THE STUDY OF COPPER BIOAVAILABILITY AND MECHANISM OF UPTAKE IN THE TYPE I METHANOTROPH METHYLOMICROBIUM ALBUS BG8

Thesis by
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In Partial Fulfillment of the Requirements for the Degree of the Doctor of Philosophy

California Institute of Technology
Pasadena, California
1996

(Defended May 30, 1996)
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Science has ‘explained’ nothing;

the more we know the more fantastic world becomes

and the profounder surrounding darkness.

– Aldous Huxley

I expect I shall be a student to the end of my days.

– Anton Chechov
Acknowledgments

I am very grateful to my advisor, Dr. Mary E. Lidstrom, who cordially invited me to work in her lab, and who introduced me to the fascinating world of microbiology, genetics and molecular biology. She gave me a lot of support and counsel with my research, career, and even parenting. Dr. Jim J. Morgan was of great assistance in discussion of the results of my experiments and advising on presenting the data for publication. I am also thankful to Dr. Sunney I. Chan for his helpful comments on my paper. Dr. Peter Green was invaluable with his willingness to ensure smooth running of ICP–MS. I cannot thank enough my fellow Russians: Mila Chistoserdova, Andrey Chistoserdov and Sergei Stolar for their support, numerous advise, and, the most important, their friendship. I also appreciate the input of all other members of Lidstrom lab into my research.

Thomas Lloyd was a pleasure to work with on the siderophore project, and I wish him a very successful completion of his thesis. Anna E. Arreola, Jennifer Miller, Elizabeth Price and Wai Kwan were all great MURF/SURF students and I appreciate their help with experiments essential for completion of this project. I must thank Fran Matzen for her inestimable assistance with paperwork.

I would like to dedicate this thesis to the people I love the most: my husband, son, parents and parents-in-law. I treasure their encouragement, I cherish their unconditional love and friendship.
Abstract

Two aspects of copper uptake by the type I methanotroph *Methylomonas albus* BG8 were investigated – the effect of copper speciation in the growth medium on copper accumulation and mechanisms of copper transport in this microorganism.

Copper accumulation in *M. albus* BG8 consisted of nonspecific sorption of copper to outer cell layers and copper internalization. Most of the copper accumulated by the cells was nonspecifically sorbed to the cellular surface and was removable by EDTA. This phenomenon was especially prominent when cultures were grown at high total copper concentrations in the growth medium. This reversible binding of copper to external sites (e.g., amino acid, carboxylic, hydroxy groups, etc.) was described by a hyperbolic model with a mean maximum binding capacity of \((1.54 \pm 0.06) \times 10^{-15}\) moles/cell and an apparent half saturation constant of \((1.43 \pm 0.05) \times 10^{-7}\) moles/l. Copper availability to *M. albus* BG8 was related to the cupric ion concentration rather than to that of total copper added to the growth medium. Total internalized copper (total copper not removable by EDTA) was relatively constant at \(1–3 \times 10^{-17}\) moles of copper per cell despite a 100-fold variation in medium total copper and cupric ion concentrations, indicating the presence of a specific homeostasis mechanism for copper.

A specific copper uptake system is expected to be copper-regulated. Several copper-regulated polypeptides were identified in both soluble and membrane cellular
fractions. The corresponding gene of one copper-repressible polypeptide, corA, was cloned and sequenced. CorA appeared to be vital for *M. albus* BG8 since an insertion mutant defective in the gene grew very poorly on plates or in liquid culture. It was suggested that CorA might be a divalent metal porin.

Three putative copper ATPase genes, *atpA, atpB, atpC*, were cloned and the complete sequence of *atpA* was obtained. The gene product of *atpA* contained a copper-binding signature motif, suggesting that AtpA does play a role in copper transport in *M. albus* BG8.

A hypothesis for copper uptake in *M. albus* BG8 was suggested by this research. A copper-repressible protein (CorA) described in Chapter Three may be an outer membrane porin that works in tandem with the putative copper ATPase(s) (Chapter Four). Such a porin would not necessarily be specific for copper, and might be overexpressed under conditions of any divalent metal limitation to facilitate the metal diffusion into periplasm.
List of Abbreviations

dDH₂O, double distilled water

GCG, Genetic Computer Group

ICP/MS, inductively coupled plasma/mass spectrometer

ISE, ion selective electrode

MMO, methane mono-oxygenase

NAD⁺, nicotinamide-adenine dinucleotide

NADH, nicotinamide-adenine dinucleotide carrying a proton

NMS, nitrate mineral salts medium

PCR, polymerase chain reaction

pMMO, particulate methane mono-oxygenase

sMMO, soluble methane mono-oxygenase

TCE, trichloroethylene
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Chapter One

Introduction and Literature Review

1.1. Introduction

Almost one-half of the population of the United States relies on aquifers for their domestic water supply (Wilson et al. 1983). This makes groundwater pollution a critical issue, especially in regard to compounds that are poorly degraded by natural microbial populations. One of such stable and widespread contaminants that is difficult to remove from the environment, is trichloroethylene (TCE). TCE is a synthetic chlorinated hydrocarbon, extensively used for degreasing and cleaning of metals, military hardware and electronic components, dry-cleaning and fumigation. Inattentive storage, inappropriate disposal, and accidental spills of TCE combined with its relatively high water solubility has made it one of the most frequently reported groundwater pollutants in the United States (Ensley 1991; Murray 1993). It is a suspected carcinogen, and it has been designated as a priority pollutant by the US. Environmental Protection Agency (EPA, 1985; Phelps et al. 1990). While strong sorption of TCE onto sediments makes pump-and-treat techniques excessively lengthy and financially unfeasible, in situ TCE bioremediation has received increasing attention as a promising contaminant-destructive approach as opposed to contaminant-relocative approach (Bouwer and McCarty 1983; Parsons et al. 1984; Vogel and McCarty 1985; Ensley 1991; Halden and Chase 1991).
Partial biodegradation of TCE and other chlorinated alkenes under anaerobic conditions has been reported (Bouwer and McCarty 1983; Parsons et al. 1984), however, one of the products of degradation that accumulates is vinyl chloride, a known stable carcinogen (Vogel and McCarty 1985; Ensley 1991).

TCE has been shown to be aerobically co-metabolized by a variety of microorganisms in the presence of their primary growth substrate. These include toluene oxidizers (Nelson et al., 1987; Folsom and Chapman, 1991), ammonia oxidizers (Arciero et al. 1989), propane oxidizers, phenol oxidizers (Nelson et al. 1987; Ensley, 1991; Folsom and Chapman, 1991; Hopkins et al. 1993), and methane oxidizers (methanotrophs) (Little et al. 1988; Halden and Chase 1991; Park et al. 1991, Higgins et al. 1980). Methanotrophs have been suggested to be one of the more favorable groups of bacteria for in situ TCE bioremediation due to their ubiquity in nature, their ability to degrade chlorinated alkenes to non-toxic compounds that can be utilized by heterotrophic bacteria, and their ability to reduce TCE levels to below drinking water standards.

1.2. Trichloroethylene Biodegradation by Methanotrophs

Methanotrophs are Gram-negative bacteria that utilize methane as a sole source of carbon and energy. They oxidize methane to carbon dioxide by a route involving two-electron oxidation steps via methanol, formaldehyde and formate (Figure 1.1) (Higgins et al. 1980). Methane mono-oxygenase (MMO), the first enzyme in the pathway, is capable of inserting an oxygen atom into a variety of hydrocarbon substrates and chlorinated
compounds, including TCE, to produce alkene epoxides, which then break down in water to intermediates that are easily utilized by heterotrophic bacteria (Figure 1.2) (Little et al. 1988, Halden and Chase 1991).

The methanotrophs are divided into three groups (type I, type II and type X) based on their morphological, biochemical, and phylogenetic characteristics (Hanson et al. 1991). MMO is found in two forms in methanotrophs: soluble (sMMO) and membrane-bound, or particulate (pMMO). The pMMO is present in all known methanotrophs, whereas sMMO is found in only a few strains, mostly type II and type X (Hanson et al. 1991; King 1992; Peltola et al. 1993). In strains that contain both pMMO and sMMO, pMMO predominates in cultures grown under conditions of copper-sufficiency, while sMMO is found only under conditions of copper limitation (Halden and Chase 1991; Park et al. 1991; King 1992; Tsien and Hanson 1992; Chan et al. 1993; Peltola et al. 1993).

Both pMMO and sMMO can oxidize TCE, but the maximum TCE oxidation rates by cells containing sMMO are 100-1000 fold higher than cells containing pMMO (DiSpirito et al. 1992). However, a recent study (DiSpirito et al. 1992) has shown that type I methanotrophs expressing only pMMO are capable of TCE removal to the level required for drinking water (5ppb).

Little is known of the structure of pMMO since it is highly unstable in vitro and until very recently (Ngyuen and Chan, unpublished) has not been reproducibly purified. However, it appears to be a copper enzyme (Chan et al. 1993; Ngyuen et al. 1994). It was suggested that the bulk of the membrane-bound copper ions exist in the form of trinuclear copper clusters serving as catalytic sites of monooxygenase activity of the
pMMO system (Chan et al. 1993; Nguyen et al. 1994). Addition of copper to growth medium in a concentration range that is neither growth-limiting nor toxic results in increased cell yield and MMO activity (Collins et al. 1991), as well as changes in pMMO kinetics (Semrau et al. 1993). This effect is even more dramatic on TCE oxidation, as cells containing pMMO that are grown at 2μM copper added to the medium show no detectable TCE degradation (Semrau 1995, Smith 1996). Therefore, these responses to copper could have a major influence on TCE bioremediation by natural populations of methanotrophs, especially the type I strains that lack sMMO. All available data suggest that natural populations of methanotrophs are not copper-limited (Lidstrom and Semrau 1995). It is important, therefore, to investigate copper bioavailability to a type I methanotroph under conditions of copper-sufficiency.

1.3. Copper Speciation in Various Environments and its Bioavailability

Copper is widely found in the environment. It is present as a component in such ores as chalcopyrite (CuFeS₂) and chalcocite (Cu₂S). Copper, released from minerals during weathering, forms soluble cupric and cuprous salts, as well as organic complexes. An extensive use of copper as an antimicrobial agent in agriculture causes its additional release into soils and groundwater (Brown et al. 1992a).
1.3.1. Copper in Seawater

Copper is a trace element in seawater. Total copper concentrations in seawater vary from 0.1-0.5 nM at the surface to 5 nM in bottom waters. Copper distribution in seawater was proposed to be controlled by its depletion at the surface and regeneration from the sediments, as well as by copper scavenging throughout the water column (Boyle and Edmond 1975; Bruland et al. 1980). It appears from numerous studies that greater than 99.7% of the total dissolved copper is present as organic complexes in seawater (Sunda and Ferguson 1983; Sunda and Hanson 1987; Coale and Bruland 1988; Coale and Bruland 1990; Moffett et al. 1990; Van den Berg et al. 1990, Bruland et al. 1991; Van den Berg and Donat 1992). At depths of 1000 m, about 50-70% of the total dissolved copper is organically bound. Two classes of copper-binding ligands were identified throughout the water column and were suggested to be of biological origin: a strong ligand class \( \log K_{\text{stab}} = 11.5 \) in the surface waters (low copper concentrations) and a weaker class of ligands at middepths (higher copper concentrations) \( \log K_{\text{stab}} = 8.5 \) (Coale and Bruland 1990). A pH, temperature and, possibly, pressure dependence of the ligand/copper association has been suggested by Coale and Bruland (1990).

Inorganic copper species contribute less than 0.3% of the total copper with free cupric ion \( [\text{Cu}^{2+}(\text{H}_2\text{O})_6] \) comprising approximately 4% of the inorganic fraction (Bruland et al. 1991). Complexation with carbonate and hydroxide dominates the inorganic copper speciation in seawater; a significant effect of sulfide complexation on
inorganic copper speciation has been also reported in some areas (Coale and Bruland 1990). The free copper concentration varies from $10^{-13.1}$ M at the surface to $10^{-9.9}$ M at 300 m depth. It appears that only a small fraction of total copper is present as Cu(I) in seawater (Moffett and Zika 1988).

**1.3.2. Copper in Soil**

Copper is distributed between the liquid and solid phase in soils. The total concentration of Cu in pore water is estimated to be on the order of $10^{-7}$ to $10^{-9}$ M. Over 99% of the Cu is complexed by ligands, predominately humic substances (Hodgson et al. 1965; Berggren 1989). Concentrations of free copper depend on the type of the soil, and are in a range between $10^{-10}$ M and $10^{-12}$ M for uncontaminated soils (Hodgson et al., 1965).

**1.3.3. Copper in Freshwater Environments**

Copper in freshwater environments can be distributed between several phases including sediment, suspended particulates, true solution (porewater and water column) and it can be adsorbed onto hydrous oxide and humic colloids in water. Copper in true solution can exist as the free aqueous species $[\text{Cu}^{2+}(\text{H}_2\text{O})_6]$, inorganic complexes, and organic complexes (Guy and Kean 1980). On the basis of studies of oxidation kinetics of Cu(I) (Moffett and Zika 1983), it has been concluded that concentrations of Cu(I) in freshwater and low salinity environments should be exceedingly small. A significant fraction of the Cu(II) in freshwater is complexed by humic substances (Hering and Morel
1988), which results in cupric ion concentrations of $10^{-11} - 10^{-18}$ M (Apte et al. 1990a, Van Den Berg et al. 1990).

1.3.4. Copper Bioavailability

Copper exhibits complex chemical speciation in the environment, however, there is a general agreement that biological uptake of copper and its toxicity depend on the level of cupric ion, not the total copper concentration (Sunda and Guillard 1976; Anderson and Morel 1978; Zevenhuizen et al. 1979; Blust et al. 1986; Coale and Bruland 1990; Bruland et al. 1991; Langford and Guzman 1992). Inorganically complexed metal species equilibrate very rapidly with free metal ion. Thus, the total inorganic complex concentrations define an upper limit for the bioavailability of the metal (Morel et al. 1991).

The significant effects of hardness, temperature, and pH on copper bioavailability have been indicated in many studies (Gadd and Griffiths 1978; Sunda and Ferguson 1983; Coale and Bruland 1990; Meador 1991). The work of Bruland and coworkers (1991) has established the importance of Cu$^{2+}$: Mn$^{2+}$ and Cu$^{2+}$: Zn$^{2+}$ competition for cellular uptake sites in overall copper bioavailability.

Several experiments reported in the literature on metal-biota interactions have shown that speciation and, thus, bioavailability of copper ions may change during growth of microorganisms. A growing microorganism excretes compounds that can form complexes with metals and change the pH of the medium causing precipitation of metal ions as oxalates or phosphates (Bird et al. 1985; Hughes et al. 1991; Cabral, 1992). It has
been found, however, that EDTA and NTA added to normal culture media overwhelm such effects (Guy and Kean 1980).

1.4. Copper Ion Selective Electrode as a Method for Measuring Cupric Ion Concentration in Fresh Water Medium

Various techniques have been used to measure free copper concentration. These methods include:

- Anodic Stripping Voltammetry (ASV) and Differential Pulse Anodic Stripping Voltammetry (DPASV) (Coale and Bruland 1988; Powell and Town 1991);
- Adsorption on an ion-exchange resin;
- Liquid/liquid extraction (Berggren 1992; Hodgson et al. 1965; Moffett and Zika 1987; Moffett et al. 1990; Nair et al. 1991);
- Chemiluminescence detection (Sunda and Huntsman 1991; Coale et al. 1992);
- Competitive equilibrium with acetyl acetone or EDTA and subsequent SEP-PAK adsorption (Sunda and Hanson 1987);
ISE presents a simple and inexpensive approach to copper speciation. It allows the direct detection of the free copper in fresh water or other media with low chloride ion activity. Chloride interference with cupric ion selective electrode measurements has been reported (Westall et al. 1979). The assay of metal uptake does not require separation of cells from the medium, thus limiting artifacts in experiments with cultures. At constant temperature and ionic strength, the potential of an ISE is directly proportional to the logarithm of the primary ion concentration. The copper selective electrode, when calibrated with an appropriate metal-buffer can be used to monitor free copper ion in bacterial medium with cupric ion concentrations as low as $10^{-19}$ M (Avdeef et al. 1983).

To obtain the best performance of an ICE, the response of the electrode should be close to the Nernstian response over the widest possible pH range, i.e. the electrode potential $E$ should obey the equation: $E = E^* + \frac{RT}{2F} \ln[\text{Cu}^{2+}]$, where $[\text{Cu}^{2+}]$ is the concentration of copper(II) in the measured solutions.

Calibration of ion-selective electrodes in the range below 1μM - 10μM with solutions obtained from salts or by dilution of more concentrated standard solutions is undesirable for the following reasons:

- The preparation of very dilute solutions is inaccurate,
- Loss or contamination of dilute solutions may create serious positive or negative errors (Avdeef et al. 1983; Hulanicki et al. 1991).

The use of metal buffers helps to avoid these difficulties. Several copper buffers have been reported to be successfully used to obtain linear calibration curves up to
pCu=12 (pCu=-log[Cu^{2+}]): Cu^{2+}/NH_{3}; Cu^{2+}/EDTA; Cu^{2+}/NTA; Cu^{2+}/en
(ethylenediamine) (Avdeef et al. 1983). Since Cu^{2+} concentration depends on pH, a pH buffer should be added also. The following buffers have been often used: acetate, maleate, borax, TRIS [tris(hydroxymethyl)-methylamine], ACES [2-(2-amino-2-oxoethyl-amino)ethanesulfonic acid, 2-(carbamoyl-methylamino)ethanesulfonic acid], HEPES [4-(2-hydroxyethyl] piperazine-1-ethanesulfonic acid]], PIPES [piperazine-1,4-bis(2-ethanesulfonic acid]], etc. (Hulanicki et al. 1991). In this study a Cu^{2+}/EDTA/borax buffer system was used to calibrate a copper ion selective electrode in the low concentration range.

1.5. Mechanisms of Metal Uptake and Transport Across the Cell Membrane Found in Microorganisms

Metal uptake by microorganisms, defined as transport of nutrient metals through membrane(s) into the cytoplasm for further assimilation, appears to occur mainly in the following ways: nonspecific binding of metal to cell surfaces, passive diffusion of lipid soluble inorganic metal complexes, and specific transport (Doyle et al. 1975; Gadd and Griffiths 1978; Blust et al. 1986).

The first method of uptake is important since most heavy metals are easily adsorbed onto the surface of microbial cells, both living and dead. Copper can be complexed by polygalacturonic acid, one of the components of the outer layers of
bacterial cells (Gadd and Griffiths 1978). The hydrophobic nature of the cell membrane prevents passive movement of polar solute molecules such as inorganic salts, and ions. However, despite careful regulation of heavy metal uptake in cells, liposoluble uncharged complexes may penetrate cell membranes by becoming dissolved in the lipid phase of the membrane and so the second mechanism of metal uptake also occurs (Doyle et al. 1975).

The third mechanism of specific transport is widespread in microorganisms, and involves various transport systems. They can be classified on the basis of the mechanism and the protein components of a given system into the following groups: active transport, secondary active transport, binding protein dependent transport, and group translocation (Neidhardt et al. 1990) (Figures 1.3 and 1.4). Active transport (Figure 1.3A) is a process involving an energy-dependent pump in which the substance being transported combines with a stereospecific, transmembrane protein which then releases the chemically unchanged substance inside the cell (Brock and Madigan 1991).

