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Effect of copper on cell division, nitrogen metabolism, morphology, and sexual reproduction in the life cycle of Closterium moniliferum (Chlorophyceae)

Erleen Svihevec Christenson
Portland State University

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EFFECT OF COPPER ON CELL DIVISION, NITROGEN METABOLISM, MORPHOLOGY, AND SEXUAL REPRODUCTION IN THE LIFE CYCLE OF CLOSTERIUM MONILIFERUM (CHLOROPHYCEAE)

by

ERLEEN SVIHOVEC CHRISTENSON

A dissertation submitted in partial fulfillment of the requirements for the degree of

DOCTOR OF PHILOSOPHY
in
ENVIRONMENTAL SCIENCES AND RESOURCES - BIOLOGY

Portland State University
1983
TO THE OFFICE OF GRADUATE STUDIES AND RESEARCH:

The members of the Committee approve the dissertation of Erleen Svihovec Christenson presented February 18, 1983.

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AN ABSTRACT OF THE DISSERTATION OF Erleen Svihovec Christenson for the Doctor of Philosophy in Environmental Sciences and Resources-Biology presented February 18, 1983.

Title: Effect of Copper on Cell Division, Nitrogen Metabolism and Sexual Reproduction in the Life Cycle of Closterium moniliferum (Chlorophyceae)

APPROVED BY MEMBERS OF THE THESIS COMMITTEE

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Additions of copper were shown to affect cell morphology, growth rates and nutrient uptake in *Closterium moniliferum*. These parameters are interrelated in the total life cycle of the organism. It was found that the timing of events in the life cycle, including sexual reproduction, could be changed when copper was added. When increasing concentrations of copper were added to the growth medium, *Closterium moniliferum* exhibited a stimulatory, inhibitory, or toxic dose-response typical of organisms to trace metals. The stimulatory effect, occurring at pCu* 14.4, was demonstrated by an early increase in cell number, increased nitrate uptake, and early onset of sexual reproduction. The inhibitory effects of greater concentrations of copper (pCu* 12.1) were a longer lag phase, decreased nitrate uptake, and later onset of sexual reproduction. Light micrographs and scanning electron micrographs of normal and aberrant cells demonstrated the effect of copper on morphology.

It was demonstrated that the sexual phase in some strains of *Closterium moniliferum* could be triggered by changes in the amount of nitrogen in the medium. Although it was expected that each cell would have a minimal cell quota (Q nitrogen/cell) before sexual reproduction occurred, work here demonstrated that Q was not the critical factor in the initiation of the process. Sexual reproduction occurred at Q = .05 micromoles nitrogen/cell up to .21 micromoles nitrogen/cell. Neither was cell density in batch
cultures the critical factor. Sexual reproduction occurred in the range of 400 cells/ml up to 6100 cells/ml. Different nitrogen sources gave different responses in the onset of sexual reproduction. Nitrate depletion, not ammonia, appears to be the critical component in induction of sexual reproduction.

The actual effect of metals like copper on the population dynamics of *Closterium moniliferum* in natural environments is yet to be determined. The stimulatory effect on nitrate uptake by copper, and its accompanying earlier formation of zygotes, may have survival value for the organism since the zygotes become thick-walled and are not as subject to environmental perturbations as vegetative cells.
DEDICATION

I dedicate this dissertation to my mom and dad, Otto and Ann Svihovec, who instilled in me a zest for learning, a desire to strive for excellence, and the North Dakota persistence that they exemplified.
ACKNOWLEDGEMENTS

I wish to thank John Rueter for imparting his enthusiasm for research and for stimulating my brain to formulate hypotheses; Ed Lippert for serving as chairperson through the bureaucratic maze; Richard Petersen for many discussions which helped convince me that it was possible to hang on to sanity while in graduate school; Joann Sanders-Loehr and Arnold Pickar for their perspectives on this work and Wolfgang Fahrenbach for his expertise with the scanning electron microscope.

Most of all, I wish to thank my family who, willingly and unwillingly, have been participants in this project these 4½ years:

my daughter Roxanne, who didn't graduate from college before I did;

my daughter Samantha, who washed more dishes than she cares to count;

my son Cy, who has been waiting to play ball.

I especially thank my husband Wes, who deserves a medal for forbearance, tolerance, solicitude and support. He deservedly shares my moment of accomplishment, as he helped me prove the resiliency of the human spirit.
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CHAPTER I

INTRODUCTION

The purpose of this research was to determine the effects of copper as an agent of environmental stress on the life cycle of Closterium moniliferum. The effects studied were cell division, sexual reproduction, zygote germination, and nutrient uptake.

Excess copper has been shown to have a toxic effect on growth and cell division (Sunda and Guillard, 1976; Anderson and Morel, 1978; Rueter, 1979; McKnight, 1981; and Petersen, 1982) and nutrient uptake (Rueter et al., 1981) in algae. Dinoflagellates, diatoms, and some members of the Chlorophycophyta have been used in this recent research. Closterium moniliferum (Chlorophycophyta, Desmidiaceae) follows a more elaborate life cycle than other algae which have been studied. This allowed the additional dimension of the complete life cycle to be examined. As well as the previously studied aspects of growth, cell division, and nutrient uptake, onset of sexual reproduction and germination of zygotes were included.

Closterium moniliferum is a member of the group of the green algae (Desmidiaceae) which is commonly found in unpolluted, pristine bodies of water and is less common in polluted or nutrient enriched water. This may indicate
that **Closterium** is an organism that is sensitive to excess pollutants, and it has been used in a rapid and reliable method for assessment of the trophic character of the plankton community (Thunmark, 1945). However, no critical studies of the specific response of this alga to metal ions have ever been conducted.

This alga offers a unique opportunity to study the effects of metal ions because of its elaborate life cycle. When exposed to metal ions, most cells show a typical dose-response relationship. The presence of low concentrations will enhance a physiological process but at higher concentrations they become inhibitory and eventually toxic (Figure 1). This research included a study of the total life cycle of **Closterium**, including sexual reproduction integrally tied to nutrient uptake and cell division in the presence of carefully selected concentrations of copper.

Growth of populations of **Closterium moniliferum** were monitored to correlate increase in cell number with uptake of nutrients. Desmids begin sexual reproduction when nitrogen amounts are low in the medium (Biebel, 1964), therefore, nitrogen was monitored in cultures with increasing amounts of copper. The results suggest that copper may be acting on the assimilatory nitrogen metabolism (Figure 2) of **Closterium** by acting on nitrate uptake and/or nitrate reduction. The timing of sexual reproduction should be affected by copper: if nitrate uptake were stimulated, sexual
Figure 1. Schematic representation of the effect of metal ions on physiological processes. These are frequently beneficial but become inhibitory and toxic at higher concentrations (based on Fraústo da Silva and Williams, 1976).
Figure 2. Suggested sites of action of copper (Cu^{2+}) on nitrate metabolism. The numbers 1, 2, 3 indicate specific sites where inhibition may be occurring:

1) Copper inhibits nitrate uptake;
2) Copper inhibits nitrate reduction by acting on the enzyme nitrate reductase;
3) Copper inhibits nitrate reduction by interfering with NAD(P) reduction.
reproduction would occur earlier; if nitrate uptake were inhibited, sexual reproduction would be delayed.

Besides the requirement for nitrogen depletion in the environment before sexual reproduction can occur, the vegetative cells must divide. Light microscopy and scanning electron microscopy were used to monitor quantitative and qualitative changes in cell morphology of dividing cells.

The morphological studies add additional information to the growth studies and may provide a key to the inhibition of cell division exemplified by the increased lag phase. Inhibition of nitrate uptake by copper parallels the inhibition of cell division. Because the nitrogen requirements of *Closterium moniliferum* are integrally tied to its sexual reproduction, it is seen that copper can be an effective tool in understanding the total life cycle of this alga.

**THE LIFE CYCLE AND MORPHOLOGY OF CLOSTERIUM MONILIFERUM**

The cells of *Closterium moniliferum* are elongate unicells, narrowed toward both poles, sometimes slightly tumid at the equator and slightly arcuate. A central nucleus lies between the two chloroplasts at the equator of the cell. After division, a cell has one semi-cell of the parent cell wall and one semi-cell which is formed by new cell wall secretions (Figure 3). When mature, the cells are symmetrical with both semi-cells of the same size (Figure 4).
Cell Division

First division

Figure 3. Diagram showing two successive divisions of a Closterium cell to illustrate the distribution of the parent cell walls to the daughter cells. The walls of the 1st generation cell (1a and 1b) are drawn with heavy longitudinal striations; those of the 2nd generation (2a and 2b) with lighter striations; while those of the 3rd generation (3a and 3b) have striations shown as dotted lines. gp = girdle portion; ap = apical portion (Brook, 1981).
Figure 4. Mature cell of *Closterium moniliferum*, strain #96. 430 X.
The cells themselves act as gametes when light is present (Ueno and Sasaki, 1978; Hogetsu and Yokoyama, 1979) and when certain physiological conditions, such as low nitrogen concentrations, exist (Biebel, 1964). The cells divide and the immature daughter cells mate to form a single zygote in one homothallic strain (#96, Figure 5) or twin zygotes in another homothallic strain (#171, Figure 6). Sexual reproduction occurs before the newly divided cells attain the symmetry characteristic of mature cells, a trait first observed in *Closterium moniliferum* by Fox (1957) and further elucidated by Lippert (1967). The cell contents from two cells travel through a conjugation tube and fuse to form a zygote, which is the only diploid stage in the life cycle.

Zygotes require a maturation time of 1-2 months in the dark (Lippert, 1967). Mature zygotes will germinate when they are exposed to light and are supplied with nutrients and moisture. These new cells, released from the germinating zygotes, begin a new life cycle and can reproduce sexually or vegetatively, depending on conditions of growth. In strain 96, sexual reproduction begins when low nitrogen concentrations exist. This phenomenon was chosen as a point of interest in the overall hypothesis of this research. The interrelationship of nitrogen and copper could influence the total life cycle (e.g. delaying sexual reproduction, inhibiting nitrogen assimilation and/or inhibiting cell division).
Figure 5. Conjugation in Closterium: the C. moniliferum-ehrenbergii type. 1) The pairing of immature daughter cells and formation of conjugation papillae; 2) Secretion of hyaline conjugation vesicle; 3) Initiation of movement of gametes; 4) Final stages of gamete fusion; 5) Mature zygospore and empty cells showing conjugation pore.

cv = conjugation vesicle; g = gametes; p = conjugation pore; pa = papillae (Brook, 1981).
Figure 6. Type of sexual reproduction in strain 171 (homothallic) of Closterium moniliferum resulting in the production of twin zygotes.
THE TRACE METAL COPPER

Copper is a required micronutrient (Price, 1970) yet is one of the trace metals that is most toxic to phyto-plank­ton (Hollibaugh et al., 1980). Its toxic effect is in large part due to its affinity for sulfhydryl groups and its ability to displace metal ion cofactors or active metal centers from metal dependent enzymes (Passow et al., 1961). The action of copper on the sulfhydryl groups may inactivate physiologically crucial SH-dependent enzymes or alter their catalytic specificity or control. Reaction of copper with side chains or replacement of other metals may result in a change in the electrostatic charge and a shift in the ionization constant of the active center to give a change in catalytic ability (Passow et al., 1961).

Mandelli (1969) suggested that copper is strongly bound in carboxyl, phosphate, amino, or sulfhydryl groups and thus its effect tends to be non-specific. Other evidence suggests, however, that copper may act more specifically on the SH groups. For example, Gurd and Wilcox (1956) found that copper oxidized SH groups forming S-S bridges and Fisher and Jones (1981) suggested a common action on SH groups for Hg, Cu, Zn, Pb, Cd, and Mn by showing the relative toxicities of the metals are correlated ($r = 0.961$) with the solubility products of the correlated metal sulfides. While copper does show specificity towards SH groups, sensitive SH groups are so ubiquitous and of importance to so
many different enzymes, that sites of action within a cell are multiple.

The addition of sulfhydryl reactants affects cell division. Hughes in 1950 reported that sulfhydryl reactants in general could block cells from entering prophase and Dustin had earlier (1947) suggested that mitotic poisons were enzyme inhibitors. Stern (1956) found that sulfhydryl groups were important in regulating plant cell division in developing anthers of the lily. Kanazawa and Kanazawa (1969) found that nuclear division in copper treated Chlorella cells was completed but cellular division did not occur. They assumed that the process suppressed by copper was probably concerned with some events related to the cell membrane rather than with those occurring inside the cell. It is probable that both events occur, with the membrane showing the primary effect since it is the location of first contact with the metal.

Evidence indicates that the membrane may include sites of action for metals. Rothstein (1959) saw a rapid reversibility of some of the effects of metal toxicity indicating the membrane as one target. Although photosynthesis inhibition and potassium excretion induced by copper and mercury followed similar dose-response curves in the green alga Chlorella pyrenoidosa, Kamp-Nielsen (1971) concluded that the primary effect of bound metals (copper and mercury) was to cause some destruction of a diffusion barrier causing an out-flow of potassium. Overnell (1975), working
with dinoflagellates, found results consistent with the suggestion of McBrien and Hassall (1965) working with *Chlorella*, that the primary toxic effect of copper is to increase the permeability of the cell. He reported that with Cu\(^{2+}\), K\(^+\) release occurred at a marginally lower concentration than did inhibition of photosynthesis. The action of copper on permeability has not been confirmed since the work of deFilippis (1979) showed little effect on potassium loss from *Chlorella*.

