The effects of altered membrane fatty acid composition on the toxic interactions of heavy metals with Saccharomyces Cerevisiae

Niall G Howlett (1998)

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THE EFFECTS OF ALTERED MEMBRANE FATTY ACID COMPOSITION ON THE TOXIC INTERACTIONS OF HEAVY METALS WITH SACCHAROMYCES CEREVISIAE

Niall G. Howlett

A thesis submitted in partial fulfilment of the requirements of Oxford Brookes University for the degree of Doctor of Philosophy

January 1998
Abstract

The effects of altered membrane fatty acid composition on the toxic interactions of heavy metals with *Saccharomyces cerevisiae* were examined. *Saccharomyces cerevisiae* was enriched with the polyunsaturated fatty acids (PUFAs) linoleate (18:2) and linolenate (18:3) by growth in 18:2- or 18:3-supplemented medium. Incorporation of the exogenous PUFAs resulted in them comprising greater than 65% and 40% of the total fatty acids in whole-cell and plasma membrane lipids, and nuclear membrane lipids, respectively. Incorporation of the exogenous PUFAs had no discernible adverse effects on cell division. However, inhibition of cell division in the presence of Cd(NO₃)₂ was accentuated by growth in the presence of the di-unsaturated fatty acid linoleate. Furthermore, susceptibility to both Cd²⁺ and Cu²⁺-induced plasma membrane permeabilisation and whole cell toxicity was markedly accentuated in PUFα-enriched cells, and increased with the degree of fatty acid unsaturation. The increased sensitivity of PUFα-enriched cells to membrane permeabilisation and whole-cell toxicity was correlated with increased levels of lipid peroxidation in these cells. Cu²⁺- and Cd²⁺-induced lipid peroxidation was rapid and associated with a decline in plasma membrane lipid order, detected by fluorescence depolarization measurements. Levels of the lipid peroxidation products thiobarbituric acid-reactive substances (TBARS) and conjugated dienes were markedly higher in PUFα-enriched cells, compared with unsupplemented cells, following exposure to cadmium or copper. Thus, lipid peroxidation was demonstrated as a major means of heavy metal toxicity in a microorganism for the first time. In addition, the effects of PUFα-enrichment on the interactions of heavy metals with cellular nucleic acids were examined. Exposure of PUFα-enriched cells to the redox-active metals chromium and copper resulted in the uncoupling of DNA synthesis from cell division, leading to sequential S phases. For example, DNA levels of up to 8C were evident in 18:3-enriched cells after only 4.5 h exposure to 100 μM Cu(NO₃)₂. Using flow cytometry, the heterogeneity in susceptibility to copper toxicity of exponential phase *S. cerevisiae* was also examined. Susceptibility towards copper toxicity was demonstrated to be cell cycle stage-dependent, whereby G₂/M phase cells were found to be the most susceptible towards copper toxicity. Staining with the oxidant-sensitive probe 2',7'-dichlorodihydrofluorescein diacetate (H₂DCFDA) revealed that the greater copper sensitivity of G₂/M phase cells correlated with elevated endogenous levels of reactive oxygen species in these cells.
Abstracts, presentations and publications


Acknowledgements

I would like to express my immense gratitude to my supervisor Dr. Simon V. Avery, and both past and present members of the Avery laboratory, in particular, Michael J. Hoptroff, Stefania Radice, Kimberly S. Babb, Kevin J. Blackwell, and Shareeka L. Smith. I also wish to thank Profs. P. C. Tai and S. A. Crow Jr. for enabling me to continue my PhD research at Georgia State University, Atlanta. In addition, I would like to thank all those in Kell Hall, Atlanta, and the Tonge Building, Oxford for helpful discussion, times well spent, and for often fanning the embers. Last, but not least, I would like to thank my family for their generosity and continued encouragement.
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<tr>
<td>ade</td>
<td>adenine sulphate</td>
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<tr>
<td>AP</td>
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<tr>
<td>ATPase</td>
<td>Adenosine triphosphatase</td>
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<td>BSA</td>
<td>Bovine serum albumin</td>
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<td>can</td>
<td>L-Canavaline</td>
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<td>OGYE</td>
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<td>TBARS</td>
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<td>YEPD</td>
<td>Yeast extract peptone dextrose base</td>
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<td>YNB</td>
<td>yeast nitrogen broth</td>
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CHAPTER 1

Literature review
The continuing flux of heavy metals in the biosphere is an ever present concern. A wide variety of activities result in the release of heavy metals into the environment. However, release of metals from anthropogenic sources generally exceeds that emanating from natural sources. Indeed, on the EPA’s list of most commonly discharged priority pollutants, five out of sixteen pollutants are heavy metals (Novotny, 1995).

Examples of natural sources of metal fluxes include emissions from hydrothermal vents, volcanic eruptions and biogenic processes (Nisbet and Fowler, 1995; Novotny, 1995). For example, it is estimated that between 500,000 and 5 million tonnes of heavy metals are released by the Broken Spur hydrothermal vent field in the North Atlantic each year (Nisbet and Fowler, 1995). Furthermore, metallic minerals are abundant elements of the earth’s crust. Natural sources of cadmium, chromium and copper include zinc carbonate and sulphide ores, chromite and chromium oxide, and copper sulphide and chalcopyrite, respectively (Novotny, 1995).

The principal anthropogenic point sources of heavy metals include industrial activities such as the smelting and refining of metal ores, and the production and fabrication of metallic commercial products (Novotny, 1995; Nriagu, 1991). Diffuse enrichment of aquatic ecosystems with heavy metals is a consequence of municipal and industrial effluents, urban runoff, e.g. combustion of lead gasoline and deterioration of lead pipes, mine effluents, sewage discharges and atmospheric fallout, often resulting in severe localised pollution (Nriagu, 1991). The enrichment of soils with heavy metals arises primarily from both industrial and household refuse, industrial solid wastes, sewage, organic wastes, coal and wood ashes, fertilisers and pesticides, as well as atmospheric fallout (Novotny, 1995; Nriagu, 1991).
Thriving biological communities exist around natural metal-rich deposits, e.g. chemolithotrophic bacterial mats on polymetal sulphide-enriched surfaces of mid-ocean ridges (Nisbet and Fowler, 1995). Iron, manganese and sulphur are essential for planktonic photosynthesis, while copper, zinc, and mercury act as essential co-factors for microbial metalloproteins (Nisbet and Fowler, 1995). However, ecosystem over-enrichment, caused by the anthropogenic mobilisation of toxic metals, disrupts the stable, biogeochemical equilibrium of trace elements. Unlike organic xenobiotic compounds, toxic metallic compounds are not subject to natural degradation. While acute ecosystem stress has been demonstrated at the local scale, the chronic effects associated with regional and global metal over-enrichment have yet to be fully discerned. The continuing mobilisation of toxic metals, organometallic compounds and radionuclides has thus resulted in considerable research attention on interactions between these compounds and biota (Gadd, 1993; Vernet, 1992; Hughes and Poole, 1989).

Metal-microbe interactions have been the focus of major research attention for several reasons. Microorganisms generally occupy the lowest trophic levels in food chains that may ultimately lead to humans. Concern has arisen because of the ubiquity of microorganisms, and their high capacities for accumulation of heavy metals, organometallic compounds, and radionuclides. Consequently, these toxic compounds may have multiple exposure pathways to humans via food, water and air. For example, $^{137}$Cs may be transferred through food chains originating from microorganisms and eventually leading to humans. Certain food chains have been identified as routes of high $^{137}$Cs transfer, e.g. Lichen-caribou-man food chain (Avery, 1995a). Heavy metal toxicity
towards humans generally involves neurotoxicity, hepatotoxicity and nephrotoxicity. For example, prolonged environmental exposure to cadmium can result in nephritis, severe kidney failure, and skeletal disease (Elinder, 1992). Furthermore, compounds of cadmium (II), chromium (VI), and nickel (II) are known human carcinogens (Dally and Hartwig, 1997; Wetterhahn and Dudek, 1996; Xu et al., 1996).

Microorganisms also play crucial roles in the biogeochemical cycling of many essential elements, e.g. carbon (C), nitrogen (N), phosphorous (P), and sulphur (S), thus ensuring their stable environmental equilibrium. However, microbial activity is severely diminished in the presence of elevated levels of heavy metals, organometallic compounds, and radionuclides. For example, exposure of soil microbial populations to elevated levels of Cd, Ni, Pb, and Zn results in significant decreases in CO₂ evolution, and specific inhibition of cyanobacterial nitrogen fixation (Trevors et al., 1986). Thus, enrichment of terrestrial and aquatic ecosystems with heavy metals could adversely affect the biogeochemical cycling of C, N, P and S, essential trace elements, and organic matter decomposition processes (Nriagu, 1991).

Furthermore, as a consequence of the high heavy metal uptake-capacities of microorganisms, metal-microbe interactions have also been the subject of considerable biotechnological research attention in the last two decades, due to the potential use of microorganisms for the bioremediation of toxic industrial effluents (Avery, 1995a; Macaskie, 1991). Microorganisms generally accumulate metals in two distinct and well characterised phases; a metabolism-independent phase involving adsorption to the cell wall and a metabolism-dependent phase of intracellular transport across the plasma membrane (Gadd, 1993). Metabolism-independent metal accumulation is facilitated by
the presence of numerous metal-coordinating ligands such as sulphydryl, carboxyl and hydroxyl groups on the cell surface, and in membrane proteins (Shumate II and Strandberg, 1985). Microbial extracellular products, e.g. phytochelatins, siderophores, and exopolysaccharides, also have the capacity to sequester and complex metals. Metabolism-dependent intracellular transport of metals in microorganisms is generally mediated by membrane-bound proteins. For example, iron transport into the yeast *S. cerevisiae* is mediated by the *FET3, FTR1* and *FET4* plasma membrane-bound transporters (Stearman *et al.*, 1996), while a high affinity copper transporter is encoded by *CTRL* (Dancis *et al.*, 1994). Current methodologies for the detoxification of metal/radionuclide-containing industrial effluents include ion-exchange techniques, e.g. natural and synthetic zeolites, and elevated pH-metal precipitation (Avery, 1995b). However, metal-removal efficiencies of zeolites can be reduced significantly as a result of competition by ions of similar charge and ionic radii, while elevated pH metal-precipitation is ineffective for certain metal species, e.g. Cs⁺ (Avery, 1995b; Macaskie, 1991; Harjula and Lehto, 1986). The use of microbial biomass for the removal of toxic heavy metals, organometallic compounds and radionuclides from industrial effluents may represent an effective and cheap alternative to existing methodologies (Avery, 1995b). Currently, microbial mats consisting predominantly of sulphate-reducing bacteria of the *Thiobacillus* genus, and cyanobacteria offer the greatest potential for the decontamination of heavy metal-enriched sediments and effluents (Bender *et al.*, 1994; Fude *et al.*, 1994).

**Transition metals are chemically defined as those with *d* electrons and incompletely filled *d* orbitals. The Groups of the *d* block contain only three elements that**
correspond to the filling of the 3d, 4d and 5d shells, respectively. Each transition Group typically divides into two parts; the lightest element and the pair of heavy elements. The chemistry of the heavy elements is very similar due to the lanthanide contraction (Mackay and Mackay, 1991). Ligand field theory is particularly confined to d-element chemistry as transition metal ions are small. The M²⁺ and M³⁺ ions are centres of high charge density and can be strongly coordinated to lone pair donors (Mackay and Mackay, 1991). The toxic effects of heavy metals can thus be attributed partly to the multiplicity of coordination complexes and clusters that they can form (Butler and Harrod, 1989).