A movement of a substrate across the cytoplasmic membrane due to a previously established ion gradient is called secondary active transport. It employs three types of proteins: symports (Figure 1.4A), antiports (Figure 1.4B) and uniports (Figure 1.4C). Uniports transport a substance from one side of the membrane to another. Antiports and symports move the substrate across the membrane along with a second substrate required for transport of the first. With antiports, the first and second substrate move in opposite
directions across the membrane, whereas with symports both substrates move in the same

Binding protein-dependent transport systems (Figure 1.3B) include a periplasmic
substrate-binding protein and three or four additional inner membrane-associated
proteins. The former serves as the primary recognition site for uptake, the latter are
involved in substrate translocation and energy coupling. This mechanism has been
proven to be ATP dependent (Higgins et al. 1990). Sensitivity to osmotic shock and
differential sensitivity to metabolic inhibitors distinguish this transport system from
others (Higgins et al. 1990). Group translocation (Figure 1.3C) is a mechanism in which
the substrate is chemically altered as it crosses the cytoplasmic membrane, usually via

A special uptake system has been identified in microorganisms grown under iron
limitation. Iron(III)-specific chelating compounds (siderophores) are responsible for
solubilization of iron(III) hydroxide aggregates and their further uptake by some
bacterial, yeast, and fungal species (Simpson and Neilands 1976; Trick et al. 1983).
Siderophores are released into the medium for subsequent transmembrane transfer of the
metal-ligand complex via specialized receptor proteins (Morel et al. 1991). Once
ferrisiderophore (siderophore with bound iron) has been linked to the receptor, iron
uptake may occur by two different mechanisms. In the first, only iron enters the
membrane while the ligand remains outside. In the alternative mechanism the
ferrisiderophore complex enters the cell and the chelate is either hydrolyzed or reused
after iron release (Martinez et al. 1990).
1.6. Copper Resistance and Metabolism in Bacteria

Copper plays a dual biological role as an essential micronutrient which is required for the synthesis of metalloproteins and as a growth inhibitor at high concentrations. Copper toxicity is associated with its ability to modify the active sites of metalloproteins and to catalyze membrane disrupting redox reactions in the cells. Therefore, the microorganisms must possess delicate mechanisms for maintaining such a restricted level of free copper within a cell that is neither limiting nor toxic (Sunda and Huntsman 1985; Lee et al. 1990; Silver and Walderhaug 1992; Brown et al. 1992b). A number of bacteria have been shown to be copper resistant due to nonspecifically produced high levels of extracellular polysaccharides (Bitton 1978), iron-binding siderophores (McKnight and Morel 1979) and other organic products (Mittelman and Geesey 1985; Hardstedt-Romeo and Guassia-Barelli 1980). However, some examples of a specific copper-inducible resistance have been identified.

Copper concentrations in bacterial cells can be regulated at two levels: metal sequestration and metal uptake. Metallothioneins, low molecular weight cysteine-rich metalloproteins are involved in sequestering intracellular copper in eukaryotes (Ecker et al. 1986; Butt et al. 1984). The marine bacterium Vibrio alginolyticus produces extracellular copper-binding proteins for copper detoxification (Harwood-Sears and Gordon 1990), while sulfate-reducing Desulfovibrio spp. tolerate high copper levels by producing sulfide that complexes copper and reduces copper toxicity (Cervantes and Gutierrez-Corona 1994). Sequestration of copper by a set of four periplasmic and outer
membrane proteins has been suggested as a resistance mechanism in *Pseudomonas syringae* (Mellano 1988; Cha and Cooksey 1991; Cooksey and Azad 1992; Cha and Cooksey 1993) (Figure 1.5). This mechanism has been well-studied. Four genes have been shown to encode these proteins, *copA, copB, copC* and *copD*. These genes have been sequenced and several copper binding motifs have been identified. Two additional genes *pcoR* and *pcoS* follow the *pcoABCD* operon and seem to be members of a two-component regulatory system involved in copper regulation (Cervantes and Gutierrez-Corona 1994). The proteins produced by *cop* genes are copper-inducible. It has been found that the periplasmic proteins CopA and CopC are able to bind 11 and 1 copper atoms per polypeptide, respectively (Figure 1.5). CopB is an outer membrane protein and CopD is a probable inner membrane protein. (Brown *et al.* 1992b; Silver 1992; Mellano 1988; Cha and Cooksey 1991; Cooksey and Azad 1992; Cha and Cooksey 1993, Cooksey 1994). It is thought that in addition to the copper sequestering function of CopC, it can also function in copper uptake, together with CopD (Cooksey 1994).

An alternative strategy for maintaining internal cupric ion concentrations is adopted by *Escherichia coli*. Four plasmid-encoded genes (*pcoARBC*) are required for copper resistance and seven chromosomal genes (*cutA-cutF, cutR*) are involved in copper metabolism in *E.coli* (Figure 1.6). Under non-toxic copper concentrations, copper enters the cell via two uptake proteins: CutA and CutB. CutA is a specific copper uptake protein, whereas CutB is also involved in Zn uptake. There are two intracellular copper-binding storage/carryer proteins (CutE and CutF) that may be responsible for preventing Cu(I) from damaging intracellular components. Export of copper that exceeds
physiological requirements is mediated by two structural proteins: CutC and CutD, which are probably copper efflux ATPases (Cervantes and Gutierrez-Corona 1994). Under conditions of high copper concentrations, plasmid-encoded pco genes are expressed. It appears that pcoA and pcoB genes interact with cutC and cutD to export extra copper from the cell, whereas pcoC is involved in cytoplasmic copper storage/transport. CutR and pcoR genes have been suggested to be involved in regulation of copper metabolism and resistance in *E. coli* and they seem to belong to the large family of bacterial two-component regulatory systems involved in sensing environmental signals (Silver and Walderhaug 1992; Brown *et al.* 1992b; Lee *et al.* 1990; Rogers *et al.* 1991; Cooksey, 1993; Gupta *et al.* 1995).

Recent work has correlated rearrangements in the gene encoding a copper-transporting P-type ATPase with Menkens syndrome, a lethal disease that involves reduced activity of numerous copper-containing proteins (Vulpe 1993). There are, as well, several examples of bacterial copper transport P-type ATPases. Two copper efflux ATPases have been identified in *Enterococcus hirae* (Oddermatt *et al.* 1993), the above-mentioned proteins CutC and CutD of *Escherichia coli* have been suggested to be copper efflux ATPases (Cervantes and Gutierrez-Corona 1994), and a *Helicobacter pylori* gene encoding a putative copper ATPase has been cloned and sequenced recently (Ge and Taylor 1995). Examples of other cation P-type ATPases, both eukaryotic and bacterial, include Na⁺, K⁺-ATPase isolated from sheep kidney (Shull *et al.* 1985), K⁺-ATPase of *Escherichia coli* (Hesse *et al.* 1984), Ca²⁺-ATPase of rabbit sarcoplasmic reticulum (Hesse *et al.* 1984; Brandl *et al.* 1986), a putative cation ATPase of *Synechococcus* sp.
(Kashiwagi et al. 1995), Cd\(^{2+}\)-efflux ATPase of Staphylococcus aureus (Nucifora et al. 1989), K\(^{+}\)ATPase of Streptococcus faecalis (Soloiz et al. 1987), and many others.

These enzymes belong to a family of membrane-bound cation translocation pumps consisting of a main polypeptide chain of about 110,000 MW that perform both catalytic and transport functions. In some cases there are additional subunits or dimerization (Cervantes and Gutierrez-Corona 1994; Inesi and Kirtley 1992). There are several common structural features in all P-type ATPases despite the variety of ions they transport. The polypeptide chain includes a relatively large extramembranous domain and eight to ten transmembrane helical structures that span the membrane bilayer, forming four to five hairpins (Figure 1.7). All ion transporting ATPases contain a phosphorylation domain with the conserved amino acid sequence DKTG(T/L)T and an aspartyl residue that is involved in the formation of a covalent intermediate (E-P) during the reaction cycle (Figure 1.8). Another common feature is two conformational forms of the phosphorylated intermediate, referred to as E1 and E2, covalently identical but differing in their affinity to substrates (Epstein 1990).

Despite the importance of copper to the physiology of methanotrophs, nothing is presently understood about the molecular mechanisms of copper acquisition from the environment. The research presented in this thesis investigates this area.
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1. Methane monooxygenase
2. Methanol dehydrogenase
3. Formaldehyde dehydrogenase
4. Formate dehydrogenase

Figure 1.1 The oxidation of methane to carbon dioxide by methanotrophs (from Murray and Richardson 1993).
Figure 1.2 Major products of TCE degradation by methanotrophic bacteria. Methanotrophs mediate steps indicated by the thick arrows, the thin arrows indicate spontaneous breakdown of TCE epoxide in water; the dashed arrows reflect processes that are stimulated by heterotrophic bacteria (from Little et al. 1988). Chloral is produced at a low level (1-5% of total products), but it does not accumulate in whole cells (Newman and Wackett 1991).
Figure 1.3 Examples of microbial nutrient transport systems (modified from Cronan et al. 1989).

Upper panel shows initial step, lower panel shows conformation after transport.

A. Active transport of a nutrient (b) involves conformational change of the stereospecific, transmembrane protein (a).

B. Binding protein-dependent transport system. b, periplasmic nutrient; c, binding protein; d-g, inner-membrane-associated proteins. f and g proteins couple ATP hydrolysis to the transport.

C. Group translocation is a mechanism in which the substrate (b) is chemically altered, usually phosphorylated, as it is transported by a protein h.
Figure 1.4 Secondary active transport through symports, antiports and uniports: schematic presentation (from Neidhardt et al. 1990).
Panel A shows a symport; panel B displays an antiport; panel C exhibits a uniport. 
A', anion; C', cation; S, uncharged substrate, H', proton.
The periplasmic proteins CopA and CopC, and the outer membrane protein CopB are involved in sequestering copper. CopC and CopD (inner membrane protein) are thought to be involved in copper transport into the cytoplasm.
Figure 1.6 A model of copper resistance and metabolism in *E. coli*.

Copper enters the periplasm through a porin X. CutA and CutB are copper uptake proteins; CutE and CutF are storage/intracellular transport proteins; CutC and CutD are copper efflux ATPases. The copper resistance proteins increase intracellular copper storage (PcoC) and increase the export of copper (PcoA and PcoB). CutR, involved in regulation of copper metabolism, is not shown on the diagram. IM, inner membrane; OM, outer membrane.
Figure 1.7 A diagram of the general structure of P-type ATPases (from Epstein, 1990).

The conserved feature is a cation channel formed by the six membrane-spanning regions I-VI. The dashed line indicates the C-terminus region where variation in the number of membrane-spanning domains is possible. The regions of highly conserved protein sequence are circled. Region 1 is the phosphatase domain, which removes the phosphate from an aspartic acid residue as part of the reaction cycle. Region 2 is the phosphorylation domain (shown phosphorylated). Region 3 is the ATP-binding domain. The external loops labeled a and b are hydrophilic in most P-type ATPases.
**Figure 1.8** A schematic diagram of the reaction cycle of the Na⁺,K⁺-ATPase (from Epstein, 1990).

Covalent binding of enzyme (E) and phosphate (P) is indicated by a line (-), tight but noncovalent binding is shown by a dot (•).

**Step 1.** E₁ with bound ATP is phosphorylated and binds three Na⁺ ions from the inside of the cell.

**Step 2.** Conformational change of P-E₁ to P-E₂ with release of Na⁺ outside of the cell.

**Step 3.** E₂ occludes two K⁺ ions from the outside and releases phosphate.

**Step 4.** E₂ releases K⁺ inside the cell, while binding ATP.
Chapter Two

Study of Copper Accumulation by the Type I Methanotroph
Methylomicrobium albus BG8

2.1. Introduction

Copper plays a significant role in the physiology of methanotrophs, Gram-negative bacteria that grow aerobically on methane as a sole source of carbon (1), by activating and stabilizing a key enzyme, particulate methane monooxygenase (pMMO). The addition of copper to the growth medium in the range that is neither growth-limiting nor toxic results in increased cell yield and pMMO activity, as well as a pronounced change in pMMO kinetics (2).

There has been considerable interest in these bacteria in recent years because of the possibility of using methanotrophs for the removal of trichloroethylene (TCE) from contaminated aquifers (3, 4). These microorganisms can degrade TCE through MMO to non-toxic compounds that can be utilized by heterotrophic bacteria (5-7). A variation in copper concentrations in the growth medium produces a significant change in the kinetics of TCE oxidation (2). Such a response to copper could have a major influence on TCE bioremediation by natural populations of methanotrophs. However, very little is presently known about the mechanism of copper uptake or the bioavailability of different copper.

species in these bacteria. One study of mutants in the type II methanotroph, *M. trichosporium* OB3b, has suggested that a specific uptake system exists in this strain (8).

Several copper species can be present simultaneously in natural waters as well as in a culture medium. These include aqueous cupric ion [Cu$^{2+}$(H$_2$O)$_6$], which we will refer to as cupric ion, inorganic complexes (e.g., hydroxide and carbonate complexes), and organic complexes (9). However, studies of copper toxicity in microorganisms have demonstrated that cupric ion is the most bioavailable chemical species and that chelators affect total copper availability by altering cupric ion concentrations or activities (10-16). Three mechanisms of trace element uptake by microorganisms have been proposed: nonspecific sorption on cell surfaces, metabolism-dependent intracellular uptake, and passive diffusion of lipid-soluble inorganic metal complexes (12, 17, 18). It is not known in methanotrophs whether the uptake of copper is mediated by an active or passive transport mechanism.

In our work, we considered three pools of copper, as depicted in Figure 2.1: copper in the growth medium ("bulk copper"), copper sorbed on the cellular surface ("surface-associated copper"), and copper inside the cells ("internal copper"). We refer to the sum of surface-associated and internal copper as "accumulated copper." "Total copper" refers to the amount of copper that was added to the medium as copper nitrate, and includes various copper species present in all three pools.

The objectives of the work presented here were two-fold. The first objective was to study the effect of a range of total copper and chelator (EDTA) concentrations on copper accumulation by *M. albus* cells and to determine the relationship between cupric
ion concentrations in the external medium and cellular copper concentrations. The second objective was to establish the existence of an active transport system for copper in *M. albus* BG8. An EDTA wash was used to remove surface-bound copper and to separate copper uptake from copper sorption. These experiments were carried out under conditions in which copper was neither growth-limiting nor toxic, since the physiological effects of copper on methanotrophs that have been observed occur under these conditions (2, 19).

**2.2. Experimental Section**

**2.2.1. Bacterial Strain and Growth Conditions.**

*M. albus* BG8 (20), a type I methanotroph, was used. The bacteria were grown on nitrate mineral salts (NMS) medium (1), under an atmosphere of 1:1 methane/air mixture at 30°C. Liquid cultures were inoculated from agar plate cultures and grown with shaking at 200 rpm. Total copper in the medium used for agar plates was 10 μM unless stated otherwise. Concentration of copper in the stock solution was verified with mass spectrometer measurements.

All chemicals used for medium preparation were of reagent grade. All solutions were made with double distilled water (dd H₂O). All glass flasks, jars, and polypropylene beakers used were presoaked in 1N HCl overnight, and then rinsed with dd H₂O three times to minimize trace metal contamination.
Copper nitrate was filter sterilized and was added aseptically to autoclaved medium to give the desired total copper concentration.

NMS medium (1) contains 11.7 μM EDTA including 11 μM Fe-EDTA complex and 0.7 μM EDTA, which is added to the medium from a preequilibrated stock of trace elements and is mostly present in the form of metal-EDTA complexes. A medium containing metal-EDTA complexes, Fe-EDTA in particular, equilibrates with cupric ion relatively slowly (9, 16). We found that in NMS medium, 24 h is needed to achieve a pseudoequilibrium, after which there is no further significant decrease in cupric ion concentration (Figure A2.1). We use the term pseudoequilibrium to refer to a state of the system in which all fast exchange reactions are completed, but true equilibrium has not been established due to the slow kinetics of exchange for Fe-EDTA against cupric ion. Therefore, in all experiments performed we allowed at least 24 h of medium incubation before inoculation with *M. albus* BG8 to avoid short-term high cupric ion concentration toxic effects on cells and to bring the system to pseudoequilibrium.

2.2.2. Ion Selective Electrode (ISE) Pretreatment and Calibration.

The Orion (Boston, MA) 942900 solid-state Copper ISE (CuISE) and the Orion 900200 double junction, half-cell electrode were used to measure cupric ion concentrations in the medium. The electrodes were connected to an Orion 710A pH/mV meter. Ionic strength adjuster (0.1M sodium nitrate, final concentration) was added to all standards and samples to swamp small variations in sample ionic strength and to keep the activity coefficient constant. The electrodes were stored and cleaned as described by Avdeef (27).
All readings were taken when the rate of change of potential had fallen below 0.2 mV/min, as was recommended by Simpson (22). All measurements were performed at 25 °C.

In the range $10^{-3} - 10^{-5}$ M cupric ion, successive dilutions of a 0.1 M standard solution of CuNO₃ in dd H₂O were used as calibrating solutions. Lower cupric ion concentrations for CuISE calibration were obtained with cupric ion buffers [Cu²⁺(H₂O)₅/EDTA with borax buffer for pH control], prepared as described by Hansen (23). The calibrations were always performed at the beginning of the series of solutions and checked every 3 h thereafter.

The response of the electrodes was tested and has been found to be linear in the range of pCu 3-10, as expected (23) (Figure A2.2). In 42 calibrations performed during the experiments, the mean slope was 27.8 mV per decade of cupric ion concentration change. The calculated ideal theoretical Nernstian slope is 29.5 mV/pCu at 25°C.

2.2.3. Mass Spectrometry.

A Perkin-Elmer (Norwalk, CT) 5000 inductively coupled plasma/mass spectrometer (ICP/MS) was used to analyze solutions for total copper concentrations. The additions calibration technique was used to compensate for sample matrix effects (24). Copper was measured as $^{63}$Cu since it is more abundant than $^{65}$Cu. Diluted solutions of 1000 ppm copper nitrate standard were used for ICP/MS calibration (Figure A2.3). A diluted yttrium nitrate standard solution served as an internal standard. Both standards were obtained from Perkin-Elmer.
The nebulizer flow was optimized before each series of measurements to minimize oxide interferences. All samples were diluted 10-fold before analysis to meet ICP/MS requirements on total dissolved solids (maximum of 2%) (24).

Parameters were set as follows: dwell time of 30 ms; 7 sweeps per reading; 7 readings per replicate; 1 point per pick. Measurements were performed on high-resolution mode.

2.2.4. Optical Density Measurement and Calibration.

Optical density was measured with a Klett - Summerson (Long Island City, NY) photoelectric colorimeter and was reported as Klett units. Optical density was calibrated to viable cell numbers by dilution plating of cell suspensions at several Klett unit values. One Klett unit was considered equal to \((6.7\pm0.3)\times10^5\) viable cells.

2.2.5. Copper Accumulation.

The accumulation of cupric ion by *M. albus* BG8 cells was determined by growing cells in NMS medium containing eight total copper concentrations: 3, 4.2, 8.4, 12.3, 17.3, 19.1, 28.1, and 37.0 \(\mu\)M as measured by ICP/MS (Table 2.1). In all experiments the pH of the medium before inoculation was 6.5, and did not change during growth of *M. albus* BG8. Cell-free NMS medium was used as an adsorption control (Figure A2.4). The medium in control and experimental flasks for each experiment was prepared exactly the same way.

After preequilibration of the medium (250 mL in each flask) for at least 24 h, the experimental flasks were inoculated with *M. albus* BG8. Just prior to BG8 addition and at
different times afterwards, 5-mL samples were collected for cupric ion concentration, pH, and ICP/MS measurements. At the same time optical density of the growing cells was measured with a Klett colorimeter. Duplicate 1-mL cell-containing subsamples were prepared from 5-mL samples by centrifugation at 16000g for 10 min, at room temperature. The supernatant phases were collected, diluted and measured for copper by ICP/MS. The pelleted cells were washed with 1 mL of dd H₂O, and centrifuged again. The washing solution was retained for ICP/MS analysis in several experiments and was found to contain a negligible amount of copper (less than 1% of the added amount). Cells then were harvested, resuspended and lysed. The obtained subsamples with cell digests and cell-free samples (controls) were diluted and measured with ICP/MS.

The average moles of copper per cell was calculated by dividing the quantity of moles of copper in 1 mL of cell pellet by the number of cells in 1 mL of pellet, assuming no cell loss during centrifugation.

The SURFEQL computer model (25) was used to predict copper speciation in NMS medium prior to inoculation (after 24-hour preequilibration). The program was run at fixed hydrogen ion concentration (pH= 6.5), corresponding to the pH of NMS medium. The equilibrium constants were from the SURFEQL database. Case-specific input included total concentrations of metals and ligands present in the medium. We considered two limiting cases. For the equilibrium case we assumed that all EDTA, including EDTA added as an iron complex, was available for complexation. In the second, nonequilibrium case, the Fe-EDTA complex was presumed to have a low chemical reactivity with respect to change in complexation (9), thus being kinetically
unavailable. A negligible amount of copper (less than 2% of the added amount) was measured to partition as solid precipitates with the cells during centrifugation.