Using isolated chloroplasts, Cedeno-Maldonado *et al.* (1972) found that copper affected the photosynthetic process itself. Shioi *et al.* (1978 a and b) showed that copper inactivated the electron transport of photosystem II and Haberman (1969) showed that both the Hill and Mehler reactions were inhibited by copper.

**Copper As A Toxicant/Nutrient**

Since the investigations of Moore and Kellerman (1904) copper has been used extensively as an algicide in algal control programs. It has recently been established that the total amount of copper in natural waters is not the crucial factor in toxicity to organisms; rather, it is the speciation (chemical form) of copper and the equilibration of those species in the environment/medium that determines the activity and toxicity of ionic copper (Sunda and Guillard, 1976; Anderson and Morel, 1978; Jackson and Morgan, 1978; Rueter *et al.*, 1979; and Sunda and Gillespie, 1979).
Different organisms respond to the activity of ionic copper with different sensitivities; the levels where copper is toxic varies (Erickson et al., 1972).

Because the biological response is dependent on free ion activities, the effect of chelators is very important. In the laboratory, Anderson and Morel (1978) found that pre-equilibration for 24 hours with the metal chelator, EDTA, was necessary to prevent a dramatic overestimate of copper sensitivity of the organisms. Sunda et al. (1981) found stimulation of phytoplankton growth by organic chelators (EDTA and NTA) appeared to result directly from the ability of these chelators to tightly complex copper without appreciably binding manganese. Fitzgerald and Faust (1963) reported that EDTA decreased algicidal effects of copper on an equivalent basis as naturally occurring metal complexing agents. Upwelling water is likely to have copper present in ionic and thus toxic form (Steemann Nielsen and Kamp-Nielsen, 1970). Sampling from upwelling seawater has shown that an emerging biological community may produce natural chelators which may act to increase phytoplankton growth similar to EDTA and NTA additions (Martin, 1967; Barber and Ryther, 1969; Murphy et al., 1976; and Van der Berg et al., 1979).

The effects of copper toxicity can be partially or completely reversed by appropriate iron, manganese, or chelator additions (Haberman, 1969; Gross et al., 1970; and Sunda et al., 1981). With the aid of thermodynamic
calculations, Sunda et al. interpreted the results as a physiological interaction between copper and manganese in which copper competes for manganese nutritional sites, thereby interfering with manganese metabolism. Experimental results of Braek et al. (1976) with copper, zinc, and manganese indicate a common route for divalent metal ions in *Phaeodactylum tricornutum*. Anderson and Morel (1978), however, did not find the results of copper sensitivity altered with manganese concentrations varied over two orders of magnitude. In some cases, there is a synergistic effect between two metals (copper and silver) to produce greater toxicity (Young and Lisk, 1972) although Petersen (1982) found no clear evidence of interaction between copper ion and zinc ion in producing toxicity.

It seems quite certain that the essentiality of copper to algae arises from its role as a prosthetic group in a number of vital enzymes and proteins, for example, being a component of the molecule plastocyanin of photosynthesis. Its ability to function depends on its intrinsic favorable chemical properties, including the ease of changing its oxidation state. As a transition metal, it has the capacity to complex effectively with a variety of proteins and porphyrins (Frieden, 1981) and with substances that are metal chelators.

Besides being a required nutrient for algae (Myers, 1962; Manahan and Smith, 1973), Rothstein (1959) suggests
a beneficial effect of copper in protecting active sites. Sato (1980) found that copper could bind with toxic compounds to return activity to the enzyme phenolase in spinach chloroplasts. Oxalate, usually toxic, acted as a chelator with the copper.

**Copper Acting on the Life Cycle of Closterium moniliferum**

Copper may act on the amount of nitrogen available to *Closterium moniliferum* and could be a factor in inducing sexual reproduction. Depleted nitrogen has been found to induce sexual reproduction in desmids (Biebel, 1964), and copper has been shown to inhibit nitrate uptake in some organisms (Harrison *et al.*, 1977). Sexual reproduction may be tuned to environmental patterns and may be a mechanism of survival at a time of physiological stress. Although the organisms may reproduce by vegetative cell division only, it may be advantageous, when in the presence of added copper, to begin a different stage in the life cycle. Zygotes are thick walled and should be less susceptible to increases in metal concentration. This research showed that copper acted on the assimilatory process of nitrate uptake and was able to influence the onset of sexual reproduction.

**Adaptation of Organisms to Copper**

An organism must contend with the problem of maintaining a minimal internal concentration of copper sufficient to meet its needs, but, at the same time, prevent the
occurrence of a toxic level of the metal. Genetic selec-
tion for higher tolerance has occurred in some algae
(Russell and Morris, 1970; Foy and Guloff, 1972; and Stokes
et al., 1973). Potential tolerance mechanisms for copper
and algae include exclusion, extracellular binding, precipi-
tation on the outside of the cell membrane, utilization
of nonsensitive intracellular sites, and metabolic shunts
(Stokes, 1979).

Secretion of algal exudates may ameliorate copper
toxicity by extracellular binding (Swallow et al., 1978
and Van der Berg et al., 1979) or changes in pH which would
change the solubility and subsequently the toxicity of copper
(Foy and Gerloff, 1972).

Exclusion of copper (Foster, 1977; Hall et al., 1979)
or differential uptake of copper ions (Jensen et al., 1976;
Bentley-Mowat and Reid, 1977; Bowen and Gunatilaka, 1977;
and Hogan and Rauser, 1981) have also been suggested as
mechanisms for copper tolerance.

There is increasing evidence in higher plants and
fungi (Neurospora and Saccharomyces) that copper tolerance
appears to be due in part to the presence of specific heavy
metal binding proteins, metallothioneins (Rauser and
Curvetto, 1980). These may act as depots for heavy metals
and in addition, serve to bind, and thus sequester, the
toxic metal (Lerch, 1981). Stokes et al. (1977) extracted
a low-molecular-weight (8000 daltons) copper-binding protein
from the cytosol of a copper-tolerant Scenedesmus. The
amount of this protein increased in response to increased copper in the medium. In 1977 Nakajima et al. described a low-molecular-weight copper-binding compound from the cells of *Chlorella regularis*. Of the soluble intracellular copper, 49% was bound to two identifiable protein fractions. These copper metallothioneins may play a role in copper metabolism in absorption, transport, storage, and excretion of the metal.

NITROGEN METABOLISM

Nitrogen metabolism is integrally tied to the cell cycle of *Closterium*. Because nitrogen depletion in the environment induces sexual reproduction (Biebel, 1964; Ichimura, 1971), the effect of copper on nitrogen metabolism, and in particular nitrogen uptake, is of great importance. Nitrate uptake kinetics depend on the species of algae, the concentration of nitrate in the environment and on the nitrogen cell quota (Eppley et al., 1969; Eppley and Renger, 1974). Nitrate is transported into the cell and is very rapidly assimilated by being reduced to nitrite with further reduction occurring to produce ammonia, which is then incorporated into amino acids (Lehninger, 1975).

Because of the requirement for an 8-electron reduction, the most generally accepted sequence for nitrate reduction was long believed to be:

\[ \text{NO}_3^- \leftrightarrow \text{NO}_2^- \leftrightarrow \text{N}_2\text{O}_2 \leftrightarrow \text{NH}_2\text{OH} \leftrightarrow \text{NH}_3 \] (Syrett, 1962).
However, more recent work with algae (Zumft et al., 1969; Aparicio et al., 1971) suggests that only two enzymes catalyze the entire reduction of nitrate to ammonia. The first enzyme is nitrate reductase (NAD(P)H:nitrate oxidoreductase) which catalyzes the reduction of nitrate to nitrite. This enzyme is a pyridine nucleotide-linked molybdoflavoprotein. The electrons are transferred from NAD(P)H to FAD, molybdenum being a required co-factor. The second enzyme is nitrite reductase (NAD(P)H:nitrite oxidoreductase) which catalyzes the reduction of nitrite to ammonia.

Nitrate reductase is believed to be readily accessible in unicellular organisms and nitrate uptake is considered to be closely correlated with nitrite assimilation. In most organisms, the enzyme is not constitutive and synthesis appears to be induced within a few hours when nitrate concentrations are increased in the medium (Harrison et al., 1977).

Most algae will preferentially take up reduced forms of nitrogen such as ammonia or urea rather than nitrate. The suppression of nitrate uptake by ammonium has been reported for natural populations (MacIsaac and Dugdale, 1969, 1972) and for laboratory cultures with Chlorella (Morris and Syrett, 1963; Syrett and Morris, 1963; Grant and Turner, 1969) and with marine phytoplankton (McCarthy and Eppley, 1972). Their work points to substantial suppression of nitrate uptake until ammonia falls below 1 microgram-atom/liter. It is believed that the end
products such as ammonia and amino acids are inhibiting the synthesis or the activity of nitrate reductase (Syrett, 1981).

In contrast, however, Harrison (1973), working with Peridinium and other dinoflagellates and diatoms from a bloom, and Topinka (1978), working with Fucus, found that under saturating nutrient conditions nitrate is utilized at significant rates even in the presence of high levels of ammonia. Bienfang (1975) argues that under nutrient-limiting conditions, with Dunaliella, the ambient ammonia never reaches concentrations high enough to inhibit an induced nitrate-utilization system. No information on use of nitrogen sources in Closterium has been reported.

Nitrate reductase can undergo rapid changes in activity as a consequence of oxidation-reduction reactions in response to the ongoing metabolism in the cell (Hewitt, 1975). Copper has been found to inhibit the activity of nitrate reductase and the uptake of nitrate (Harrison et al., 1977). Shioi et al. (1978b), from their work with Ankistrodesmus, suggest that this may occur because copper inactivates the electron transport of photosystem II and there is not an adequate supply of reductants for nitrate reduction.

Previous work has demonstrated that copper does decrease photosynthesis and chlorophyll content in Chlorella (McBrien and Hassall, 1965); Steemann Nielsen et al., 1969; Rosko and Rachlin, 1977) and in Dunaliella and Phaeodactylum (Overnell, 1975). In Chlorella, Hassal (1963) showed that
photosynthesis was more sensitive to copper additions than was the specific growth rate. No previous work had been done on copper's effects on Closterium.

Closterium is a slower growing organism than many unicellular algae and is not commonly used as a research organism. It has been used in this research because of its metabolic trigger for sexual reproduction. Whether copper directly affects nitrogen availability or indirectly affects photosynthesis, chlorophyll production, or cell division is a complex question. If the effect is mainly on nitrogen metabolism, copper as an environmental pollutant is particularly significant in the life cycle of Closterium moniliferum.

ENVIRONMENTAL SIGNIFICANCE

Since Closterium has been used as an indicator organism, the information on copper's effect gained in the laboratory may be helpful in assessing natural populations.

Copper is one of the chemicals that man cycles more rapidly than nature. The recycling of copper occurs at a ratio of 12:1 (4460 X 10^3 metric tons man-induced rate: 375 X 10^3 metric tons geological rate) (Ketchum, 1972). Anthropogenic effects can be noted in the significant increase this century in the level of copper production and release of copper to the atmosphere (Table I). The level of copper in natural waters is approximately 10^{-7} M or less
### TABLE I

HISTORICAL TRENDS IN GLOBAL COPPER PRODUCTION AND RELEASE OF COPPER TO THE ATMOSPHERE

<table>
<thead>
<tr>
<th>Period</th>
<th>Copper Production Consumption ($10^{12}$ g)</th>
<th>Atmosphere Copper Emission ($10^{10}$ g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pre-1900</td>
<td>58</td>
<td>41.1</td>
</tr>
<tr>
<td>1900-1910</td>
<td>7.5</td>
<td>5.3</td>
</tr>
<tr>
<td>1910-1920</td>
<td>11.3</td>
<td>8.0</td>
</tr>
<tr>
<td>1920-1930</td>
<td>13.5</td>
<td>9.6</td>
</tr>
<tr>
<td>1930-1940</td>
<td>16.3</td>
<td>11.7</td>
</tr>
<tr>
<td>1940-1950</td>
<td>23.8</td>
<td>17.0</td>
</tr>
<tr>
<td>1950-1960</td>
<td>32.4</td>
<td>23.0</td>
</tr>
<tr>
<td>1960-1970</td>
<td>61.4</td>
<td>43.5</td>
</tr>
<tr>
<td>1970-1980</td>
<td>82.5</td>
<td>58.5</td>
</tr>
<tr>
<td>1980-1990</td>
<td><strong>136</strong></td>
<td><strong>97.4</strong></td>
</tr>
<tr>
<td><strong>Total, 3800 B.C.-A.D. 1980</strong></td>
<td><strong>307</strong></td>
<td><strong>315</strong></td>
</tr>
</tbody>
</table>

(from Nriagu, 1979)
One of the effects of copper additions to natural waters is decreased growth and species shifts in the natural community. Sunda et al. (1981) showed a shift from diatoms to green flagellates and Thomas and Seibert (1977) showed a decrease in the proportion of centric diatoms and dinoflagellates.