Microbial cells provide a ligand-rich environment in which heavy metals can exert their deleterious effects through a variety of mechanisms. For example, cadmium reacts with polythiol groups on cellular macromolecules, and both cobalt and cadmium can substitute for zinc in Zn-containing enzymes, e.g. carboxypeptidases (Price and Morel, 1990).

While the redox-active metal copper is an essential cellular redox component; copper is required for catalysis by enzymes such as Cu, Zn superoxide dismutase, lysyl oxidase, cytochrome c oxidase, and dopamine β hydroxylase (Knight et al., 1996), at elevated concentrations copper is extremely cytotoxic. Among other mechanisms of toxicity, copper indiscriminately binds to thiolate moieties and interacts with enzyme active sites (Cervantes and Gutierrez-Corona, 1994). Moreover, the interaction of both cadmium and copper, along with other transition metals, with cellular and organellar membranes can result in the rapid loss of membrane stability and impairment of membrane function (Fodor et al., 1995; Ohsumi et al., 1988). Cadmium- and copper-induced plasma-membrane permeabilisation with associated cellular K⁺ efflux has been extensively reported in the yeast Saccharomyces cerevisiae, and is considered a major mechanism of
toxicity in this organism (Ohsumi et al., 1988; De Rome and Gadd, 1987; Kessels et al., 1985; Gadd and Mowll, 1983). However, the underlying mechanism of heavy metal-induced membrane permeabilisation has yet to be clearly discerned.

One general cause for loss of membrane integrity in biological systems, in response to a variety of stimuli, can arise through reactive oxygen species (ROS)-mediated damage (Halliwell and Gutteridge, 1989). Redox-active metals, such as copper, are known to be capable of inducing ROS-generation through redox-cycling activity. Furthermore, nonredox-active metals, such as cadmium, may indirectly induce the accumulation of ROS through depletion of ROS-scavenging molecules such as glutathione and protein-bound sulhydryl groups (Figueiredo-Pereira et al., 1998; Li et al., 1997; Hassoun and Stohs, 1996; Stohs and Bagchi, 1995).

Free radicals, which includes the vast majority of reactive oxygen species, are chemically defined as highly unstable molecules that have an unpaired electron(s) in an outer orbital (Davies, 1995; Dix and Aikens, 1993). Free radicals are stabilised through abstraction of electrons from donor molecules, with the concomitant destabilisation of the donor molecule. In biological systems ROS are generated endogenously via normal metabolic processes. Primarily, ROS are generated during electron transport events such as mitochondrial respiration, or the respiratory burst of neutrophils, monocytes, and macrophages (James et al., 1998; Pongracz and Lord, 1998; Akaike et al., 1996). ROS are also generated during steroidogenesis and the β-oxidation of high-molecular weight fatty acids (Rapoport et al., 1998; Davidge et al., 1995). In addition to ROS generation during normal metabolism, exposure to numerous exogenous chemical agents such as the
monofunctional alkylating agent methyl methanesulfonate (MMS), methyl viologen (paraquat), mitomycin C, menadione, phenylhydrazine, and cumene hydroperoxide, as well as heavy metals, may result in the generation of ROS (Flattery O’Brien and Dawes, 1998; Brennan et al., 1994; Hama-Inaba et al., 1994).

The major ROS that occur in biological systems include superoxide anion (O$_2^-$) (a mono-radical), hydroxyl radical (OH$^-$), hydrogen peroxide (H$_2$O$_2$), and dioxygen (O$_2$) (a bi-radical; two unpaired electrons in the $\pi^*$ orbital). The non-enzymatic, univalent, reduction of molecular oxygen results in the generation of several highly reactive intermediates (1) (Davies, 1995).

\[
\begin{align*}
e^- & \quad \quad \quad \quad \quad e^- \\
\vdots & \quad \quad \quad \quad \quad \vdots \\
O_2 & \rightarrow O_2^- \rightarrow H_2O_2 \rightarrow HO^- \rightarrow H_2O
\end{align*}
\]

In mitochondria, superoxide anion can be generated at complex I (NADH-Ubiquinone) or at complex II (cytochrome c) of the electron transport chain. Furthermore, iron-sulphur clusters within the respiratory chain mediate the generation of superoxide anions (Keyer and Imlay, 1996). Superoxide anion can also be generated through auto-oxidation via interactions with cellular reductants, such as ascorbate, glutathione and NADH (Park et al., 1998; Minasi and Willsky, 1991). In addition, phagocytic cells, such as monocytes, neutrophils, and macrophages, utilise an NADPH oxidase enzymatic system to generate superoxide anion as part of their defence against pathogenic microorganisms (James et al., 1998; Pongracz and Lord, 1998). While the superoxide anion itself is not highly
reactive, it can give rise to highly reactive ROSs. For example, protonation of the superoxide anion generates the highly active oxidising agent, the perhydroxyl radical (HO$^\cdot$) (Valentine et al., 1998; Dix et al., 1996). Hydrogen peroxide is generated through the β-oxidation of high molecular weight fatty acids, and upon superoxide anion dismutation (Van der Leij et al., 1992; Seaman et al., 1982). Despite being a ROS, hydrogen peroxide is not a free radical as it has no unpaired electrons. However, hydrogen peroxide efficiently crosses biological membranes and leads to the generation of the highly reactive hydroxyl radical. Hydrogen peroxide undergoes both a slow, spontaneous decomposition, and a more rapid, metal-dependent decomposition, via the Fenton reaction (2), to yield the hydroxyl radical.

$$\text{Fe}^{2+} + \text{H}_2\text{O}_2 + \text{H}^+ \rightarrow \text{Fe}^{3+} + \text{HO}^\cdot + \text{H}_2\text{O} \quad (2)$$

Significantly, many redox-active transition metals, including Cu$^+$, Ni$^{2+}$ and Ti$^{3+}$, facilitate the generation of the hydroxyl radical, via Fenton-type reactions. Furthermore, the reduction of Cu$^{2+}$ and Fe$^{3+}$ by the superoxide anion, via the Haber-Weiss reaction (3), results in the regeneration of the reduced metal form which can subsequently undergo another Fenton reaction.

$$\text{Fe}^{3+} + \text{O}_2^- \rightarrow \text{Fe}^{2+} + \text{O}_2 \quad (3)$$

Thus, while transition metals play essential roles as cellular enzymatic redox components, excess levels of transition metals in biological systems promote the formation of several highly reactive oxygen species.

Reactive oxygen species, primarily O$_2^\cdot$, HO$^\cdot$, HO$_2^\cdot$, H$_2$O$_2$, and O$_2$ are capable of eliciting cellular damage through interactions with DNA, protein and lipids. The
interactions of ROS with DNA results in the formation of a wide variety of DNA lesions, including strand breaks, base loss or damage, and fragmentation of the deoxyribose moiety (Demple and Harrison, 1994; Ramotar et al., 1991). Indeed, DNA double-strand breaks may lead to genome rearrangements, such as deletions, duplications, and translocations, which have been implicated in carcinogenesis (Brennan and Schiestl, 1998; Manivasakam and Schiestl, 1998; Brennan et al., 1994). The hydroxyl radical has been implicated as a key ROS responsible for eliciting DNA damage, generating several base lesions in double-stranded DNA, including thymine glycol, 8-oxoguanine and formamido-pyrimidine among other base oxidation products (Demple and Harrison, 1994). Hydrogen peroxide-mediated cell toxicity results predominantly from the generation of hydroxyl radical via the Fenton reaction (2) (Flattery O’Brien and Dawes, 1998; Brennan et al., 1994). The DNA damage elicited by ionizing radiation, the antitumour antibiotic bleomycin, and peroxides is mediated via the formation of ROS, with the subsequent generation of apurinic/apyrimidinic (AP) sites and single-strand breaks with 3’-deoxyribose fragments (Teoule, 1986). Exposure to the oxidative mutagens methyl viologen (paraquat), mitomycin C, phenylhydrazine, and cumene hydroperoxide also induces a variety of DNA lesions, both directly (N7-methylguanine, O6-methylguanine, 3-methyladenine), and as a result of lesion processing (abasic sites, nicks, gaps, double-strand breaks). Oxidative mutagens also increase the frequency of both intrachromosomal and interchromosomal recombination (Brennan et al., 1994). A role for the hydroxyl radical in H2O2-induced gross chromosomal rearrangements has been implicated by the ability of the free radical scavenger dimethyl sulphoxide (DMSO)
to significantly inhibit the induction of both intrachromosomal and interchromosomal recombination by H$_2$O$_2$ (Brennan et al., 1994).

ROS-mediated protein damage occurs via oxidation of the amino acids histidine, lysine, arginine, tyrosine, phenylalanine, tryptophan, proline, methionine and cysteine, and through the formation of protein-crosslinks (Iwai et al., 1998; Luikenhuis et al., 1998; Moskovitz et al., 1997). Oxidative damage to proteins leads to increased proteolytic susceptibility; oxidation of amino acids to carbonyl derivatives signals proteins for proteasome degradation (Iwai et al., 1998; Jungmann et al., 1993a). For example, oxidation of arginine, lysine, proline, and threonine residues of the iron-dependent degradation domain of iron regulatory protein 2 (IRP2), to their corresponding carbonyl derivatives, results in ubiquitination and subsequent proteasomal degradation (Iwai et al., 1998). While oxidation of amino acids is primarily mediated by the hydroxyl radical, the superoxide anion also exhibits considerable protein reactivity; O$_2^-$ reacts with proteins that contain transition metal-prosthetic groups, such as haem moieties or iron-sulphur clusters (Keyer and Imlay, 1996; Prasad et al., 1989).

In addition to causing severe DNA and protein damage, ROS can also induce membrane lipid damage via lipid peroxidation. The cumulative effects of lipid peroxidation have been implicated in several pathological conditions including atherosclerosis, haemolytic anaemias and ischaemia reperfusion injuries (Davies, 1995; Steinbrecher et al., 1990). ROS-induced lipid peroxidation proceeds via the abstraction of hydrogen atoms from methylene groups separating unsaturated fatty acid double bonds (Glende and Recknagel, 1994; Dix and Aikens, 1993). The ROS O$_2^-$, in equilibrium with HO$_2^-$ (perhydroxyl radical), HO$_2^-$ alone, HO', O, and H$_2$O$_2$ in the presence of O$_2^-$, and
trace quantities of transition metals, are capable of initial H atom abstraction from methylene groups separating unsaturated fatty acid double bonds, thus initiating the lipid peroxidative chain reaction (Dix and Aikens, 1993). Initiation of the lipid peroxidative chain results in the generation of carbon centered radicals (L'), with the concomitant formation of conjugated dienes. Propagation of the lipid peroxidative chain proceeds via radical coupling, whereby L' reacts with O₂ resulting in the formation of lipid peroxyl radicals (LOO'). Lipid peroxyl radicals can then abstract further bis-allylic H atoms from adjacent unsaturated fatty acids in a reaction classified as an atom transfer reaction (Porter et al., 1995; Dix and Aikens, 1993). Propagation is cycled through rounds of LOO' abstraction of bis-allylic H atoms in the presence of O₂ to generate new LOO', which ultimately results in the conversion of lipids to lipid hydroperoxides (LOOH) (Dix and Aikens, 1993). LOOHs are subsequently degraded to a variety of compounds including alkanals, alkenals, hydroxyalkenals, ketones, aldehydes and alkanes (Coudray et al., 1995). Termination of lipid peroxidative chains occurs via the Russel mechanism, whereby two LOO's react to form a nonradical product, or where one LOO' reacts with another terminating radical, or free radical inhibitor, to generate self-quenching, nonpropagating radical species (Dix and Aikens, 1993).

