pH in the flasks ranged from 6.85 for the control to 6.7 for the flask with increased pCO$_2$.

<table>
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<tr>
<th>pCO$_2$ (%)</th>
<th>0.03</th>
<th>0.33</th>
<th>1.0</th>
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<tr>
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<td>8</td>
<td>15</td>
<td>5</td>
<td>7</td>
</tr>
<tr>
<td>Empty cysts</td>
<td>7</td>
<td>35</td>
<td>90</td>
<td>93</td>
</tr>
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</table>

**Table 1. Excystment on proline agar in varying CO$_2$ atmospheres**

Washed cysts were spread at 2 x 10$^6$ cysts per plate on agar plates buffered at pH 6.8 with 0.02 M-potassium phosphate and containing 10$^{-8}$ M-L-proline. The plates were placed in desiccators, the atmospheres adjusted to various CO$_2$ concentrations, and the desiccators incubated at 34°C. Excystment was evaluated after 4 hr incubation.
CO₂-induced excystment of *naegleria*

CO₂ or proline, excystment regularly occurred sooner in response to CO₂ than to proline.

Measurement of the response to CO₂ depended on having a measure of the relative rate of excystment. If excystment of a population were synchronous, there would be an abrupt transition from full cysts to transitional cysts and then to empty cysts. Excystment under the conditions we used was asynchronous, and a plot of the % full cysts against time of incubation gave a sigmoid curve. If such a curve represented a summed normal distribution of excystment times, the probit transformation would convert it to a linear curve (Finney, 1962). Probability curves for *naegleria* excystment approximate linearity (Fig. 3). The mean time of excystment (the time for 50% excystment) varied with the pCO₂ (Fig. 8). The reciprocal of the mean excystment time was used as a measure of the relative rate of excystment (this approach, used with a similar morphological change, will be discussed in Fulton & Dingle, to be published). Though the mean excystment time varied with pCO₂, the slopes of the probability curves, which represent the standard deviation did not vary greatly, so the heterogeneity of the response of the cyst populations, to increased pCO₂ was not markedly affected by the relative rate of excystment.

The effect of CO₂ concentration on the relative rate of excystment was determined by multiple measurements under standardized conditions (Fig. 4). The response of cysts was a function of pCO₂ in the range from 0·15% to about 1%. Excystment at any pCO₂ less than 0·15% was too slow and variable to obtain valid
measurements of rate. Thus we do not know whether a threshold level of pCO₂
(i.e. above atmospheric) is necessary to induce excystment. Cysts can be stored
on agar plates in atmospheric pCO₂ for prolonged periods without significant
excystment; this suggests that excystment does not occur unless pCO₂ exceeds
0.03%.

A portion of the CO₂ dissolving in an aqueous solution ionizes to increase the
H⁺, HCO₃⁻, and CO₃²⁻ content of the solution. Is molecular CO₂ the stimulus for
excystment, or is one of its ionic products the effective agent? Acid pH values
have been reported to induce excystment (Darby, 1929), but naegleria cysts
placed in phosphate buffers from pH 3.3 to 7 did not excyst until CO₂ was added.

In the buffered solutions used the addition of CO₂ did not markedly alter the pH
value (e.g. Fig. 2), so changes in pH value are not responsible for excystment.
Bicarbonate ions also did not induce excystment (Fig. 5); at high concentrations
the slight excystment observed was probably due to molecular CO₂ produced in the
solution. The addition of CO₂ to the bicarbonate solutions induced excystment
(Fig. 5), showing that bicarbonate neither stimulated nor inhibited the process.
Carbonate ions, as well as carbonic acid, are produced in very small amounts at the
neutral pH value used for excystment. Carbonate should be produced in similar
or greater amounts in the bicarbonate solutions, which did not induce excystment.
Also, changing the pH value of the excystment environment, and thus markedly
altering the proportions of carbonic acid and carbonate (see Scott, 1939), did not
markedly affect the rate of excystment in response to CO₂. For example, the relative
rate of excystment in response to 0.6% pCO₂ in phosphate buffered solutions
**CO₂-induced excystment of naegleria**

at pH 5 and pH 9 was about 70% of the rate at pH 7. Thus by elimination we conclude that dissolved molecular CO₂ gives the signal for excystment.

Effect of cyst population density. In a study of the effect of cyst population density on the relative rate of excystment in response to CO₂, we found that increasing the population density in liquid eventually led to excystment at atmospheric pCO₂. The relative rate of population density-induced excystment was a function of the population density (Fig. 6). Considerable variability was encountered in the value of the population density required to obtain excystment with different batches of cysts.