Some algae have been known to react to nonoptimal environmental conditions by beginning a different stage in their life cycles. An example of the effect of nonoptimal environmental conditions is the resting spore formed by Leptocylindrus denicus when nitrate level was reduced below 0.5 micromoles in CEPEX experiments (Davis et al., 1980). Under conditions of copper stress, the marine dinoflagellate Gonyaulax tamarensis was found to become immotile with no increase in size or cell division (Anderson and Morel, 1978). These cells were termed a "temporary resting stage in the life cycle" which could be revived if the stress due to copper had not been too severe and normal culture conditions were restored.

This thesis shows that a different series of events occurs with the life cycle of Closterium moniliferum. The zygotes produced in the presence of copper do not differ from the zygotes produced without additional copper present. Copper does not affect the germination of the zygotes, which
are surrounded by a thick layered cell wall. Rather, copper stress is significant in the life cycle in the time of onset of sexual reproduction. The occurrence of earlier sexual reproduction as a result of added copper may allow the organisms to survive pulses of toxicants in the environment by reducing the amount of time the cells spend in the vegetative state. This may, however, reduce the utility of Closterium moniliferum as an indicator organism of unpolluted waters.

PURPOSE OF THE RESEARCH

The purpose of this research was to study the sensitivity of the life cycle of Closterium moniliferum to copper. Closterium has an elaborate life cycle which is dependent on environmental changes. Environmental changes for this study were effected by adding copper or by using different sources of nitrogen in the medium.

The parameters of the life cycle that were measured were increase in cell number, onset of sexual reproduction, zygote germination, and uptake of nitrogen. The effect that copper had on each of these parameters was the essence of this research.

Cell division is always critical in a population but it is especially critical here since the process of cell division must occur before sexual reproduction and is therefore closely tied to that stage in the life cycle.

The uptake of the nutrient nitrogen is also closely tied to sexual reproduction; depletion of nitrogen in the
medium, in the form of nitrate, appears to be a trigger for sexual reproduction. A stimulatory or inhibitory response of nitrate uptake or of cell division could alter the timing of the life cycle.

The ultimate question, then, is if the timing of the life cycle can be changed by copper additions in the environment, can this lead to survival of *Closterium* in natural waters by production of resistant zygotes? The success of this laboratory study may give direction to field studies in which phytoplankton are able to survive pulses of toxicants in natural waters.
CHAPTER II

MATERIALS AND METHODS

THE ORGANISMS

The organisms used in this study were strains 96 and 171 of *Closterium moniliferum* from the Portland State University algal culture collection. Both strains are presently stored in the Culture Collection of Algae at the University of Texas (Starr, 1978). Strain 96 was used for all experiments; both strains were used for the scanning electron microscopy work to compare morphology and for the long term growth/uptake experiments to compare the time of onset of sexual reproduction.

**Strain 96**

Strain 96 is a homothallic strain that produces a single zygote in sexual reproduction. It was originally isolated by B.E. Lippert from a soil sample collected by Melvin Goldstein from a marshy area near Shulman's Equipment Co. Rt. 46 in New Jersey on August 21, 1959. The average size of a mature cell of strain 96 is 29 micrometers x 147 micrometers; zygote measurement is 45 x 45 micrometers. It is stored in algal culture at the University of Texas at Austin as LB 2302 *Closterium moniliferum* (Bory) Ehr.
Strain 171

Strain 171 is a homothallic strain that produces twin zygotes in sexual reproduction. It was originally isolated and collected by B.E. Lippert from a farm pond, 2 miles south of a sewage disposal plant, on Fern Hill Road, south of Forest Grove, Oregon in August, 1959. The average size of a mature cell of strain 171 is 43 micrometers x 214 micrometers; zygote measurement is 48 x 54 micrometers. It is stored in algal culture at the University of Texas at Austin as LB 2303 Closterium moniliferum (Bory) Ehr.

ENVIRONMENTAL CONDITIONS

Cultures were grown at 21° C in an Environator under continuous light (cool-white fluorescent) of 4000 lux measured with a GE Triple Range 214 Light Meter. Since Whitton (1968) reported that toxicity of metals, particularly copper, was decreased by light, the cultures were grown in continuous light to eliminate the variable of light/dark cycles. Cultures were manually swirled every 24 hours to facilitate the availability of nutrients to all cells and to prevent the cells from adhering continuously to the glass containers. Fitzgerald (1975) found that continuous shaking of cultures of Scenedesmus resulted in no improved maximum rates of growth as long as relatively low volumes of liquid per flask were used. Also, Bentley-Mowat and Reid (1977) reported that there was not much difference in sensitivity
of species to copper in batch cultures as compared with continuous culture.

PREPARATION OF GLASSWARE AND SOLUTIONS

Glassware

Glassware used for all experimental work was washed in 95% dodecyl sodium sulfate, rinsed 5 times in tap water, and put in an acid wash of 10% HCl solution for 24 hours. After the acid wash, the glassware was rinsed 5 times with tap water and 5 times with deionized water from a Sybron/Barnstead system (nanopure water). After air drying, the glassware was covered with parafilm for storage.

Medium

The organisms were grown in batch cultures in the defined medium FRAQUIL (Morel et al., 1979). (See Appendix for quantities of components and method of preparation.) The water used in the medium (and in all solutions) was nanopure deionized water from a Sybron/Barnstead system. A supplement to FRAQUIL is a vitamin solution, including thiamine HCl, biotin, and B-12. Preliminary work demonstrated that there was no significant increase in growth or zygote production with the vitamins present so vitamins were not added to the growth medium for experimental work. The lack of a vitamin requirement may have been due to the fact that cultures were not continually axenic and the bacteria were supplying the growth factors. Cells were periodically
reisolated to eliminate any bacterial contamination, although axenic cultures were difficult to maintain. Moss (1973) however feels that exponentially growing cultures are probably not affected by bacterial interferences. In his work, he found that algae required the addition of vitamins even with bacteria present and therefore accepted growth measurements obtained with vitamin utilization in non-axenic cultures as valid. Also, Tassigny (1971) considers the results with unialgal desmid cultures in mineral medium to be equally valid to those in axenic culture.

Batch cultures of 150 ml in 250 ml flasks or 250 ml in 500 ml flasks were used to maintain a similar surface to volume ratio and prevent carbon limitation. For short term (2 days) nitrate uptake experiments, 1 liter of culture was maintained in a 2800 ml flask.

To minimize contamination, the medium was filter sterilized. The autoclaving process may add metals and change the activity level of the copper so filtration was the sterilization method preferred. To obtain copper activity close to the predicted value, a Chelex column was eluted with FRAQUIL (without trace metals added) to remove contaminant trace metals (Morel et al., 1978). The trace metals required in FRAQUIL were then added and the medium was filter sterilized (0.45 micrometers Millipore filter). Glassware was not autoclaved due to metal contamination; some bacterial contamination may have occurred with the
acid wash procedure but testing for bacterial growth showed it was minimal.

Copper sulfate was added to FRAQUIL in the experimental flasks and the medium was allowed to equilibrate 24 hours before the organisms were added (when EDTA was the buffer). When TRIS was used as a buffer in systems where copper was added, equilibration was considered to occur almost instantaneously and the organisms were added within a half hour after the copper had been added.

**Preparation of Copper Solution**

Copper sulfate (CuSO\textsubscript{4}·5 H\textsubscript{2}O) was added to sterile deionized water to give a 10\textsuperscript{-1} M solution. This stock solution was stable and could be stored in the refrigerator indefinitely.

**COPPER CONCENTRATION AND SPECIATION**

The toxic effect of copper on organisms is determined by the amount of copper in the ionic form, Cu\textsuperscript{2+}, rather than total concentration of all forms of copper (lab results of Sunda and Guillard, 1976; calculations of Jackson and Morgan, 1978; and field results of McKnight, 1981). EDTA acts as chelator for copper and was present in FRAQUIL at 5 x 10\textsuperscript{-6} M. If the total concentration of copper is less than the total concentration of EDTA there is a defined system with no problems of copper precipitation.
The final dilutions of copper in the experimental flasks in the long term growth experiments were $1.3 \times 10^{-6}$ M, $10^{-7}$ M, and $10^{-8}$ M with EDTA as the buffer. When TRIS was added as a buffer in short term experiments, to a final concentration of $10^{-3}$ M, copper concentrations were increased to $10^{-5}$ M.

Copper activity (concentration of Cu$^{2+}$) was calculated by means of the MICROQL program using constants of components in the experimental medium FRAQUIL (Westall et al., 1976). Computed copper activity (expressed as pCu*, the negative log of cupric ion concentration) in long term experiments with EDTA as the buffer ranged from 12.1 to 15.4. (See Table II and Figure 7.) pCu* as referred to in this work is always calculated and not measured. When the medium was eluted through the Chelex column, the pCu actual values are assumed to be very close to the values calculated with MICROQL (pCu*). This calculation of copper activity does not include organic complexation that may occur during growth, therefore it represents maximum toxicity, i.e. minimum values of pCu* (Sunda et al., 1981).

Swallow et al. (1978) used an ion electrode to measure the depression of cupric ion concentration by phytoplankton exudates and found that only one in 8 algal species excreted copper complexing materials. McKnight (1979) also measured organic products but earlier attempts to measure excretion products of algae in laboratory conditions were not successful because of their low concentration (Fogg, 1977). Darling
<table>
<thead>
<tr>
<th>Copper Molarity</th>
<th>pCu*</th>
</tr>
</thead>
<tbody>
<tr>
<td>1E-3</td>
<td>3.6</td>
</tr>
<tr>
<td>1E-5</td>
<td>5.5</td>
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<tr>
<td>9E-6</td>
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<td>10.8</td>
</tr>
<tr>
<td>2E-6</td>
<td>11.8</td>
</tr>
<tr>
<td>1.3E-6</td>
<td>12.1</td>
</tr>
<tr>
<td>1E-6</td>
<td>12.3</td>
</tr>
<tr>
<td>1E-7</td>
<td>13.4</td>
</tr>
<tr>
<td>1E-8</td>
<td>14.4</td>
</tr>
<tr>
<td>9.97E-10</td>
<td>15.4</td>
</tr>
</tbody>
</table>

[EDTA] = 5 \times 10^{-6} \text{ M}
Figure 7. Negative log of copper concentration in the medium FRAQUIL vs. calculated copper activity. Arrow indicates EDTA concentration in the medium.
(1979) found extracellular release of organic products of *Selenastrum capricornutum* had a small but significant effect on ionic copper.

**pH MEASUREMENT**

pH was routinely measured with an Ionalyzer specific ion meter model 404. The beginning pH of autoclaved FRAQUIL and of Chelexed FRAQUIL with trace metals added was the same (7.05). When the various amounts of copper were added to the experimental flasks, the pH did not vary significantly. The pH at the end of the experiments was not alkaline indicating the flasks were not carbon limited. The effects seen in the experiments were considered to be the result of copper treatment, not of pH changes.

The pH increased during zygote maturation (up to 7.87) and remained high during germination (7.94).

The solubility diagram of copper (Figure 8) shows the cupric ion to be available at the experimental pH.

**MONITORING CELL GROWTH**

**Cell Counts**

A Sedgwick-Rafter counting chamber was used for cell counts; the cells present in the total milliliter were counted so non-uniform distribution within the counting chamber would not contribute to counting error (McAlice, 1971). For long range (3 weeks) experiments, initial cell density ranged from 16 to 112 cells/ml; final cell density reached
Figure 8. Solubility diagram of copper. The solid line surrounding the shaded area gives the total solubility of Cu(II) which up to pH value of 6.96, is governed by the solubility of malachite, $\text{Cu}_2(\text{OH})_2\text{CO}_3(s)$. In the low pH region azurite, $\text{Cu}_3(\text{OH})_2(\text{CO}_3)_2(s)$ is metastable but may become stable at higher $C_T$. Above pH 7 the solubility is controlled by the solubility of CuO (tenorite). The predominant species with increasing pH are $\text{Cu}^{2+}$, $\text{CuCO}_3(\text{aq})$, $\text{Cu(CO}_3)_2^{2-}$ and hydroxo copper(II) anions. $C_T = 10^{-2}$ (Stumm and Morgan, 1970).
a maximum of 6100 cells/ml. Dilutions were made for counts when cell number exceeded 600 cells/ml. The counting chamber and cover glass were cleaned with water and rinsed in 95% ethanol after each count.

**Cell Morphology**

Quantitative assessment of changes of gross cell morphology was done during long term growth experiments by grouping the cells into three categories: those cells of normal size and shape (Figures 9 and 10); those of abnormal size and shape that completed cytokinesis (Figures 11 and 12); and those incapable of completing cell division (Figures 13 and 14). Those cells incapable of finishing cell division were recognized when 2 mature symmetrical cells were contained within a continuous cell wall. The latter two categories of cells are considered to be under stress due to copper.

**Sexual Reproduction**

The onset of sexual reproduction in long term growth experiments was monitored. The frequency of sexual reproduction was determined by counting the number of zygotes and the number of potential zygotes (abortions) and comparing that number to the total number of cells with this equation:

\[
% \text{ Sexual Reproduction} = \frac{2(\# \text{ Zygotes} + \text{ Abortions})}{\# \text{ Cells} + 2(\# \text{ Zygotes} + \text{ Abortions})} \tag{2}
\]
Figures 9 and 10. Photomicrographs of strain #96 (Top) and strain #171 (Bottom) without copper treatment. The cells shown here are immature cells pairing before sexual reproduction. LM 430 X.
Figures 11 and 12. Photomicrographs of abnormal cells of strain #96. LM 430 X.
Figures 13 and 14. Photomicrographs of strain #96 that have not completed cytokinesis. LM 430 X.
Determination of Cell Chlorophyll

The amount of chlorophyll was determined by the fluorometric method of Strickland and Parsons (1972). There was no significant difference in chlorophyll amounts using the processes of filtration or centrifugation, so, for convenience, the pellet from a sample centrifuged for nitrate analysis was used.