The generation of lipid hydroperoxides within the hydrophobic core of membranes results in several deleterious biophysical effects. Principally, oxidized lipids alter the packing behaviour of membrane phospholipids. The accumulation of polar residues within the membrane induces a repulsive effect between adjacent oxidised fatty acyl chains, increasing their spatial separation (Van Ginkel and Sevanian, 1994).
hydroperoxide groups and short-chain oxidation products into membrane phospholipids, results in deterioration of membrane integrity, with the subsequent loss of membrane function (Van Ginkel and Sevanian, 1994; Mehlhorn, 1986).

Biological systems have evolved numerous antioxidant activities to sense, protect, and repair ROS-mediated damage to cellular constituents. The molecular defences against ROS in the model organism *Saccharomyces cerevisiae* have been well characterised (Santoro and Thiele, 1997; Moradas-Ferreira *et al.*, 1996; Jamieson 1995). Yeast antioxidant activities include enzymatic systems, non-enzymatic systems comprising low molecular weight antioxidants, heavy metal-chelators, and damage repair systems (Santoro and Thiele, 1997; Moradas-Ferreira *et al.*, 1996; Jamieson, 1995). Furthermore, *S. cerevisiae* possesses distinct protective oxidative stress responses to both hydrogen peroxide, and superoxide-generating compounds such as menadione (Jamieson, 1992).

The yeast enzymatic oxidative stress defence system consists primarily of cytosolic copper/zinc-containing superoxide dismutase (*SOD1*), mitochondrial manganese-containing superoxide dismutase (*SOD2*), peroxisomal catalase A (*CTA1*), cytosolic catalase T (*CTT1*), mitochondrial cytochrome-c peroxidase (*CCP1*), and thiol-specific antioxidant (*TSA*) (Santoro and Thiele, 1997; Moradas-Ferreira *et al.*, 1996; Jamieson, 1995). Superoxide dismutases (SODs) catalyse the disproportionation of the superoxide anion to hydrogen peroxide and dioxygen (Chang *et al.*, 1991; Gralla and Valentine, 1991). Exposure of yeast to the superoxide anion-generating compounds menadione and paraquat has been reported to result in increased *SOD1* and *SOD2* mRNA
transcript levels (Galiziano and Labbe-Rois, 1993). Furthermore, mutations in either SOD1 or SOD2 results in elevated sensitivity to oxygen and free-radical generating compounds. In addition, sod1Δ mutants exhibit a variety of abnormal phenotypes, including auxotrophies for the amino acids lysine, cysteine and methionine (Liu and Culotta, 1994; Gralla and Valentine, 1991). However, the oxygen sensitivity, and lysine and methionine auxotrophies of sod1Δ sod2Δ double mutants can be suppressed by mutations in either the BSD1 or BSD2 genes (Liu and Culotta, 1994). Significantly, both the BSD1 and BSD2 genes encode proteins involved in metal ion-homeostasis. The BSD1 gene is identical to the PMRI gene, which encodes a golgi-located P-type ATPase homologue (Rudolph et al., 1989). It has been proposed that BSD1/PMRI functions in maintaining low cytosolic levels of calcium and manganese. Thus, it is thought that pmr1 mutants suppress oxygen toxicity in sod1 mutants as a result of the hyper-accumulation of manganese ions (Lapinskas et al., 1995). The BSD2 encoded transmembrane protein is thought to play a role in the intracellular transport and sequestration of copper and cadmium, into cellular compartments such as the vacuole and golgi apparatus (Liu and Culotta, 1994). In addition to suppressing oxygen toxicity in sod1 mutants, bsd2 mutants display increased resistance to, and accumulation of, both cadmium and copper. Both manganese and copper have known SOD mimetic activities. Therefore it is thought that alterations in intracellular levels of both copper and manganese may generate bioavailable metal complexes which can serve as SOD mimetics. The interactions of SOD1, SOD2, BSD1, BSD2 and PMRI further highlight the important biological relationship between metal ion homeostasis and oxidative stress in S. cerevisiae.
Peroxisomal catalase A, cytosolic catalase T, and glutathione peroxidase catalyse the dismutation of hydrogen peroxide to water and molecular oxygen (Lapinskas et al., 1993; Cohen et al., 1985). Peroxisomal catalase A (CTA) serves primarily to remove hydrogen peroxide formed from the β-oxidation of high-molecular weight fatty acids. While the exact function of cytosolic catalase T (CTT) is as yet unknown, CTT is important for survival under conditions of severe osmotic stress (Schüller et al., 1994). Under conditions of osmotic stress HOG (high osmolarity glycerol response)-dependent CTT transcription is mediated via the stress response element (STRE) (Schüller et al., 1994). Post-translational modification of an STRE-binding transcription factor, mediated by the MAP kinase products of the HOG1 and PBS2 genes, most likely facilitates increased CTT mRNA transcription (Schüller et al., 1994). Expression of both CTT and CTA is induced by oxygen via positive control by heme (Hortner et al., 1982). Regulation of CTT and CTA gene expression by heme is mediated by the transcriptional activator Hap1p (Ruis and Hamilton, 1992). CTT transcription is also induced by Cu^{2+} via the MAC1 transcription factor (Jungmann et al., 1993b; Lapinskas et al., 1993; Thiele, 1992). Acatalasaemic double mutants (cttlΔ cta1Δ) have similar growth rates, and comparable susceptibility to hydrogen peroxide in the exponential growth phase, to wild-type cells under non-oxidative stress conditions. However, stationary phase acatalasaemic mutants are much more sensitive to hydrogen peroxide than wild-type cells, indicating that catalase plays an important role in the yeast hydrogen peroxide-adaptive stress response (Izawa et al., 1996).

The product of the TSA4 gene, thiol-specific antioxidant, is a cytosolic enzyme that protects cellular components against oxidation systems in which a thiol functions as a
reducing equivalent, e.g. di-thiothreitol (DTT)/Fe$^{3+}$/O$_2$ (Kim et al., 1988). It is thought that Tsap protects cells from oxidative damage by catalysing the removal of thiyl radicals (Yim et al., 1994). Exposure to 100% O$_2$ or Fe$^{3+}$ results in an increase of Tsap levels. Furthermore, tsa mutants grow more slowly under aerobic conditions, and in the presence of peroxides and paraquat, indicating that Tsap has an important physiological antioxidant role (Yim et al., 1994; Chae et al., 1993).

The yeast non-enzymatic defence system consists primarily of metallothionein (CUP1), glutathione (GSH1) and thioredoxin (TRX1 and TRX2). Metallothioneins are highly conserved, small cysteine-rich metal binding proteins that are essential for the detoxification of metals. The yeast CUP1 metallothionein gene is transcriptionally activated by copper, superoxide anion-generating chemicals, particularly menadione, and by growth on non-fermentable carbon sources (Liu and Thiele, 1996). Over-expression of CUP1 can suppress the cysteine and methionine auxotrophies of sod1A mutants, and can also restore the growth of sod1Asod2A double mutants on non-fermentable carbon sources (Tamai et al., 1993). Copper activates CUP1 via the copper metalloregulatory transcription factor ACE1. However, induction of CUP1 by menadione requires the yeast heat shock transcription factor (Hsf1p) which acts at the CUP1 promoter heat shock element (Liu and Thiele, 1996). CUP1 transcriptional activation is not observed in the presence of hydrogen peroxide, suggesting that CUP1 transcriptional activation is not a consequence of ROS exposure in general.

The tripeptide γ-L-glutamyl-L-cystinylglycine (glutathione) is one of the major and most abundant antioxidant molecules of S. cerevisiae. Glutathione possesses a redox-active sulphydryl group which reacts with oxidants to produce oxidised glutathione.
(GSSG) (Stephen and Jamieson, 1996). The GSHI gene product \( \gamma \)-glutamylcysteine synthetase catalyses the first and rate-limiting step in the biosynthesis of glutathione. GSHI is more strongly induced by the superoxide anion generating compounds menadione and plumbagin than by hydrogen peroxide (Stephen and Jamieson, 1996). GSHI transcription is controlled by the transcription factor encoded by the YAP1 gene (Stephen et al., 1995). The YAP1 and YAP2 gene products are leucine zipper-containing transcription factors similar to the c-Jun mammalian transcriptional activators. The expression of numerous oxidative stress response genes in \emph{S. cerevisiae}, including GSHI, GLR1 (glutathione reductase), TRX2 (thioredoxin), TPS2 (trehalose-6-phosphate phosphatase), and YCF1 (Mg-ATP-dependent transporter of glutathione-conjugated moieties) has been shown to be Yap1p-dependent. High level expression of Yap1p and Yap2p confers resistance to alkylating agents, cycloheximide, iron chelators and zinc (Stephen et al., 1995; Hussain and Lenard, 1991). While menadione-mediated induction of GSHI is only slightly reduced in yap1A mutants, yap1A and yap2A mutants are considerably more sensitive to hydrogen peroxide, indicating that YAPI and YAP2 are critical for the hydrogen peroxide-adaptive stress response.

Thioredoxin is a dithiol, sulphhydryl-rich protein (two redox-active cysteines in its active site) with antioxidant activity. \emph{Saccharomyces cerevisiae} contains two genes encoding thioredoxins, \emph{TRX1} and \emph{TRX2} (Gan, 1991). The precise molecular antioxidant mechanism of thioredoxin is unknown, however \emph{TRX2} is induced by hydrogen peroxide but not by menadione. In addition, \emph{trx1Atrx2A} mutants cannot grow in the absence of methionine or cysteine, and contain elevated levels of GSSG (Muller, 1991; Muller 1996). As stated previously, \emph{TRX2} is transcriptionally activated by the Yap1p.
transcription factor (Kuge and Jones, 1994). Deletion of both \textit{TRX1} and \textit{TRX2} is a non-lethal event in \textit{S. cerevisiae}, however \textit{trx2A} mutants are hypersensitive to hydrogen peroxide and the alkyl hydroperoxide, \textit{tBOOH}, indicating a role for \textit{TRX2} in hydrogen peroxide dismutation (Kuge and Jones, 1994).

Direct repair of ROS-induced AP sites and 3'-damage in yeast is mediated primarily by the class II AP endonuclease/3'-diesterase, encoded by the \textit{APNI} gene. Yeast strains containing mutations in the \textit{APNI} gene display an elevated spontaneous mutation rate (Ramotar et al., 1991). In addition, DNA repair enzymes encoded by genes belonging to the \textit{RAD52} epistasis group repair double-strand breaks by homologous recombination (Guzder et al., 1994; Klein, 1988). Re-reduction of oxidised protein sulphhydril groups is thought to be mediated by enzymes such as disulphide reductase, glutaredoxin (\textit{GRXI} and \textit{GRX2}) (Luikenhuis et al., 1998), and methionine sulfoxide reductase (\textit{MSRA}) (Moskovitz et al., 1997). Both \textit{GRXI} and \textit{GRX2} have been found to contain putative STREs, and their expression is increased in response to hydrogen peroxide, menadione, heat and osmotic shock (Luikenhuis et al., 1998). In addition, oxidatively modified soluble proteins in \textit{S. cerevisiae} are recognised by ubiquitin-conjugating enzymes, encoded by \textit{UBC1}, \textit{UBC4}, \textit{UBC5} and \textit{UBC7}, and are thus tagged for degradation by the proteasome (Jungmann et al., 1993a). Phospholipase enzymes are responsible for the hydrolysis of lipid hydroperoxides within lipid bilayers, while lipid hydroperoxides that have been released into the cytosol are thought to be cleaved by glutathione peroxidase (Tran et al., 1993). Indeed, a membrane-bound glutathione peroxidase from the yeast \textit{Hansenula mrakii} has been purified, and displayed high
substrate specificity for the lipid peroxidation by-products oleic acid hydroperoxide and phosphatidylcholine hydroperoxide in vitro (Tran et al., 1993).