2.2.6. Copper Accumulation in M. albus BG8 under Different Complexation Conditions.

Four complexation experiments were performed with different total EDTA concentrations (11.7, 15.7, 21.7, 31.7 μM). It was not possible to use EDTA concentrations outside of this range because of poor growth of the culture, presumably due to altered bioavailability of required trace elements under these conditions. In each experiment the cells were exposed to eight total copper concentrations in the range from 1 to 100 μM. Cellular copper concentrations for these experiments were measured at the end of logarithmic growth, but no later than 30 h. after inoculation. Ionic copper concentrations were measured with ISE before M. albus BG8 inoculation in all experiments (Table 2.1). The samples were prepared and analyzed as described in the previous section, except that three replicates instead of two were prepared for ICP/MS measurements.

2.2.7. Sorption Estimates.

Cells were grown on agar plates containing either 10 μM or 2 μM copper. The cells from plates containing 10 μM copper were exposed to a range of total copper concentrations from 1.2 to 54 μM as measured by ICP/MS. The cells from plates containing 2 μM copper were inoculated into liquid NMS medium containing total copper in the range from 0 to 7 μM. The total EDTA concentration in the medium was
11.7 µM except for one experiment where it was increased to 15.7 µM and 31.7 µM, while copper concentrations ranged from 0 to 2.8 µM. Ionic copper concentrations were measured with ISE prior to *M. albus* BG8 inoculation in all experiments. All samples were prepared in duplicate.

Sorption experiments were started by inoculation of *M. albus* BG8 into preequilibrated NMS medium. At time intervals, the optical density of the growing cells was measured with a Klett colorimeter, and 1-mL samples were removed for either further ICP/MS analysis (untreated samples), to assess copper accumulated by the cells (sorbed plus internal) or for an EDTA wash (washed samples) to remove sorbed copper. The untreated samples were prepared and analyzed as described above. The samples intended for EDTA wash were centrifuged at 16000g for 10 min, at room temperature. The pelleted cells were washed with 1 mL dd H₂O, and centrifuged again. The collected cells then were resuspended in dd H₂O containing 0.1M EDTA (final concentration) for 24 h. The cells were then harvested by centrifugation, washed with 1 mL dd H₂O, and prepared for ICP/MS measurements as described above.

### 2.2.8. Study of Uptake of Cell-Surface Associated Copper.

*M. albus* BG8 was exposed to five total copper concentrations: 2.3, 3.5, 7.5, 39.0 and 55 µM as measured by ICP/MS. The total EDTA concentration in the medium was 11.7 µM. Within 1 h after inoculation, 1-mL samples were withheld for ICP/MS analysis with or without EDTA wash to determine the amount of sorbed copper. After 1.5 or 3.5 h of incubation (at 30°C, with shaking at 200 rpm) the remaining cells were harvested by
centrifugation of the flask contents (150 mL) at 16000g for 10 min, at room temperature.
The pelleted cells were washed with 10 mL of 1mM PIPES buffer (pH=6.5) to remove
loosely-bound copper, and centrifuged again. The collected cells then were transferred
into modified NMS medium, which differed from the regular NMS by the absence of
copper, Fe-EDTA, and EDTA. Otherwise growth conditions were normal.

At time intervals, the optical density of the growing cells was measured with a
Klett colorimeter, and 1-mL untreated and washed samples were prepared as described
above. Two replicates were made on each sample.

2.3. Results And Discussion

2.3.1. Copper Accumulation.

Because of the importance of cupric ion to copper bioavailability in other systems,
the relationship between copper accumulation and cupric ion concentration in the
medium was studied in actively growing cultures of M. albus BG8. The cells were grown
in medium containing eight different total copper concentrations. No significant change
in the growth rate of M. albus BG8 was observed within the total copper concentration
range from 3 to 37 μM (Figure 2.2), showing that copper was both sufficient and non-
toxic to M. albus BG8 in these experiments.

Cupric ion concentration in NMS medium was determined during growth of M.
albus BG8 for all eight initial copper concentrations tested. Since the same general trends
were observed in all eight experiments, the results for only the lowest (3 μM) and highest
(37 μM) total copper concentrations tested are presented in Figure 2.3A (all the other results are shown in Figures A2.4-A2.6). The cupric ion concentration decreased with time, while cell-free controls showed no significant cupric ion (Figure 2.3A) or total copper loss (Figure A2.7) within the time of the experiments. This allowed us to attribute the change in cupric ion concentration to the presence of *M. albus* BG8, and to assume that the effect was not due to further copper complexation by EDTA present in the medium or to copper adsorption to the walls of the flasks.

In all eight experiments, the same pattern of cupric ion concentration in the medium was observed: an initial rapid depletion within 10 h after inoculation, which was followed by a slow but detectable decrease. A different pattern was seen for the cellular copper accumulation. In this case, there was rapid accumulation within the first hour after inoculation, and only a small further accumulation occurred (Figure 2.3B). The difference in the patterns could be explained if the rapid initial cellular accumulation of copper disturbs the equilibrium between cupric ion and other copper species in the medium. SURFEQL predictions indicate that the two main contributors to the pool of bulk copper are Cu-EDTA complexes and Cu₃(PO₄)₂ precipitates, and establishment of equilibria of cupric ion with these pools is expected to be slower than the rapid cellular accumulation rates observed (9, 16) (Figure 2.1). It is possible that the two phases of cupric ion decrease in the medium reflect two different processes in which equilibria are being reestablished, perhaps in combination with ongoing accumulation in the first phase. In order to examine the dependence in *M. albus* BG8 of the cellular copper content on
cupric ion concentration in the medium, accumulation experiments under different complexation conditions were carried out.

2.3.2. Copper Accumulation in *M. albus* BG8 under Different Complexation Conditions.

*M. albus* BG8 cells were grown in media containing different total copper concentrations and different EDTA concentrations. Before inoculation, initial cupric ion concentrations were determined (Table 2.1). At the end of growth the copper content per cell and the copper in the medium were determined as described in the Experimental Procedures. Metal recovery (sum of cell digest and filtrates) based on the measured total copper added was 101% (average). All replicates showed very good reproducibility: most of the error bars on Figures 2.4 and 2.5 are within the symbols that represent the averaged data. However, cupric ion concentration measurements showed more variability (Table 2.1).

When the copper per cell from these experiments and from the experiments shown in Figure 2.3 was plotted against the negative logarithm of the total copper concentration (Figure 2.4), a decrease in copper content per cell was seen as the EDTA concentration increased. The results of experiments with the two highest EDTA concentrations (21.7 and 31.7 μM) are similar, presumably because in these cases copper complexation by EDTA produced almost the same initial cupric ion concentrations, as measured with ISE (Table 2.1).
In contrast to Figure 2.4, the copper content of *M. albus* BG8 cells plotted as a function of pCu for all experiments falls on one curve (Figure 2.5), suggesting that cellular accumulation of copper depends on the cupric ion concentration. A hyperbolic relationship between cell copper content and cupric ion activity was observed in a similar study with an alga (10). We found that the cellular copper content of *M. albus* BG8 also demonstrates a hyperbolic dependence with cupric ion concentration (Figure 2.5). The following model was used, based on that presented by Sunda and Guillard (10):

\[
\text{Cu/cell} = \frac{1.54 \times 10^{-15} [Cu^{2+}]}{[Cu^{2+}] + 1.43 \times 10^{-7}},
\]

where \([Cu^{2+}]\) is the cupric ion concentration, \((1.54 \pm 0.06) \times 10^{-15} \text{ mol/cell}\) is an estimation of a mean maximum binding capacity, and \((1.43 \pm 0.05) \times 10^{-7} \text{ mol/L}\) reflects an apparent half-saturation constant.

The observed hyperbolic dependence is consistent with two main hypotheses. The first assumes reversible binding of copper to several similar cellular ligand sites with their eventual saturation (10). The second involves carrier-mediated transport with saturation of transport molecules (26).

### 2.3.3. Sorption versus Uptake (Surface-Associated versus Internal Copper).

In order to determine what portion of the copper accumulated by *M. albus* BG8 cells was due to nonspecific sorption, experiments were carried out in which cells were washed with EDTA. It is assumed that most of the copper removable by EDTA is nonspecifically sorbed to external sites. Cells were grown first on agar plates containing 2 μM or 10 μM copper and then in a medium containing different total copper
concentrations. Samples were withdrawn at different time points during growth, and the copper content in cells that were untreated and cells incubated with 0.1 M EDTA for 24 h and then washed was analyzed and reported as moles of copper per cell. Figure 2.6 shows the results of three experiments with NMS medium containing 1.2, 12, and 43 μM total copper. The other experiments demonstrated the same pattern of copper incorporation (Figure A2.8).

In all cases the amount of copper per cell increased significantly after exposure of the cells to the copper-containing medium, and then as cells increased in number, the average amount of copper per cell decreased and leveled out for both untreated and washed samples (Figure 2.6). The pattern of total copper accumulation by the biomass in these cultures was similar to that shown in Figure 2.3B (Figure A2.9). These data suggest that in both untreated and washed cells no significant copper accumulation occurs after the initial accumulation on a per cell basis.

A significant difference in the total copper per cell was observed between the treated and untreated samples, with as much as a 40-fold decrease after EDTA treatment at the higher copper concentrations. These data suggest that most of the copper accumulated by the cells in these experiments was sorbed to the cellular surface at all copper concentrations tested. The pattern of copper accumulation seen in untreated samples suggests that cells inoculated from agar plates rapidly sorb copper, but as they grow exponentially in liquid medium, new copper sorption does not occur at a significant level. This may be due to changes in cellular surface characteristics, for instance amounts
of extracellular polysaccharide synthesized, under these two different physiological conditions.

In all of these experiments, rapid saturation of cellular accumulation of copper in both EDTA-treated and non-treated samples occurred (Figure 2.6). However, while the amount of copper measured in untreated cells showed the expected dependence on the pCu of the medium, the amount of copper found in the cell digests after EDTA wash was essentially constant, at about $3 \times 10^{-17}$ mol of copper per cell for experiments that had been inoculated from plates containing 10 $\mu$M copper, and at about $1 \times 10^{-17}$ mol of copper per cell for inoculation from plates containing 2 $\mu$M copper, despite a 100-fold change in both cupric ion and total copper concentrations (Figure 2.7). This constancy was seen both for samples collected within 1h of inoculation of cells into copper-containing medium (Figure 2.7) and for samples prepared at the end of growth, 25-30 h after inoculation (Figure A2.10).

Copper that is not removable by a strong chelating agent like EDTA during a 24-h exposure and wash treatment may be either inside the cells or complexed on the outside of the cell at sites with higher affinity for copper than EDTA. However, the literature indicates that microorganisms are not normally capable of copper uptake from metal-EDTA complexes present in the medium (9, 12, 27, 28) and, therefore, the existence of copper-binding sites with such high affinity seems unlikely, suggesting that the copper not removable by EDTA may be inside the cells (28). If this copper does represent copper inside the cell, then an analysis of Figure 2.7 shows that the saturation of copper
uptake into the cells could be achieved at approximately 1-3\times10^{17} moles of copper per cell.

In phytoplankton, the uptake of essential trace metals is believed to occur via binding to a surface ligand and subsequent transfer across the cell membrane (29). This process usually follows Michaelis-Menten kinetics. Our results are also consistent with such a mechanism (Figure 2.1). However, we were unable to estimate Michaelis-Menten parameters from our data, because the rapid uptake precluded measuring the initial uptake rates in these experiments by our protocol.

Since copper uptake into the cells was saturated in our experiments and because most of the copper accumulated by the cells was sorbed to the cellular surface, the hyperbolic model presented above reflects mainly the reversible binding of copper to cellular sites with a mean binding capacity of 1.54\times10^{-15} moles/cell, and an apparent half saturation constant of 1.43\times10^{-7} moles/l. The rapid adsorption saturation seen in our experiments (untreated samples) probably reflects saturation of two types of sites: a relatively small number of specific, possibly membrane-bound, sites associated with a putative copper transport system, and a relatively large number of nonspecific sites, possibly reflecting various functional groups such as amino acid, carboxylic, hydroxy groups, etc. (Figure 2.1). Since in all cases the amount of copper removable by EDTA was greater than that which was not EDTA removable, the kinetics of sorption observed would be expected to be dominated by the nonspecific sites.
2.3.4. Desorption of Cell-Surface Associated Copper and Uptake into the Cells.

To study the fate of the cell surface-bound copper (presumably on non-specific sites), a series of experiments was conducted in which *M. albus* BG8 cells were incubated in copper-containing NMS medium for a short period of time (1.5 or 3.5 h) to allow accumulation of copper and then were transferred to copper-free medium. Samples were then taken at various times and copper was measured in four pools: total cellular copper accumulated by the cells, cellular copper not removable by EDTA, copper removed with the EDTA treatment, and copper in the filtrate (medium from which cells had been removed). Since the results were the same in both sets of experiments (1.5 or 3.5 h of incubation), data from only the experiments with 1.5 h. incubation are shown.

In all experiments, cells grew slower than usual after transfer to copper-free medium. Doubling time increased on the average by a factor of 2 as compared to growth in regular NMS medium. This is probably due to the absence of EDTA in the copper-free medium, which is important in providing solubility and availability of trace elements essential for growth. EDTA was excluded from the copper-free NMS medium in these experiments to prevent stripping of copper from the cellular surface.

A comparison of the data for the untreated and EDTA-treated cells shows that the protocol gave good recovery and good reproducibility. The measured copper accumulated by the cells was approximately equal to the sum of the EDTA-removable and nonremovable copper (Figure 2.8). An insignificant amount of copper was found in the filtrate in the experiments with 2.3, 3.5, or 7.5 μM total copper concentration (Figures
2.8A and 2.8B) suggesting that little copper was desorbed from the cell surface during this time period. The total amount of copper accumulated by the cells and the copper not removable by EDTA both remained approximately constant in these experiments. When cells were grown initially with higher copper (39 μM), the amount of copper lost to the medium was substantial (Figure 2.8C), and was correlated with the loss of copper from the cells. However, in this case, the copper not removable by EDTA was approximately constant (Figures 2.8A-C). This implies that under these conditions, the copper species sorbed on non-specific sites was not further internalized by *M. albus* BG8 cells either directly, or indirectly after re-entering the medium. This lack of availability of desorbed species for further uptake during the tested time period may be due to kinetic factors: an imbalance between rates of desorption, reequilibration in the medium, and copper uptake (Figure 2.1).

The combined results of our experiments suggest the scenario shown in Figure 2.1, in which the cupric ion is directly available for accumulation including both sorption and internalization by *M. albus* BG8, and suggest that the rates of copper accumulation are greater than the dissociation and medium reequilibration rates. Therefore, cupric ion concentration in the medium is probably kinetically controlled by a balance between all sources and sinks (Figure 2.1) and this determines overall copper availability to *M. albus* BG8. Moreover, the results of sorption versus uptake experiments demonstrated saturable kinetics of copper uptake and independence of cellular uptake from a 100-fold extracellular copper concentration variation. This suggests that cupric ion is probably taken up by *M. albus* BG8 via a specific transport system (Figure 2.1).
The data presented here suggest that the growth of M. albus BG8 will not be copper-limited in most natural environments due to their efficient copper uptake system(s), which are yet to be studied in detail. This is supported by previous reports suggesting that natural populations of methanotrophs are not copper-limited (2, 8). If we wish to employ a type I methanotroph in TCE removal from a contaminated site, it may not be necessary to fertilize soil with copper to encourage high TCE degradation rates.
2.4. References


Table 2.1  Results of ICP/MS and ISE Measurements of the Initial Concentrations of Total Copper and Cupric Ion.

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Copper Pools  

1) Bulk  
2) Surface-Associated  
3) Internal

Figure 2.1 Model for copper accumulation by *M. albus* BG8.

Three pools of copper are assumed: (1) copper in the growth medium (bulk copper), including CuL\textsubscript{inorg} (soluble copper complexes with inorganic ligands, such as CuCO\textsubscript{3}, CuHPO\textsubscript{4}, CuSO\textsubscript{4}, (CuOH)\textsuperscript{+}, etc.), Cu\textsubscript{3}(PO\textsubscript{4})\textsubscript{2} precipitate, CuEDTA and Cu\textsuperscript{2+}(H\textsubscript{2}O)\textsubscript{6}; (2) surface-associated copper including transport site complexes (CuX) and copper sorbed on non-specific surface sites removable by EDTA (CuY); and 3) internal copper (Cu\textsubscript{in}). Arrow widths show the probable relative rates of each reaction when transport system is not saturated.
Figure 2.2 Specific growth rate observed within a total copper concentration range from 3 to 37 μM.

Total copper is the amount of copper nitrate added to the growth medium.
Figure 2.3 Cupric ion concentration and copper accumulation by *Methylococcus albus* BG8 during growth.

Cells were inoculated into the medium at time 0.

A. **Cupric ion concentration in the medium.** Filled symbols denote 3μM, open ones denote 37μM total copper added to NMS. Circles show optical densities as measured by the Klett colorimeter. Squares indicate cupric ion concentration measured by CuISE in the control flasks (no *M. albus* BG8). Triangles show the time course of cupric ion disappearance from the medium in the flasks inoculated with *M. albus* BG8 as measured with CuISE.

B. **Change in relative amount of cell-associated copper during the time course of the experiment.** Filled diamonds indicate 3μM and open ones indicate 37 μM total copper added to NMS.
Figure 2.4 Cellular copper content of *M. albus* BG8 versus negative logarithm of total copper added to the medium.

○ , [EDTA]=11.7 μM; ▼ , [EDTA]=15.7 μM; □ , [EDTA]=21.7 μM; 
△ , [EDTA]=31.7 μM;
○ , data from Figure 2.3 ([EDTA]=11.7 μM).
Figure 2.5 Cellular copper content of *M. albus* BG8 versus negative logarithm of the cupric ion concentration (pCu) in the medium at different EDTA concentrations.

Symbols are as in Figure 2.4.
The curve is a plot of the hyperbolic model.
Figure 2.6 Copper accumulation in cells either untreated or incubated with EDTA and then washed.

Filled symbols represent untreated samples; open ones indicate washed samples (incubated in 0.1M EDTA for 24 h and then washed).

Total copper concentrations in the medium:
circles - 1.2 \mu M, triangles - 12 \mu M, squares - 43 \mu M.

Growth curves were similar to those shown in Figure 2.3A.
Figure 2.7 Copper accumulated in cells within 1h of inoculation during sorption experiments versus total copper concentration and cupric ion concentration.

Circles, experiments were inoculated from agar plates containing 10 μM copper; triangles, experiments were inoculated from agar plates with 2 μM copper.
Figure 2.8 Fate of cell-surface associated copper.

Averaged data (from two replicates) are shown in all figures. Filled bars, copper measured in the cell digests (mol); hatched bars, copper measured in the filtrates (mol); open bars, copper measured in EDTA-treated fractions (mol). Plots on the left represent measurements in untreated samples; plots on the right reflect measurements in washed samples. Panels A-C show data for three experiments with the following total copper concentrations used for growth before transfer to copper-free medium: 2.3 μM, 7.5 μM, and 39 μM.
Appendix

In this appendix, figures that are relevant to Chapter Two but were not included in the original paper (Berson and Lidstrom 1996) are presented.
**Figure A2.1** The decrease in free copper concentration during media preequilibration.

The time required to achieve a pseudoequilibrium in NMS media at various total copper concentrations was studied. Open circles, 3 μM total copper; solid circles, 7 μM total copper; triangles, 12 μM total copper in NMS media.
**Figure A2.2** One of the cupric ion selective electrode calibrations obtained with cupric ion buffers [CuNO₃/EDTA/borax].

This particular calibration gave a slope of 28 mV/pCu in the range from $10^{-10}$ to $10^{-3}$ M. The calculated ideal theoretical Nernst slope is 29.5 mV/pCu at 25 °C.
Figure A2.3 An example of ICP/MS addition calibration with copper nitrate standards.
Figure A2.4 Cupric ion concentration and copper accumulation by *Methylomicrobiun albus* BG8 during growth; 4μM (filled symbols) and 28 μM (open symbols) total copper added to NMS.

Cells were inoculated into the medium at time 0.

A. **Cupric ion concentration in the medium.** Circles show optical densities as measured by the Klett colorimeter. Triangles indicate cupric ion concentration measured by CuISE in the control flasks (no *M. albus* BG8). Squares show the time course of cupric ion disappearance from the medium in the flasks inoculated with *M. albus* BG8 as measured with CuISE.

B. **Change in relative amount of cell-associated copper during the time course of the experiment.**
Cells were inoculated into the medium at time 0.