One capillary drop of saturated MgCO$_3$ was added to prevent acid conditions which would decompose the chlorophyll to give phaeophytin pigments. Ten ml of 90% acetone was the total amount used for extraction. In dim light, a few ml of 90% acetone were added to the pellet in the centrifuge tube; this was transferred to a grinding tube. The centrifuge tube was rinsed with more acetone and the rinse was added to the grinding tube. Cells were ground for 2 minutes at the highest speed on an Eberbach Power-Stir. The ground cells were transferred back to the centrifuge tube for overnight extraction. The grinding tube and pestle were rinsed until the total 10 ml of 90% acetone was added. The tube was capped and covered with aluminum foil and refrigerated for 24 hours.

The amount of chlorophyll was determined by use of a Turner fluorometer which was calibrated using a Bausch and Lomb Spectronic 100. On the Spectronic 100, readings were taken at wavelengths of 665, 645, and 630 nm for use in the Strickland and Parsons formula:
chlorophyll a = 11.6E665 -1.31E645 -0.14E630 (3)

where E = extinction at these wavelengths.

Using this same sample, an F factor was determined on the fluorometer:

\[
F \text{ factor} = \frac{\text{chlorophyll a (calculated from (3))}}{\text{reading on fluorometer}}
\] (4)

Using this F factor value, subsequent chlorophyll amounts were determined by a single reading on the fluorometer.

The filters used in the fluorometer were # 47B as the primary filter and # 2A15 as the secondary filter (Turner). Cell counts were taken of the same samples so the amount of chlorophyll per cell could be calculated.

Carbon Uptake as a Function of Copper

The cultures contained TRIS so equilibration would occur quickly (within the first 5 minutes) when copper was added (Anderson and Morel, 1978). Copper was added to the test flasks to give a final concentration of $10^{-5}$ M. To the copper treated flask and a control flask (100 ml of culture) were added 100 microliter of $^{14}$carbon (1 micro-curie/ml). For background count, a third flask without $^{14}$carbon or copper added was run in parallel. Triplicate samples were taken at 16 hours and 40 hours. Ten ml of sample were filtered onto a Whatman GF/C glass microfibre filter (24 mm). The filter was placed in a grinding tube for chlorophyll extraction. The method for chlorophyll determination was the same as above. After overnight
extraction, the samples were centrifuged and chlorophyll readings were done of the supernatants. The pellet and supernatant were then recombined in a scintillation vial and the acetone was allowed to evaporate under an intense light. When dry, 8 ml of scintillation liquid (Scinti Verse I, Fisher Scientific Company) were added. Radioactivity counts were taken for 5 minutes on a Beckman Scintillation Counter.

The filtrates of the samples were analyzed for fixed $^{14}$C carbon as an indicator of excreted organics. The pH of the filtrate was lowered to pH 2 by adding 2 drops concentrated HCl. Air was bubbled through the liquid continuously for a few hours, with an occasional shaking of the sample to wash the sides of the scintillation vial. Eight ml of scintillation liquid were added and samples were read as above.

Gloves were used throughout the procedure and all materials were soaked after use for 24 hours in Count-off (New England Nuclear) and thoroughly rinsed with water.

NITROGEN UPTAKE AS A FUNCTION OF COPPER

Long Term Nitrate Uptake and Growth

Growth experiments were run for three weeks to examine the relationship between nitrate taken up by the cells and the time of induction of sexual reproduction. Control and experimental flasks were run in duplicate. Experimental flasks had pCu* of 12.1 - 14.4. Control flasks had pCu*
of 15.4. Inocula were added to 250 ml of FRAQUIL containing 100 micromoles/l nitrate. Cell counts and samples for nitrate determination were taken every 2 days until sexual reproduction or the stationary phase of growth occurred.

Short Term Nitrate Uptake

When cultures were depleted of nitrate, a spike of nitrate (final concentration of 10-20 micromoles/l) was added. Subcultures of 75 ml were taken and dispensed into flasks designated as controls and experimental flasks. A cell count was taken and pH was measured. To the experimental flasks, copper concentrations were added. The flasks were swirled left to right and right to left for 15 minutes. The completion of this mixing was considered zero time and a sample was taken for nitrate amount. The amount of nitrate was found to be consistent in all flasks at 0 time. Samples were taken periodically for 48 hours (at 5, 19, 24, and 48 hours) until nitrate was virtually depleted in the control flasks.

Other short-term uptake experiments were done with the nitrate spike being added 5 hours before the copper was added. It was necessary to use TRIS as a buffer to provide rapid equilibrium.

Long Term Ammonium Uptake and Growth

To determine if copper more directly affected nitrate uptake than ammonia uptake, long term growth experiments were also done with ammonia as the nitrogen source. To
test for ammonia uptake, nitrate was not added to the media; 
\( \text{NaN}_3 \) was replaced by the same molarity \( (10^{-4} \text{ M}) \) solution of \( \text{NH}_4 \text{Cl} \). The pH of both solutions were adjusted to 6.96. The final molarity of copper added was \( 1.35 \times 10^{-6} \) which gives a \( \text{pCu}^* \) in the nitrate media of 12.1. Less than 1 micromole nitrate was present in the FRAQUIL with \( \text{NH}_4 \text{Cl} \). The procedure here was the same as for long term nitrate uptake.

**Measurement of Nitrate**

Each time a sample was taken for nitrate determination, a cell count was also done. The sample for nitrate determination was centrifuged on a Universal Model U Centrifuge at approximately 750 g for 20 minutes. Nitrate determination of the supernatant was done immediately or it was frozen for later testing. Nitrate was measured using the method proposed by Strickland and Parsons (1972) and later modified by Eppley (1978). It is assumed that all nitrate not in the medium has been taken into the cells.

A cadmium-copper reduction column was used to reduce the remaining nitrate in the supernatant to nitrite; nitrite was then measured spectrophotometrically. When a sample was reduced, part of it was sent through the column to act as a pre-wash to assure a more accurate reading. Although the nitrate reduction column used was a scaled down model for use with small samples (Rueter, 1979), a 5 ml wash was not adequate to completely wash out the previous sample.
Reducing samples of 0.1 micromole nitrate after a 100 micromole sample gave readings of 25 x the expected value due to carryover. Washes of 20 ml with a 5 ml sample collected for nitrate testing resulted in consistent readings.

All samples to be run through the column were made basic with concentrated ammonium chloride (100 microliters per 5 ml). This was done because basic solutions tend to clear the column more thoroughly (Strickland and Parsons, 1972). As soon after reduction as possible, 100 microliters of sulfanilamide were added to the 5 ml sample to form a diazonium salt with the nitrite. After 3 minutes reaction time, 100 microliters of NEDA (N-l-napthyl-ethylene-diamine dihydrochloride) solution were added to form the red azodye. Complete color development occurred within 10 minutes and samples needed to be read within 2 hours. Samples were read at 543 nm on a Bausch and Lomb Spectronic 100. Dilutions were made to read samples above 30 micromoles NO₃⁻ (extinction levels above 0.900).

Because of the difference between columns, a standard curve (with standards of 1, 5, 12.5, 25, 50, and 100 micromoles nitrate/l) was run on each column each time (Figure 15).

FRAQUIL had 100 micromoles of nitrate present before inoculation of cells. The amount of nitrite in the medium was measured by testing for nitrite without the medium passing through the column. The amount of nitrite in FRAQUIL was negligible.
Figure 15. Absorbance (543 nm) vs. concentration of nitrate. The two lines represent standard curves from two different nitrate reducing columns.
An ideal rate of passing through the column was 5 ml in less than 5 minutes. If a column became sluggish, moving the liquid through it by using forced air cleared the column temporarily to obtain a faster rate. When sluggish, or after 100 samples were reduced, the columns were reactivated by placing the cadmium filings in 5% v/v HCl, rinsing them in distilled water until the pH was greater than 5, and then saturating the filings in a 2% copper solution. The column was then repacked (Strickland and Parsons, 1972).

A solution of dilute NH₄Cl was used to pass through the column between samples that had great differences of nitrate concentration. The dilute NH₄Cl was also used on the columns in storage; parafilm over the open top of the glass columns assured that the columns would not evaporate to dryness when not in use.

Measurement of Ammonia

The amount of ammonia present in the medium was measured with an Orion ammonia electrode (model 95-10). While being constantly stirred with a magnetic stirrer, a 25 ml sample was treated with 2.5 ml of 10 M NaOH. The reading was recorded when it became stable.

Determination of Nitrate Reductase Activity

The basic procedure for determining activity of nitrate reductase (ferrocytochrome:nitrate oxidoreductase EC. No. 1.9.6.1) was as follows. Samples were harvested by
centrifugation at 2800 g for 20 minutes or by the process of filtration. The supernatant was discarded and the cells were homogenized with a glass homogenizer in ice in a homogenizing solution consisting of 3 ml 0.15 M KPO₄ buffer (pH 7.0), 0.6 ml sodium dithionite or dithiothreitol, and 20 mg polyvinylpyrrolidone. The sodium dithionite (or dithiothreitol) was used to maintain integrity of the enzyme (Lewis et al., 1982) and the polyvinylpyrrolidone was used to eliminate interference from phenols (Klepper and Hageman, 1969; Stock and Lewis, 1982).

The assay is that of Lowe and Evans (1964) wherein the reaction velocity was determined by measuring the production of nitrite in a dithionite/methyl viologen system. One unit produces one micromole of nitrite/minute at 30° C and at pH 7.0 under specified conditions.

Test tubes were placed in a 30° C bath. To these tubes were added 0.1 ml phosphate buffer, 0.1 ml 0.02% methyl viologen, 0.1 ml 23 mM sodium dithionite prepared in 48 mM sodium bicarbonate solution, 0.1 ml 0.10 M NaNO₃ and 0.1 ml 0.05 M MgSO₄. One tenth of a milliliter of copper sulfate (10⁻³ to 10⁻⁹ M) was added to the experimental tubes. After 5 minutes of temperature equilibration, 0.1 ml enzyme or cell extract (0.1 ml H₂O for blank) was added and incubated for 30 minutes. The reaction was stopped by vigorous aeration until blue color was completely removed. 0.5 ml sulfanilamide (58 mM in 3N HCl) and 0.5 ml NEDA solution (0.39 mM N-1-napthyl-ethylene-diamine dihydrochloride) were
quickly added to the test tube. 1.5 ml glass distilled water was added and 10 minutes time allowed for color development at room temperature. Samples were read at 543 nanometers.

Using the known activity of purified enzyme (E.coli) as a reference, the filtered sample yielded activity of $2.9 \times 10^{-5}$ units cell$^{-1}$ minute$^{-1}$ whereas the centrifuged sample yielded activity of $1.7 \times 10^{-5}$ units cell$^{-1}$ minute$^{-1}$. When copper was added in preliminary experiments to give a final molarity of $10^{-3}$ M copper, no enzyme activity was produced. This indicates that copper does directly affect the activity of nitrate reductase, which was indicated in the long term and short term nitrate uptake experiments.

Enzyme activity from cell extracts of Closterium moniliferum did not always occur, however, so modifications of the basic procedure above were tried. Assuming the lack of activity was due to the enzyme being attached to the membrane, drastic steps were taken to disrupt the cells. Besides homogenization, the French press, Triton X, and glass beads in a mortar and pestle in various combinations were used. To determine if the French press procedure was too harsh a procedure and was destroying the activity of the enzyme, purified enzyme (obtained from E.coli, Sigma) was run through the French press twice. There was a loss of 44% of the activity. With the homogenization process, the purified enzyme activity decreased by 42%.
NADH was not an acceptable electron donor for purified enzyme or cell extract; both methyl viologen and benzyl viologen were suitable electron donors. The addition of FMN did not affect enzyme or extract activity.

The solution of sodium dithionite needed to be prepared fresh just before use. Its freshness was tested by observing a deep blue color formed when combined with methyl viologen which indicated a reducing environment.

To determine if the experimental procedure was proper, tests with purified enzyme were always run when testing cell extract.

Since copper was to be added to the test tubes to test its effect on enzyme activity, TRIS buffer was necessary for rapid equilibration. One-tenth of a milliliter of $10^{-3}$ TRIS buffer was added to the reaction tubes. This addition did not significantly decrease enzyme activity.

Other variables that might have contributed to the inconsistency of the procedure were tested. These included temperature, time (from 1 minute up to 24 hours), excess and minimum sodium dithionite or dithiothreitol, and use of cultures that were nitrate depleted (with nitrate spike added to induce enzyme production) or cultures that were actively growing. Sodium salts were removed since Tischner (1976) reported that nitrate reductase activity did not occur in a K-Na-PO$_4$ buffer. These changes did not produce activity in the supernatant; often, some nitrate reductase activity was present in the cells. Supernatants were tested...
after each step in the procedure. Simple homogenization of a mixed algal culture produced enzyme activity however. This indicates that there is a special problem in releasing the enzyme from Closterium moniliferum cells or in maintaining its activity.