The lipid composition of the plasma membrane of *S. cerevisiae* is complex and tightly regulated. The major lipid classes of the yeast plasma membrane are glycerophospholipids, sphingolipids, and sterols. Glycerophospholipids consist of two fatty acyl chains ester-linked to glycerol-3-phosphate; various substituents such as choline [in phosphatidylcholine (PC)], ethanolamine (in PE), serine (in PS), myoinositol (in PI), and glycerol [in phosphatidylglycerol (PG)], can be linked to the phosphoryl group. The phospholipid component of the inner leaflet of the *S. cerevisiae* plasma membrane is enriched in phosphatidylethanolamine (PE), phosphatidylinositol (PI) and phosphatidylserine (PS). Diphosphatidyl glycerol or cardiolipin, the dimeric form of PG, is also present in yeast cells. Sphingolipids have a ceramide backbone which is composed of a long-chain base phytosphingosine that is N-acylated with a hydroxy C_{26} fatty acid. *Saccharomyces cerevisiae* contains only three major sphingolipids: inositol phosphate ceramide, mannosyl-inositolphosphate-ceramide, and mannosyl-diinositolphosphate-ceramide (Van der Rest et al., 1995). Sterols are compact rigid hydrophobic molecules with a polar hydroxyl group. In contrast to higher eukaryotes, in which cholesterol is the most abundant sterol, the yeast plasma membrane contains mainly ergosterol and minor amounts of zymosterol (Paltauf et al., 1992; Zinser et al., 1991; Rattray et al., 1975). Between 70-80% of the fatty acids in membrane phospholipids of *S. cerevisiae* consist of the monounsaturates palmitoleic acid (16:1) and oleic acid (18:1). Δ7-cis-Heptenoic acid (14:1) is also present (particularly in PS and
PE), but in much smaller quantities (Wagner and Paltauf, 1994). The remaining fatty acids are saturated and consist of palmitic acid (16:0) and lesser amounts of stearic acid (18:0) and myristic acid (14:0). Yeast Δ-9 fatty acid desaturase (encoded by the OLE1 gene) catalyses the formation of the initial double bond between the 9th and 10th carbons of both palmitoyl (16:0) and stearoyl (18:0) coenzyme A (CoA) substrates to form 16:1 and 18:1. *Saccharomyces cerevisiae* however does not have the ability to synthesise polyunsaturated fatty acids (PUFAs) (Van der Westhuizen *et al.*, 1994; Wagner and Paltauf, 1994).

 Whereas the molecular basis of both constitutive and inducible elements of the yeast antioxidant and heavy metal defence systems have been well characterised (Li *et al.*, 1997; Stephen and Jamieson, 1996; Liu and Culotta, 1994; Chang *et al.*, 1991), no studies have sought to investigate membrane compositional modifications in response to oxidative stress, or how membrane composition influences susceptibility to heavy metal toxicity. This is surprising in view of the large variability in microbial membrane composition which exists. In order to maintain a relatively constant membrane environment, and thus maintain correct membrane functionality, microorganisms rapidly alter their lipid metabolism and membrane composition during changes in external physico-chemical parameters, e.g. ethanol concentration, temperature, pressure and salinity (Schulthess and Hauser, 1995; Alexandre *et al.*, 1994; Hosono, 1992; Hazel and Williams, 1990; Suutari *et al.*, 1990). Genetically determined intrinsic inter-species differences in microbial membrane lipid composition are also widespread (Van der Westhuizen *et al.*, 1994; Vestal and White, 1989). The biochemical effects of altered yeast plasma membrane phospholipid composition have been investigated. For example,
increased affinity constants for amino acid transport have been observed for yeast choline and ethanolamine auxotrophs, with plasma membranes enriched with PC or PE, respectively (Triveldi et al., 1982). PI-enrichment of the yeast plasma membrane has also been associated with enhanced H⁺-ATPase activity (Triveldi et al., 1987). Also, the effects of specific enrichment of the plasma membrane of S. cerevisiae with exogenously-supplied PUFAs and sterols on growth dynamics, and amino acid and pyrimidine transport have been described (Parks et al., 1995; Bossie and Martin, 1989; Keenan et al., 1982; Watson and Rose, 1980). Hoptroff et al. (1997) recently demonstrated an elevated rate of Cs⁺-accumulation in S. cerevisiae exogenously enriched with the di-unsaturated fatty acid linoleate. Thus, maintenance of membrane homeoviscosity appears to be critical for the regulation of plasma membrane protein activity. In addition, the effects of modification of membrane fatty acid composition on the heat shock and H₂O₂-stress responses have also been examined (Polla et al., 1997; Carratu et al., 1996; Steels et al., 1994). Incorporation of an exogenous PUFA into membrane phospholipids of anaerobically grown S. cerevisiae resulted in decreased intrinsic and induced thermotolerance, and an increase in sensitivity towards H₂O₂ (Steels et al., 1994). Furthermore, incorporation of a saturated fatty acid into membrane phospholipids of a temperature-sensitive H. capsulatum strain (up-regulated OLE1 promoter) resulted in an increase in heat shock protein (hsp82 and hsp70) mRNA transcription at 37 °C (Carratu et al., 1996). Here, it was postulated that a primary membrane-associated heat shock sensor (which initiates the signal cascade leading to phosphorylation of heat shock factor (HSF)) is regulated in part through active modification of membrane lipid composition. However, while Polla et al. (1997) also
demonstrated the increased H$_2$O$_2$-sensitivity of PUFA-enriched human premonocytic
cells, no discernible differences in HSP synthesis were observed following heat shock.

Microorganisms, particularly yeast and fungi, are highly amenable to
manipulation of membrane lipid composition, and thus serve as ideal experimental
organisms for modelling membrane lipid-dependent functions in higher organisms.
Enrichment of the yeast plasma membrane with PUFAs has no effect on growth rate, thus
circumventing the major problem associated with blunt-tools such as temperature-
dependent manipulations of membrane lipid composition (Guerzoni et al., 1993). PUFA-
enrichment in S. cerevisiae is known to result in sharp repression of OLE1 transcription,
resulting in decreased synthesis of 16:1 and 18:1 (Bossie and Martin, 1989).

Incorporation of long chain PUFAs into membrane phospholipids alters the
packing behaviour of phospholipid moieties, and ultimately leads to a decline in
membrane stability (Vossen et al., 1995; De Vos et al., 1993). However, to date no
reports have examined how heavy metal-induced membrane permeabilisation is related to
initial membrane phospholipid composition. Furthermore, the susceptibility of
membrane phospholipids to ROS-induced lipid peroxidation is known to increase with the
degree of phospholipid fatty acyl chain unsaturation (Van Ginkel and Sevanian, 1994).
The latter point is of particular relevance to the present studies as heavy metals can
promote the generation of ROS through redox-cycling activity via the Fenton (2) and
Haber-Weiss (3) reactions, and by depletion of free radical scavengers (Figueiredo-
Pereira et al., 1998; Li et al., 1997; Hassoun and Stohs, 1996; Stohs and Bagchi, 1995).
However, the relationship between membrane fatty acid composition, ROS-mediated
cellular damage, and heavy metal toxicity towards microorganisms has yet to be elucidated.

Thus, the objectives of this research were to examine the effects of altered membrane fatty acid composition on the toxic interactions of heavy metals with the model organism *Saccharomyces cerevisiae*. 
CHAPTER 2

Investigation of the Relationship Between Cadmium Toxicity and
Degree of Plasma Membrane Fatty Acid Unsaturation in Saccharomyces
cerevisiae
2.1. Introduction

The toxicity of cadmium, along with several other heavy metals, towards microorganisms has attracted considerable research attention in recent years as a result of the continuing anthropogenic mobilisation of heavy metals in the environment. The principal anthropogenic point sources of cadmium in the environment include the electroplating, galvanising, zinc and lead mining, and smelting industries. In addition, nickel-cadmium alkaline batteries are extensively used throughout the aircraft industry and in numerous portable devices, e.g. cellular phones. Cadmium is a potent toxin to living cells, for which limited biological function has been defined to date (Price and Morel, 1990). Cadmium is a nephrotoxin and potential carcinogen, with a long biological half-life in humans (Dally and Hartwig, 1997; Brennan and Schiestl, 1996).

Cadmium, along with Zn and Hg, is a d-block element that forms M$^{2+}$ ions of high charge density which can be strongly coordinated to lone pair donors. As a consequence, the toxic effects of these transition metal ions can be attributed partly to the multiplicity of coordination complexes and clusters that they can form (Mackay and Mackay, 1991; Butler and Harrod, 1989). Most heavy metal ions have a strong affinity for ligands such as phosphates, purines, pteridines, porphyrins, and cysteiny1 and histidyl side chains of proteins, which are all abundant in microbial cells. Thus, microbial cells provide an environment in which heavy metals can exert deleterious effects through a variety of mechanisms (Gadd, 1993). Cadmium reacts with polythiol groups on cellular macromolecules, particularly protein sulphhydr1 groups, and can substitute for zinc in Zn-
containing enzymes, e.g. carboxypeptidases (Li et al., 1997; Price and Morel, 1990). In addition, cadmium resistance in yeast can be mediated by ubiquitin-dependent proteolysis (Jungmann et al., 1993a), suggesting that cadmium exerts its toxic effects in part through the formation of abnormal proteins. Furthermore, as stated previously, cadmium, along with other transition metals, causes perturbations in the integrity of certain structural components, e.g. cellular and organellar membranes (Abemann et al., 1996; Kessels et al., 1985). Plasma-membrane permeabilisation with associated cellular K⁺ efflux has been reported in a variety of microorganisms following cadmium exposure, and is considered a major mechanism of cadmium toxicity in the yeast Saccharomyces cerevisiae (Aβmann et al., 1996; Kessels et al., 1985; Gadd and Mowll, 1983).

Recent studies with higher organisms have indicated that susceptibility to heavy metal toxicity may be partly dependent on the lipid composition and physical properties of cellular membranes (Vossen et al., 1995; De Vos et al., 1993). These observations are of considerable pertinence to metal-microbe interactions as many microorganisms exhibit large fatty acid compositional changes during environmental acclimation (Hazel and Williams, 1990). Intrinsic inter-species differences in microbial fatty acid profiles are also widespread (Van der Westhuizen et al., 1994; Vestal and White, 1989). Many membrane-dependent functions are sensitive to such alterations (Avery et al., 1995a; Hazel and Williams, 1990; Murata, 1989). However, there is currently a notable lack of information relating to how this variation might influence microbial susceptibility to plasma membrane-targeting toxic agents.

Recently, a link between plasma membrane composition and metal sensitivity was reported in yeast. It was found that copper-induced plasma membrane disruption in S.
*cerevisiae* was markedly accelerated in cells enriched with the di-unsaturated fatty acid linoleate (18:2) (Avery et al., 1996). Because polyunsaturated fatty acids are particularly susceptible to free radical attack, these results were also consistent with a role of oxidative stress in copper-toxicity towards yeast, as has been indicated for higher organisms (Stohs and Bagchi, 1995; Shinar et al., 1989). A role of free radicals has been further supported by studies in other laboratories which have demonstrated copper hypersensitivity in *S. cerevisiae* strains which carry mutations in genes encoding components of the oxidative stress response (Culotta et al., 1995; Tamai et al., 1993; Greco et al., 1990). Brennan and Schiestl (1996) have recently demonstrated the elevated Cd$^{2+}$-sensitivity of *sod1Δ sod2Δ* and *gsh1Δ S. cerevisiae* strains. In addition, Cd$^{2+}$-induced toxicity and recombination were inhibited in the presence of the free radical scavenger N-acetyl cysteine (Brennan and Schiestl, 1996). In the present study we address the question of whether the nonredox-active nature of cadmium (unlike copper) therefore precludes a role of plasma membrane lipid composition in determining its toxicity to *S. cerevisiae*.