Comparison of inducers of excystment

The addition of proline and an increase in the cyst population density both appeared to induce excystment by increasing the amount of CO₂ in the cyst environment. In both cases the *Aerobacter aerogenes* present in washed cyst suspensions are believed responsible for the CO₂ production. Proline can be decarboxylated or metabolized to CO₂ by *A. aerogenes*. If uninduced cysts are metabolically inactive, the aerobacter would be responsible for the metabolism of proline to CO₂. In liquid media enough CO₂ to induce excystment is produced from the stimulation of aerobacter metabolism by proline, but on agar plates the CO₂ produced diffuses away and excystment occurs only when gas exchange is prevented. Increasing the population density of cysts increases the number of aerobacters/ml., and thereby the amount of CO₂ produced/ml. by the bacteria. If these assumptions are correct, CO₂ is the active agent in all cases. Elimination of the bacteria, or removal of the CO₂ produced by them, should prevent excystment in response to proline or density.

The pCO₂ of the cyst environment can be decreased by absorbing CO₂ from the gas phase of the flask with KOH. CO₂-induced excystment did not occur in the presence of alkali absorbant. Similarly, excystment in response to increased cyst population density or added proline was inhibited by removal of CO₂ from the gas phase. KOH inhibition of excystment was dependent on sufficient shaking to ensure rapid gas exchange between liquid and gas phases. No excystment occurred in response to any inducers when KOH was present, whereas simultaneous controls without KOH did excyst.

A small amount of CO₂ (e.g. atmospheric concentration) might be essential for excystment in response to the environmental alteration produced by proline or density. This possibility, as well as the role of the aerobacter, can be tested by allowing the pCO₂ to remain at atmospheric levels but preventing the production of additional CO₂ by the bacteria. Ideally this could be achieved by studying the excystment of sterile cysts (free of metabolizing as well as colony forming bacteria), but many attempts to sterilize cysts or to obtain viable cysts by sterile cultivation of amoebae were unsuccessful. (Cysts were formed in sterile cultures on autoclaved aerobacter, but they had abnormal morphology and excysted in liquid without any stimulus.) Because of these difficulties, partial sterilization was accomplished with streptomycin, to which *Aerobacter aerogenes* is sensitive. Overnight incubation of washed cysts on NM containing 200 μg. streptomycin/ml. lowered the population of viable aerobacter from about 10⁷ to less than 10³ per 10⁷ cysts plated. These cysts did not appear to be affected by streptomycin, and excysted
normally in response to CO₂. They did not, however, excyst at population densities which normally induce the excystment of untreated cysts. Streptomycin treatment also inhibited proline-induced excystment, though when streptomycin and proline were added to washed cysts simultaneously, slow excystment still occurred, suggesting that the bacteria, while unable to form colonies, were still able to metabolize proline to CO₂.

The inhibition by KOH of CO₂-, proline-, and density-induced excystment produces strong support for the hypothesis that increased pCO₂ is the stimulus for excystment in all cases. The inhibition of proline- and density- but not CO₂-induced excystment by streptomycin treatment of cysts indicates that the bacteria which contaminate the cyst suspensions are responsible for the CO₂ production.

![Fig. 7. Temporal sequence of morphological changes during excystment.](image)

Fig. 7. Temporal sequence of morphological changes during excystment. Ten ml. of washed cysts at 5 x 10⁶ per ml. in phosphate buffer (pH 7.7) were shaken at 30° in 2.4 % pCO₂. At intervals samples of 100 cells were classified under phase contrast into the arbitrary stages described in the text: ○, stage 1; ●, stage 2; △, transitional and empty cysts.

![Fig. 8. Influence of duration of exposure to CO₂ upon subsequent excystment.](image)

Fig. 8. Influence of duration of exposure to CO₂ upon subsequent excystment. Washed cysts at 3 x 10⁶ per ml. in phosphate buffer were distributed in 10 ml. samples to six flasks. One flask was left open (●), and the remaining flasks adjusted to 0.8 % pCO₂. The flasks were shaken at 30°, and at intervals a flask containing CO₂ was removed, its contents centrifuged and resuspended in 10 ml. phosphate buffer, and shaking continued at atmospheric pCO₂. In each case the transfer took about 3 min. The cysts were in elevated pCO₂ for: ○, 30; △, 45; △, 55; ▼, 80; and ▼, 150 min.

**Response of cysts to CO₂**

Phase contrast observation of living organisms revealed extensive cytoplasmic changes before excystment. In an uninduced or stage 1 cyst the cytoplasm was filled with small particles in active Brownian motion, and no contractile vacuoles were present. After exposure of cysts to CO₂, large phase-dense granules appeared in the cytoplasm (stage 2). Under standard excystment conditions, half of the cysts changed from stages 1 to 2 within about 20 min. (Fig. 7). Soon thereafter contractile vacuoles were formed, and amoeboid movement began. About an hour
after a cyst changed from stages 1 to 2, an amoeba emerged through one of the pores in the cyst, leaving behind an empty cyst wall (Fig. 7). Cysts can recognize CO$_2$ in their environment within 20 min. and begin the sequence of changes which lead to emergence.