A. Cupric ion concentration in the medium. Circles show optical densities as measured by the Klett colorimeter. Triangles indicate cupric ion concentration measured by CuISE in the control flasks (no *M. albus* BG8). Squares show the time course of cupric ion disappearance from the medium in the flasks inoculated with *M. albus* BG8 as measured with CuISE.

B. Change in relative amount of cell-associated copper during the time course of the experiment.

**Figure A2.5** Cupric ion concentration and copper accumulation by *Methylomicrobiunm albus* BG8 during growth; 12μM (open symbols) and 19 μM (filled symbols) total copper added to NMS.
Cells were inoculated into the medium at time 0.

A. **Cupric ion concentration in the medium.** Circles show optical densities as measured by the Klett colorimeter. Triangles indicate cupric ion concentration measured by CuISE in the control flasks (no *M. albus* BG8). Squares show the time course of cupric ion disappearance from the medium in the flasks inoculated with *M. albus* BG8 as measured with CuISE.

B. **Change in relative amount of cell-associated copper during the time course of the experiment.**
Figure A2.7 Total copper concentration in adsorption controls (cell-free NMS medium) versus time.

The following total copper concentrations were studied:

- ▼, 3.0 µM; □, 4.2 µM; ▲, 8.4 µM; ●, 12.3 µM;
- ◇, 17.3 µM; ▼, 19.1 µM; ■, 28.1 µM; △, 37.0 µM.
Figure A2.8 Copper accumulation in cells either untreated or incubated with EDTA and then washed; 6μM (circles) and 54μM (triangles) total copper concentration in the medium.

Filled symbols represent untreated samples; open ones indicate washed samples (incubated in 0.1M EDTA for 24 h and then washed).
Growth curves were similar to those shown in Figure 2.3A.
Figure A2.9 Copper accumulation by *Methylomicrobium albus* BG8 during sorption experiments.

Cells were inoculated into the medium at time 0.

**A.** Filled squares, 1.2 μM total copper in the medium; filled triangles, 12 μM total copper in the medium; open triangles, 43 μM total copper in the medium

**B.** Filled diamonds, 6μM total copper in the medium; open diamonds, 54 μM total copper in the medium.
Figure A2.10 Copper accumulated in cells by the end of growth (25-30 h) during sorption experiments versus total copper concentration and cupric ion concentration.

Growth media were inoculated from agar plates containing 10 μM copper.
Chapter Three

Cloning and Characterization of corA, a Gene Encoding a Copper-repressible Polypeptide Presumably Involved in Copper Uptake by *Methylomicrobium albus* BG8.

3.1. Introduction

The initial step in the oxidation of methane by methanotrophic bacteria involves methane monooxygenase (MMO). It has been found that the membrane-associated (particulate) form of MMO (pMMO) is a copper-containing enzyme with copper arranged in trinuclear clusters serving as the sites of the enzymatic activity (Chan *et al.* 1993). Copper is found widely in the environment (see Chapter One: 1.3.1.-1.3.3.). The total concentration of Cu in various environments is estimated to be on the order of $10^{-7}$ - $10^{-9}$ M (Hodgson *et al.* 1965; Boyle and Edmond 1975; Bruland *et al.* 1980). However, a significant fraction of copper is associated with various organic ligands present in sea water (Sunda and Hanson 1987; Coale and Bruland 1988; Coale and Bruland 1990; Moffett *et al.* 1990; Van den Berg *et al.* 1990, Bruland *et al.* 1991; Van den Berg and Donat 1992), fresh water (Hering and Morel 1988) and soils (Hodgson *et al.* 1965; Berggren 1989). Consequently, the reported concentrations of free copper are very low: $10^{-9.9}$ - $10^{-13.1}$ M for sea water (Coale and Bruland 1990), $10^{-11}$ - $10^{-18}$ M in fresh water (Apte *et al.* 1990, Van Den Berg *et al.* 1990), and $10^{-13}$ - $10^{-16}$ M for soils (Hodgson *et al.* 1965). As described in Chapter Two, our copper accumulation experiments with the type
I methanotroph *Methylomonas albus* BG8 have shown that copper availability to the cells is related to the cupric ion concentrations, not simply its total concentration (Berson and Lidstrom, 1996). Copper uptake by methanotrophs, therefore, is expected to involve a high-affinity copper transport system capable of overcoming low cupric ion concentrations in the environment.

Very little is presently known about the mechanisms by which copper enters cells in methanotrophs. Fitch and co-authors (1993) have indicated the existence of specific copper uptake and metabolism in the type II methanotroph, *M. trichosporium* OB3b. Our recent work (Berson and Lidstrom 1996) has established that a rapid saturation of copper uptake occurs in *M. albus* BG8 cells at 1-3x10^{17} moles of copper per cell despite a 100-fold variation in medium total copper and cupric ion concentrations. This suggests that cupric ion is taken up via a specific transport system.

Many transport mechanisms for uptake of trace metals are presently known (Hughes and Poole 1989; Silver and Walderhaug 1992; Brown *et al.* 1992b; Lee *et al.* 1990; Rogers *et al.* 1991; Cooksey, 1993). In some, multiple transporters for a single ion are found, in others, one transporter is able to transfer several different ions across the membrane (Silver and Walderhaug, 1992). Since copper uptake is central to methane oxidation by the pMMO, it is expected that methanotrophs might employ multiple systems. However, it is possible to obtain some understanding of a specific transport process by identifying and studying putative copper uptake proteins. Such proteins are expected to be copper-repressible, as at high levels of copper they would be needed in lower amounts, and under copper-limitation they should be overexpressed. However,
proteins involved in copper efflux might be copper inducible. In this chapter a screen for copper-regulated polypeptides was undertaken, and the cloning and characterization of corA, a gene encoding a copper-repressible protein are presented.

3.2. Materials and Methods

3.2.1. Growth of *M. albus* BG8.

*Methylocrobium albus* BG8 (Whittenbury et al. 1970), a type I methanotroph, was grown on nitrate mineral salts (NMS) medium (Whittenbury et al. 1981) in batch culture with shaking at 200 rpm. Filter sterilized copper nitrate and separately autoclaved phosphate buffer (Whittenbury et al. 1981) were added aseptically to autoclaved medium to avoid formation of precipitates. The media then was incubated for 24h with shaking before inoculation to avoid short-term toxic effects of the copper (Berson and Lidstrom 1996). The cells were grown to the late exponential phase at 30°C under a methane/air atmosphere (approximately 1:3 vol/vol) and harvested. Cells were grown at three total copper levels: under conditions of copper limitation (no copper added to the medium), in medium with optimal physiological levels of copper (10 μM and 20 μM), or in medium with high copper concentration, but still not toxic to *M. albus* BG8 under these growth conditions (40 μM). Mutants grew poorly in liquid cultures and, therefore, they were maintained on NMS plates. For chromosomal DNA isolation from mutants, they were grown on the plates for 2 to 3 weeks and harvested by resuspension in a small volume of NMS medium.
3.2.2. Isolation of a Copper-repressible Protein and Sequence Determination.

Membrane and cytoplasmic/periplasmic fractions of *M. albus* BG8 were prepared as follows. Cells were harvested by centrifugation at 5000g for 10 min. and then washed three times with 20 mM PIPES [piperazine-\(N,N'\)-bis(2-ethanesulfonic acid)] buffer (pH 7.0). Cells were then resuspended in the same buffer (approximately 3 g (wet weight) of bacteria was resuspended in 5 ml of PIPES). The cell suspension was passed three times through a French pressure cell at 137 Mpa. Remaining whole-cell debris was removed by centrifugation at 10,000g for 20 min. The supernatant from this centrifugation was recentrifuged at 150,000g for 1 h. The resulting supernatant consisted of periplasm plus cytoplasm (soluble fraction); the pellet contained outer and inner membranes (membrane fraction). The membrane fraction was resuspended 1:2 [wt/vol] in 20 mM PIPES buffer and homogenized. This suspension was diluted in a dissociation buffer 1:4 vol/vol, and incubated for 2 h at room temperature. The soluble fraction was boiled for 10 min. in a dissociation buffer solution added as 1:1 [vol/vol]. The dissociation buffer was prepared as follows: 250 mM Tris-HCl, 5% [wt/vol] SDS, 3.3% [vol/vol] β-mercaptoethanol, 20% [vol/vol] glycerol, 0.2% [wt/vol] bromophenol blue.

The SDS-PAGE screening of copper-repressible proteins was performed with an Hoefer Mighty Small™ Gel Electrophoresis Unit (Hoefer Scientific Instruments, San Francisco, Ca). Various concentrations of acrylamide/bis-acrylamide in base and stacker gels were tested, but the best resolution was obtained on the 14% (membranes) and 10% (soluble fraction) gels. The gels were cast as follows. The base gel contained 14% (10%)
[wt/vol] acrylamide, 0.37% (0.27%) [wt/vol] bis-acrylamide, 2M Tris-Base, pH 8.6, 20%
[wt/vol] SDS, 10% [wt/vol] ammonium persulfate, 5μl of TEMED per 10 ml of the
solution. The stacker gel consisted of 7.2% (5.1%) [wt/vol] acrylamide, 0.19% (0.14%)
[wt/vol] bis-acrylamide, 0.5M Tris-Base, pH 6.2, 20% [wt/vol] SDS, 10% [wt/vol]
ammonium persulfate, 5μl of TEMED per 5ml of the solution. The running buffer
contained 200mM glycine, 0.1% [wt/vol] SDS, and 20 mM Tris-Base. The gels were run
at a constant current of 20mA per gel until the tracking dye reached the bottom of the gel
(approximately 1h). Depending on sample concentration, 10 to 20 μL of membrane or
soluble fraction was loaded. The gels were stained with FastStain™ (Zoion Biotech,
Newton, Ma) to visualize polypeptide bands.

After the polypeptide bands of interest were identified in the course of the initial
screening, SDS-PAGE was carried out with the Xcell II gel system (Novex, Inc., San
Diego, Ca). Precast 1-mm-thick 8-16% polyacrylamide gradient Tris-glycine gels were
used for protein separation. The gels were prerun with the dissociation buffer to remove
any residual free radicals that commonly cause blocked amino termini of polypeptides.
Then, the gel was run again with samples prepared from cultures grown without copper
added to the medium and a control sample prepared from the cultures grown with 10μM
copper added. The electrophoresis was performed at 4°C to minimize any heat damage to
proteins and under a constant current of 40 mA. The running buffer contained 250 mM
glycine, 0.1% [wt/vol] SDS, 25 mM Tris, and 0.1 mM thioglycolate (added fresh just
before use). Polypeptide bands were blotted from the SDS-polyacrylamide gel onto an
Immobilon P membrane (Millipore Corp., Bedford, Ma) at 4°C. A Bio-Rad Trans-Blot
cell was used to transfer polypeptide bands overnight at 100 V in a transfer buffer. The transfer buffer consisted of 12mM Tris, 96mM glycine, 10% [vol/vol] methanol, and 0.5 mM dithiothreitol. The membranes were stained in Coomassie blue (0.2 [wt/vol] Commassie blue R-250 in 45% [vol/vol], 10% [vol/vol] acetic acid). The membranes were destained in 45% [vol/vol] and 10% [vol/vol] acetic acid. The membranes were rinsed several times in water to decrease the glycine and Tris concentrations. Appropriate bands were excised and sequenced by Edman degradation on ABI 476 pulsed liquid protein sequencer (Applied Biosystems, Inc., Foster City, Ca) by the Protein/Peptide Micro Analytical Facility at California Institute of Technology. One of these sequences was used to design an oligonucleotide probe using an eight-amino acid sequence fragment, Gly-Thr-Phe-Phe-Asp-Lys-Asn-Asn. The oligonucleotide probe OB1, 5’-GG(A, G, C, or T)-AC(A, G, C, or T)-TT(C or T)-TT-(C or T)-GA-(C or T)-AA(A or G)-AA(C or T)-AA-3’, with 512-fold redundancy, was synthesized at the Microchemical Facility at the California Institute of Technology.

3.2.3. DNA Purification.

Chromosomal DNA was isolated from *M. albus* BG8 by a slightly modified procedure of Satio and Miura (1963). The cells were grown to the late exponential phase, harvested by centrifugation at 7,500g for 10 min., washed once with saline-EDTA (0.15 M NaCl and 1 M EDTA, pH 8.0), and resuspended in 10 ml of saline-EDTA to which 4 mg per ml of lysozyme was added. The suspensions were placed in a -70°C freezer for 1h. 10 ml of Tris-SDS buffer (0.1 M Tris, 1% SDS, 0.1 M NaCl, pH 9.0) was
added to the frozen samples, and the tubes with the suspensions were swirled in a 50°C water bath until the solution became clear or nearly clear. The DNA was extracted with neutral phenol-saline-EDTA solution (phenol was saturated with 2 volumes of 0.1 M Tris [pH 9.0], 1 volume TE buffer [10 mM Tris-HCl, pH 7.4, 1 mM EDTA, pH 8.0] and then saturated with the saline-EDTA solution until pH 7.0). The extracted DNA was precipitated with isopropanol and washed in ethanol until no phenol smell could be detected. This DNA was resuspended in TE buffer.

Plasmid DNA isolation was carried out as follows. 3 ml of Luria-Bertani (LB) medium (1% [wt/vol] bacto-tryptone, 0.5% [wt/vol] bacto-yeast extract, 1%[wt/vol] NaCl) containing an appropriate antibiotic was incubated with a single colony clone overnight at 37°C with shaking. Cells were harvested and resuspended in 250 µl of TE buffer to which 4 mg per ml lysozyme was added. The cell suspension then were diluted with 50 µl of 10% [wt/vol] SDS solution and boiled at 100°C for 2 min. To precipitate proteins, 150 µl of 7.5 M ammonium acetate solution was added to the suspensions, they were incubated on ice for 10 min and then centrifuged at 16000g for 10 min. The supernatant containing DNA was collected, and DNA was precipitated with isopropanol, pelleted, and washed with ethanol. This plasmid DNA was resuspended in TE buffer.

All enzymes used were purchased from New England Biolabs, Inc. (Beverly, Ma). Restriction enzyme digestions were performed as described by Maniatis (et al., 1982). DNA fragments obtained from restriction enzyme digestions were separated by electrophoresis (150 V, approximately 1h) through 1% agarose gels in a Tris-borate buffer (0.089 M Tris-borate; 0.089M boric acid; 0.002M EDTA). A 1Kb DNA Ladder
(Boehringer, Mannheim, Germany) was used for sizing both chromosomal and plasmid DNA fragments. The gels were stained by adding the fluorescent dye ethidium bromide (0.5 μg/ml) to the buffer. The gels intended for hybridizations with a nucleotide probe were denatured in a solution of 150 mM NaCl and 0.5 N NaOH for 30 min. on a shaker at room temperature. The denaturing solution was then replaced with a neutralizing solution, 0.5MTris-HCl (pH 8.0), and the gel was shaken for 30 min. at room temperature. Finally, the gel was dried at 80°C in a vacuum gel dryer (Bio-Rad, Hercules, CA).

3.2.4. Cloning and Sequencing of corA.

The oligonucleotide probe OB1 was end labeled with 5'-[γ^{32}P]ATP as described below. 20 pmoles (1μl) of the synthesized oligo was resuspended in 38 μl ddH₂O and put into a boiling water bath for 10 min. 10x phosphorylation buffer (5 μl), 20 units of T4 polynucleotide kinase (1μl), and 150 μCi of 5'-[γ^{32}P]ATP were mixed with the denatured oligo and incubated at 37°C for 45 min. To stop the reaction, 50 μl of 0.1 M EDTA was added to the mix after the incubation. 10x phosphorylation buffer is composed of 0.5M Tris-HCl, 0.1M MgCl₂, 50mM dithiothreitol, 1mM spermidine, 1mM EDTA.

The labeled oligo OB1 was denatured and hybridized to the dried gels overnight at 39°C with shaking. The temperature of the hybridization with OB1 was chosen based on the following formula (Maniatis et al. 1982): \( T_{hybr} = T_m - 10^\circ C = [69.3 + 0.441(GC\%) - 650/L] - 10^\circ C \), where \( T_{hybr} \) temperature of hybridization; \( T_m \), oligo melting temperature; GC%, the percent of G and C in the oligo; L, number of base pairs in the oligo. The
hybridization solution consisted of 1M sodium chloride, 0.1M sodium citrate, 0.56 % [wt/vol] SDS, 200 μl 0.5M EDTA (pH 8); 0.5% [wt/vol] dry powdered milk. After hybridization the gels were washed 3 times with 75 mM sodium chloride, 7.5 mM sodium citrate, and 0.1% SDS to remove any residual non-specifically bound radioactivity. Air dried gels were then exposed to X-ray film. To increase the sensitivity of the film, intensifying screens were always used, and the exposure was carried out at -75°C (Maniatis et al. 1982).

DNA fragments excised from agarose gels were purified using GeneClean II® kit (Bio 101 Inc., La Jolla, Ca). Transformations into competent Escherichia coli DH5α™ (Life Technologies, Gaithersburg, MD) were carried out following the protocol provided by the manufacturer. The transformants were grown on LB agar plates to which 2.5% [wt/vol] X-gal (5-bromo-4-chloro-3-indoly1-β-D-galactoside), 2.5% [wt/vol] IPTG (Isopropyl-β-D-thiogalactopyranoside), and 100 mg/ml [wt/vol] ampicillin (Amp) were added (Maniatis et al. 1982). White colonies (carrying recombinant plasmids) were transferred onto ME25 82 mm nitrocellulose filters (Schleicher & Schuell, Los Angeles, CA) placed on LB agar plates with Amp and onto master agar plates with Amp. All plates were grown overnight at 37 °C. The next day DNA from the colonies grown on filters was liberated, bound onto the filters and hybridized at 39°C with OB1 probe using the procedures described by Maniatis (et al. 1982).

Plasmid DNA for sequencing was isolated using a Qiagen-tip 20 kit (Qiagen, Los Angeles, CA ). DNA sequencing was performed with an Applied Biosystems automated sequencer at the California Institute of Technology, Pasadena, Sequencing Facility, for
both strands. Initial sequencing was performed with M13 Forward and M13 Reverse sequencing primers (New England Biolabs, Inc., Beverly, Ma). Synthetic oligodeoxynucleotide primers complementary or identical to previously sequenced DNA fragments were synthesized at the Microchemical Facility at the California Institute of Technology, and they were used for later sequencing. The DNA sequence translation and analyses of DNA and polypeptide sequences were performed by using Genetic Computer Group (GCG), University of Wisconsin programs.

3.2.5. Construction of an Insertion Mutation in corA

An insertion mutant in corA was constructed by homologous recombination as described elsewhere (Ruvkun and Ausubel 1981; Chistoserdova and Lidstrom 1994). A 1.4-kb DNA fragment containing a kanamycin resistance gene (Km\(^r\)) was excised from plasmid pUC4K (Pharmacia, Uppsala, Sweden) using HincII sites to produce blunt ends and purified. The plasmid containing the targeted corA gene was digested with BstEII. BstEII can not be inactivated by heat, and, therefore, the plasmid DNA was precipitated from the restriction reaction by adding 10 mM EDTA, 0.4 volumes of 5M ammonium acetate, and 2 volumes of isopropanol as recommended by Maniatis (et al. 1982). The BstEII 5\(^\prime\)-protruding ends were made flush with Klenow fragment and deoxynucleoside triphosphates (dNTPs). The Kan\(^r\) gene was ligated into BstEII sites of the plasmid, and the construction was used to transform competent E.coli DH5\(\alpha\)™ cells as described by Maniatis (et al. 1982). The resulting construction (pOB17) was transformed into E.coli S17-1 (Simon et al. 1983) and the transformants were plated on LB plates with Amp,
Km, and tetracyclin (Tc). The disrupted corA was introduced into the M. albus BG8 chromosome by conjugation with an E. coli strain containing pOB17 and selection on NMS agar plates containing Km. The biparental matings were carried out as follows. Two full loops of M. albus BG8 cells, freshly grown on plates, and one full loop of E. coli containing pOB17 were resuspended into 1 ml of NMS each, and 600 μl of M. albus suspension was mixed with 60 μl of E. coli suspension. The mixture was centrifuged, resuspended in 50 μl of NMS, and placed on a NMS agar plate (10μM total copper concentration) as a thin spot about 1 inch in diameter. The plates were incubated either under methane/air atmosphere, or with 0.1% [vol/vol] methanol for 2 days at 30°C. The biomass was washed off each plate with 2 ml of NMS. The volume of the suspensions was brought down to 100 μl by centrifugation and resuspending in 100μl of NMS, and they were spread on NMS plates to which Km was added. The plates then were grown either with methane or methanol at 30°C for one to three weeks.