Interestingly, cadmium toxicity was also found to be highly dependent on plasma membrane composition and showed increases with the degree of fatty acid unsaturation; maximal Cd$^{2+}$ sensitivity was evident in cells enriched with the tri-unsaturated fatty acid, linolenate. The dependence of yeast sensitivity to metals on plasma membrane composition appears to be more widespread than originally anticipated.
2.2. Materials and Methods

2.2.1. Organism and culture conditions. *S. cerevisiae*NCYC 1383 (MATa, his3-Δ1, leu2-3, leu2-112, ura3-52, trpl-289) was routinely maintained on solid YEPD medium, comprising 2% (w/v) neutralised bacteriological peptone, 1% (w/v) yeast extract (Oxoid, U.K.), 2% (w/v) glucose, and 1.6% (w/v) agar (no. 3, technical, Oxoid, U.K.), and solid YNB selective medium comprising 0.67% (w/v) yeast nitrogen base without amino acids, 1.6% (w/v) technical agar, 60 μg ml⁻¹ leucine, 40 μg ml⁻¹ tryptophan, and 20 μg ml⁻¹ histidine and uracil. For experimental purposes, *S. cerevisiae* was grown in 100 ml of YEPD broth, of the same composition as solid YEPD, but lacking agar and supplemented with 1% (w/v) of the non-ionic surfactant tergitol (NP-40, Sigma, U.K.), in 250 ml Erlenmeyer flasks (tergitol supplementation, to help solubilise fatty acids, was found to have no discernible effect on the fatty acid composition of *S. cerevisiae*). Experimental flasks were inoculated to OD₅₅₀~0.1 from 48 h starter cultures, and incubated at 25 °C with orbital shaking at 120 rev. min⁻¹. Where indicated, linoleate or linolenate (Sigma, U.K.) (final concentrations, 1 mM) were added from filter-sterilised 20 mM stock solutions, solubilised with 5% (w/v) tergitol. When specified, Cd(NO₃)₂ was aseptically added from a stock solution, previously sterilised by autoclaving. Cell numbers were determined using a modified Fuchs-Rosenthal haemocytometer slide after appropriate dilution with distilled deionised water: more than 400 cells were counted in each sample.
2.2.2. Preparation of cell homogenates. Cells were harvested by centrifugation at 1,500 g for 5 min and washed twice with distilled deionised water at 4 °C to restrict further lipid metabolism. Cells were disrupted by vigorous shaking with 0.5 mm-diameter glass beads (Sigma, U.K.) for 15 min at 4 °C using a Mickel homogeniser (Mickle Laboratories, Guildford, U.K.). The beads were removed by vacuum filtration through a glass-sintered filter and washed with distilled deionised water.

2.2.3. Plasma membrane isolation. It was found that the method used for isolation of plasma membranes from unsupplemented and linoleate-supplemented cells did not yield membranes of sufficient purity from linolenate-enriched cells [possibly for the reasons suggested by Alterthum and Rose (1973)], thus, two methods for preparation of plasma membranes were used. Unsupplemented and linoleate-enriched plasma membranes were purified from whole-cell homogenates by differential and density-gradient centrifugation, using a modified method of Serrano (1988). Late-exponential/early-stationary phase cells were harvested by centrifugation at 1,500 g for 5 min and washed twice with distilled deionised water at 4 °C. The pellet was resuspended in 5 mM Tris.Cl (pH 7.5), 700 mM sorbitol to give O.D.₈₀₀ ~ 6.0. The suspension was then mixed with ¼ volume of 300 units ml⁻¹ lyticase (Sigma, U.K.), prepared in the above buffer, and DTT was added to a final concentration of 6.5 mM. The suspension was then incubated for 1-2 h at 30 °C, with occasional shaking to aid cell wall digestion. Spheroplast formation was followed by examining for birefringence using phase-contrast microscopy (Rose and Veazey, 1992). Approximately 80-90% spheroplast yield was usually observed after 2 h incubation. Spheroplasts were harvested by centrifugation at 3,000 g for 10 min and
washed twice with lyticase-free buffer. The spheroplast pellet was finally resuspended in 10-15 ml of 15 mM Mes-Tris (pH 6.5), 500 mM sorbitol, 100 mM glucose, and incubated at 30 °C for 10 min to activate the plasma membrane H⁺-ATPase. Osmotic lysis was achieved by dilution with two volumes of 25 mM Mes-Tris (pH 6.5), 5 mM EDTA, 0.2% (w/v) BSA, 0.2% (w/v) casein hydrolysate, 1 mM DTT and 1 mM PMSF, at 4 °C. All subsequent steps were carried out at 4 °C. To aid spheroplast homogenisation the suspension was subjected to gentle mechanical disruption with five slow strokes of a hand-held homogeniser (BDH, U.K., 0.15-0.25 mm clearance). The suspension was centrifuged at 700 g for 10 min to remove cell-wall debris. The resulting supernatant was then centrifuged at 35,000 g for 15 min, yielding a total-membrane pellet. The pellet was resuspended in 20% (v/v) glycerol, 10 mM PMSF. The resuspended total-membrane fraction was applied to a discontinuous sucrose gradient comprising 2 parts 43% (w/w) sucrose and 1 part 53% (w/w) sucrose. After centrifugation for 6 h at 120,000 g in a Beckman SW40Ti swinging bucket rotor, the plasma membrane-enriched fraction was collected at the 43/53 interface with a Pasteur pipette (narrow tip). The plasma membrane-enriched fraction was diluted with 4 volumes of distilled deionised water and centrifuged at 80,000 g for 20 min. After centrifugation, the final pellet was resuspended in 20% (v/v) glycerol, 10 mM PMSF, flash frozen in liquid nitrogen and stored at -70 °C until used.

Isolation of linolenate-enriched plasma membranes was achieved using a method of two-phase partitioning, adapted from Menendez et al. (1995). Purification of the total-membrane pellet was as described above. The total-membrane pellet was resuspended in buffer (5 mM KH₂PO₄ (pH 7.8), 330 mM sucrose, 1 mM DTT) to give a total weight of
4.0 g. The resuspended total-membrane fraction (4.0 g) was then added to a 12.0 g dextran/polyethylene glycol polymer mixture, to generate a 16.0 g two-phase system of final composition 5.7% (w/w) dextran T500 (Pharmacia Biotech, U.K.), 5.7% (w/w) polyethylene glycol 3350 (Sigma, U.K.), 5 mM KH₂PO₄ (pH 7.8), 330 mM sucrose, 1 mM EDTA, 1 mM DTT. The two-phase system was mixed thoroughly, transferred to a 25 ml separating funnel and allowed to settle for 2 h at 4 °C, until a sharp interface separating the two phases was clearly observable. The upper phase was removed using a Pasteur pipette (wide tip). The remaining lower phase, with interface, was then diluted with nine volumes of 15 mM Mes-Tris (pH 6.5), 330 mM sucrose, 1 mM DTT, and centrifuged at 60,000 g for 30 min. After centrifugation, pellets were resuspended in fresh buffer, divided into 100 μl aliquots, flash frozen in liquid nitrogen and stored at -70 °C until used.

2.2.4. Enzyme assays. The purity of plasma membranes and whole-cell homogenates in fractions obtained after sucrose density-gradient centrifugation or phase-partitioning were determined by assaying for vanadate-sensitive (plasma membrane) and azide-sensitive (mitochondrial membrane) ATPase activities, using the methods described by Widell and Larsson (1990). Optimal detergent activation of plasma membrane ATPase, after sucrose density-gradient centrifugation, was achieved according to the method of Serrano (1988). Protein determination was by the method of Bradford (1976), using BSA (Sigma, U.K.) as a standard.
2.2.5. **Lipid extraction and fatty acid analysis.** Lipids were extracted from whole-cell homogenates or membrane fractions using the method of Bligh and Dyer (1959) as modified by Griffiths and Harwood (1991), with the following modifications. Lipids from whole-cell homogenates or membrane fractions were extracted in to a methanol/chloroform/water (2:1:0.8) monophasic solvent mixture. Chloroform and distilled deionised water were added, resulting in the formation of a biphasic solvent mixture. After centrifugation at 1,000 g for 5 min the upper (aqueous) phase was removed and the lower (organic) phase, containing the lipids, was transferred to a clean glass tube. The chloroform was evaporated under N₂, and for fatty acid analysis methyl esters were generated by acid-catalysed esterification [2.5% (v/v) H₂SO₄ in dry methanol] at 70 °C for 2 h. Methyl esters were extracted with redistilled petroleum spirit (b.p. 60-80 °C) and subsequently analysed by gas-liquid chromatography. Pentadecanoate was used as an internal standard. Separations were routinely achieved using 10% SP-2330 on 100/120 Chromosorb-WAW (Supelco, Saffron Waldon, Essex, U.K.) packed into a stainless steel column [1.8 m x 0.3 mm (outer diameter)]. Fatty acids were identified by comparison with authentic standards.

2.2.6. **Potassium efflux.** Cells from the late-exponential/early-stationary phase were harvested by centrifugation at 1,500 g for 5 min and washed twice with distilled deionised water. Washed cells were suspended to a density of approximately 5 x 10⁷ ml⁻¹ in 40 ml of 20 mM Mes buffer (pH 6.0), and the suspension agitated by magnetic stirring. After 2 min, glucose was added to a final concentration of 1% (w/v). After a further 10 min equilibration, Cd(NO₃)₂ was added to the desired concentration. Extracellular K⁺ was
measured continuously using an EE-K K\(^+\)-selective electrode (EDT Instruments, Dover, U.K.) coupled to a Jenway 3045 ion-analyser. Cd\(^{2+}\)-induced K\(^+\) efflux measurements were corrected for small non-cadmium-induced changes in extracellular K\(^+\) [which never exceeded 0.1 nmol K\(^+\) (10\(^6\) cells\(^{-1}\)) over the time courses examined] that occurred in control suspensions in the absence of Cd(NO\(_3\))\(_2\).

2.2.7. Cell viability. Cell viability was determined as the ability to produce colony-forming units. At specified intervals after the addition of Cd(NO\(_3\))\(_2\) to cell suspensions (see above), aliquots were removed and, after appropriate dilution with sterile, distilled deionised water, plated on yeast-selective OGYE agar (Lab M, U.K.), containing 0.1% (w/v) oxytetracycline (Sigma, U.K.). Colonies were enumerated after 4 d incubation at 25 °C. Colony counts did not change with prolonged incubation up to 20 d.