PHOTOGRAPHY

Light Microscopy

Gross cell morphology was observed with a Zeiss microscope and recorded with a Zeiss c35 camera. Film used was black and white Plus-X-Pan ASA 125/22 DIN. Magnification of the cells was 100 X or 430 X.

Scanning Electron Microscopy (SEM)

The cells were grown in liquid medium and therefore needed to be attached to a substrate for SEM work. Polycarbonate filters were chosen because they were not soluble in the ethanol used for dehydration purposes and they were able to survive the critical drying process. They were also found to be an unobtrusive background for the organisms as compared to glass filters. While the cells were filtered, they were rinsed with FRAQUIL and never were allowed to become completely dry. The filter, with cells attached, was placed in a fixative composed of 3-4% formalin, 0.75% glutaraldehyde, and phosphate buffer of pH 7.0 (Fahrenbach, personal communication). The preparation of the fixative and all manipulations with the fixative were
done under a hood because of the toxicity of the formalin and glutaraldehyde.

After overnight fixation at room temperature, the samples were rinsed with buffer and sequentially dehydrated through 20%, 40%, 60%, and 80% ethanol solutions in 5 minute intervals. The samples were then transferred to absolute alcohol until they could be placed in the transitional fluid, Freon. The transitional fluid was necessary to prevent morphological changes in the cells by eliminating the liquid-gas interfaces as the samples were processed through the critical point drying. The critical point dryer used was a Model Hummer 1 by Technics. When dry, the filters were attached to SEM studs with double sticky tape and then coated with gold. Coating with gold makes the sample conductive for observation. Photographs were taken with Polaroid 55 positive/negative film. Magnifications ranged from 100 X to 5000 X on an AMR Model 1000 scanning electron microscope using 20 kv.

PRESENTATION OF DATA

Graphs were plotted on an interactive Digital Plotter using a Tektronix 4051 with a program for X-Y plot and the calibration curve for nitrate standards modified by Jim Sweet (personal communication).
CHAPTER III

INHIBITION OF GROWTH AND CELL DIVISION BY EXCESS COPPER

INTRODUCTION

Cell division was definitely inhibited by the addition of excess copper to batch cultures of *Closterium moniliferum*. There was an increased lag phase in long term experiments and delayed onset of sexual reproduction. The increased lag phase indicates the effect of copper is on the process of cell division itself. Chlorophyll measurements and ¹⁴C uptake show, however, that these aspects of metabolism were also affected.

A toxic sublethal amount of copper causes an increased lag phase, which can eventually be overcome with the culture reaching a normal growth rate. Smaller amounts of copper may be stimulatory to growth and beneficial to the organisms. The evidence for copper's effect is presented in this chapter as growth patterns, growth rate, cell division and morphology, chlorophyll content and ¹⁴C carbon uptake.

The effect of copper was examined by long term experiments of 3 weeks duration where cell division, nitrogen (nitrate or ammonia) remaining in the medium, and onset of sexual reproduction were monitored. It was shown here
that copper affected all of these processes. The first process is discussed in this chapter; the latter two are presented in Chapter IV.

Micrographs were used to examine aberrant morphology to determine if there was a correlation between changes in cell number and morphological changes. Further study of morphology is presented in Chapter V.

RESULTS AND DISCUSSION

Growth Patterns

When copper was added to batch cultures of *Closterium moniliferum*, initial cell division was inhibited and an increase in lag phase was observed at pCu* below 14.4 (Figure 16). FRAQUIL medium, with no added copper, has a pCu* of 15.4. (pCu* is calculated activity of copper ion = -log [Cu_{aq}^{2+}].) Cultures with lower beginning pCu* were able to overcome this increased lag phase and were able to attain high maximum cell number. In some cases, the cultures with increased lag phase at pCu* 12.1, surpassed the maximum cell number of those cultures with no copper added (Strain 96, Figure 17). This suggests an adaptability of the cells with time with the effect of copper occurring during the lag phase. It is possible that the lag phase may be overcome by exudation from living cells or leaching from dead cells to render the heavy metal less toxic by chelation (Braek *et al.*, 1976).
Figure 16. Number of cells per ml vs. time (days) in strain 96 of *Closterium moniliferum*. Growth was inhibited at pCu* 12.1 in the first days of the growth curve.
Figure 17. Number of cells per ml vs. time. Strain 96 of Closterium moniliferum.
The stimulatory effect of pCu* 14.4 and the similar responses at pCu* 13.4 and 15.4 in strain 96 exemplify the typical dose response of biological organisms to a trace metal (See Figure 1).

In strain 171, the addition of small amounts of copper did not inhibit growth. The final cell density was highest in this instance at pCu* 14.4 (Figure 18).

Figure 19 includes error bars on the growth curve; these may appear to be rather large in some instances but this is due to the fact that one of the duplicate cultures sometimes lagged a day behind the other but with the same pattern (as represented in Figure 20).

Increasing the copper concentration from a pCu* of 15.4 to 14.4 generally stimulated cells and gave an increase in cell division (Figure 21). Further increases in ionic copper above pCu* of 14.4 were inhibitory and decreased the rate of cell division. These effects are most apparent during the first week of growth and are particularly dramatic by day 7 (Figure 22). Figure 22 again demonstrates the typical dose-response relationship mentioned in Chapter I (Figure 1) where low concentration of a metal ion enhances a physiological process but a higher concentration becomes inhibitory and eventually toxic.

Once sexual reproduction begins in a batch culture, the growth rate, defined in terms of cell number over time, slows as shown in Figure 17. This slow down is in part attributed to the stationary phase and in part to induction
Figure 18. Number of cells per ml of strain 171 of Closterium moniliferum vs. time. The addition of small amounts of copper did not inhibit growth; the final cell density was highest at pCu* 14.4.
Figure 19. Number of cells per ml vs. time (days) in strain 96 of Closterium moniliferum. Error bars represent duplicate cultures.
Figure 20. Number of cells per ml vs. time (days) in strain 96 of *Closterium moniliferum*. Duplicate cultures @ pCu* = 15.4.
Figure 21. Number of cells vs. time (days) in strain 96 of Closterium moniliferum. This demonstrates the typical dose-response of an organism to a trace metal. Cell growth was stimulated with the addition of small amount of copper but became inhibited at greater copper activities.
Figure 22. Number of cells per ml at day 7 vs. pCu* in strain 96 of Closterium moniliferum. Small additions of copper appear to be stimulatory while larger amounts become inhibitory. This is expected from the suggested dose-response of organisms to trace metals (see Figure 1).
of sex precluding vegetative cell division in any given pair of cells.

The stimulation of cell division at pCu* 14.4 varied in different experiments, (e.g. Figure 16 versus Figure 17), suggesting a narrow range where stimulation can occur or possible contamination of a sensitive system. The lengthening of the lag phase at pCu* 12.2 was reproducible in both strains however. When stimulation of cell division occurred, nitrate uptake increased and sexual reproduction occurred earlier.

The longer lag phase evident at low pCu* is a result of fewer cell divisions, but it has not been determined if this occurs because copper is acting on the cell division process itself or because copper affects nutrient uptake. Nutrient uptake inhibition in turn slows growth since cells must reach a minimum size before cell division can occur. Conversely, slower growth could slow nutrient uptake.

Growth Rate

*Closterium moniliferum* cells are unicellular and are rather large (maximum size of one strain used was 214 micrometers in length); therefore, they do not have a particularly rapid metabolism or growth rate. Calculating the growth rate is complicated because of the occurrence of sexual reproduction occurring. Once zygotes begin to be formed, those cells involved as gametes do not divide vegetatively so cell number decreases or at least cell number increase is slowed.
Within the first days of these growth experiments, no sexual reproduction is involved but the lag phase is included so growth rate would be low in all cases. Calculating the growth rate during different stages of the growth curve does show that cultures treated at \( pCu^* \) 12.1 can reach an equivalent growth rate and maximum cell number as cultures at \( pCu^* \) 15.4 (Table III). The specific growth rate was computed using natural log (\( \log_e 2 \)) so the rate corresponds to the number of divisions per day.

The growth rate and final cell densities again suggest adaptability to the presence of added copper. Steemann Nielsen and Kamp-Nielsen (1970) reported that normal growth rates were re-established in *Chlorella pyrenoidosa* after extended lag phases due to sub-lethal doses of copper. Work with toxic metals has generally shown decreased growth of algae (Sunda, 1975; Sunda and Guillard, 1976; and Jensen *et al.*, 1976). Sunda (1975) reported that the specific growth rate was a function of \( pCu \) and Morel *et al.* (1978) found a prolonged lag phase in their work with *Skeletonema costatum*. The work here shows the prolonged lag phase due to copper's effect as well as the re-establishment of the normal growth rate.

In looking for a genetic change to copper tolerance, Darling (1979) expected that cells would grow better when subcultured at the same \( pCu \). However, he found that subcultures of *Selenastrum capricornutum* had a lower specific
### TABLE III

SPECIFIC GROWTH RATE (DAY$^{-1}$)

<table>
<thead>
<tr>
<th>pCu*</th>
<th>DAYS 1-7</th>
<th>DAYS 8-13</th>
<th>$\mu_{max}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>15.4</td>
<td>.46</td>
<td>.32</td>
<td>.76</td>
</tr>
<tr>
<td>14.4</td>
<td>.55</td>
<td>.08</td>
<td>1.4</td>
</tr>
<tr>
<td>13.4</td>
<td>.44</td>
<td>.36</td>
<td>.80</td>
</tr>
<tr>
<td>12.1</td>
<td>.35</td>
<td>.54</td>
<td>.75</td>
</tr>
</tbody>
</table>
growth rate. He postulated, then, that the lower growth rate may have been due to an increase in internal copper concentration or the accumulation of copper on the outside of the cell wall preventing cell division as suggested by Steemann Nielsen and Wium-Andersen (1970). The lag phase shown here may be due to this action of copper.

Cell Division and Morphology

In 1950, Hughes reported that copper interfered with the process of mitosis at prophase and the work of Kanazawa and Kanazawa with Chlorella (1969) showed that cell division was inhibited by copper at cytokinesis. Further research since then indicates that this effect of copper is a rather general phenomenon. The fact that copper affects cell division is particularly important here because cell division and sexual reproduction are closely linked in the strains of C. moniliferum studied. Sexual reproduction takes place before vegetative cells mature, so cell division must occur shortly before sexual reproduction begins.

In Closterium moniliferum, cell division is definitely blocked by copper. In Closterium, cells normally separate when immature and non-symmetrical. Since sexual reproduction in these strains only occurs between immature cells, sexual reproduction is also blocked. When cells are exposed to copper, (pCu* 12) light micrographs (Figures 23-27) and scanning electron micrographs (Figures 28-30) show mature cells still joined together.
Figures 23 and 24. Strain 96 of Closterium moniliferum. Cells unseparated after treatment with copper. LM 430 X.
Figures 25 and 26. Strain 96 of *Closterium moniliferum*. Cells unseparated after treatment with copper. LM 430 X.
Figure 27. Strain 96 of Closterium moniliferum. Enlargement of unseparated cells after treatment with copper. LM approximately 800 X.
Figure 28. Strain 96 of *Closterium moniliferum*. Cells unseparated after treatment with copper. SEM 500 X.
Figure 29. Cells treated with copper. Cell on the bottom left did not divide after cell growth. Cell at middle right shows distortion at cell isthmus. SEM 300 X.
Figure 30. Strain 96 of Closterium moniliferum. Cells unseparated after treatment with copper. SEM 500 X.
Another morphological effect that results from uncoupling of cell growth and cell division is abnormal cells. It is improbable that these distorted aberrant cells (Figures 12 and 31-34) would be able to divide; if this is so, it would further decrease the growth rate and the ability to have sexual reproduction. The scanning electron micrograph of Figure 31 illustrates the old semi-cell that is normal and the new semi-cell that results from abnormal growth. The cell shown in Figure 32 indicates that the abnormal semi-cell may be able to divide to give a totally distorted cell with no normal morphology evident. This would also indicate that mitosis is not the process affected by excess copper.

Kiermayer (1970) in his work with the desmids Cosmarium and Micrasterias found that cell wall deposition continued in the absence or interruption of cell expansion and most significantly, that the new wall material was deposited in a predetermined pattern. The accumulation of wall material at any region may be related to whether that particular region of wall would have continued to expand under normal circumstances.

Kiermayer also found that if cells were prevented from undergoing any expansion at all after cytokinesis, the septum itself developed a pattern of uneven wall thickening, again related to the potential form of the expanding semi-cell. This appears to be the effect (Figures 33 and 34) in Closterium treated with copper.
Figure 31. Strain 96 of Closterium moniliferum. Note how the new semi-cell is distorted and does not give a symmetrical appearance. SEM 500 X.
Figure 32. Abnormal cell of *Closterium moniliferum*. SEM 500 X.
Figures 33 and 34. Strain 96 of Closterium moniliferum. Cells that have been treated with copper. Note the distorted cell shape particularly the bulbous shape at the isthmus. LM 430 X.
Chlorophyll Content as a Function of Copper

The total amount of chlorophyll as produced by a population of cells was decreased when the cells were in the presence of increased copper. Although the total amount of chlorophyll produced in 48 hours was less in those populations treated with more copper, the cells also did not divide so chlorophyll/cell was higher (Table IV). Copper may have affected cell division directly or the effect may have been indirect by less chlorophyll production and therefore less photosynthesis and energy production.