3.2.6. Analysis of Insertion Mutation in corA.

The corA insertion mutants were streaked on NMS plates containing Amp to determine their phenotype and to distinguish between single and double crossover events. One mutant (OB12.7), that seemed to be a result of a double crossover, was grown on an NMS plate with Km (it was not possible to grow the mutant in liquid culture), characterized by PCR amplification with primers PR12-F6 (5’-ATG-TAT-CCC-TGC-ATG-GCA-CTG-3’) and PR12-R1 (5’-TTA-CGG-GAT-ACT-GAC-TTC-TAC-3’). The primer PR12-F6 was chosen from a region located 110 bp downstream from the start of
corA; the primer PR12-R1 was selected from the sequence located at the very end of corA. The PCR products obtained from OB12.7 and from wild-type M. albus (w.t.) with the above primers were compared. The PCR reactions were carried out in 100 µl volumes under the following conditions: approximately 20ng template DNA, PCR buffer (10 mM Tris-HCl, 50 mM KCl), 1 mM MgCl₂, 0.2 mM of each dNTP, 1U of Taq polymerase (Boehringer, Germany), 100 pmol of each primer, and 5% [vol/vol] DMSO. A Hybaid thermal cycler (Combi TR-2) was used for 30-cycle amplification with following reaction conditions: denaturation, 94°C for 1 min.; annealing, 55°C or 50°C for 1 min.; polymerization, 72°C for 2 min.

To analyze the growth response of the OB12.7 mutant to copper concentration in the medium, the mutant and the w.t. were grown at three levels of total copper concentration in NMS medium: copper-limited conditions (no copper added to NMS), physiological level of copper (10 µM copper added to NMS), and high, but still allowing the growth of w.t. copper concentration (50 µM of copper added). Each experiment was performed in triplicate. After 24-h preequilibration of the medium (10 ml in each flask), the flasks were inoculated with w.t. or the mutant strain. At different times, the optical density of the growing cells was measured with a Klett - Summerson (Long Island City, NY) photoelectric colorimeter and was reported as Klett units. Optical density was calibrated to viable cell numbers as was described in 2.2.4.
3.3. Results and Discussion

3.3.1. Purification and Sequencing of the 29-kDa Polypeptide.

*M. albus* BG8 cells were screened for the presence of copper-regulated polypeptides, under the assumption that these should be involved in copper metabolism and/or transport. The screening was carried out as follows. Cells were grown at 4 copper levels, copper limitation (no copper added to NMS medium), 10 μM, 20 μM, and 40 μM total copper added. These cultures were harvested during the late exponential phase of growth and fractionated to produce soluble (cytoplasm and periplasm) and membrane (outer and inner membranes) fractions. These samples were screened by SDS gel electrophoresis for the presence of polypeptides found at different relative levels in the cells grown under copper limitation than in those grown with copper added to the medium. Three copper repressible (Cu\textsubscript{1}\textsuperscript{rep}, Cu\textsubscript{2}\textsuperscript{rep}, Cu\textsubscript{3}\textsuperscript{rep} found at higher levels in copper-limited cells), and one copper inducible (Cu\textsubscript{1}\textsuperscript{ind}, found at lower levels in copper-limited cells) polypeptides were identified in both membrane (Figures 3.1a, A3.1a-A3.3a) and soluble (Figures 3.1b, A3.1b-A3.3b) fractions. Cu\textsubscript{1}\textsuperscript{rep} was a strong band, while the others were minor. An attempt was made to transfer all of these polypeptides onto Immobilon P membranes for sequencing, by optimizing conditions of blotting for each protein band. To increase protein elution efficiency, gel composition was varied (both gradient and single gel concentrations were used), as well as several transfer buffers, transfer times and power settings. However, only the Cu\textsubscript{1}\textsuperscript{rep} (membranes) and Cu\textsubscript{3}\textsuperscript{rep} band (soluble fraction), found in samples prepared from cells grown under copper-limitation, were successfully
blotted onto membrane and sequenced. It is possible that the other polypeptides did not elute well onto membranes because they are present at lower levels. In some cases poor growth was obtained with cells grown with added copper (Figure 3.1a), but overexpression of CorA always correlated with lack of copper, not with poor growth, which implied that it is copper-repressible rather than a general starvation polypeptide. Therefore, Cu$_1^{rep}$ was termed CorA, for Copper-repressible polypeptide A. The following N-terminal sequence was obtained for CorA: Ala-Thr-Ala-Ile-Ser-Gly-Thr-Phe-Phe-Asp-Lys-Asn-Thr-Ser-Ala-Asp-Met-Thr-Val-Arg-Ala-Tyr-Ser-(Ser)-Tyr-Asn-Leu-Ser-(Ser). The sequence was not significantly homologous to any protein sequences in SwissProt database.

Cu$_3^{rep}$ was not consistently expressed under copper limitation and, therefore, copper effect on the expression of this polypeptide was questioned. Indeed, its N-terminal sequence, Met-Lys-Lys-Ile-Leu-Asp-Val-Val-Lys-Pro-Gly-Val-Val-Thr-Gly-Glu-Asp-Val-Gln, showed high homology (76.5% to 84.2% in 19 aa overlap) to N-termini of fructose bisphosphate aldolases (Figure 3.2). Fructose bisphosphate aldolases are not involved directly or indirectly in metal uptake, consequently, the cloning of the gene, corresponding to polypeptide Cu$_3^{rep}$, was not undertaken.

**3.3.2. Cloning and Sequencing of corA**

Chromosomal DNA was isolated from *M. albus* BG8, digested with *SacI, PstI, Sphi, SalI, XbaI*, and *BamHI*, separated on 1% agarose-Tris-borate gel and hybridized with the oligonucleotide probe OB1 (Figure 3.3). Bands of 6, 4, 6, 4.2, 0.5, and 8 kb,
respectively were identified. The probe was based on the Gly-Thr-Phe-Phe-Asp-Lys-Asn-Asn fragment of the N-terminal sequence (see 3.2.2.). A 4.2-kb SalI fragment was chosen for cloning. A fraction of SalI-digested M. albus BG8 chromosomal DNA containing DNA fragments sized from 3.5 to 5 kb was excised from an agarose gel and purified. The purified DNA was then ligated with SalI-digested pUC19 (New England Biolabs, Inc., Beverly, MA) and used to transform E. coli.

Plasmid DNA was isolated from the colonies that hybridized with the OB1 probe and screened for insert size. All of these clones appeared to have the same SalI insert of approximately 4.2 kb in either the forward or reverse orientation. Plasmid DNA from one of the clones with the forward insert (pOB10a) and one with the reverse insert (pOB10b) was isolated for further subcloning (Figure 3.4). Six clones per each insert, randomly chosen, were examined for the size and the orientation of the inserts. Three clones containing the 0.9-kb SalI-SphI fragment (pOB11) and three clones containing the 3.3-kb SalI-SphI fragment (pOB12) were identified (Figure 3.4). One of each of these clones was used for sequencing. The subclones pOB11 and pOB12 were then sequenced on both strands and the combined sequence is presented in Figure 3.5.

3.3.3. Analysis of the Sequence Data.

The sequence (Figure 3.5) was analyzed by identifying start and stop codons and by using Testcode program from the GCG Package (Figure 3.6). The Testcode program analyzes local nonrandomness at every third base in sequence in a frame-independent way, and it produces a plot that allows gene localization. Both methods revealed three
open reading frames (orf2, orf3 (corA), orf4), two partial open reading frames at the ends of the insert (orf1 and orf5), and 984-bp, 205-bp, and 247-bp regions without a distinguishable area of polypeptide coding (Figure 3.4).

The 695-bp fragment, orf3, was recognized as corA, since it contained a region coding for the N-terminal sequence of CorA. Based on the DNA sequence and its translation, it appears that CorA contains 695 residues, and it has a molecular mass of 21,860 Da. The discrepancy between the mass, estimated based on the amino acid sequence (21,860 Da), and the approximate mass of the polypeptide, predicted from the SDS-PAGE screening (28,500 Da), is probably because addition of the dye to polypeptides for SDS-PAGE electrophoresis sometimes causes them to migrate differently than their true molecular weights. Therefore, the mass of 21,860 Da was presumed to be the true mass of CorA. CorA was found in the membrane fraction of the cells (Figure 3.1) and this finding is supported by the presence of a leader sequence of 30 residues (Figure 3.5) and by hydropathy plots that indicate that CorA should be a membrane polypeptide (Figure 3.7). Eight possible membrane-spanning regions were identified in CorA by using the residue specific hydrophobicity index of Kyte and Doolittle (Figure 3.7). A search of both DNA and protein data bases revealed some homology of CorA (17.5% identity in 126 aa overlap) to rabbit and human calcium release channel protein (Takeshima et al. 1989) (Figure 3.8). No specific copper-binding domains were found in CorA. It is possible, however, that CorA is a metal porin (which can be involved in the transport of several metals, including copper), and it is induced
under conditions of a nutrient metal starvation. To provide more direct information about the function of CorA, mutants defective in corA were generated (see below).

The sequences of the other open reading frames (Figure 3.4) and their corresponding polypeptides were compared with DNA and protein data banks (SwissProtein Bank, GenBank). Only one of these, the 221-amino-acid-long polypeptide encoded by orf1 (665 bp), demonstrated high similarity to a known protein. A 29-kDa extragenic suppressor protein (268 amino acids) that is involved in facilitating the function of heat shock proteins (Yano et al. 1990) had 50.7% identity in 221 aa with the product of orf1 translation (Figure 3.9). No significant amino acid sequence similarity was found between the products of orf2, orf4, orf5 and any known proteins (SwissProtein Bank, GenBank). Hydropathy analysis (Figure 3.10) suggests that all open reading frames in the insert encode membrane polypeptides, however a recognizable leader sequence was found only in orf2 (22 residues). The second open reading frame, orf2, of 539 bases is located downstream of orf1, and it encodes a polypeptide of 180 amino acids. The fourth, 455-bp, open reading frame, orf4, encodes a polypeptide of 152 aa. The fifth, partial, open reading frame, orf5, (329 bp) codes for 113 residues of the N-terminus of a polypeptide. The open reading frames orf1 and orf5 are transcribed in the same direction as corA, while orf2 and orf4 are transcribed in the opposite direction.

3.3.4. Mutant Characterization

To obtain information on the function of corA, an insertion mutant defective in the gene was constructed. Mutants were generated by homologous recombination
between *M. albus* BG8 chromosome and the mutated gene in a plasmid. A 3.3-kb *SalI*-SphI fragment containing the corA gene (695 bp) was cloned into plasmid pUC19 to produce pOB12. The unique BstEII site is located 463 bp upstream from the end of corA and 780 bp upstream from the end of the 3.3-kb insert in pOB12 (Figure 3.4). The BstEII site was used to introduce Km\(^\text{r}\) cassette into pOB12. The plasmid, carrying the Km\(^\text{r}\) gene transcribed in the same direction as corA (pOB16), was identified by plating the transformants on LB plates containing both Amp and Km, and by the restriction analysis of these clones with SphI and XhoI. The EcoRI site was used to ligate pOB16 with the suicide vector pAYC61 (Chistoserdov *et al.* 1994). The size of the constructed plasmid (pOB17) in selected clones was checked by digestion with EcoRI (Figure 3.11). The phenotype of these mutants was checked by growing them on NMS plates containing Amp or Kan. The double crossovers should be Km\(^{\text{r}}\)/Amp\(^{\text{s}}\), whereas single crossovers are expected to be Km\(^{\text{r}}\)/Amp\(^{\text{r}}\), since they carry an insertion of the entire plasmid (Figure 3.12). DNA from one mutant (OB12.7) with a Kan\(^{\text{r}}\)/Amp\(^{\text{s}}\) phenotype was isolated and analyzed by PCR amplification with primers PR12-F6 and PR12-R1 located at the ends of the gene. The PCR product amplified from OB12.7 mutant was about 1.4 kb longer than those generated from wild type (Figure 3.13). These data suggested that this mutant contained the disrupted corA in the chromosome and was the result of the desired double crossover recombination event. This mutant grew very poorly on agar plates.

An attempt was made to determine the relationship between copper concentrations in the medium and growth of the OB12.7 mutant. The mutant and w.t. were inoculated into preequilibrated NMS medium with no copper added, 10 \(\mu\text{M}\), or 50
μM copper added. While w.t. cells grew best at 10μM and grew with longer lag phases at
no copper added and 50 μM copper added, the mutant did not grow at all under these
conditions, after 6 days of incubation. This result implies that corA is vital for growth of
M. albus BG8.

3.4. Conclusions

Circumstantial evidence suggests that CorA might be involved in copper uptake.
The polypeptide was isolated as a copper-repressible polypeptide, which implies a
regulatory effect of copper on CorA. The product of translation of corA shows similarity
with calcium release channel proteins, which suggests that CorA might belong to a family
of divalent metal porins induced under conditions of metal starvation. It is possible that
such a porin is not specific for copper, but rather works with a range of divalent metals.
The mutant analysis indicates the importance of corA for normal physiology of M. albus
BG8. However, the inability to characterize copper uptake in the CorA mutant due to it’s
poor growth makes it impossible to draw a definite conclusion about the role of CorA in
copper transport/metabolism.
3.5. References


Figure 3.1 Identification of copper-repressible and copper-inducible polypeptides in particulate (a) and soluble (b) fractions of *Methylomicrobium albus* BG8 by sodium dodecyl sulfate-polyacrylamide [14% (a) and 10% (b)] gel electrophoresis. Cell extracts of *M. albus* BG8 were grown with 10 μM (lane 1); 20 μM (lanes 2, 3, 5), 40 μM (lane 4), and no copper (lanes 6-9) added to the medium. Samples in lanes 2-5 were prepared from poorly growing cells; samples in lanes 1, 6-9 were prepared from cultures with normal growth. Two copper-repressible polypeptides were identified in the particulate fractions: Cu1rep (~28.5 kDa) and Cu2rep (~33 kDa). A copper-repressible polypeptide Cu3rep (~49.5 kDa) and a copper-inducible polypeptide Cu1ind (~49 kDa) were identified in the soluble fraction.
A.

\[
\begin{array}{ccc}
\text{Cu}_3\text{rep} & \text{MkkILDVVKPGVVTGEVDVQ} \\
10 & 19 \\
\text{Alf}\_\text{Ha} & \text{MAKLLDVKPGVVTGEDVQKVFAAY} \\
10 & 20 \\
\end{array}
\]

84.2% homology in 19 aa overlap

B.

\[
\begin{array}{ccc}
\text{Nu}_3\text{rep} & \text{MkkILDVVKPGVVTGEVDVQ} \\
10 & 19 \\
\text{Alf}\_\text{Ec} & \text{MSKIFDFVKPGVITGDVQKVFOQVAKENF} \\
10 & 20 & 30 & 40 & 50 \\
\end{array}
\]

76.5% homology in 17 aa overlap

**Figure 3.2.** Comparison of the N-terminal amino acid sequence of Cu$_3^\text{rep}$ with the N-terminal part of fructose bisphosphate aldolases.

A. Alf\_Ha, fructose bisphosphate aldolase from *Haemophilus influenzae*; SwissProt Data Base Accession number P44429.

B. Alf\_Ec, fructose bisphosphate aldolase from *Escherichia coli*; SwissProt Data Base Accession number P11604.
Figure 3.3 Hybridization of oligonucleotide probe OB1 to restriction enzyme digests of *Methylocrobium albus* BG8.

Lanes: 1, molecular weight standard (1kb ladder, not labeled); 2, *SacI* digest; 3, *PstI* digest; 4, *SphI* digest; 5, *SalI* digest; 6, *XbaI* digest; 7, *BamHI* digest; 8, molecular weight standard (32P-labeled λ DNA/ *HindIII* fragments). A 4.2-kb band of *SalI*-digested chromosome is circled.
Figure 3.4 Physical map of the 4.2-kb *Methylomicrobium albus* BG8 chromosomal region containing *orf1*, *orf2*, *orf3* (*corA*), *orf4*, *orf5*. Plasmids containing the whole region (pOB10a and pOB10b) or its fragments (pOB11 and pOB12) are shown below the map. The directions of the orfs transcription are indicated by the arrows above the orfs, and the direction of cloning with respect to direction of *corA* transcription is shown by the arrows on the inserts. The asterisk shows the beginning of the sequence hybridizing with OB1 probe.
Figure 3.5 Nucleotide sequence and deduced amino acid sequence of the 4.2-kb *M. albus* BG8 chromosome region containing *orf2, orf3* (*corA*), *orf4*, and partially *orf1, orf5*.

Asterisks indicate stop codons. Putative Shine-Dalgarno sequences are shown in bold, putative leader sequences are underlined. The amino acids that were determined by sequencing of the blotted protein band are shown in bold italic.
Figure 3.5 (continued)
TCGGCTGTACAAlAACGAGTTTATTCGCTGAAATCCGAGCATCTGAACTCCTGCTGCC
1141 +------------------------------------------+++++++
AAGCGCAATGTTTTTCTTTTTCACAATAGGCGAGTTTAGGCCTCGTAGACTTGGCAGGAC
N P Q V F S F N I G D F G S C R F R G T
ATGTTCCCCCTACGTTGAGCCCGCGGGCCGGAAAAGGAAAAGACAGCAAAAGGATCGGTA
1201 +------------------------------------------+++++++
TACAAGGGCACTCACAAGGCGGGGGCGCCCTTCTTTTTCTCTCTCTCTCTCTCTCTTAGGAT
W T G T L N V A G S F S F L L L I P
AAACCATAAAGGCAGCTGATTATTCATCAACTTTCTTCCCTAAATCATGCTCAACTTCTTTTG
1261 +------------------------------------------+++++++
TTTGTATTTCCGCTACATAAAGTATTTGAGGAGGAGTTTTGTTAGTAGCAGATGGAGAAC
L V M F R L S K M ≤ orf2

1321 TCAGAGTCCGCCATATATATCTCGTGCGGGTTTGGGAGTCAACTGATAGCGAC
1381 AAGCGCAATCAGCGGTGCTGGTCTGGGAAAGGCCGCTGCTGGGCGCGGGCGCCTCAAC
1441 GCCCGCGCGCTTCATTGATCCGCTCAGGCTCGTCCGCCAACAGCCCGCGCTGCC
1501 CAGTGATTTCCAACCAGTTCAGGTAAGGCTAGCGATTAAATAGGAGTTAGGA
1561 AAGCGCAATTCGCTGTGAAAGCCCTCTATATATAAACCCCCCTGCCCCCCTTTTTCAAG
1621 GGGGAGGCGCGCATTTTCCTTTTTATAGCAGTATGCAAATAAACCAGGGTTATTTTCTCAGGGAG
1681 CTTGTGTGAATTATACCGTTACATGATCCCAGGTCTACGTTGCCAAGGATAAATAA
1741 TGCATGGGTGATCTTATAAGGAAAGAACCCGGAAGACTGTTGACGGGTGAGGGGCTTA
1801 GCCGCTGCTGCTATTCATTTCTCAACAGGAAAGCAGACATCTCCTCTCTTCGGTAAAAAGGGGGA
1861 TTTGCTCTGCAAAACAGTGGATGAGCAACGCTGTTACAACGCTCAAATCGGATTAAACAG
1921 TGCCGATGGTTAAAAACCCTGCCTGGCGGATATTCACCATAGCGCGATTGCCTGGCAAA
1981 CTCGATTTGAAATTCTGCCGAGCTCTGGGAGTCCGGCTCGCTGCTCGCTGCTGCTGGA
2041 AGCCTTTCAACACACAGAAATCTGGAATGACACACTTTTAAATATCCCTATATATGTTTAT
2101 ATGTTGTAACAAAAATTATAATATAGGCTATAATATCGCTATAATGATTATTCCCTCAGAT
2161 GACATGTTTATAGGTGTACTTGGCTCTTGTGGAATTTCTCCTGGATACGGCGACAGCG
2221 CATCGGGGAGCTACGCGATAACGTTAAAAATTTTTATTTTTTTAAGGATACCCCATGAAAATACAC

orf3 (corA) ⇒ M K Y N

2281 AATAAATTTAAAAAGTTTGGCATTGATTTCGATGAGCTGGCTGCTGCTGCTGCTGCTGGCACC
N K F K S L A L I S V A A G L F F A N P
2341 CTCCAAGTGCCTACAGCTATATCTCGCCACTTTTTTTTGACAAAAAACATACGTACGGGAC
L Q A A T A I S G T F F D K N N T S A D
2401 ATGACCTGTCCGCCCTATTCTCTCGGTACAACCCTAGTATGGGCTATCTGGGCTGGAAGCAT
M T V R A Y S W Y N L S M G Y L G W T H

Figure 3.5 (continued)
2461 CACTGGAATCGGGGATTTGCTCAAGCTGAAAGAAAGGCAAGCCCGTGCACCATTGCGCTGACT
Figure 3.5 (continued)
Figure 3.5 (continued)
Figure 3.5 (continued)
Figure 3.6 Frame-independent gene localization using the TestCode Program from the GCG Package.
The plot produced by the program is shown. X axis shows the number of residues in a sequence. The top horizontal line shows bias in the composition of bases found in the third positions of the codons. The lower horizontal line is the codon preference curve. Larger values of the testcode statistic (above the top horizontal line) correspond to regions with a greater likelihood of encoding a protein. The predicted open reading frames are indicated by the bars.
Figure 3.7 Predicted structure of corA gene product.