2.2.8. Cadmium uptake. Late exponential/early stationary phase cells were harvested by centrifugation at 1,500 g for 5 min and washed twice with 20 mM Mes (pH 6.0). The final pellet was resuspended in 150 ml of the buffer to give an approximate cell density of 5 x 10\(^7\) ml\(^{-1}\). 50 ml-portions of this suspension were incubated with shaking at 120 rev. min\(^{-1}\) in 100 ml Erlenmeyer flasks at room temperature. After 10 min, glucose was added to a final concentration of 1% (w/v). After a further 10 min equilibration, Cd(NO\(_3\))\(_2\) was added. At specified intervals (up to 2 h), samples were removed, harvested by centrifugation at 1,500 g for 5 min and washed twice with distilled deionised water at 4 °C. Final cell pellets were digested by addition of 0.5 ml of 6 M HNO\(_3\) and immersion of tubes in a water bath at 100 °C for 2 h. Cell debris was removed by centrifugation and,
after appropriate dilution with deionised water, the cadmium concentrations of the supernatants determined using a Baird alpha-2 atomic absorption spectrophotometer, with reference to appropriate standard solutions.
2.3. Results

2.3.1. Influence of linoleate-supplementation on growth of *S. cerevisiae* NCYC 1383 in the absence and presence of Cd(NO$_3$)$_2$. Linoleate (1 mM) supplementation was found to have no discernible effect on the growth of *S. cerevisiae* in YEPD medium. Cell doubling times during the exponential phase were approximately 2.5 h, and final cell yields after 24 h approximately $1.5 \times 10^8$ ml$^{-1}$, in both unsupplemented and linoleate (18:2)-supplemented media (Fig. 2.1). Growth in the presence of 80 μM Cd(NO$_3$)$_2$ was associated with extended lag phases, longer doubling times and reduced final cell yields. However, these effects were considerably more marked for 18:2-supplemented cultures. For example, cell yields after 24 h growth in the presence of Cd(NO$_3$)$_2$ were approximately $4 \times 10^7$ ml$^{-1}$ and $1.8 \times 10^7$ ml$^{-1}$ in unsupplemented and 18:2-supplemented cultures, respectively; these represented respective 74% and 88% decreases compared to the cell yield of cultures incubated without Cd(NO$_3$)$_2$ (Fig. 2.1).

2.3.2. Fatty acid composition of *S. cerevisiae* NCYC 1383 during growth in unsupplemented, linoleate- and linolenate-supplemented media. In order to confirm that growth of *S. cerevisiae* in polyunsaturated fatty acid (PUFA)-supplemented medium resulted in incorporation of exogenous fatty acids into total cellular lipids, lipid extracts from cells grown in unsupplemented and PUFA-supplemented media were analysed. *Saccharomyces cerevisiae* readily incorporated exogenous PUFAs from its growth medium (Fig. 2.2; Table 2.1).
Fig. 2.1. Influence of linoleate-supplementation on growth of *Saccharomyces cerevisiae* NCYC 1383 in the absence and presence of cadmium. *S. cerevisiae* from a 48 h starter culture was inoculated to *A*$_{550}$ ~ 0.1 in unsupplemented (○,●) and linoleate-supplemented (□,■) medium. Cell numbers were determined during growth in the absence (○,□) and presence (●,■) of 80 μM Cd(NO$_3$)$_2$. Points represent means from three replicate counts (more than 400 cells total); Standard error of the mean (SEM) values are smaller than the dimensions of the symbols.
Fig. 2.2. Gas-chromatograph trace of the fatty acid profiles of *S. cerevisiae* NCYC 1383 previously grown in unsupplemented (a) and linoleate-supplemented (b) medium.
In agreement with other reports (Bossie and Martin, 1989; Stukey et al., 1989), analysis of cell lipid extracts from *S. cerevisiae* grown in unsupplemented medium revealed only trace quantities of 18:2, while 18:3 was not detected. However, cells grown in the presence of 18:2 or 18:3 readily incorporated high levels of these fatty acids. For example, after 16 h growth (late-exponential/early-stationary phase), when incorporation of exogenous fatty acids was at a maximum, 18:2 constituted approximately 68% (Fig. 2.2) and 18:3 approximately 69% of total fatty acids in cells from 18:2- and 18:3-supplemented cultures, respectively (Table 2.1). Increases in PUFA content, as a proportion of total cellular fatty acids, during growth in supplemented media were contemporaneous with decreases in the monounsaturated fatty acids palmitoleate (16:1) and oleate (18:1); the fatty acid composition at 0 h (not shown) was similar to that of cells in unsupplemented medium at 16 h. The unsaturation index (average number of double bonds per fatty acid) was found to increase during exponential growth, and was maximal during the late-exponential/early-stationary phase (Fig. 2.3; Table 2.1). At any time, values for cellular unsaturation index decreased in the order 18:3-supplemented > 18:2-supplemented > unsupplemented (Fig. 2.3; Table 2.1).

The fatty acid composition of *S. cerevisiae* grown in the presence of 80 μM Cd(NO₃)₂ showed differences to that of cells grown without Cd(NO₃)₂ (Table 2.2). Between 0 and 16 h, cells growing in the presence of Cd²⁺ generally displayed a lower content of PUFAs, as a proportion of total cellular fatty acids, than cells growing in the absence of Cd²⁺ after corresponding periods of growth. This was most evident as lower proportions of 18:2 and 18:3 during early growth in 18:2- and 18:3-supplemented
**Table 2.1.** Changes in cellular fatty acid composition during growth of *S. cerevisiae* NCYC 1383 in unsupplemented, linoleate- and linolenate-supplemented media

<table>
<thead>
<tr>
<th>Medium</th>
<th>Fatty acid</th>
<th>Age of culture (h)</th>
<th>4</th>
<th>8</th>
<th>12</th>
<th>16</th>
<th>20</th>
<th>24</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unsupplemented</td>
<td>16:0</td>
<td></td>
<td>29.1±0.3</td>
<td>25.5±1.6</td>
<td>24.1±0.3</td>
<td>20.2±0.7</td>
<td>17.1±0.1</td>
<td>17.0±0.3</td>
</tr>
<tr>
<td></td>
<td>16:1</td>
<td></td>
<td>38.2±0.2</td>
<td>46.0±6.6</td>
<td>51.7±0.2</td>
<td>55.7±1.4</td>
<td>57.4±0.1</td>
<td>54.4±1.1</td>
</tr>
<tr>
<td></td>
<td>18:0</td>
<td></td>
<td>15.9±0.1</td>
<td>10.7±3.9</td>
<td>4.6±0.1</td>
<td>4.2±0.9</td>
<td>3.1±0.1</td>
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<tr>
<td></td>
<td>18:1</td>
<td></td>
<td>16.8±0.7</td>
<td>17.8±1.2</td>
<td>19.5±0.2</td>
<td>19.9±0.3</td>
<td>22.4±0.1</td>
<td>24.5±tr.</td>
</tr>
<tr>
<td></td>
<td>18:2</td>
<td></td>
<td>tr.</td>
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<td>tr.</td>
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<td>tr.</td>
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<tr>
<td></td>
<td><em>U.I.</em></td>
<td></td>
<td>0.55</td>
<td>0.64</td>
<td>0.71</td>
<td>0.76</td>
<td>0.80</td>
<td>0.79</td>
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<tr>
<td>18:2-supplemented</td>
<td>16:0</td>
<td></td>
<td>21.9±0.1</td>
<td>20.5±0.9</td>
<td>20.6±0.1</td>
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<td>20.1±0.2</td>
<td>19.7±0.7</td>
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<tr>
<td></td>
<td>16:1</td>
<td></td>
<td>22.9±3.1</td>
<td>9.8±0.3</td>
<td>7.4±0.1</td>
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<td>6.3±tr.</td>
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<tr>
<td></td>
<td>18:0</td>
<td></td>
<td>9.4±0.7</td>
<td>6.0±tr.</td>
<td>5.0±0.1</td>
<td>4.6±tr.</td>
<td>4.6±tr.</td>
<td>5.5±0.2</td>
</tr>
<tr>
<td></td>
<td>18:1</td>
<td></td>
<td>10.7±0.3</td>
<td>5.0±tr.</td>
<td>3.2±0.1</td>
<td>2.4±tr.</td>
<td>2.8±0.1</td>
<td>3.4±0.3</td>
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<td></td>
<td>18:2</td>
<td></td>
<td>35.1±2.8</td>
<td>58.7±0.5</td>
<td>63.8±1.2</td>
<td>67.7±0.8</td>
<td>66.2±0.3</td>
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<tr>
<td></td>
<td><em>U.I.</em></td>
<td></td>
<td>1.04</td>
<td>1.32</td>
<td>1.38</td>
<td>1.44</td>
<td>1.42</td>
<td>1.38</td>
</tr>
<tr>
<td>18:3-supplemented</td>
<td>16:0</td>
<td></td>
<td>23.9±0.3</td>
<td>20.4±1.0</td>
<td>19.6±0.4</td>
<td>18.8±tr.</td>
<td>20.6±0.4</td>
<td>18.5±tr.</td>
</tr>
<tr>
<td></td>
<td>16:1</td>
<td></td>
<td>27.0±1.2</td>
<td>10.3±1.8</td>
<td>6.9±0.5</td>
<td>5.2±0.2</td>
<td>6.2±0.4</td>
<td>7.7±0.6</td>
</tr>
<tr>
<td></td>
<td>18:0</td>
<td></td>
<td>9.8±0.4</td>
<td>6.4±0.3</td>
<td>5.0±0.1</td>
<td>4.9±tr.</td>
<td>5.5±0.2</td>
<td>5.2±tr.</td>
</tr>
<tr>
<td></td>
<td>18:1</td>
<td></td>
<td>15.1±0.8</td>
<td>5.0±0.7</td>
<td>3.1±0.2</td>
<td>2.4±tr.</td>
<td>3.3±0.2</td>
<td>4.1±0.2</td>
</tr>
<tr>
<td></td>
<td>18:2</td>
<td></td>
<td>tr.</td>
<td>tr.</td>
<td>tr.</td>
<td>tr.</td>
<td>tr.</td>
<td>tr.</td>
</tr>
<tr>
<td></td>
<td>18:3</td>
<td></td>
<td>24.2±0.9</td>
<td>57.9±4.9</td>
<td>65.4±1.8</td>
<td>68.7±1.0</td>
<td>64.4±1.6</td>
<td>64.5±0.2</td>
</tr>
<tr>
<td></td>
<td><em>U.I.</em></td>
<td></td>
<td>1.15</td>
<td>1.89</td>
<td>2.06</td>
<td>2.14</td>
<td>2.03</td>
<td>2.05</td>
</tr>
</tbody>
</table>

Fatty acids are abbreviated with the first figure representing the number of carbon atoms in the fatty acyl chain and the second representing the number of double bonds. *U.I.* refers to the unsaturation index (average number of double bonds per fatty acid). Values for percentage fatty acid composition are means from two replicate determinations ± SD. (tr. < 0.1).
**Table 2.2.** Changes in cellular fatty acid composition during growth of *S. cerevisiae* NCYC 1383 in unsupplemented, linoleate- and linolenate-supplemented media in the presence of 80 μM Cd(NO₃)₂

<table>
<thead>
<tr>
<th>Medium</th>
<th>Fatty acid</th>
<th>Age of culture (h)</th>
<th>U.I.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>4</td>
<td>8</td>
</tr>
<tr>
<td><strong>Unsupplemented</strong></td>
<td>16:0</td>
<td>34.7±1.0</td>
<td>39.9±2.1</td>
</tr>
<tr>
<td></td>
<td>16:1</td>
<td>34.1± tr.</td>
<td>35.8±0.7</td>
</tr>
<tr>
<td></td>
<td>18:0</td>
<td>11.4±1.1</td>
<td>7.2±1.4</td>
</tr>
<tr>
<td></td>
<td>18:1</td>
<td>19.8±0.2</td>
<td>17.1±1.4</td>
</tr>
<tr>
<td></td>
<td>U.I.</td>
<td>0.54</td>
<td>0.53</td>
</tr>
<tr>
<td><strong>18:2-supplemented</strong></td>
<td>16:0</td>
<td>32.2±5.0</td>
<td>28.1±0.2</td>
</tr>
<tr>
<td></td>
<td>16:1</td>
<td>20.0± tr.</td>
<td>19.8±3.6</td>
</tr>
<tr>
<td></td>
<td>18:0</td>
<td>8.8±1.0</td>
<td>6.2±0.3</td>
</tr>
<tr>
<td></td>
<td>18:1</td>
<td>11.5±0.7</td>
<td>8.8±0.7</td>
</tr>
<tr>
<td></td>
<td>18:2</td>
<td>27.5±3.0</td>
<td>37.1±4.8</td>
</tr>
<tr>
<td></td>
<td>U.I.</td>
<td>0.87</td>
<td>1.03</td>
</tr>
<tr>
<td><strong>18:3-supplemented</strong></td>
<td>16:0</td>
<td>28.1±0.4</td>
<td>23.6±2.5</td>
</tr>
<tr>
<td></td>
<td>16:1</td>
<td>20.3±1.5</td>
<td>14.4±1.3</td>
</tr>
<tr>
<td></td>
<td>18:0</td>
<td>8.7±1.1</td>
<td>5.8±0.3</td>
</tr>
<tr>
<td></td>
<td>18:1</td>
<td>10.2±0.3</td>
<td>8.3±1.7</td>
</tr>
<tr>
<td></td>
<td>18:2</td>
<td>5.3±1.2</td>
<td>3.4±1.6</td>
</tr>
<tr>
<td></td>
<td>18:3</td>
<td>27.4±1.4</td>
<td>44.5±7.9</td>
</tr>
<tr>
<td></td>
<td>U.I.</td>
<td>1.23</td>
<td>1.63</td>
</tr>
</tbody>
</table>