14Carbon Uptake as a Function of Copper

The amount of 14carbon taken up per cell was decreased when cells were in the presence of increased copper (pCu* 5.5). Cells without excess copper had nearly twice the assimilation ratio minute⁻¹ mg chlorophyll⁻¹.

CONCLUSION

A toxic sublethal amount of copper causes an increased lag phase, which can eventually be overcome with the culture reaching a normal growth rate. This suggests adaptability of the organisms to the presence of copper. Smaller amounts of copper may be stimulatory to growth and beneficial to the organisms as suggested by the dose-response of physiological processes to metal ions.

Cell division was definitely inhibited by the addition of excess copper to batch cultures of Closterium
## TABLE IV

**AMOUNT OF CHLOROPHYLL/CELL AS A FUNCTION OF pCu**

<table>
<thead>
<tr>
<th>pCu*</th>
<th>mg chlorophyll/cell before treatment</th>
<th>mg chlorophyll/cell 48 hours after copper was added</th>
</tr>
</thead>
<tbody>
<tr>
<td>15.4</td>
<td>6.2 x 10^{-4}</td>
<td>7.9 x 10^{-4}</td>
</tr>
<tr>
<td>14.4</td>
<td>6.8 x 10^{-4}</td>
<td>12 x 10^{-4}</td>
</tr>
<tr>
<td>12.1</td>
<td>6.3 x 10^{-4}</td>
<td>12 x 10^{-4}</td>
</tr>
</tbody>
</table>
moniliferum. Evidence for this is the growth patterns at different pCu*, with the longest lag phase at pCu* 12.1. Light micrographs and scanning electron micrographs show distorted and unseparated cells which indicate copper affects cell wall deposition or mitosis. Inhibition of either of these processes would increase the lag phase.

The evidence for adaptability is presented both in the growth patterns and in the specific growth rates. After an increased lag phase, the cultures growing at low pCu* were able to overcome the toxic effects and reach a growth rate equal to that at a higher pCu*.

The initial decreased growth rate could be due to a number of processes. Nutrient uptake may be inhibited and therefore slow growth. It is unlikely that chlorophyll production is the limiting factor since chlorophyll/cell remains high with reduced cell division. The micrographs suggest uncoupling of growth and cell division. The effect could be on the cell's outer surface or on the cell membrane. Chapter IV addresses the uptake of nitrogen in growth experiments and Chapter V addresses other morphological changes.
CHAPTER IV

INHIBITION OF NITROGEN METABOLISM BY EXCESS COPPER

INTRODUCTION

Growth rate has been shown to be a function of the intracellular concentration of the macronutrients of silicon (Paasche, 1973), phosphorus (Fuhs, 1969), and nitrogen (Caperon and Mayer, 1972a and b). The toxic effect of metals (in this work copper) and the limitation of macronutrients (in this work nitrogen) interact to affect the total metabolism of the cell.

The life cycle of Closterium moniliferum is influenced by its external environment in the media and in natural waters. Depletion of nitrate in the media caused populations of strain #96 of Closterium moniliferum to begin sexual reproduction. Thus, if the trace metal copper acts on the nitrogen metabolism of Closterium, it could have an ultimate effect on the inducement of the sexual stage of the life cycle.

Since both cell division and nitrate depletion are necessary for sexual reproduction, the hypothesis proposed was that with fewer opportunities for cell division and inhibited nitrate uptake due to the effect of copper, there would be less or delayed sexual reproduction at a lower
pCu. This hypothesis was generally found to be true. The exception was where copper appeared to be stimulatory in its effect on cell division and nitrate uptake (pCu* = 14.4) which ultimately led to more frequent and earlier sexual reproduction.

This chapter examines long term (3 week) and short term (48 hours) nitrate uptake and the timing of the onset of sexual reproduction as a result of changes in the rate of uptake due to added copper. Rate of zygote germination and abortions are also examined.

RESULTS AND DISCUSSION

In this work, copper inhibited uptake of nitrate. When this uptake was delayed, sexual reproduction was delayed. Previously, (1977) Harrison et al. showed that copper inhibited nitrate uptake and the activity of the assimilatory enzyme, nitrate reductase, in natural algal populations. This action of copper on the nitrogen metabolism could affect sexual reproduction here in Closterium, but it is also probable (Chapter III) that sexual reproduction may also be affected by copper acting on the process of cell division itself.

Nitrate Uptake in Long Term Experiments

Most evidence points to the membrane as the primary site of copper toxicity (Rothstein, 1959). This indicates that nutrient uptake could be the primary process affected.
Although the uptake of many nutrients could be affected by added copper, nitrate was chosen to be the measured parameter because of its close relationship with sexual reproduction in desmids (which was demonstrated by Biebel in 1964) and shown in my early work with *Closterium moniliferum*.

Nitrate uptake experiments confirmed the inhibition of nitrate at low pCu* and the stimulation at pCu* 14.4 with strain #96. When pCu* = 14.4, uptake of nitrate (Figure 35) was more rapid than at pCu* of 12.1, 13.4, or 15.4, resulting in a medium depleted of nitrate earlier in the experiment. This apparent stimulation of nitrate uptake and subsequent early sexual reproduction must be sensitive because not all repeat experiments showed these results. This may be due to a slight difference in pCu due to autoclaving of the media in the first experiments, to different general cell conditions, or to contamination of a sensitive system. Inhibited nitrate uptake at the lower pCu* was reproducible however.

At pCu* 12.1, inhibition of nitrate uptake delayed the onset of sexual reproduction whereas stimulation at pCu* 14.4 caused early sexual reproduction in strain #96 (Figure 36). The number of cells per milliliter at the time of onset of sexual reproduction was not a critical factor.

Strain #171 did not display this expected nitrate depletion before sexual reproduction. Other researchers
Figure 35. Nitrate remaining in the medium vs. time (Days). Sexual reproduction occurred when nitrate was depleted in strain 96 of *Closterium moniliferum.*
Figure 36. Cell number and nitrate uptake vs. days. Note the correlation between nitrate depletion and onset of sexual reproduction.
working with _Closterium sp._ did not find it essential for the nutrient medium to be nitrogen-deficient for sexual reproduction to occur. Enhanced carbon dioxide supply induced conjugation (Starr, 1955; Dubois-Tylski and Lacoste, 1970; and Lippert, 1973), although Lippert (1969) found greater CO₂ (10%) increased cell lysing and abortion rate.

Although strains #171 and #96 are both in the _C. moniliferum_ group, the reason that their responses may be different is that one of them (#171) may be mis-classified (Lippert, personal communication).

**Nitrate Quota**

This research was approached with the hypothesis that cells would enter the sexual reproduction phase of the life cycle when the nitrogen cell quota became critical (minimal) within the cell. It was thought that copper might inhibit nitrate uptake and thereby decrease the amount of nitrogen in each cell which would lead to sexual reproduction. This was not found to be the case. With sub-lethal concentrations of copper added, nitrate uptake was not completely prevented. At pCu* 12.1 nitrate uptake in strain #96 was inhibited and sexual reproduction was delayed until nitrate was depleted in the medium. In the beginning of the experiment, cells at pCu* 12.1 had large nitrogen quotas because nitrate uptake continued while cell division was slowed. When sexual reproduction occurred, however, cell quota of nitrate was as low as those cells growing at pCu* 15.4 (Table V).
TABLE V
NITROGEN QUOTA AND ONSET OF SEXUAL REPRODUCTION AS A FUNCTION OF COPPER

<table>
<thead>
<tr>
<th>pCu*</th>
<th>Day of Nitrate Depletion</th>
<th>Day of Sexual Reproduction</th>
<th>N Quota at time of sex (micromoles)</th>
</tr>
</thead>
<tbody>
<tr>
<td>15.4</td>
<td>12</td>
<td>14</td>
<td>0.045</td>
</tr>
<tr>
<td>14.4</td>
<td>6</td>
<td>8</td>
<td>0.205</td>
</tr>
<tr>
<td>13.4</td>
<td>11</td>
<td>13</td>
<td>0.045</td>
</tr>
<tr>
<td>12.1</td>
<td>13</td>
<td>14</td>
<td>0.045</td>
</tr>
</tbody>
</table>

*N Quota = \frac{\text{micromoles NO}_3 \text{ taken from the medium}}{\# \text{ of cells} + 2 (\# \text{ of zygotes} + \text{aborted cells})}
It is apparent, however, that minimum cell quota is not the factor affecting the onset of sexual reproduction since sexual reproduction occurred earlier when nitrate uptake was rapid (pCu* 14.4) and cell numbers were low; these cells had a high cell quota at the time of sexual reproduction. Also, in earlier experiments with strain #96, sexual reproduction occurred sooner when beginning nitrate concentrations were less (Figure 37). Q, the nitrate quota, was high in these cells when early sexual reproduction occurred.

Although Eppley (1978) reported nitrate uptake kinetics depend on the concentration of nitrate in the medium and nitrogen cell quota, this work with Closterium showed sexual reproduction to be a function of the depletion of nitrate in the medium, not nitrogen cell quota.

Onset of Sexual Reproduction in Strain 96

Sexual reproduction in this strain occurred shortly after nitrate depletion of the media (Table V) and was not dependent on cell number (Figures 36 and 37). When copper stimulated cell division and nitrate uptake (populations at pCu* 14.4), sexual reproduction occurred earlier than in populations without added copper. Both sexual reproduction and nitrate uptake were delayed at pCu* 12.1.

It is reasonable to assume that if the FRAQUIL medium had less nitrogen, sexual reproduction would occur earlier. This was found to be true experimentally (Table VI). It
Figure 37. Number of zygotes per total cell number $\times 100$ vs. initial concentration of nitrate.

Total number of cells per ml at the time of sexual reproduction vs. initial concentration of nitrate.
<table>
<thead>
<tr>
<th>Nitrate Concentration At Beginning of Growth Experiment (micromoles/liter)</th>
<th>Onset of Sexual Reproduction (Days)</th>
<th>Maximum Cell Number (per ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>25</td>
<td>7</td>
<td>430</td>
</tr>
<tr>
<td>50</td>
<td>9</td>
<td>1000</td>
</tr>
<tr>
<td>100</td>
<td>16</td>
<td>3230</td>
</tr>
<tr>
<td>200</td>
<td>16</td>
<td>6100</td>
</tr>
</tbody>
</table>
is interesting to note that when sexual reproduction occurred early in a culture, due to beginning nitrogen limitation (Figure 37), the percent of culture undergoing sexual reproduction was greater than in those with larger cell number and later sex. The percent of culture undergoing sexual reproduction was calculated:

\[
\text{% Sexual Reproduction} = \frac{2(\# \text{ Zygotes} + \# \text{ Abortions})}{\text{Cell #} + 2(\# \text{ Zygotes} + \# \text{ Abortions})}
\]

Although the populations stimulated by copper (pCu\* = 14.4) had higher frequency of sexual reproduction early in the growth curve (Figure 38), the populations at other pCu\* values were able to achieve as high a frequency later. This again indicates the cells can adapt to the presence of copper and overcome its effects.

Although it is difficult to quantify, an effect of copper on sexual reproduction, independent of nitrate, may be inferred by the morphology of copper treated cells. At day 7 of the long-term experiment, 11.2% of the cells in the population were aberrant at pCu\* 12.1 compared to less than 1% at pCu\* 15.4 (Table VII). Aberrant cells include cells that are distorted (Figures 31-34), unseparated in the process of cell division (Figures 23-30), with less chlorophyll than normal (Figures 39-42) or with many vacuoles (Figures 43 and 44). It is assumed that these aberrant cells are unlikely to be able to initiate or complete the process of cell division (which must precede sexual reproduction).
Figure 38. Percent sexual reproduction @ day 15 of the growth curve vs. pCu*. 
<table>
<thead>
<tr>
<th>pCu*</th>
<th>Day 7</th>
</tr>
</thead>
<tbody>
<tr>
<td>15.4</td>
<td>0.95%</td>
</tr>
<tr>
<td>14.4</td>
<td>1.5%</td>
</tr>
<tr>
<td>13.4</td>
<td>3.7%</td>
</tr>
<tr>
<td>12.1</td>
<td>11.2%</td>
</tr>
</tbody>
</table>
Figure 39. Strain 96 of Closterium moniliferum. This scanning electron micrograph shows evidence of a shrunken protoplast inside the cell. SEM 500 X.

Figure 40. Strain 96 of Closterium moniliferum. Cell with less than normal amount of chlorophyll and with protoplast pulled away from the cell wall. LM 430 X.
Figures 41 and 42. Strain 96 of *Closterium moniliferum* treated with copper. These cells are examples of cells that were treated with copper and had less chlorophyll per cell than those cells that did not have extra copper added to the medium. LM 200 X.
Figure 43. Strain 96 of *Closterium moniliferum*. This cell demonstrates the presence of large vacuoles formed throughout the cell when added copper is in the medium. p = pyrenoid; v = vacuole. LM 430 X.