The hydrophobicity of CorA was predicted by the Kyte-Doolittle algorithm. Thick bars show potential membrane-spanning domains. Below is a plot which indicates probability of finding an amino acid residue on the surface of the protein molecule. The third panel shows flexibility of the peptide backbone. Next is antigenic index, surface probability and chain flexibility combined. The probability of the peptide chain to form secondary structures (α-helixes, β-sheets, β-turns) are shown next.
Figure 3.8 Comparison of the deduced amino acid sequence of CorA with the sequence of calcium release protein (SwissProt Data Base Accession number P05838).

Identical residues are indicated by vertical lines, conserved substitutions are shown by dots.
**Figure 3.9** Sequence comparison of predicted amino acid sequence of orf1 and suhB, extragenic suppressor protein (SwissProt Data Base Accession number P22783).

Identical residues are indicated by vertical lines, conserved substitutions are shown by dots.
**Figure 3.10** Predicted hydropathy plots of orf1, orf2, orf4 and orf5 gene products by the Kyte-Doolittle algorithm. Probable membrane-spanning domains are indicated with bars. Scales above the plots show the number of residues in a sequence.
Figure 3.11 Construction of plasmids pOB16 and pOB17 carrying the mutated corA gene.

First, the 3.3-kb *M. albus* BG8 DNA fragment containing corA was inserted into pUC19, then, corA was disrupted by the insertion of a kanamycin resistance gene (Km') to produce pOB16. pOB16 was ligated with the suicide vector pAYC61 to yield pOB17.
Figure 3.12 The phenotypes of wild-type (w.t.), single and double crossover corA mutants of *M. albus* BG8.

p, a promoter; corA, gene encoding copper-repressible polypeptide; corA^{dt}, corA deleted by an insertion of kanamycin resistance cassette (Km).
Figure 3.13 Characterization of OB12.7 mutant.

PCR-amplified products were separated on an agarose gel. Lanes 1 and 8, 1kb DNA Ladder DNA size standard. The following PCR products were loaded: lanes 2, 3, 5, 6, PCR with wild type *M. albus* BG8; lanes 4 and 7, PCR with OB12.7. Lanes 2-3, annealing at 50°C; lanes 5-8, annealing at 55°C. Samples of w.t. DNA were prepared from two batches: 2 and 5, cells grown under copper limitation, 3 and 6, cells grown with sufficient copper.
Appendix

To optimize separation of the polypeptide bands by electrophoresis several polyacrylamide concentrations in the gels were tested. In this appendix the results of sodium dodecyl sulfate-polyacrylamide gel electrophoresis with 10% (membranes) and 14% (soluble fraction) (Figure A3.1), 12% (Figure A3-2), and 16% (Figure A3-3) polyacrylamide in the base gels are shown. However, the best separation of the membrane bands was achieved with 14% polyacrylamide in the base gel, while the soluble fraction was better separated on a 10% polyacrylamide gel (Figure 3.1).
Figure A3.1 Identification of copper-repressible and copper-inducible polypeptides in particulate (a) and soluble (b) fractions of *Methylomicrobium albus* BG8 by sodium dodecyl sulfate-polyacrylamide [10% (a) and 14% (b)] gel electrophoresis.

Cell extracts of *M. albus* BG8 were grown with 10 μM (lane 1); 20 μM (lanes 2, 3, 5), 40 μM (lane 4), and no copper (6-9) added to the medium. Lanes 2-5, cells grew poorly; lanes 1, 6-9, normal growth of cultures. A copper-repressible polypeptide Cu$_3^{\text{rep}}$ (∼49.5 kDa) and a copper-inducible polypeptide Cu$_1^{\text{ind}}$ (∼49 kDa) were identified in the soluble fraction.
Figure A3.2 Identification of a copper-repressible polypeptides in particulate (a) and soluble (b) fractions of *Methylomicrobium albus* BG8 by sodium dodecyl sulfate-polyacrylamide [12%] gel electrophoresis.

Cell extracts of *M. albus* BG8 were grown with 20 μM (lanes 1, 6, 9); 10 μM (lane 5), and no copper (lanes 2-4, 7, 8) copper added to the medium. Samples in lanes 3 and 8 were grown in medium with 20 μM and 40 μM EDTA added. Samples in lanes 1 and 6 were from poorly growing cultures, all other samples were from normally growing cultures. A copper-repressible, Cu_{1}^{rep} (~28.5 kDa), polypeptide was identified in the particulate fraction.
Figure A3.3 Identification of copper-repressible and copper-inducible polypeptides in particulate (a) and soluble (b) fractions of *Methylo bacterium albus* BG8 by sodium dodecyl sulfate-polyacrylamide [16%] gel electrophoresis.

Cell extracts of *M. albus* BG8 were grown with 10 μM (lane 1); 20 μM (lanes 2, 3, 5), 40 μM (lane 4), and no copper (6-9) added to the medium. Lanes 2-5, cells grew poorly; lanes 1, 6-9, normal growth of cultures. A copper-repressible, Cu$_{1}^{\text{rep}}$ (~28.5 kDa), polypeptide was identified in the particulate fraction. A copper-inducible polypeptide, Cu$_{1}^{\text{ind}}$ (~49 kDa), was identified in the soluble fraction.
Chapter Four

Cloning and Sequencing of a Putative Copper Transporting P-type ATPase.

4.1. Introduction

Copper is an essential nutrient due to its role as a cofactor of many oxygenases and electron-transport proteins. Yet in excess, copper can cause extensive cellular damage. Consequently, the regulation of intracellular copper activity is required in all cells. Two mechanisms for the homeostasis of copper have been studied in bacteria: copper resistance/uptake in Escherichia coli and in Pseudomonas syringae. In E. coli four plasmid-encoded genes (pcoARBC) are required for copper resistance and seven chromosomal genes (cutA-cutF) are involved in copper metabolism. In P. syringae four structural plasmid-encoded genes (copABCD) constitute the copper-resistance system. Both copper resistance systems are regulated by a pair of regulatory genes (pcoSR (E. coli) and copCR (P. syringae)) (Butt et al. 1984; Ecker et al. 1986; Mellano 1988; Harwood-Sears and Gordon 1990; Lee et al. 1990; Cha and Cooksey 1991; Rogers et al. 1991; Cooksey and Azad 1992; Silver and Walderhaug 1992; Cha and Cooksey 1993; Cervantes and Gutierrez-Corona 1994; Gupta et al. 1995, etc.). However, recent studies have identified P-type ATPases as a new class of copper transporting proteins involved in the control of intracellular copper both in prokaryotic and eukaryotic cells (Odermatt et
al. 1993; Silver et al. 1993; Tanzi et al. 1993; Kanamaru et al. 1994; Rad et al. 1994; Vulpe et al. 1994; Fu et al. 1995; Ge et al. 1995). All P-type ATPases are involved in translocation of a specific cation ($H^+$, $Na^+$, $K^+$, $Ca^{2+}$, $Mg^{2+}$, $Cd^{2+}$ or $Cu^{2+}$), including uptake (movement into the cell), e.g., $K^+$-ATPase (Hesse et al. 1984), efflux (movement out of the cell), e.g., $Ca^{2+}$-ATPase (Brandl et al. 1986), and cation exchange (one moving in, one moving out), e.g. $Na^+$/K$^+$-ATPase (Shull et al. 1985). All known P-type ATPases, from bacteria to human, contain several highly conserved functional domains, such as a phosphatase motif, TGES (Figure 4.1A); a conserved proline in the transduction region, XPX (Figure 4.1B); a phosphorylation region, DKTGT(L or I)T (Figure 4.1C); and an ATP-binding domain, GDG(I or V)ND(A or S)P(A or S)L (Figure 4.1D). It is postulated that the conserved aspartate residue in the ATP-binding domain is phosphorylated by ATP during an intermediate step of cation transport, then, the phosphatase domain removes the phosphate from the aspartic acid during the reaction cycle (see Chapter One for details). The transduction domain is thought to transfer intracellular protein-bound cation to the membrane channel region (Silver and Ji 1994).

There are several unique features found only in copper and cadmium transporting ATPases that set them apart from the rest of the P-type ATPases. One is the cysteine residues flanking the conserved proline in the transduction domain (except for one of the two copper transporting ATPases in Enterococcus hirae, CopB, that has only one cysteine next to the proline) and the presence of a second proline six to eight positions downstream (Figure 4B). Another unique feature is a putative metal binding motif, GMXCXXC. This motif is repeated six times in the copper efflux ATPase that is thought
to be defective in the copper-related Menkes disease, Mc1 (Vulpe et al. 1993), and it is repeated five times in WD, a copper-ATPase associated with Wilson’s disease of copper deficiency (Tanzi et al. 1993). The motif is present in CopA, the copper uptake ATPase of Enterococcus hirae (Odermatt et al. 1993), and CadA, the cadmium-transporting ATPase of Staphylococcus aureus (Nucifora et al. 1989) (Figure 4.1E). The same motif was previously noted in other proteins that transport or bind heavy metals, such as MerA, mercuric reductase, and in MerP, a periplasmic mercury binding protein (Silver et al. 1989), and can be a general heavy metal binding site.

Methanotrophs, aerobic bacteria that grow on methane as a sole source of carbon and energy, are expected to have a tightly regulated copper uptake/efflux system since the key enzyme of methane oxidation, particulate methane monooxygenase (pMMO), requires copper, but on the other hand, high levels of copper (above 50 μM) are toxic to methanotrophs. The presence of a specific copper uptake system(s) in methanotrophs is supported by the results of copper accumulation experiments with a type I methanotroph M. albus BG8 (Berson and Lidstrom 1996). In analogy to other bacteria, it seemed possible that a P-type copper-transporting ATPase(s) could be involved in copper transport in methanotrophs. This chapter describes cloning of three putative copper ATPase genes, atpA, atpB and atpC, their sequencing (complete sequence of atpA has been obtained, atpB and atpC were sequenced only partially), and an attempt to make a mutant in atpB.
4.2. Materials and Methods

4.2.1. Bacterial Strains, Culture Conditions, and Preparation of DNA.

*Methylocrobium albus* BG8 (Whittenbury *et al.* 1970), a type I methanotroph, was grown with 10μM copper nitrate added to nitrate mineral salts (NMS) medium (Whittenbury *et al.* 1981) as described in Chapter Three (3.2.1) to the late exponential phase and harvested. Chromosomal DNA was extracted from *M. albus* BG8 by the slightly modified procedure of Satio and Miura (1963) as described in detail in Chapter Three (3.2.3). Plasmid DNA was prepared by the method outlined in Chapter Three (3.2.3).

All enzymes used were purchased from New England Biolabs, Inc. (Beverly, Ma) unless stated otherwise. Restriction enzyme digestions were performed as described by Maniatis (*et al.*, 1982). DNA separation by electrophoresis, all ligations, transformations, DNA-DNA hybridizations with dried gels and with nitrocellulose filters, oligonucleotide probe labeling were carried out following the protocols described in Chapter Three.

4.2.2. Polymerase Chain Reaction.

The polymerase chain reaction (PCR) method was used to generate a probe for screening fragments of *M. albus* BG8 chromosome obtained by digestion with restriction enzymes. Amino acid sequences of CopA, a copper uptake protein of the Gram-positive bacterium *Enterococcus hirae* (Odermatt *et al.* 1993) and MC1, a putative human copper-ATPase (Vulpe *et al.* 1993) were aligned (Figure 4.1) to identify
consensus regions. Conserved regions, DKTGTIT (positions 425-431 in the amino acid sequence of CopA and positions 1044-1050 in the amino acid sequence of MC1) (Figure 4.1 C) and MVGDGIND (positions 618-625 in CopA and positions 1298-1305 in MC1) (Figure 4.1D) were identified. Primers P1-A and P2-A were generated, which were identical to the fragments of DNA sequence from copA of E. hirae that corresponded to the two consensus amino acid sequences. The other primers were backtranslated from the amino acid sequences based on M. albus BG8 codon usage frequencies (Table 4.1). The following degenerate primers were designed and synthesized at the Microchemical Facility at the California Institute of Technology: forward primers P1-A (5'-GAT-AAA-ACT-GGA-ACG-ATT-AC-3'), P1-B (5'-GAC-AAA-ACC-GGC-ACC-ATC-AC-3'), P1-C (5'-GA(C or T)-AA(A or G)-AC(C or G or T)-GG(A or C or T)-AC(C or G or T)-AT (C or T)-AC (C or G or T)-3'), and reverse primers, which were designed as reverse complements to the translated amino acid sequence MVGDGIND, P2-A (5'-TCA-TTG-ATT-CCA-TCA-CCG-ACC-3'), P2-B (5'-TCG-TTG-ATG-CCG-TCG-CCG-ACC-3').

For each primer, the temperature of hybridization was calculated based on the following formula (Maniatis et al. 1982): 

\[ T_{hybr} = T_m - 10 \degree C = [69.3 + 0.441(GC\%) - (650/L)] - 10\degree C, \]

where \( T_{hybr} \), hybridization temperature; \( T_m \), oligo melting temperature; \( GC\% \), the percent of G and C in the oligo; \( L \), number of base pairs in the oligo. The following hybridization temperatures were obtained: P1-A, 42.2\degree C; P1-B, 51.1\degree C; P1-C, 46.6\degree C; P2-A, 49.3\degree C; P2-B, 57.7\degree C. Each primer was end labeled with 5'-[\( \gamma \)\(^{32}\)P]ATP, denatured and hybridized with dried agarose gels that contained M. albus BG8 chromosomal DNA digested with the following restriction enzymes: PstI, SphI, SalI, EcoRI, BamHI, HindIII. Only P1-C
and P2-B gave positive hybridization with the chromosomal DNA [up to 7 hybridizing bands were identified for each restriction (results are not shown due to high background)] and, therefore, this pair was used in PCR reactions.

The PCR reactions were carried out in 100 µL volumes under the following conditions: approximately 20 ng template DNA, PCR buffer (10 mM Tris-HCl, 50 mM KCl), 0.2 mM each dNTP, 1 U of Taq polymerase (Boehringer, Germany), 100 pmol each primer, and in some cases, 5% [vol/vol] DMSO. The PCR reactions were performed with five concentrations of MgCl₂: 0.5 mM, 1 mM, 1.5 mM, 2 mM, or 2.5 mM. A Hybaid thermal cycler (Combi TR-2) was used for 25-cycle amplification with the following reaction conditions: denaturation, 94°C for 1 min.; annealing, 55°C or 50°C or 44°C for 1 min.; polymerization, 72°C for 2 min.

PCR products obtained were separated by electrophoresis on 1.4% agarose gels. The combination of 1mM MgCl₂, DMSO, and an annealing temperature of 44°C gave the best resolution of three major PCR products: pcrA, 340 bp, pcrB, 620 bp and pcrC, 900 bp) on the gel (Figure 4.2). It appeared that the presence of DMSO increased production of the main PCR products, while bringing down the background (Figure 4.2), so DMSO was added to all subsequent PCR reactions. Two additional PCR reactions were carried out with only one primer added, as controls. Two of the major PCR products pcrB and pcrC, were in the approximate size range expected, and they were chosen for further study.

The optimized PCR reaction was run once more, PCR products were separated on 1.4% agarose gels, DNA bands of pcrB and pcrC were excised from the gel and purified
using a Geneclean II © kit (Bio 101 Inc., La Jolla, Ca). The purified DNA was then cloned into the pCR™II (Invitrogen, San Diego, CA) vector that was used to transform INVαF™ E. coli cells (Invitrogen, San Diego, CA) using the protocol recommended by the manufacturer. The transformants were plated on Luria-Bertani (LB) agar with 50 µg/ml ampicillin (Amp) and with 2.5% [wt/vol] X-gal (5-bromo-4-chloro-3-indolyl-β-D-galactoside) added. X-gal is a color indicator for the cells containing plasmids with inserts, which remain white on plates with X-gal. To determine the orientation of the inserts, 12 white clones per each insert were picked, grown overnight in LB broth containing 50µg/ml of Amp, and were used for plasmid DNA isolation. The plasmid DNA samples obtained were digested with EcoRI, which cuts the vector 9 bp up- and downstream of the PCR insert, and electrophoresed on 1.6% agarose gels. Two bands were observed in each case: one, of approximately 4 kb, corresponding to the pCR™II vector, and another band, of either about 600 bp or 900 bp. Four different sizes of the second band were identified among the 620-bp inserts, 2.3, 2.5, 2.8, and 2.9, and three sizes of the second band were seen among 900-bp inserts, 3.1, 3.3, and 3.4 (not shown). These clones were grown overnight in LB with Amp, and the plasmid DNA was isolated using a Qiagen-tip 20 kit (Qiagen, Los Angeles, CA). DNA sequencing was performed with an Applied Biosystems automated sequencer at the California Institute of Technology, Pasadena, Sequencing Facility, for both strands with M13 Forward and M13 Reverse sequencing primers (New England Biolabs, Inc., Beverly, MA).

Clones 2.5 and 2.9 were found to contain plasmids with the same DNA fragment inserted in either the forward or reverse direction. The recombinant plasmids from E.coli
strains 2.3, 2.5, and 2.8 were called pOB18, pOB19, and pOB20 correspondingly, and were assumed to encode portions of ATPase-like genes. In order to obtain more complete information, these plasmids were used as probes for screening *M. albus* BG8 genes.

**4.2.3. Cloning and Sequencing of ATPase genes**

DNA was isolated from pOB18, pOB19, and pOB20 and digested with *EcoRI* and the DNA fragments obtained were separated on agarose gels. Two bands were observed in each case: one, of 3.9 kb, corresponding to the pCR™II vector, the other, approximately 620-bp, of the inserts. The 620-bp band was excised from each agarose gel, purified using a GeneClean II™ kit (Bio 101 Inc., La Jolla, Ca), and randomly labeled with [α-³²P]dCTP as described by Maniatis (Maniatis *et al.* 1982).

Chromosomal DNA of *M. albus* BG8 was digested with *PstI*, *SphI*, *SalI*, *EcoRI*, *BamHI*, and *HindIII*. The DNA fragments obtained were separated by electrophoresis on 0.7% agarose gels (three such gels were prepared). The gels were dried and hybridized with the labeled 620-bp fragment of the 2.3, 2.5, and 2.8 clones at 68°C as described in Chapter Three. These fragments will be called by the name of the clone from which they were isolated, e.g. clone 2.3 will mean 620-bp DNA fragment isolated from clone 2.3. A 1.8-kb *EcoRI* band, hybridizing to clone 2.5, and a 2.5-kb *SalI* band, hybridizing to clone 2.8 were chosen for subsequent cloning. Hybridization with clone 2.3 did not produce any convenient fragments for cloning (not shown), and, therefore, restriction of *M. albus* BG8 chromosomal DNA with different enzymes (*Apol*, *KpnI*, *SaeI*, *XbaI*, and *BspMI*) was
performed. The hybridization of the dried gel with clone 2.3 identified a 3-kb *Kpnl* fragment that was used for further cloning.