Footnotes as for Table 2.1.
media (although 18:2 comprised between 2% and 5% of fatty acids in 18:3-supplemented Cd\(^{2+}\)-exposed cells), and a lower proportion of 16:1 in cells grown in unsupplemented medium. However, whereas no further incorporation of exogenous fatty acids was evident during stationary phase in the absence of Cd\(^{2+}\) (Table 2.1), 18:2 and 18:3 incorporation continued up to 24 h in Cd\(^{2+}\)-exposed fatty acid-supplemented cultures (Table 2.2). After 24 h growth, unsaturation indices of the latter cultures were approximately 22% higher than those of corresponding cultures grown in the absence of Cd\(^{2+}\) (Fig. 2.3). Despite these differences, the results presented in Figure 2.3 and Table 2.2 confirm that cultures grown in the presence of Cd\(^{2+}\) still incorporate exogenous fatty acids to an extent that can markedly alter their overall degree of fatty acid unsaturation. Thus, the cultures that were most inhibited by Cd\(^{2+}\) in Fig. 2.1 were those that had the higher fatty acid unsaturation index (Fig. 2.3; Table 2.2).

2.3.3. Fatty acid composition of plasma membrane-enriched fractions. The incorporation of exogenous linoleate and linolenate was maximal after 16 h growth (Fig. 2.3; Table 2.1), when cultures were in the late-exponential/early-stationary phase. Hence, cells from 16 h cultures were used for all subsequent short-term experiments. However, as this study was primarily concerned with cadmium toxicity exerted at the plasma membrane, it was initially necessary to confirm incorporation of exogenous 18:2 and 18:3 into plasma membrane lipids. In order to assess purity of isolated plasma membrane fractions (see Section 2.2.4), vanadate-sensitive ATPase (plasma membrane marker) and azide-sensitive ATPase (mitochondrial membrane marker) activities were assayed. Plasma membrane-enriched fractions from cells grown in unsupplemented and
**Fig. 2.3.** Fatty acid unsaturation of *S. cerevisiae*NCYC 1383 during growth in unsupplemented and PUFA-supplemented medium, in the absence and presence of cadmium. Cells from a 48 h starter culture (lacking fatty acid supplements) were inoculated into fresh medium to a concentration of $2 \times 10^6$ cell ml$^{-1}$. Unsaturation indices were determined during growth at 25 °C in unsupplemented (○,●), linoleate- (□, ■) or linolenate- (Δ, ▲) supplemented medium in the absence (○, □, Δ) or presence (●, ■, ▲) of 80 μM Cd(NO$_3$)$_2$. Because of the large differences in unsaturation index values, the data for unsupplemented and linoleate-supplemented cells (a), and linolenate-supplemented cells (b), are presented separately.
18:2-supplemented media displayed vanadate-sensitive ATPase activities 15-fold and 26-fold greater than those of respective whole-cell homogenates (Table 2.3). In contrast, azide-sensitive activities were higher in the homogenates. The lower phase of the aqueous two-phase polymer system, used for purification of plasma membranes from 18:3-supplemented cells, displayed vanadate-sensitive ATPase activity approximately 7-fold greater than that of the upper phase and 16-fold greater than that of the homogenate. However, whereas azide-sensitive activity was highest in the upper phase, enrichment of the mitochondrial ATPase was evident in both upper and lower phases when compared to homogenate activity, suggesting some contamination of the 18:3-enriched plasma membrane fraction. Repeated partitioning resulted in no further improvement in plasma membrane purity. However, in view of the greater relative enrichment of the vanadate-sensitive ATPase (16-fold compared to 8-fold for azide-sensitive activity), and, most importantly, that all plasma membrane fractions displayed high vanadate-sensitive activities, similar to those of plasma membranes isolated by other workers (Serrano, 1988; Schmidt et al., 1983), it was considered that the fractions obtained were of sufficient purity for lipid analysis.

The fatty acid compositions of plasma membrane-enriched fractions from cells grown for 16 h in unsupplemented, 18:2- and 18:3-supplemented media (Table 2.4) were very similar to those of their corresponding whole-cell homogenates (Table 2.1). The proportions of 18:2 and 18:3 in plasma membrane fractions from supplemented cells were approximately 5% higher and 5% lower, respectively, than those of whole cells. In unsupplemented cells, the relative proportion of 16:1 was slightly lower in plasma membranes than in whole-cell homogenates. In spite of these minor differences, fatty
Table 2.3. Purification of plasma membranes from *S. cerevisiae* NCYC 1383 grown for 16 h in unsupplemented, linoleate- and linolenate-supplemented media

<table>
<thead>
<tr>
<th>Fraction</th>
<th>Specific ATPase activity [μmol (mg protein)$^{-1}$ min$^{-1}$]</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Vanadate-sensitive</td>
</tr>
<tr>
<td><strong>Unsupplemented</strong></td>
<td></td>
</tr>
<tr>
<td>Homogenate</td>
<td>0.055</td>
</tr>
<tr>
<td>Plasma membrane</td>
<td>0.799</td>
</tr>
<tr>
<td>18:2-supplemented</td>
<td></td>
</tr>
<tr>
<td>Homogenate</td>
<td>0.026</td>
</tr>
<tr>
<td>Plasma membrane</td>
<td>0.679</td>
</tr>
<tr>
<td>18:3-supplemented</td>
<td></td>
</tr>
<tr>
<td>Homogenate</td>
<td>0.066</td>
</tr>
<tr>
<td>Upper phase</td>
<td>0.145</td>
</tr>
<tr>
<td>Lower phase</td>
<td>1.078</td>
</tr>
</tbody>
</table>

The results shown are typical values from one of at least three experiments.
Table 2.4. Fatty acid composition of plasma membrane-enriched fractions from *S. cerevisiae* NCYC 1383 grown for 16 h in unsupplemented, linolate- and linolenate-supplemented media

<table>
<thead>
<tr>
<th>Fatty acid</th>
<th>Medium</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Unsupplemented</td>
</tr>
<tr>
<td>16:0</td>
<td>23.4±1.1</td>
</tr>
<tr>
<td>16:1</td>
<td>50.7±0.8</td>
</tr>
<tr>
<td>18:0</td>
<td>5.3±0.9</td>
</tr>
<tr>
<td>18:1</td>
<td>20.6±1.1</td>
</tr>
<tr>
<td>18:2</td>
<td>tr.</td>
</tr>
<tr>
<td>18:3</td>
<td>n.d.</td>
</tr>
<tr>
<td>U.I.</td>
<td>0.71</td>
</tr>
</tbody>
</table>

Values for percentage fatty acid composition are means from two separate experiments ± SD. (n.d. = not detectable; tr. < 0.1).
Acid unsaturation indices calculated for plasma membrane-enriched fractions were similar to those of whole cells and again decreased in the order 18:3-supplemented > 18:2-supplemented > unsupplemented (Table 2.4).

2.3.4. Cd\(^{2+}\)-induced K\(^+\) efflux from unsupplemented and PUFA-supplemented *S. cerevisiae* NCYC 1383. Cadmium can cause perturbations in the structural integrity of microbial organellar and cellular membranes (Aßmann *et al.*, 1996; Gadd and Mowll, 1983). Here, release of intracellular K\(^+\) was used as a means of assessing cadmium-induced impairment of plasma membrane integrity. Following Cd(NO\(_3\))\(_2\) addition, K\(^+\) efflux from *S. cerevisiae* previously grown in PUFA-supplemented media was markedly higher than that from cells grown in unsupplemented medium (Fig. 2.4). Rates of K\(^+\) release were maximal between 5 and 10 min after Cd\(^{2+}\) addition. After 12 min exposure to 50 μM Cd(NO\(_3\))\(_2\), amounts of K\(^+\) released from 18:2- and 18:3-enriched cells were approximately equal at 1.5 nmol (10\(^6\) cells)\(^{-1}\), while that from cells grown in unsupplemented medium was only 0.6 nmol (10\(^6\) cells)\(^{-1}\) (Fig. 2.4a). At 100 μM Cd\(^{2+}\), K\(^+\) efflux was greater, indicating more rapid plasma membrane permeabilisation at this higher Cd\(^{2+}\) concentration (Fig. 2.4b). However, relative rates of K\(^+\) release showed a similar pattern to those at 50 μM Cd\(^{2+}\), although a difference between 18:2- and 18:3-supplemented cells also became apparent. Thus, amounts of K\(^+\) released after 12 min exposure to 100 μM Cd\(^{2+}\) were approximately 1.9, 5.0 and 5.9 nmol (10\(^6\) cells)\(^{-1}\) from cells grown in unsupplemented, 18:2- and 18:3-supplemented media, respectively. The results indicate that the susceptibility of *S. cerevisiae* to Cd\(^{2+}\)-induced plasma membrane-disruption increases with the degree of plasma membrane fatty acid unsaturation.
**Fig. 2.4.** Cadmium-induced potassium efflux in unsupplemented and PUFA-supplemented *S. cerevisiae* NCYC 1383. Cells were grown for 16 h in unsupplemented (⌀), linoleate- (●) and linolenate- (☐) supplemented medium, harvested, washed and resuspended in 20 mM Mes (pH 6.0). After addition of glucose (1% w/v), Cd²⁺ was added to a final concentration of (a) 50 μM and (b) 100 μM. Values for K⁺ release are means from three replicate determinations ± SEM where these values exceed the dimensions of the symbols.
2.3.5. Effect of cadmium on the viability of unsupplemented and PUFA-supplemented \textit{S. cerevisiae} NCYC 1383. In order to determine whether increases in the susceptibility of cells grown in unsaturated fatty acid-supplemented media to Cd\textsuperscript{2+}-induced K\textsuperscript{+} efflux were reflected in whole-cell toxicity measurements, changes in cell viability were determined after Cd\textsuperscript{2+} exposure (Fig. 2.5). Cd\textsuperscript{2+} exposure resulted in losses of cell viability that were particularly sharp during the first 1-2 h of incubation. Reductions in viability were greater in PUFA-supplemented cells than in unsupplemented cells. For example, after 1 h incubation in the presence of 50 \textmu M Cd(NO\textsubscript{3})\textsubscript{2}, only 51\% and 37\% of cells previously grown in 18:2- and 18:3-supplemented media, respectively, were still viable (capable of colony formation), whereas 87\% viability was maintained in cells from unsupplemented medium (Fig. 2.5a). Between 2 and 5 h incubation in the presence of 50 \textmu M Cd\textsuperscript{2+}, small increases in the percentage viability of unsupplemented and 18:2-supplemented cells were evident, possibly representing cell division among \textit{S. cerevisiae} sub-populations of lower cadmium-sensitivity. Thus, after 5 h incubation, the percentage viability of unsupplemented cells was approximately seven-fold higher than that of 18:3-supplemented cells. Incubation in the presence of 100 \textmu M Cd(NO\textsubscript{3})\textsubscript{2} had a more marked effect on cell viability, which in all cases was reduced to less than 20\% within 2 h (Fig. 2.5b). Again, the decline in viability was most rapid in PUFA-supplemented cells. After 5 h, percentage viability was reduced to approximately zero in 18:2- and 18:3-supplemented \textit{S. cerevisiae}, whereas 14\% viability was maintained in unsupplemented cells. Similar results and trends to those described above were obtained using alternative methods for determining viability, e.g. dye (methylene violet) exclusion (results not shown).
**Fig. 2.5.** Effect of cadmium on the viability of unsupplemented and PUFA-supplemented *S. cerevisiae* NCYC 1383. Cells were grown for 16 h in unsupplemented (○), linoleate- (●) and linolenate- (□) supplemented medium, harvested, washed and resuspended in 20 mM Mes (pH 6.0). After addition of glucose (1% w/v), Cd²⁺ was added to a final concentration of (a) 50 μM and (b) 100 μM. Percentage viability was calculated by reference to colony counts from samples obtained just prior to Cd²⁺ addition. Values are means from three replicate determinations from single experimental flasks ± SEM. Results from one of at least three experiments are shown.
2.3.6. **Cd²⁺ uptake by unsupplemented and PUFA-supplemented *S. cerevisiae* NCYC 1383.**