In order to test whether the continual presence of increased pCO$_2$ is required for excystment, or whether CO$_2$ induced a reaction after which excystment can continue to completion independently of the presence of CO$_2$, cysts were exposed to an inducing concentration of CO$_2$ for varying times, and then shaken at atmospheric CO$_2$. Thirty minutes of exposure to CO$_2$ was sufficient to induce excystment of about 50% of a cyst population; the extent of excystment was a function of the time the cyst population was exposed to CO$_2$ up to about an hour (Fig. 8). Exposure to CO$_2$ therefore initiates a sequence of events which can continue to completion in the absence of excess CO$_2$.

After about 80 min. exposure to CO$_2$, half the cysts had reached morphological stage 2 (Fig. 7) and could complete excystment independently of increased pCO$_2$ (Fig. 8). This correlation suggests that the critical effect of CO$_2$ is to cause some early event, associated with the appearance of large granules, which results in a commitment to excystment.

**DISCUSSION**

Our major conclusion is that the only stimulus in the environment that naegleria cysts recognize as a signal for excystment is molecular CO$_2$. The fitness of this signal as a means for a phagotroph to recognize an environment in which food is available is obvious. The level of pCO$_2$ in an environment such as soil is a function of the rate of production of CO$_2$—a measure of the population density of organisms, the temperature, and other conditions influencing growth and metabolism—and the rate of loss of CO$_2$ by diffusion. Because of this, CO$_2$ would seem to be one of the few molecules whose concentration in soil would indicate when overall conditions would be favourable for naegleria amoebae to survive and grow. The water content of soil, its pH or temperature, or many of the stable organic molecules soil contains would be less sensitive signals of such an environment. This hypothesis, though it certainly does not consider the complexities of soil ecology, has a simplicity which makes it attractive. Furthermore, the levels of CO$_2$ found in topsoils are sufficient to induce the excystment of naegleria (see Thimann, 1968).

Ballard (1958) found that CO$_2$ induces germination of clover seeds. Loomis (see 1959, 1964), in a study of the role of increased pCO$_2$ in controlling sexual differentiation of hydra, has drawn attention to the many biological processes that are influenced by CO$_2$.

Other environmental variables induce the excystment of naegleria, but these appear to act by increasing the amount of CO$_2$ produced by bacteria in the cyst environment. Proline- and density-induced excystment in the laboratory display the effect of environmentally produced CO$_2$. Although cyst population density was the observed variable in density-induced excystment, it is the simultaneously altered bacterial population density that is responsible for excystment. With some cyst preparations, less responsive to density-induced excystment, excystment could be obtained by adding additional aerobacters. On the nutrient agar for naegleria, NM, it is the growth of aerobacter—and presumably the CO$_2$ micro-
environment created by this growth—which induces excystment of naegleria. In all cases, increased pCO₂ accounts for excystment. From this point of view, as long as bacteria are present anything which increases their rate of CO₂ production—proline, other chemicals, a temperature increase, etc.—would lead to excystment. In this connexion, it is of interest that the washed cyst suspensions used in our experiments are contaminated to the extent of about one viable aerobacter per cyst; one Aerobacter has roughly one thousandth the volume of a cyst. This relatively small quantity of aerobacter, when stimulated by proline or increased in population density, is able to produce enough CO₂ to induce excystment.

Many published studies of excystment of other organisms could be explained in the same way—i.e. that the addition of bacteria, organic acids, etc., induces excystment via CO₂. For example, Crump (1950), in a study of the ability of various bacteria to induce the excystment of a limax amoeba, stated that ‘The unsuccessful attempts already recorded to induce excystment by placing cysts in culture fluid in which Aerobacter... had grown, but from which they were removed by centrifuging, suggests that the substances concerned may be so transient that they disappear as fast as they are formed, and only when there are living bacteria in the medium is enough of the stimulating material present to act successfully on the cysts’. In studying the induction of excystment of didinium by bacteria, Beers (1946) noted that ‘the effective substances are highly unstable. They are probably produced at the surface of the bacteria by an enzyme system. They diffuse a short distance into the medium, where they induce excystment, but only if present in relatively high concentrations—i.e. the concentration of the bacteria must be high to induce excystment. Once in the medium the substances are rapidly altered or destroyed’. Such statements might apply to excystment induced by bacteria-produced CO₂.

Most of this study is included in a dissertation submitted to the graduate faculty of Brandeis University by M.A. in partial fulfilment of the requirements for the degree of Doctor of Philosophy. This work was done during the tenure of a Jack Cohn predoctoral fellowship, and completed while M.A. held a postdoctoral traineeship from training grant T1HD22 of the Institute of Child Health and Infant Development of the National Institutes of Health. The research was supported by a grant from the National Science Foundation.

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