The 1.8-kb *EcoRI*, 2.5-kb *SalI*, and 3-kb *Kpnl* fragments were cloned into the pUC19 vector (New England Biolabs, Inc., Beverly, Ma) using the protocols described in Chapter Three to generate pOB21 (contains partial *atpB*), pOB22 (contains partial *atpC*), and pOB31 (contains complete *atpA*). These recombinant plasmids were used for sequencing of the *EcoRI*, *SalI*, and *Kpnl* fragments on both strands at the Sequencing Facility at the California Institute of Technology, Pasadena. At the beginning, sequencing was performed with M13 Forward and M13 Reverse sequencing primers (New England Biolabs, Inc., Beverly, Ma). After the initial sequences were obtained, synthetic oligodeoxynucleotide primers complementary or identical to previously sequenced DNA fragments were synthesized at the Microchemical Facility at the California Institute of Technology, and they were used for sequencing. To translate and to analyze the DNA sequence programs supplied by the Genetic Computer Group (GCG), University of Wisconsin, were used.

### 4.2.4. Construction of Insertion Mutations in *atpB*.

An insertion mutant defective in *atpB* was obtained by homologous recombination as described in Chapter Three using the kanamycin resistance cassette (Km\(^\dagger\)) as a selecting inactivating marker. In short, the 611-bp fragment from pOB19 (pCR\(^\text{TM}\)II vector containing Km\(^\dagger\) and ampicillin resistance (Amp\(^\dagger\))) was recloned into pAYC63 (Chistoserdov, personal communication) containing Amp\(^\dagger\) and chloromphenicol
resistance (Cm\(^r\)) using LB agar plates with 50 mg/ml [wt/vol] Cm (in ethanol) to select recombinants. This clone was called pOB24. The AvaI site in pOB24 appeared to be the most promising for the insertional mutation since it is located almost in the middle of the 611-bp fragment, 270 bp downstream from the beginning of the insert. However, the site was not unique in pOB24, since pAYC63 (3.3 kb) also contains an AvaI site in the polylinker. Therefore, prior to the introduction of the Km\(^r\) cassette, the site was removed from the vector by standard methods described by Maniatis (1982) to generate pOB25 with the unique AvaI site in the middle of atpB. The plasmid carrying the Km\(^r\) gene transcribed in the same direction as atpB (pOB26) was ligated with the suicide vector pAYC61 (Chistoserdov et al. 1994), and the resulting plasmid (pOB30, Figure 4.3) was transformed into *E. coli* S17-1 (Simon et al. 1983). The resulting *E. coli* strain was employed as a donor in homologous recombination with *M. albus* BG8. The mutants were selected on NMS agar plates with 15\(\mu\)g/ml Km added grown under a methane/air atmosphere.

### 4.3. Results and Discussion

#### 4.3.1. Isolation and Cloning of Three Putative Atpase Genes, atpA, atpB and atpC.

In order to determine whether *M. albus* BG8 might contain one or more genes encoding P-type ATPases, the consensus amino acid sequences, DKTGTIT and MVGDGIN, found in two known copper transporters, *CopA*, the copper uptake protein of the Gram-
positive bacterium *Enterococcus hirae* (Odermatt et al. 1993) and MC1, a putative human copper-ATPase (Vulpe et al. 1993) (Figure 4.1 C and D), were used to generate three forward primers and two reverse primers for PCR amplification. The redundancy of these primers was minimized by choosing the most prevalent of *M. albus* BG8 codons (Table 4.1). Only two of these primers, the forward primer P1-C and the reverse primer P2-B, hybridized to *M. albus* BG8 DNA. Several (up to 7) bands were seen in each case, suggesting the possibility that multiple ATPase-like genes might be present. This hypothesis was studied further by cloning and sequencing putative ATPase genes. Three PCR products, *pcrA*, *pcrB*, and *pcrC* of approximately 340, 620, and 900 nucleotides correspondingly, were generated by PCR. A PCR product of approximately 600 bp was expected from the PCR reaction with primers P1-C and P2-B since the primers were created based on the highly conserved regions separated by approximately 600 bp in bacterial ATPases. The PCR reaction with only one primer did not give any products in case of P1-C, but the reaction with P2-B produced two small DNA fragments, one of which was the same size as *pcrA* (Figure 4.2), which implied that *pcrA* was the result of an incomplete PCR reaction, and, therefore, it was not studied further.

The PCR products *pcrB* and *pcrC* were cloned into pCRTMII (Invitrogen, San Diego, CA) vector. The restriction analysis of the generated clones revealed three independent clones containing inserts of approximately 620-bp, and three independent clones containing inserts of approximately 900-bp. DNA fragments purified from all six clones were sequenced on both strands. The sequences obtained (not shown) were translated into amino acid sequences, and the obtained polypeptides were compared with
known proteins from the data base (SwissProt) by using GCG (University of Wisconsin) programs. All three sequenced 620-bp fragments demonstrated high homology to cation transporting ATPases, with especially high identity to copper and cadmium ATPases: 31.9% identity in a 207 aa overlap for pOB20, 41.2% identity in a 204 aa overlap for pOB18, and 41.7% identity in a 204 aa overlap for pOB19 (not shown). The 900-bp fragments, however, were not significantly homologous to any of the genes encoding known proteins. These, probably, were generated as side-products of PCR reactions due to non-specific binding of the primers to the DNA template. They were not pursued further.

Radioactively labeled 620-bp DNA fragments from pOB18, pOB19, and pOB20 were used to screen chromosomal DNA of M. albus BG8 digested with restriction endonucleases. 1.8-kb EcoRI, 2.5-kb SauI, and 3-kb KpnI fragments were identified by clones 2.5, 2.8, and 2.3, respectively. These fragments were cloned into the pUC19 vector (New England Biolabs, Inc., Beverly, MA) to produce pOB21, pOB22, and pOB31 respectively and the inserts were sequenced on both strands.

4.3.2. Sequencing Results And Sequence Comparison With Known Copper ATPases.

The sequence analysis revealed the presence of one open reading frame (atpA), 2231 bp in length, in pOB31 (Figure 4.4), and partial open reading frames atpB (1866 bp) and atpC (689 bp) in pOB21 (Figure 4.5) and pOB22 (Figure 4.6) correspondingly. Both atpB and atpC encoded the C-terminal part of the polypeptides AtpB and AtpC.
The amino acid sequence derived from \textit{atpA}, \textit{atpB} and \textit{atpC} were compared with each other and with sequences in the SwissProt data base. It appeared that the three polypeptides are closely related to each other: AtpA/AtpB, 48.5\% identity and 71.8\% similarity (when conserved substitutions are considered); AtpA/AtpC, 48.3\% identity and 67.8\% similarity; AtpB/AtpC, 44.3\% identity and 69.1\% similarity. The AtpA/AtpB/AtpC alignment results (Figure 4.7) show especially high homology at the C-terminus of the polypeptides. All three polypeptides exhibited strong homology to P-type ATPases, especially the copper-ATPase family (Table 4.2, results only for AtpA are shown). Since only \textit{atpA} was completely sequenced and because \textit{atpB} and \textit{atpC} are very similar, further discussion will be focused on AtpA.

The product of \textit{atpA} has the highest homology (45.7\% identity in 742 aa overlap) to a cation-transporting ATPase found in the thylakoid membrane of the cyanobacterium \textit{Synechococcus} sp. PCC7942, which was recently postulated to be a copper-transporting ATPase (Kanamaru \textit{et al.} 1994). The next four proteins most closely related to AtpA were CopA from \textit{Enterococcus hirae} (Odermatt \textit{et al.} 1993), WD, human Wilson disease related ATPase (Tanzi \textit{et al.} 1993), Mc1, human Menkes disease, associated ATPase (Vulpe \textit{et al.} 1993), and CtaA, another P-type ATPase from the cyanobacterium \textit{Synechococcus} 7942 (Phung, \textit{et al.} 1994). These are all copper-transporting ATPases (Table 5.2). Potassium/copper transporting ATPase, CopA, is thought to be a copper-uptake ATPase, because disruption of the \textit{copA} gene does not affect copper-sensitivity of the cells but causes dependency on higher added copper concentrations for growth (Odermatt \textit{et al.} 1993). Wilson and Menkes diseases are human hereditary disorders of
copper transport. It has been suggested that Wilson’s disease is related to a defect in the copper-efflux ATPase, WD, that results in an inability to excrete an excess of copper and causes toxic accumulation of copper in the liver and brain (Tanzi et al. 1993). Menkes disease is associated with copper deficiency caused by a disruption in the McI gene encoding a copper ATPase, which is believed to be involved in copper transport to tissues containing copper-requiring proteins (Tanzi et al. 1993, Vulpe et al. 1993). A mutation in the ctaA gene of Synechococcus 7942, encoding a P-type ATPase, increases tolerance of the cells to higher copper concentrations while it does not cause higher resistance to other cations, suggesting that the product of ctaA is a copper ATPase (Phung et al. 1994).

A lower level of identity was observed between AtpA and cation transporting ATPases A and B with unknown specificity from Mycobacterium leprae (Fsilu and Cole, 1995), CCC2, an uptake copper ATPase from the yeast Saccharomyces cerevisiae (Fu et al. 1995), and CopB, a potassium/copper efflux ATPase from Enterococcus hirae (Odermatt et al. 1993) (Table 4.2). Outside of the copper ATPase subfamily, significant similarity was seen between AtpA and CadA, a cadmium ATPase of Staphylococcus aureus (30.2% identity) (Nucifora et al. 1989). AtpA is not similar to other P-type ATPases outside of the highly conserved regions of phosphorylation and ATP binding, which results in overall lower similarity between them (21 to 28% identity in the 742 aa overlap). The sequence alignment tree presented in Figure 4.8 summarizes similarity relationships of P-type ATPases, and places AtpA in the copper ATPase subfamily.

All key motifs of P-type ATPases such as those involved in phosphatase function, ion-transduction, phosphorylation, and ATP-binding are conserved in AtpA (Figures 4.1
and 4.6). The ion transduction region is highly conserved within the copper ATPase subfamily (Figure 4.1B). Both AtpA and AtpB show more similarity to copper ATPases than other ATPases in this domain (Figures 4.1B and 4.7). The motif, Cys-Pro-Cys-X(6)-Pro, which is thought to participate in cadmium and copper coordination (Phung et al. 1994) is also conserved in AtpA and AtpB (Figures 4.1B and 4.7). A putative copper/cadmium binding motif, Gly-Met-X-Cys-X-X-Cys previously noted in CopA, the copper ATPase of Enterococcus hirae (Odermatt et al. 1993), Mc1, Menkes copper ATPase (Vulpe et al. 1993), WD, Wilson’s Disease copper ATPase (Tanzi et al. 1993), the copper ATPase from Synechococcus sp. PCC7942 (Kanamaru et al. 1994), and CadA, the cadmium ATPase of Staphylococcus aureus (Nucifora et al. 1989) was identified in the N-terminus of AtpA (Figure 4.1E), strongly suggesting a role of AtpA in copper transport.

Hydropathy plots of AtpA, AtpB, and AtpC were generated with the GCG package using the residue-specific hydrophobicity index of Kyte and Doolittle (1982) and a span of 20 residues (Figures 4.9-4.11). Nine highly hydrophobic regions were identified in AtpA (Figure 4.9). The same regions were identified within sequenced fragments of AtpB, 3rd-9th, (Figure 4.10) and AtpC, 8th and 9th, (Figure 4.11). It is thought that a hydrophobic but uncharged sequence has a high probability to be in a transmembrane domain (Tanzi et al. 1993). To identify membrane-spanning regions, the hydrophobicity profile of AtpA was overlapped with a profile of residue charges (Figure 4.12, only hydrophobic regions identified in Figure 4.9 are shown). All nine hydrophobic peaks corresponded to the regions of no or little charge (zero or one charged residue per 14 to
20 residues) and they were predicted (Peptidestructure Program, GCG, Chou-Fasman algorithm) to be α-helices. Therefore, it is plausible to conclude that there are nine transmembrane domains in AtpA, including the metal-binding domain, 1, and the ion transduction domain, 7. A similar membrane topology was proposed for other copper ATPases (Tanzi et al. 1993, Odermatt et al. 1993, Vulpe et al. 1993).

4.3.3. Mutant Characterization

To confirm a role of AtpA, AtpB and AtpC in copper transport in *M. albus* BG8, an attempt was made to construct insertion mutants by homologous recombination between the *M. albus* BG8 chromosome and plasmids containing insertions in the genes of the interest. No mutants were obtained in *atpA* and *atpC*. The phenotype of the *atpB* insertion mutants was checked on NMS plates containing Amp, and it was found to be Km'/Amp' for all colonies obtained. Therefore, only single crossovers, containing an insertion of the entire suicide plasmid, and generating an intact gene were identified among *atpB* mutants (Figure 4.13). The inability to obtain double crossover mutants suggests that *atpB* is required for growth of *M. albus* BG8.

In conclusion, the presence of a copper or cadmium binding motif in AtpA and the high identity of AtpA, AtpB and AtpC with the subfamily of copper ATPases suggests that the cloned genes might encode copper pumps. Further studies, including more mutant construction and copper uptake experiments, will be needed to clearly define the role of this protein copper uptake.
4.4. References


Fu, D., T. J. Beeler, and T. Dunn. 1995. Sequence,mapping and disruption of *CCC2*, a gene that cross complements the Ca^{2+}-sensitive phenotype of csg1 mutants and encodes a P-type AYPase belonging to the Cu^{2+}-ATPase subfamily. Yeast 11:283-292.


Table 4.1 Codon frequency statistics for *Methylobacterium albus* BG8 calculated using the CodonPreference program of GCG Package.

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Table 4.2 The results of sequence comparison of AtpA with some other cation transporting ATPases.

AtpA sequence was compared with other ATPase sequences using the FASTa (GCG) algorithm. In the table below, sequences are displayed in the order of decreasing similarity to AtpA. Gaps were introduced into sequences to enhance their alignment, and the reported length of overlapping regions include these gaps.

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<tr>
<td>ATC2</td>
<td>Ca²⁺</td>
<td><em>Saccharomyces cerevisiae</em></td>
<td>723</td>
<td>29.9% (720)</td>
<td>P38360</td>
</tr>
<tr>
<td>KdpB</td>
<td>K⁺</td>
<td><em>Escherichia coli</em></td>
<td>682</td>
<td>27.96% (740)</td>
<td>P03960</td>
</tr>
<tr>
<td>AtnA</td>
<td>Na⁺/K⁺</td>
<td><em>Artemia salina</em></td>
<td>996</td>
<td>21.3% (580)</td>
<td>P17326</td>
</tr>
<tr>
<td>MgtB</td>
<td>Mg²⁺</td>
<td><em>Salmonella typhimurium</em></td>
<td>908</td>
<td>20.3% (650)</td>
<td>P22036</td>
</tr>
</tbody>
</table>
### A. Phosphatase Region

| Mgtb | A G D L V P A D V R L L A S R D L F I S Q S I L G E S L P V E K | 232 |
| Atna | P G D R I P A D I R T S C O S M K V D N S S L T G E S E P - - - | 205 |
| Cadd | P G E K I A M D G I I N G V S A - V N Q A A I T G E S V P V A K | 357 |
| Cada-Bac | P G K I A M D G V V G S Y S A - V N Q T A I T G E S V P V E K | 277 |
| Mc1 | P G K K F F V D G R V I E G H S M - V D E S L I T G E A M P V A K | 888 |
| Ccc2 | P G D K I P A D G I T R G E S E - T D E S L M T G E S I L V P K | 477 |

**Figure 4.1** Conserved regions found in P-type ATPases.

The deduced amino acid sequence of the *M. albus* BG8 Atpase, AtpA, was aligned with the following ATPases (from top to bottom, SwissProt data base accession numbers are shown in the parentheses): Mgtb, Mg$^{2+}$ transport ATPase, *Salmonella typhimurium* (P22036); Atna, Na$^+$/K$^+$ ATPase, *Artemia salina* (brine shrimp) (P17326); Cada, Cd$^{2+}$ ATPase, *Staphylococcus aureus* (P20021); Cadd, Cd$^{2+}$ ATPase, *Staphylococcus aureus* (P37386); Cada-Bac, Cd$^{2+}$ ATPase, *Bacillus firmus* (P30336); WD, Cu$^{2+}$ efflux ATPase associated with human Wilson disease (P35670); Mc1, Cu$^{2+}$ uptake ATPase associated with human Menkes disease (Q04656); Atpa, cation ATPase from *M. albus* BG8, this study; Pacs, probably Cu$^{2+}$-transporting ATPase, *Synechococcus* sp. (P37279); Copa, K$^+$/Cu$^{2+}$-transporting ATPase A, probably involved in copper intake, *Enterococcus hirae* (P32113); Syna, Cu$^{2+}$-transporting ATPase, *Synechococcus* sp. (P37385); Ccc2, Cu$^{2+}$-transporting ATPase, *Saccharomyces cerevisiae* (P38955); Copb, K$^+$/Cu$^{2+}$-transporting ATPase B, probably involved in copper efflux, *Enterococcus hirae* (P05425); Atc2, Ca$^{2+}$-transporting ATPase, *Saccharomyces cerevisiae* (P38360); Atkb, K$^+$-transporting ATPase, *Escherichia coli* (P03960).

The putative metal binding motif (Part E) is shown only for copper and cadmium ATPases. The sequences were aligned and arranged according to amino acid similarity by Pileup program of GCG Group. Gaps (-) were introduced when needed for better alignment. The regions with identical sequences are boxed. The signature motifs are marked with asterisks.
B. Ion Transduction Domain

<table>
<thead>
<tr>
<th></th>
<th>Mgtb</th>
<th>Attna</th>
<th>Cada</th>
<th>Cadd</th>
<th>Cada-Bac</th>
<th>Wd</th>
<th>Mc1</th>
<th>Atpa</th>
<th>Pacs</th>
<th>Copa</th>
<th>Syna</th>
<th>Ccc2</th>
<th>Copb</th>
<th>Atc2</th>
<th>Atkb</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>VAVGLTPEMLPIMVSSN</td>
<td>GIIVAKGLELATVTVCL</td>
<td>AVLVCPCALVISPTSI</td>
<td>AVLVCPCALVITPTSI</td>
<td>AVLVCPCALVISPTSI</td>
<td>TVLCIACPCSGLGLATPTAV</td>
<td>TVLCIACPCSGLGLATPTAV</td>
<td>TVLVICPACGLGLATPTISV</td>
<td>VMIACPCALGLATPTSI</td>
<td>SVLVICPACGLGLATPTAI</td>
<td>SVLVICPACGLGLATPTAI</td>
<td>SVLVICPACGLGLATPTAI</td>
<td>TVFIICPHALGLATPLLV</td>
<td>TVLVICPACGLGLATPTAI</td>
<td>ALVLCCPITGGLLSAIG</td>
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<td></td>
<td>347</td>
<td>321</td>
<td>383</td>
<td>460</td>
<td>380</td>
<td>964</td>
<td>1012</td>
<td>400</td>
<td>402</td>
<td>393</td>
<td>444</td>
<td>595</td>
<td>408</td>
<td>871</td>
<td>275</td>
</tr>
</tbody>
</table>

C. Phosphorylation Site

<table>
<thead>
<tr>
<th></th>
<th>Mgtb</th>
<th>Attna</th>
<th>Cada</th>
<th>Cadd</th>
<th>Cada-Bac</th>
<th>Wd</th>
<th>Mc1</th>
<th>Atpa</th>
<th>Pacs</th>
<th>Copa</th>
<th>Syna</th>
<th>Ccc2</th>
<th>Copb</th>
<th>Atc2</th>
<th>Atkb</th>
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<tr>
<td></td>
<td>LCCTDKTGTLTDQD</td>
<td>ICSDKTGTLTQRNMTV</td>
<td>VAFDKTGTLTKGVPPV</td>
<td>VAFDKTGTLTKGVPPV</td>
<td>VAFDKTGTLTKGVPPV</td>
<td>VMFDKTGTLTIHGVPVRV</td>
<td>VFDKTGTLTIHGVPVRV</td>
<td>LVDKGTGTVTGKPKV</td>
<td>VILDKGTGTLTQGQPSV</td>
<td>ILDKGTGTLTQGRPV</td>
<td>FVFDKTGTLTTGFGMV</td>
<td>FVFDKTGTLTTGEGKLTV</td>
<td>LLDKTGTLITNLGNRQA</td>
<td>392</td>
<td>371</td>
</tr>
</tbody>
</table>