Figure 44. Strain 96 of *Closterium moniliferum*. This result is seen in some cells in the presence of added copper. p = pyrenoid; v = vacuole. LM 600 X.
Onset of Sexual Reproduction in Strain 171

Sexual reproduction was not correlated with nitrate depletion in this strain. Sexual reproduction began when nitrate was in adequate supply and cell numbers were very small (less than 100/ml). Other researchers have induced conjugation in *Closterium* without nitrogen depletion (Starr, 1955; Dubois-Tylski and Lacoste, 1970; Lippert, 1973); they stressed the importance of an enhanced CO₂ supply. Pickett-Heaps and Fowke (1971), however, did not find an increased CO₂ supply a prerequisite for conjugation in *Closterium littorale*. As pointed out by Coesel (1974), conjugation would seem to occur most readily when illumination, temperature and CO₂ tension are at an optimum for mitotic activity. This would seem to be particularly true in these strains of *Closterium moniliferum* where sexual reproduction takes place between newly divided immature cells.

Zygote Germination

There was no significant difference in the percent germination in cultures at any pCu* when copper was added to cells when in their vegetative asexual stage. When copper was added to mature cultures at the same time as light and nutrients required for germination, again there was no significant difference in percent germination.

It may be beneficial for populations to begin sexual reproduction when copper is present in the environment since
evidence shows growing cells are susceptible. The thick-walled zygotes may act as a survival stage for the organism. During the maturation process, the zygote may provide protection. In a natural environment, the time needed for maturation may be enough time for dilution of the toxicants to occur so vegetative cells could survive. Research here has shown that toxic amounts of copper present do not prevent germination; newly formed vegetative cells are subject to the toxic effect.

Added copper does not appear to have any effect on the germination of *Closterium moniliferum*. The germination process was not inhibited but the newly formed products of germination were killed by a low pCu*.

**Abortions**

Although preliminary experiments indicated that with increased zygote production, there was also increased abortion rate, subsequent experiments did not substantiate this. The percent of abortions varied from 1 to 25% of the zygotes formed and did not correlate with pCu* or nitrate in the medium.

**Nitrate Uptake in Short Term Experiments**

The effects of copper on nitrate uptake were analyzed by nitrate depletion in cultures over short time intervals from 2 to 69 hours. In nearly all cases, nitrate uptake was most rapid when no added copper was present. The most rapid uptake occurred in the first 5 hours after nitrate
was added to the nitrogen-depleted cultures with an average uptake rate (without added copper present) of $7.27 \times 10^{-4}$ micromoles nitrate cell$^{-1}$ hour$^{-1}$. The maximum during that five hour period without added copper in any one experiment was $22 \times 10^{-4}$ micromoles cell$^{-1}$ hour$^{-1}$ and the minimum was less than $1 \times 10^{-4}$ micromole cell$^{-1}$ hour$^{-1}$.

The exception to the most rapid uptake occurring without copper being added was one experiment where added copper to give a final molarity of $10^{-7}$ M caused a more rapid uptake of $5.1 \times 10^{-4}$ micromole cell$^{-1}$ hour$^{-1}$ vs. $4.7$ micromole cell$^{-1}$ hour$^{-1}$ (with no added copper) at the end of 22 hours. However, early in the experiment (5 hour sampling) the nitrate uptake rate was not as great with copper as it was without it. In this same experiment, added copper to give a final molarity of $5 \times 10^{-8}$ M caused inhibition in comparison to the $10^{-7}$ M. This concentration of copper ($10^{-7}$ M) could be analogous to the sensitive point (pCu* 14.4) in the long term experiments with strain 96 where uptake rate was increased and sexual reproduction occurred earlier. Generally, the lesser concentrations of copper exerted less toxicity in terms of NO$_3$ uptake on the cells (Figure 45).

It was generally found that a decrease in nitrate uptake rate occurred over time (Table VIII) when no copper was added. After a rapid uptake in the beginning, there is a decrease, followed by a slight increase. At the very end of the experiments, this decrease may be due to the
Figure 45. Nitrate uptake rate vs. final copper concentration. Time factor is 24 hours. TRIS buffer was present in the test media.
TABLE VIII

NITRATE UPTAKE RATE (Micromole x 10^{-4} cell^{-1} hour^{-1})
(No copper added)

<table>
<thead>
<tr>
<th>TIME* (hours)</th>
<th>NITRATE UPTAKE RATE</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Exp. 1</td>
<td>Exp. 2</td>
<td>Exp. 3</td>
</tr>
<tr>
<td>5</td>
<td>7.1</td>
<td>7.8</td>
<td>6.8</td>
</tr>
<tr>
<td>21</td>
<td>4.5</td>
<td>7.1</td>
<td>4.7</td>
</tr>
<tr>
<td>28</td>
<td>5.0</td>
<td>7.4</td>
<td>4.9</td>
</tr>
<tr>
<td>45</td>
<td>5.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>51</td>
<td>5.3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>69</td>
<td>3.9n</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

^Nitrates may be limited at this sampling time.
lack of nitrate availability, but through most of the experi-
ment, nitrate was in adequate supply. In contrast, inhibi-
tion of nitrate uptake due to added copper generally lessened
as the experiment proceeded (Table IX). Again, this demon-
strates the adaptability of the organisms to the presence
of copper.

The cells appear to be rapidly adaptable to the
presence of copper since they are able to increase nitrate
uptake rate after initial contact with the metal. Although
lack of nitrate uptake sometimes occurred for up to 26 hours
when sub-lethal concentrations of copper were added, these
cells were subsequently able to take up nitrate. This
adaptability may partially account for a lag in nitrate
uptake rather than complete blockage. This may help explain
why there is delayed nitrate depletion and delayed sexual
reproduction characteristic of the effect of copper in long
term experiments, rather than total blockage.

Examining nitrate uptake is an indirect way of examining
the synthesis and activity of nitrate reductase since uptake
is closely linked to the presence of nitrate reductase.
Nitrate reductase is an inducible enzyme produced in the
presence of nitrate so it is assumed that there is little
or no nitrate reductase present in nitrogen depleted cultures.
When nitrate was added to these nitrogen depleted cultures,
copper was added at zero time to some flasks and 5 hours
later to other flasks. Differences in uptake were not
significant so it is not possible to say if synthesis
<table>
<thead>
<tr>
<th>TIME (hour)</th>
<th>FINAL MOLARITY OF COPPER</th>
<th>NITRATE UPTAKE RATE (Micromole x 10^{-4} cell^{-1} hour^{-1})</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>9.9 x 10^{-10}</td>
<td>10^{-6}  3.5 x 10^{-6}  7 x 10^{-6}</td>
</tr>
<tr>
<td>5</td>
<td>7.8</td>
<td>4.0  0.6  0.6</td>
</tr>
<tr>
<td>24</td>
<td>7.1</td>
<td>5.7  2.8  0.9</td>
</tr>
<tr>
<td>29</td>
<td>7.4</td>
<td>6.0  2.3  1.5</td>
</tr>
</tbody>
</table>

*0 hour is the time nitrate and copper were added to nitrate depleted cultures.
of nitrate reductase or activity is affected more by copper.

**Ammonia Uptake in Long Term Experiments**

To determine if copper equally affected all aspects of the nitrogen metabolism of *Closterium moniliferum* or if copper was more toxic to the process of nitrate uptake, long term growth experiments were run with ammonia and with nitrate as separate nitrogen sources.

A representation of nitrogen uptake and assimilation in unicellular organisms and possible sites of copper's action is shown in figure 46. Although nitrite is an intermediate in nitrate reduction, it is usually not considered to be a significant factor (Eppley and Rogers, 1970).

If copper is acting on the uptake of nitrogen into the cell as well as cell division, the effects would be shown by different growth patterns when different nitrogen sources are used. If copper influences the amount of nitrogen taken into the cells, metabolism may be affected which may ultimately affect growth. Copper may have more than one site of action, acting both on cell division and nutrient uptake. Copper may specifically affect nitrate uptake which would slow the amount of nitrogen reduced to ammonia and ultimately available for incorporation, or copper may specifically affect ammonia uptake, directly affecting the amount of ammonia available.
Figure 46. Metabolic sites that may be blocked by the action of copper. Copper may be acting on any of the 7 sites shown with a ■ interrupting the arrow.
With few exceptions, algae can apparently utilize either ammonium salts or nitrate when these are supplies at a suitable concentration. Some Chlamydomonas and Gloeocystis species (Cain, 1965) and Haematococcus (Proctor, 1957; Stross, 1963) prefer utilization of nitrate to ammonium. Nitrate reduction, however, requires energy, and if the energy supply is limited more growth will occur on ammonium-N (Lewin, 1962). In batch culture experiments, the presence of greater than micromolar levels of ammonium prevented the utilization of other nitrogen sources (Morris, 1974). Ammonium is known to both inhibit the activity of and repress the synthesis of nitrate reductase, thus preventing assimilation of nitrate (Hattori, 1962a,b); Morris and Syrett, 1963; Syrett and Morris, 1963).

The maximum inhibition by copper was seen when nitrate was the nitrogen source (Figure 47). The lag phase was prolonged and these cultures were never able to overcome the toxicity and attain the maximum cell density. The cultures with ammonium, with and without copper, were able to reach the same maximum cell number as those with nitrate and no copper. Cells grew slightly better (with the shortest lag phase) when nitrate was the nitrogen source and no added copper was present; this may have been due to the acclimatization of the cells since they had previously been grown in FRAQUIL with nitrate.

The fact that copper inhibits nitrate uptake indicates that the site of copper's action may be the inducible enzyme
Figure 47. Cell number/ml vs. time with copper and different nitrogen sources. FQ = FRAQUIL; FQ/Cu = FRAQUIL with copper (1.3 x 10^{-6} M); FQ/NH_{3} = FRAQUIL with ammonia substituted for nitrate; FQ/Cu/NH_{3} = FRAQUIL with ammonia with copper (1.3 x 10^{-6} M).
nitrate reductase. Copper has been found to inhibit the activity of nitrate reductase and the uptake of nitrate and ammonium uptake (Harrison et al., 1977). Shioi et al. (1978b), from their work with Ankistrodesmus suggest that this may occur because copper inactivates the electron transport of photosystem II and there is not an adequate supply of reductants for nitrate reduction. This would help explain why copper has more of an effect on nitrate uptake than on the reduced form of nitrogen, ammonia. Previous work has also demonstrated that copper does decrease photosynthesis and chlorophyll content (McBrien and Hassall, 1965; Steemann Nielsen et al., 1969; Overnell, 1975; and Rosko and Rachlin, 1977). Nitrogen availability may be affected by the direct effect of copper as well as the indirect effects of photosynthesis in providing reductants. The form of nitrogen that is present becomes increasingly important to the cells when metals such as copper are in the environment to affect their availability.

CONCLUSION

The onset of sexual reproduction is linked to depletion of nitrate in the medium in one of the strains (#96) tested. When copper delayed nitrate uptake, sexual reproduction was delayed; when copper stimulated nitrate uptake, sexual reproduction occurred earlier. Copper did not have this effect when ammonia was the nitrogen source. The key to sexual reproduction in Closterium is not total nitrogen
in the cell, but the supply outside the cell and this must be in the form of nitrate.

Copper generally inhibited nitrate uptake in long and short term experiments. The dosage response was demonstrated also by the stimulation of nitrate uptake (and cell division) at $pCu^*$ 14.4.

The increase in cell number parallels nitrate depletion in batch cultures. Although cells must make contact before sexual reproduction can occur, cell number is not critical in the process. The trigger is metabolic, from the depletion of nitrate.

After sexual reproduction has occurred, the zygotes need maturation time in the dark. Upon being returned to light conditions and having nutrients added, germination occurs. The presence of added copper did not affect the percent germination. Abortion rate was not affected by copper.

The following chapter (Chapter V) shows morphological effects due to excess copper.
CHAPTER V

EXCESS COPPER AFFECTS CELL MORPHOLOGY

INTRODUCTION

A commonly observed effect of heavy metal poisoning is changes in cell size and morphology (Thomas et al., 1980). This phenomenon has been seen in Chlorophyceae (Rosko and Rachlin, 1977); Chrysophyceae (Davies, 1974); and Bacillariophyceae (Nuzzi, 1972; Sunda and Guillard, 1976; Berland et al., 1977; and Morel et al., 1978).

Morel et al. (1978) reported morphological aberrations in Skeletonema costatum at the same copper activities that began to increase the lag phase of growth in batch cultures. With Closterium, however, these aberrant forms, particularly the unseparated cells, can be found in nature and in untreated cultures. Ichimura and Watanabe (1976) reported morphological variation among three related species of Closterium. What is significant here is the increase in the number of aberrant cells.

The effect of copper on morphology is probably not a direct effect but is due to its effect on the growth and metabolism of the cell. The uncoupling of cell growth and cell division by the action of trace metals has been suggested by Davies (1976) as an explanation of the
phenomenon of changes in morphology. When treated with trace metals, the cells may continue to grow but the normal separation does not occur. Favorable growth conditions may also cause this uncoupling as shown by Playfair (1910) when cell division in Closterium was accelerated and occurred before new semi-cells were completely formed.

RESULTS AND DISCUSSION

**Closterium moniliferum** displaying typical morphology is shown with light microscopy (Figures 4 and 48) and with the scanning electron microscope (SEM) (Figures 49 and 50). Pickett-Heaps (1973) suggests the use of SEM in taxonomic studies of desmids. SEM was used here not so much for an increase in magnification as for observation of changes that might occur with copper treatment.

From the increased lag phase in growth of copper-treated cells, (See Chapter III) it is apparent that completion of cell division is inhibited by copper. The inability of the cells to complete cell division is confirmed with photomicrography (Figures 13, 14, 23-27) and with SEM (Figures 28-30). Irregularities and loss of symmetry of cells are evident in this stereo pair of photographs (Figures 51 and 52).

Gross distortions occur as cells continue to grow and deposit cell wall without dividing or expanding (Figures 29 and 30) or appear to repeatedly begin the division process (Figures 31, 32; 53 and 54).
Figure 48. Typical morphology strain 96 of *Closterium moniliferum*. Cell with typical morphology with one semi-cell that has not yet attained maximum size for the cell to be considered mature. LM 430 X.
Figure 49. Mature symmetrical healthy cells of strain 96 of *Closterium moniliferum*. SEM 500 X.
Figure 50. Dynamic focusing with the scanning electron microscope. Strain 96 of Closterium moniliferum. Mature symmetrical cells that have not been treated with copper. SEM 500 X.
Figures 51 and 52. Stereo pair of electron micrographs of strain 96 of *Closterium moniliferum* treated with copper. Cells should be viewed through a stereo-viewer or by briefly staring cross-eyed at the two micrographs until the two images merge. SEM 500 X.
Figure 53. Copper treated cell of strain 96 of *Closterium moniliferum*. SEM 500 X.

Figure 54. Enlargement of rectangle in Figure 53. SEM 5000 X.
With added copper, the total amount of chlorophyll produced was decreased. When cells fail to divide, the amount of chlorophyll/cell may be higher (See Table IV) but shrunken protoplasts are visible with light microscopy (Figures 40-44) and with SEM, lack of vigor is indicated by the flaccid cell membrane (Figure 39).

Another morphological effect observed in vegetative cells was the formation of large vacuoles throughout the cell (Figures 43 and 44).

In long term growth experiments, an increase in number of aberrant cells did not begin to appear until day 6 or 7. Aberrant cells consist of those cells that did not complete separation (Figures 23-30), those with shrunken protoplasts (Figures 39-44) or vacuoles (Figures 43 and 44), and those of distorted morphology (Figures 31-34). In the first days of the experiment the highest percent of aberrant cells occurred in the cultures with pCu* = 12.1 (Table X). In cultures where sexual reproduction occurred earlier (pCu* = 14.4) with strain #96 the percent of aberrant cells increased greatly throughout the experiment with a high in one experimental flask of 56.7% of the total cell number aberrant. However, this was probably not due to copper's immediate effect, but due to senescence and/or nutrient depletion since those cells took up nitrate (and presumably other nutrients) more quickly.

At pCu* = 5.5, cultures were not viable. In growing cultures, the number of cells counted as dead (Figures 55-
<table>
<thead>
<tr>
<th>pCu*</th>
<th>Day 7</th>
<th>Average (3 weeks)</th>
</tr>
</thead>
<tbody>
<tr>
<td>15.4</td>
<td>0.95%</td>
<td>5.9%</td>
</tr>
<tr>
<td>14.4</td>
<td>1.5%</td>
<td>10.9%</td>
</tr>
<tr>
<td>13.4</td>
<td>3.7%</td>
<td>9.3%</td>
</tr>
<tr>
<td>12.1</td>
<td>11.2%</td>
<td>14.3%</td>
</tr>
</tbody>
</table>
when treated with sub-lethal concentrations of copper was not significantly different from that of cultures without added copper. In other words, although growth (cell number increase) was inhibited at sub-lethal concentrations of copper, it was due to less cell division, not death of individual cells.

It is known that *Closterium* species produce mucilage; in fact, that is the mechanism for movement in this organism. Quantitative differences in mucilage production due to copper were not measured nor were differences obvious with SEM (Figures 58 and 59).

The two strains of *Closterium moniliferum* studied differed in their sexual reproduction in that #171 produces twin zygotes (Figures 10 and 60-68), whereas #96 produces single zygotes (Figures 69-72). In 1963, Cook proposed that sexual morphology be an essential feature in identification within *Closterium* species so it is important to continue to monitor both sexual and asexual phases for morphological changes due to copper.

When the sexual process did not go to completion, the zygotes aborted (Figures 73-76). The difference in the number of abortions in populations treated with copper and those without was not significant. This indicates that although copper affects sexual reproduction by delaying its onset, once the process begins copper is not a factor in affecting it.
Figures 55 and 56. Lysed non-viable cells of *Closterium moniliferum*. LM 430 X.
Figure 57. Abnormal cell of strain 96 of Closterium moniliferum. Appearance of a cell in a culture treated with excess copper. SEM 500 X.
Figure 58. Mucilage in strain 96 of Closterium moniliferum. Note the mucilage between two cells. $pCu^* = 15.4$ SEM 1000 X.

Figure 59. Mucilage in strain 96 of Closterium moniliferum. Mucilage when cells were grown at $pCu^* = 12.1$. SEM 2000 X.
Figure 60. Papillae in strain 171 of Closterium moniliferum. This is the beginning of sexual reproduction. SEM 500 X.
Figure 61. Strain 171 of Closterium moniliferum. Twin zygotes forming. LM 430 X.

Figure 62. Twin zygotes of strain 171 of Closterium moniliferum with 4 empty cells. LM 430 X.
Figures 63 and 64. Twin zygotes of strain 171 of Closterium moniliferum. LM 600 X.
Figure 65. Zygotes of strain 171 of Closterium moniliferum. Twin zygotes in culture that had been treated with added copper. SEM 500 X.

Figure 66. Zygotes of strain 171 of Closterium moniliferum. Twin zygotes in culture that had been treated with added copper. SEM 1000 X.
Figure 67. Twin zygotes of strain 171 of Closterium moniliferum. Empty cells remain after sexual reproduction. SEM 500 X.

Figure 68. Twin zygotes of strain 171 of Closterium moniliferum. Empty cells remain after sexual reproduction. SEM 500 X.
Figure 69. Two immature cells of strain 96 of *Closterium moniliferum* pairing before conjugation. LM 430 X.

Figure 70. Scanning electron micrograph of zygote of strain 96 of *Closterium moniliferum*. Empty cells are nearby. SEM 500 X.
Figure 71. Zygotes of strain 96 of *Closterium moniliferum* that are still covered with the mucilagenous sheath from the conjugation vesicle. SEM 2000 X.

Figure 72. Zygote of strain 96 of *Closterium moniliferum* without mucilagenous material. SEM 2000 X.
Figures 73 and 74. Abortion in strain 96 of *Closterium moniliferum*. Cell contents are being released before the conjugation vesicle formed (Top); Non-simultaneous release of cell contents (Bottom). LM 430 X.
Figure 75. Abortion in strain 96 of *Closterium moniliferum*. On the right is a zygote formed; on the left and at the bottom of the micrograph are two abortions. SEM 500 X.

Figure 76. Enlargement of cell in upper left of figure 75. SEM 2000 X.
CONCLUSION

The typical symmetrical cell shape of *Closterium* is affected by additions of copper. After cell division, the cell grows and re-establishes the symmetry lost when it divided. When copper disrupts this growth process or the deposition of cell wall, distorted cells may result.

Copper affects cell morphology by acting on the cell division (mitosis or cytokinesis) itself. Although growth was occurring, it is uncoupled from cell division and mature cells remained unseparated from each other.

The unseparated cells, not being able to finish cell division, will not be involved in sexual reproduction since vegetative cell division is a prerequisite. It is improbable that the distorted cells could initiate or complete a normal conjugation. Thus, the main significance of copper's effect on morphology in regard to this work is the fact that frequency of sexual reproduction will be reduced due to lack of cell division.
CHAPTER VI

CONCLUSION

This study examined the total life cycle of an organism and its response to copper. The growth of Closterium moniliferum with copper exemplifies the typical beneficial, inhibitory, and toxic dose-response to trace metals suggested in Chapter I. The stimulatory effect was demonstrated by an early increase in cell number, increased nitrate uptake, and early onset of sexual reproduction. The inhibitory effect, with greater concentrations of copper, was demonstrated by a longer lag phase, decreased nitrate uptake, and later onset of sexual reproduction. Since these cells are adaptable with time, one of the final effects of sub-lethal concentrations of copper within the medium was to affect the timing of the life cycle.

The sexual phase in some strains of Closterium moniliferum can be triggered by nitrate depletion in the media. Although it was expected that each cell would have a minimal cell quota of nitrogen before sexual reproduction occurred, work with strain 96 demonstrated that Q (nitrogen/cell) was not the critical factor in initiation of the process. It was also demonstrated the cell number in batch cultures was not the critical factor. When ammonia was the only nitrogen source and it became depleted in the
medium, sexual reproduction did not occur. Nitrate depletion appears to be the critical component in induction of sexual reproduction.

When copper was added to batch cultures with different nitrogen sources, cells growing with nitrate were inhibited more than cells growing with ammonia. This information gives direction to further research in using copper as a tool in understanding nitrogen metabolism in microorganisms, particularly in desmids where sexual reproduction is integrally tied to the nitrogen requirements of the cell. Figure 46 shows possible sites of action of copper on nitrogen metabolism. Further work should include an examination of any of these sites that may be blocked. For example, the action of copper on nitrate uptake could be directly on the enzyme nitrate reductase. Since nitrate reductase is an inducible enzyme, antibiotics inhibiting protein synthesis could be used in combination with copper to help understand whether copper affects synthesis or activity of nitrate reductase.

The rate of germination of zygotes was not correlated with pCu*. After germination, however, the vegetative cells would again be susceptible to copper's effect. Abortion rate was not correlated with pCu*. It appears that the effect of copper occurs early in cell growth or on induction of sexual reproduction. Once the process of sexual reproduction is allowed to begin, copper's effect is not evident.
In natural waters, conjugation in desmids is most frequent in very shallow waters which can absorb carbon dioxide from the atmosphere or possibly from respiration from bottom muds (Brook, 1981). Such conditions prevail in bogs and temporary pools with changing water levels and where, clearly, the production of resistant zygospores would have considerable survival value. Similarly, the stimulatory effect of copper with its accompanying earlier sexual reproduction could have survival value for *Closterium moniliferum*. 


Fahrenbach, Wolfgang. Personal communication.


Lippert, B.E. Personal communication.


Sweet, Jim. Personal communication.


APPENDIX

FRAQUIL MEDIUM

(from Morel et al., 1979)
### Fraquil Salts (S.P.W.)

<table>
<thead>
<tr>
<th>Mass</th>
<th>Substance</th>
<th>Gram Formula Weight</th>
<th>Initial Stock Molarity</th>
<th>Initial Stock Volume</th>
<th>Dilution Factors</th>
<th>Final Molarity</th>
<th>-log M</th>
</tr>
</thead>
<tbody>
<tr>
<td>36.8g</td>
<td>CaCl₂·2H₂O</td>
<td>147.02</td>
<td>1.50 x 10⁻¹</td>
<td>500 x 2</td>
<td>2.50 x 10⁻⁴</td>
<td>3.60</td>
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<tr>
<td>37.0g</td>
<td>MgSO₄·7H₂O</td>
<td>246.48</td>
<td>1.50 x 10⁻¹</td>
<td>500 x 2</td>
<td>1.50 x 10⁻⁴</td>
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<tr>
<td>12.6g</td>
<td>Na₂HCO₃</td>
<td>84.01</td>
<td>1.50 x 10⁻¹</td>
<td>500 x 2</td>
<td>1.50 x 10⁻⁴</td>
<td>3.82</td>
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### Nutrients

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<th>Mass</th>
<th>Substance</th>
<th>Gram Formula Weight</th>
<th>Initial Stock Molarity</th>
<th>Initial Stock Volume</th>
<th>Dilution Factors</th>
<th>Final Molarity</th>
<th>-log M</th>
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<tbody>
<tr>
<td>1.74g</td>
<td>K₂HPO₄</td>
<td>174.18</td>
<td>1.00 x 10⁻²</td>
<td>500 x 2</td>
<td>1.00 x 10⁻⁵</td>
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<td>8.50g</td>
<td>NaNO₃</td>
<td>84.99</td>
<td>1.00 x 10⁻¹</td>
<td>500 x 2</td>
<td>1.00 x 10⁻⁴</td>
<td>4.00</td>
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<tr>
<td>3.55g</td>
<td>Na₂SIO₃·9H₂O</td>
<td>284.20</td>
<td>1.25 x 10⁻²</td>
<td>500 x 2</td>
<td>1.25 x 10⁻⁵</td>
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### Trace Metals

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<th>Initial Stock Volume</th>
<th>Dilution Factors</th>
<th>Final Molarity</th>
<th>-log M</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.249g</td>
<td>CuSO₄·5H₂O</td>
<td>249.68</td>
<td>9.97 x 10⁻⁴</td>
<td>10² x 10³</td>
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<tr>
<td>0.265g</td>
<td>(NH₄)₂SO₄·2H₂O</td>
<td>1235.86</td>
<td>1.50 x 10⁻³</td>
<td>10² x 10³</td>
<td>1.50 x 10⁻⁹</td>
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<tr>
<td>0.59g</td>
<td>CoCl₂·6H₂O</td>
<td>237.95</td>
<td>2.50 x 10⁻³</td>
<td>10³ x 10³</td>
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<td>8.60</td>
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<td>287.54</td>
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<td>10³ x 10³</td>
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<td>0.122g</td>
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<td>1.86g</td>
<td>Na₂EDTA</td>
<td>372.24</td>
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<td>6.18g</td>
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<td>61.83</td>
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<td>10³ x 10³</td>
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</tbody>
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