Figure 4.1 (continued)
### D. ATP-binding Region

|       | Q    | K    | N    | G    | H    | T    | V    | G    | F    | L    | G    | D    | G    | I    | N    | D    | A    | P    | A    | L    | R    | D    | A    | D    | V    | G    | I    | S    | V    |       |
|-------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|-------|
| Mgtb  | Q    | K    | N    | G    | H    | T    | V    | G    | F    | L    | G    | D    | G    | I    | N    | D    | A    | P    | A    | L    | R    | D    | A    | D    | V    | G    | I    | S    | V    | 667   |
| Atta  | R    | Q    | R    | G    | E    | F   | V    | A    | V    | T    | G    | D    | G    | V    | N    | D    | S    | P    | A    | L    | K    | K    | A    | D    | I    | G    | V    | A    | M    | 709   |
| Cadd  | K    | A    | E    | -    | H    | G    | N    | V    | A    | M    | I    | G    | D    | G    | V    | N    | D    | A    | P    | A    | A    | A    | S    | T    | V    | G    | I    | A    | M    | 714   |
| Cada-Bac | R    | S    | E    | -    | Y    | G    | N    | V    | A    | M    | I    | G    | D    | G    | V    | N    | D    | A    | P    | A    | A    | A    | S    | T    | V    | G    | I    | A    | M    | 634   |
| Mc1   | Q    | E    | B    | G    | -    | K    | R    | V    | A    | M    | V    | I    | G    | D    | G    | V    | N    | D    | S    | P    | A    | L    | A    | Q    | A    | D    | V    | G    | I    | 1318  |
| Atpa  | Q    | A    | Q    | G    | -    | E    | T    | V    | G    | M    | V    | I    | G    | D    | G    | V    | N    | D    | A    | P    | A    | L    | A    | Q    | A    | D    | V    | G    | L    | 648   |
| Pacs  | Q    | S    | R    | G    | -    | Q    | V    | A    | M    | V    | I    | G    | D    | G    | V    | N    | D    | A    | P    | A    | L    | A    | Q    | A    | D    | V    | G    | I    | 651   |
| Copa  | Q    | K    | A    | G    | -    | K    | V    | G    | M    | V    | I    | G    | D    | G    | V    | N    | D    | A    | P    | A    | L    | K    | L    | A    | D    | V    | G    | I    | A    | M    | 638   |
| Syna  | Q    | S    | G    | G    | -    | D    | A    | V    | A    | M    | I    | G    | D    | G    | V    | N    | D    | A    | P    | A    | A    | A    | V    | G    | I    | S    | L    | 686   |
| Ccc2  | Q    | K    | G    | G    | N    | K    | V    | A    | M    | V    | I    | G    | D    | G    | V    | N    | D    | A    | P    | A    | L    | A    | S    | D    | L    | G    | I    | A    | 855   |
| Atc2  | S    | S    | S    | K    | R    | P    | V    | V    | F    | C    | G    | D    | G    | T    | N    | D    | A    | I    | G    | L    | T    | Q    | A    | T    | I    | G    | V    | H    | I    | 1124  |
| Atkb  | Q    | A    | E    | G    | R    | L    | V    | A    | M    | T    | G    | D    | G    | T    | N    | D    | A    | P    | A    | L    | A    | Q    | A    | D    | V    | A    | V    | A    | M    | 535   |

### E. A Putative Heavy Metal Binding Motif

|       | Q    | G    | F    | T    | C    | A    | N    | C    | A    | G    | K    | F    | E    | K    | N    | V    | K    | K    | I    | P    | G    | V    | Q    | D    | A    | K    | V    | N    | F    |       |
|-------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|-------|
| Cada-Bac | V    | O    | G    | F    | T    | C    | A    | N    | C    | A    | G    | K    | F    | E    | K    | N    | V    | K    | Q    | L    | S    | G    | V    | E    | D    | A    | K    | V    | N    | 48    |
| Wd    | I    | T    | G    | M    | T    | C    | A    | S    | C    | V    | H    | I    | E    | S    | K    | L    | T    | R    | T    | N    | G    | I    | T    | Y    | A    | S    | V    | A    | L    | 569   |
| Mc1   | V    | R    | G    | M    | T    | C    | A    | S    | C    | V    | H    | I    | E    | S    | S    | L    | T    | K    | H    | R    | G    | I    | L    | Y    | C    | S    | V    | A    | L    | 600   |
| Atpa  | I    | L    | G    | M    | S    | C    | A    | G    | C    | V    | S    | V    | V    | E    | S    | A    | L    | N    | G    | V    | P    | G    | V    | T    | E    | V    | S    | V    | N    | 44    |
| Pacs  | L    | G    | M    | G    | C    | A    | C    | A    | A    | G    | R    | I    | E    | A    | L    | I    | Q    | A    | L    | P    | G    | V    | Q    | E    | C    | S    | V    | N    | F    | 39    |
| Copa  | I    | T    | G    | M    | T    | C    | A    | N    | C    | A    | R    | I    | E    | K    | E    | L    | N    | E    | Q    | P    | G    | V    | M    | S    | A    | T    | V    | N    | A    | 42    |
| Syna  | V    | E    | G    | M    | K    | X    | C    | A    | G    | C    | V    | A    | A    | V    | E    | R    | R    | L    | Q    | Q    | T    | A    | G    | V    | E    | A    | V    | S    | V    | N    | 50    |
| Copb  | -    | -    | -    | -    | -    | -    | -    | -    | -    | -    | -    | -    | -    | -    | -    | -    | -    | -    | -    | -    | -    | -    | -    | -    | -    | -    | -    | -    | M    | N    | N    | G    | 4     |

Figure 4.1 (continued)
Figure 4.2 PCR-amplified products in an agarose gel.
1 kb ladder was used as DNA sizing standard, lanes 1 and 7. The reaction conditions were: denaturation, 94°C for 1 min.; annealing, 55°C or 50°C or 44°C for 1 min.; polymerization, 72°C for 2 min. The PCR reactions were carried out as follows: Lane 2, MgCl₂=1mM, +DMSO; lane 3, MgCl₂=1mM, -DMSO; lane 4, MgCl₂=2.5mM, -DMSO; lane 5, MgCl₂=1mM, +DMSO, only one primer added to the reaction, P1-C; lane 6, MgCl₂=1mM, +DMSO, only one primer added to the reaction, P2-B. Three major PCR products, *pcrA*, *pcrB*, *pcrC* are marked on the gel.
Figure 4.3 Construction of plasmid pOB30 carrying mutated atpB gene.

The 611-bp M. albus BG8 DNA fragment containing the middle part of atpB was cloned into pCR™II to generate pOB19, then the fragment was recloned into pAYC63 to create pOB24. The AvaI site was removed from the polylinker to produce pOB25 with a unique AvaI site in atpB. Then, atpB was disturbed by the insertion of a kanamycin resistance gene (Km') to produce pOB26. pOB26 was ligated with the suicide vector pAYC61 to yield pOB30.
Figure 4.4 Nucleotide sequence and deduced amino acid sequence of 2.2-kb M. albus BG8 chromosome region containing atpA.

Asterisks indicate stop codons. The nine putative transmembrane domains as predicted by a hydrophobicity plot are underlined. The signature motifs in phosphatase, ion transduction, phosphorylation, and ATP-binding regions are shown in bold. A putative copper binding motif is shown in bold italic. A putative Shine-Dalgarno sequence is double underlined.
Figure 4.4 (continued)
Figure 4.4 (continued)
Figure 4.4 (continued)
Figure 4.5 Nucleotide sequence and deduced amino acid sequence of the 1.96-kb M. albus BG8 chromosome region containing a fragment of atpB.

Asterisks indicate stop codons. The six putative transmembrane domains as predicted by a hydrophobicity plot are underlined. The signature motifs in phosphatase, ion transduction, phosphorylation, and ATP-binding regions are shown in bold.
Figure 4.5 (continued)
Figure 4.5 (continued)
Figure 4.6 Nucleotide sequence and deduced amino acid sequence of 1.34-kb *M. albus* BG8 chromosome region containing a fragment of atpC and the beginning of an identified open reading frame *orf1*. Asterisks indicate stop codons. The two putative transmembrane domains as predicted by a hydrophobicity plot are underlined. The signature motif of the ATP-binding domain is shown in bold.
Figure 4.6 (continued)
Figure 4.7 Sequence comparison of AtpA and partially sequenced AtpB and AtpC. The conserved regions are boxed. The ion transduction motif conserved in all heavy metal ATPases is indicated by asterisks. Dashes represent areas not sequenced. The amino acid sequences targeted by the primers are underlined.
Figure 4.8 Graphic representation of sequence similarities between cation transporting ATPases.

The diagram was generated by the program Pileup of GCG Group. The shorter the branch length, the more closely related the sequences. The accession numbers for the sequences are the same as in Figure 4.1. Copper ATPase subfamily is boxed. AtpA (this study) is underlined.
Figure 4.9 Predicted hydropathy plot of the *atpA* gene product using the Kyte and Doolittle algorithm (1982) over a span of 20 residues. The numbered bars indicate the polypeptide sequence fragments that have peaks of hydrophobicity and a length sufficient to form a putative transmembrane domain.
Figure 4.10 Predicted hydropathy plot of the C-terminal fragment of the *atpB* gene product using the Kyte and Doolittle algorithm (1982) over a span of 20 residues. The numbered bars indicate possible transmembrane domains.
Figure 4.11 Predicted hydropathy plot of the C-terminal fragment of the atpC gene product using the Kyte and Doolittle algorithm (1982) over a span of 20 residues. The numbered bars indicate possible transmembrane domains.
Figure 4.12 Charge (top) and hydrophobicity (bottom) profiles of AtpA.
Hphobic, hydrophobic; Hphilic, hydrophilic. Putative membrane domains are underlined and numbered. Only the fragments of the sequence containing such domains are shown.
Figure 4.13 The phenotypes of wild-type (w.t.), single and double crossover *atpB* mutants of *M. albus* BG8.

p, a promoter; *atpB*, gene encoding a putative copper P-type ATPase; *atpB<sup>dt</sup>, *atpB* deleted by an insertion of kanamycin resistance cassette (Km).
Appendix

Polymerase Chain Reaction (PCR) Method for DNA Amplification *in vitro*.

The PCR method utilizes thermostable Taq polymerase, which catalyzes the addition of nucleotides to the growing chain of preexisting DNA. To use the method for obtaining sufficient amounts of desired gene(s) for cloning or other purposes it is necessary to know a portion of the gene sequence to create primers. Primers are 18-22 nucleotide sequences that are complementary to sequence of gene(s) of interest. A PCR cycle involves three steps: (1) heat denaturation of double-stranded DNA to split it into two separate molecules; (2) cooling of the PCR reaction to allow hybridization of the primers to the DNA template; (3) primer extension by synthesis of a new DNA molecule complementing the template; this step is catalyzed by Taq polymerase (Figure A4.1). A PCR cycle is usually repeated 20 to 30 times, yielding up to a billion-fold increase in the target DNA concentration (Brock and Madigan 1991)
Figure A4.1
PCR for amplification of target genes.
Chapter Five

Conclusions

The possible future application of methanotrophs in biodegradation of trichloroethylene (TCE) depends on our understanding of all the major factors involved in the process. It is known that particulate methane monoxygenase (pMMO), found in all methanotrophs, can oxidize TCE and remove it to levels below the drinking water standard (DiSpirito et al. 1992). Recent studies (Chan et al. 1993; Nguyen et al. 1994) have identified trinuclear copper clusters in pMMO as the catalytic sites of the enzymatic activity. Moreover, it has been shown that the kinetics of TCE oxidation by pMMO are strongly affected by copper concentrations in the growth medium (Semrau 1995; Smith 1996). Therefore, an understanding of copper bioavailability and uptake in methanotrophs is important for optimization of in situ TCE bioremediation. This project has made significant progress in assessing the effect of copper speciation in the growth medium on copper accumulation by the type I methanotroph *Methylomicrobium albus* BG8 (Chapter Two), as well as in recognizing and evaluating possible copper uptake mechanisms in this microorganism (Chapters Three and Four).

5.1. Copper Speciation Affects Copper Uptake by *M. albus* BG8.

Copper is widely present in the environment, however, over 99% of the total copper is complexed by organic ligands, which results in cupric ion concentrations of
$10^{-10}$ to $10^{-18}$ M (Hodgson *et al.* 1965; Sunda and Ferguson 1983; Sunda and Hanson 1987; Coale and Bruland 1988; Hering and Morel 1988; Berggren 1989; Coale and Bruland 1990; Moffett *et al.* 1990; Van den Berg *et al.* 1990, Bruland *et al.* 1991; Van den Berg and Donat 1992). Although several copper species can coexist in an environment, there is a general agreement in the literature on a key role of cupric ion in toxicity and in biological uptake of copper in all studied organisms, from bacteria to animals (Sunda and Guillard 1976; Anderson and Morel 1978; Zevenhuizen *et al.* 1979; Blust *et al.* 1986; Coale and Bruland 1990; Bruland *et al.* 1991; Langford and Guzman 1992).

In this research the relationship between copper accumulation and cupric ion concentration in the medium was investigated in the type I methanotroph *M. albus* BG8 at copper concentrations that were neither growth limiting nor toxic (Chapter Two). The disappearance of cupric ion from the growth medium was studied using a copper-selective electrode, and was compared with total copper accumulation by the cells assayed by inductively coupled plasma mass spectrometry. The amount of copper accumulated by the cells was related to the cupric ion concentration rather than to that of total copper added to the growth medium, as has been seen previously in other organisms. Copper accumulation demonstrated a hyperbolic dependence on cupric ion concentration, suggesting saturation of copper binding sites on the cell surface. A mean maximum binding capacity was estimated as $(1.54\pm0.06)\times10^{-15}$ moles/cell, and an apparent half saturation constant as $(1.43\pm0.05)\times10^{-7}$ moles/l. Sorption experiments established that most of the copper accumulated by the cells was nonspecifically sorbed
to external sites (e.g., amino acid, carboxylic, hydroxy groups, etc.). Copper that was not removed by EDTA during a 24-h treatment was assumed to be either inside the cells or bound to specific copper transport sites. The copper concentration measured in the cells after the EDTA wash was essentially constant at $1-3 \times 10^{-17}$ moles of copper per cell despite a 100-fold variation in medium total copper and cupric ion concentrations. This result implies the presence of a specific copper uptake system in *M. albus* BG8 regulated by the copper concentration in the medium.

5.2. Copper Uptake in *M. albus* BG8 Involves Membrane Proteins.

Several transport mechanisms described in Chapter One (1.5. and 1.6.) could be involved in copper uptake by methanotrophs, such as a periplasmic binding-protein dependent mechanism (Figure 5.1A); an active transport via a copper P-type ATPase (Figure 5.1B); a siderophore-type high-affinity system (similar to the one in iron uptake) (Figure 5.1C); and copper accumulation in a special periplasmic storage protein for pMMO assembly (Figure 5.1D). All except a siderophore type mechanism might involve an outer membrane component, such as a porin, which facilitates copper transport from the outside into the periplasm. It is not uncommon for the cells to have more than one transport system with different affinities for a substrate, therefore, a combination of several transport systems from Figure 5.1 might be employed in methanotrophs.

By analogy to other uptake systems in bacteria, it is expected that a specific copper uptake system would be copper-regulated. Therefore, membrane and soluble fractions of *M. albus* cells were screened for the presence of copper-regulated
polypeptides (Chapter Three). One such copper-repressible membrane polypeptide, CorA, was identified, and the corresponding gene, corA, was cloned and sequenced. An insertion mutant defective in the gene was constructed, but it grew very poorly on plates or in liquid culture. This result confirms the vital significance of CorA to M. albus cells. DNA and protein data base analysis revealed some homology of CorA to rabbit and human calcium release channel protein. This, together with the fact that the expression of CorA seems to be regulated by copper (it was isolated as a major copper-repressible polypeptide), makes it plausible that CorA might be a divalent metal porin. It was not possible to confirm the specificity of CorA because of the poor growth of the CorA mutant.

To study the hypothesis that P-type ATPases might be involved in copper uptake by M. albus BG8 cells, a partial clone library of M. albus BG8 chromosome was screened with a labeled PCR product obtained with a pair of primers designed from the consensus sequences of two known copper ATPases (Chapter Four). Three putative copper ATPase genes, atpA, atpB, atpC, were cloned and sequenced (atpA was completely sequenced, while atpB and atpC were sequenced only partially). From the sequences analysis and comparison with DNA and protein data bases it appears that all three cloned genes are closely related to each other and that they belong to the group of heavy metal transporting ATPases, possibly they are members of the copper ATPases subfamily. The fact that only single crossover mutants were obtained when an insertion mutation was attempted in one of the genes, suggests that this gene is required for normal growth of M. albus BG8.
A project in progress is focusing on the identification and study of extracellular compounds excreted by _M. albus_ BG8 under conditions of copper limitation. The siderophores identified so far in the literature (mainly in iron acquisition) have been chemically classified as hydroxamates and catechols (Martinez _et al._ 1990). Copper has a high affinity for these ligands and possibly can be accumulated by some bacteria in the same way as iron (Hider 1984). Preliminary data from potentiometric titration experiments with _M. albus_ BG8 have suggested that some extracellular copper-complexing material is excreted by the cells (Lloyd and Kwan, personal communication). The chemical nature of these compounds is currently under investigation, but preliminary qualitative tests identified the presence of hydroxamate functionality in media in which cells were grown under copper limitation (Lloyd and Kwan, personal communication). The possibility of siderophore-type copper uptake in _M. albus_ BG8 is being pursued by Dr. Morgan's group.

The following hypothesis for copper uptake in _M. albus_ BG8 is suggested by the data presented in this research. One or more copper P-type ATPase(s) appears to be involved in copper uptake and/or copper efflux in _M. albus_ BG8. Since in all gram-negative bacteria P-type ATPases are situated in the inner membrane (Figure 5.1B), it is possible that the copper-repressible protein (CorA) described in Chapter Three may be an outer membrane porin that works in tandem with the putative copper ATPase(s) (Chapter Four). Such a porin would not necessarily be specific for copper, and might be overexpressed under conditions of any divalent metal limitation to facilitate the metal diffusion into periplasm. Such a scenario, however, does not exclude the possibility that
other uptake mechanisms presented in Figure 5.1 might be involved in copper transport in *M. albus* BG8.

This project has suggested a number of further studies into the mechanisms of copper transport in methanotrophs. First, the cellular location of CorA and the putative copper ATPases should be confirmed, by expression of the cloned genes in an alternative host, and the possible regulation of CorA by other metals should be investigated. Next, the sequences of *atpB* and *atpC* should be completed, and mutants should be generated that are defective in AtpA and AtpC, to test their phenotype. These genes should be tested for their ability to complement *E. coli* CopA and CopB mutants, to further investigate their functions. Finally, the existence of copper-binding siderophore-like compounds should be addressed, as well as the role of such compounds in copper uptake under different copper-related growth conditions.
5.3. Summary

- Cu accumulation by *M. albus* BG8 depends on the concentration of cupric ion \([\text{Cu}^{2+} (\text{H}_2\text{O})_6]\) in the growth medium.

- At high total copper concentrations in the growth medium, most of the accumulation is apparently due to non-specific sorption to outer cell layers.

- Total Cu/cell that is not removable by EDTA, presumably inside the cell, is relatively constant, indicating the presence of a specific homeostasis mechanism for copper.

- A membrane protein (CorA) that is over expressed in the absence of copper (copper-repressible) is required for significant growth of *M. albus* BG8, and may be a divalent metal porin.

- *M. albus* BG8 contains at least three genes whose products show homology to the protein subfamily of P-type copper ATPases.

- One of these genes was sequenced in its entirety, and it contains the P-type copper ATPase copper-binding signature sequence, suggesting that it does play a role in copper transport.
References


Figure 5.1 Schematic representation of the different types of transport systems that might be employed in copper uptake by Methylophilus albus BG8.

A. Binding-protein dependent mechanism. a, periplasmic substrate-binding protein; b,c,d,e, inner membrane-associated proteins. d,e, proteins that couple ATP hydrolysis to the transport. f, outer membrane porin

B. Copper transport via P-type ATPase (g) exposed to both cytoplasm and periplasm. Copper bound on the periplasmic side can be carried through the membrane by a conformational change of the protein.

C. A high-affinity system, consisting of low-molecular-weight copper chelator (h) and a specific membrane receptor (i).

D. Passive diffusion through porin (f) with further copper accumulation in a special periplasmic storage protein (j) for pMMO (m) assembly.