In order to determine whether differences in Cd²⁺-induced K⁺-efflux and whole-cell toxicity could be accounted for by differences in levels of cellular Cd²⁺ uptake, the latter was examined in cells previously grown in unsupplemented, 18:2- and 18:3-supplemented media (Fig. 2.6). Cd²⁺ uptake was characterised by an initial rapid phase (complete within 10 min), which probably represented metabolism-independent cell-surface adsorption, followed by a slower phase most likely corresponding to intracellular Cd²⁺ transport, or plasma membrane permeabilisation and inward Cd²⁺ diffusion (Gadd and Mowll, 1983). Cd²⁺ uptake during both phases was similar in the presence of 50 μM and 100 μM Cd²⁺, and did not appear to be affected by the fatty acid composition of *S. cerevisiae*. Thus, after 4 h incubation, cellular Cd²⁺ levels were approximately 0.4 nmol (10⁶ cells)⁻¹ in all cases (Fig. 2.6). Differences in Cd²⁺-induced toxicity in cells grown in unsupplemented, 18:2- and 18:3-supplemented media were, therefore, not attributable to differences in cellular Cd²⁺ uptake.
**Fig. 2.6.** Cadmium uptake by unsupplemented and PUFA-supplemented *S. cerevisiae* NCYC 1383. Cells were grown for 16 h in unsupplemented (◯), linoleate- (●) and linolenate- (□) supplemented medium, harvested, washed and resuspended in 20 mM Mes (pH 6.0). After addition of glucose (1% w/v), CD\(^{2+}\) was added to a final concentration of (a) 50 μM and (b) 100 μM. Values for cellular CD\(^{2+}\) are means from three replicate determinations ± SEM where these values exceed the dimensions of the symbols.
Cadmium toxicity towards *S. cerevisiae* NCYC 1383 was accentuated when cells were enriched with the polyunsaturated fatty acids (PUFAs) linoleate and linolenate. *Saccharomyces cerevisiae* does not normally synthesise PUFAs but can readily incorporate exogenous unsaturated fatty acids into cellular lipids during growth, and this has no adverse effect on cell division (Van der Westhuizen *et al.*, 1994; Wagner and Paltauf, 1994; Bossie and Martin, 1989). The consequent advantages of such a model system for determining physiological effects directly related to altered fatty acid composition have been highlighted (Avery *et al.*, 1996; Maresca and Cossins, 1993). Conventional systems used to manipulate fatty acid composition, which include anaerobic culturing (Watson and Rose, 1980), low-temperature shift (Suutari *et al.*, 1990), or the use of cerulenin (specific inhibitor of sterol and fatty acid synthesis) (Awaya *et al.*, 1975), often adversely affect growth parameters and a range of physiological characteristics.

The very high incorporation of linoleate and linolenate observed here, which was maximal during the late-exponential/early-stationary growth phase at > 65% of total fatty acids, was similar in whole-cell homogenates and plasma membrane-enriched fractions. Therefore, the effects of cadmium on the plasma membrane integrity of cells enriched with different fatty acids could be correlated with their plasma membrane fatty acid compositions. Observed decreases in the relative proportions of monounsaturated fatty acids in linoleate- and linolenate-enriched cells were in keeping with a sharp repression.
of Δ-9 fatty acid desaturase activity in PUFA-supplemented *S. cerevisiae*, as reported elsewhere (Bossie and Martin, 1989; Stukey *et al.*, 1989).

Exposure of *S. cerevisiae* to toxic concentrations of Cd$^{2+}$ generally results in plasma membrane permeabilisation, which is readily detectable as a rapid non-stoichiometric loss of cellular K$^+$ (Gadd and Mowll, 1983; Norris and Kelly, 1977). Extensive K$^+$ efflux (such as the approximately 50% loss within 12 min at 100 μM Cd$^{2+}$ reported here) is generally irreversible, although, in mixed populations, K$^+$ released by sensitive cells may subsequently be taken up by non-sensitive cells (Gadd and Mowll, 1983). In the present study, the slight lag (~2 min) in K$^+$ efflux may have represented the time taken for Cd$^{2+}$ to traverse the plasma membrane; no membrane permeabilisation was evident in the absence of metabolism-dependent cadmium uptake (where glucose was not included in incubations) (results not shown). As described elsewhere (e.g. Avery *et al.*, 1996), reductions in yeast viability were correlated with the relative extent of plasma membrane permeabilisation; a one-to-one relationship between percentage viability and percentage K$^+$ retention was not evident, although such comparisons are complicated by all-or-none effects (Kuypers and Roomans, 1979). Cd$^{2+}$ uptake did not vary with fatty acid composition. Thus, the elevated susceptibility of linoleate- and particularly linolenate-enriched cells to Cd$^{2+}$ toxicity could not be related to altered Cd$^{2+}$ transport activity [it is well documented that the activity of certain membrane transport proteins can be altered by changes in plasma membrane fatty acid composition (Hazel and Williams, 1990; Serrano, 1978). Indeed, increased $V_{\text{max}}$ and $K_m$ values for Cs$^+$ transport in linoleate-supplemented *S. cerevisiae* have recently been reported (Hoptroff *et al.*, 1996).]
Hence, this effect was, therefore, likely to be directly related to plasma membrane properties.

Under non-stressed conditions (e.g. in the absence of toxic metals) the introduction of unsaturated fatty acyl chains into plasma membrane lipid bilayers tends to result in reduced membrane order, or increased membrane 'fluidity' (De Vos et al., 1993; Hazel and Williams, 1990). Large increases in membrane disorder can disrupt plasma membrane functions, including impermeability to the passive diffusion of ions such as K\(^+\) and Na\(^+\) (Hazel and Williams, 1990). Such effects alone could not account for the rapid Cd\(^{2+}\)-induced losses in cellular K\(^+\) observed here, but they may be accentuated through oxidation of unsaturated fatty acids (Tyson and Frazier, 1994). Reactive oxygen species (e.g. superoxide anions and hydroxyl radicals), which may be generated during normal cellular respiratory metabolism, promote lipid peroxidation (Dix and Aikens, 1993). This can result in destabilisation of hydrophobic membrane bilayers by the need to accommodate polar hydroperoxide groups, with a subsequent deterioration of membrane integrity (Van Ginkel and Sevanian, 1994; Dix and Aikens, 1993), such as that indicated here. Studies with higher organisms have shown that certain metal species can actively promote the generation of reactive oxygen species in cells, leading to saturation of cellular protective mechanisms [e.g. glutathione, superoxide dismutases] at toxic concentrations (Stohs and Bagchi, 1995). Furthermore, recent evidence suggests that the toxicity of the redox-active metal copper towards *S. cerevisiae* may also linked to oxyradical generation (Avery et al., 1996; Culotta et al., 1995; Greco et al., 1990).

Redox-cycling activity cannot account for effects induced by the non-redox-active metal cadmium in this study. However, metals without redox capacity may promote lipid
peroxidation by depleting glutathione and protein-bound sulphhydryl groups (Figueiredo-Pereira et al., 1998; Li et al., 1997; Stohs and Bagchi, 1995), or by interacting with membranes to stimulate Fe$^{3+}$/Cu$^{2+}$-initiated lipid peroxidation indirectly (Verstraeten and Oteiza, 1995; Halliwell and Gutteridge, 1989) [it should be noted that enhancement of Cd$^{2+}$ toxicity in the presence of the latter ions may not be evident in intact S. cerevisiae because of simultaneous alleviation of Cd$^{2+}$-induced perturbations of Fe metabolism (Lesuisse and Labbe, 1995)]. Indeed, the yeast cadmium factor, encoded by YCF1, specifically catalyzes the MgATP-energized uptake of bis(glutathionato)cadmium (Cd$^2$GS$_2$) by vacuolar membrane vesicles (Li et al., 1997). The increasing susceptibility of lipids with increasing degrees of fatty acid unsaturation to such oxy-radical induced lipid peroxidation (Halliwell and Gutteridge, 1989) could thus account for the results from K$^+$-efflux and whole-cell toxicity experiments presented here. Variations in cellular levels of a single PUFA (linoleate) may also give corresponding variations in metal sensitivity (Avery et al., 1996).

Oxidative degradation of unsaturated fatty acids was supported by the lower cellular fatty acid unsaturation indices evident during exponential growth in the presence of Cd$^{2+}$; depletion of unsaturated fatty acids is a useful marker of lipid peroxidation (De Vos et al., 1993). That this effect could not simply be attributed to reduced incorporation of exogenous unsaturated fatty acids, was supported by the lower degree of fatty acid unsaturation in the presence of Cd$^{2+}$ also evident in cells grown in the absence of a fatty acid supplement. The increased unsaturation index apparent at later stages of growth was probably a consequence of the elevated anti-oxidant status and stress-resistance of stationary phase yeast cells (Morodas-Ferreira et al., 1996; Steels et al., 1994).
Thus, cadmium toxicity towards *S. cerevisiae* was found to be markedly dependent on plasma membrane fatty acid composition. These findings are of pertinence to metal toxicity in the natural environment, where the fatty acid composition of microorganisms displays considerable heterogeneity (Pennanen *et al.*, 1996; White, 1993; Hazel and Williams, 1990). While several other determinants (constitutive and inducible) of microbial resistance to cadmium have already been described (Li *et al.*, 1997; Jungmann *et al.*, 1993a), this is the first report suggesting a possible passive role of cellular or plasma membrane fatty acid composition and further investigation of this relationship is warranted.
CHAPTER 3

Induction of Lipid Peroxidation during Heavy-Metal Stress in

Saccharomyces cerevisiae and Influence of Plasma Membrane Fatty

Acid Unsaturation
In chapter 2 the relationship between plasma membrane fatty acid unsaturation and susceptibility towards the toxicity of the nonredox-active metal cadmium was investigated. *Saccharomyces cerevisiae* NCYC 1383 was demonstrated to readily incorporate high levels of the exogenously-supplied polyunsaturated fatty acids (PUFAs) linoleate (18:2) and linolenate (18:3) into whole-cell and plasma membrane lipids. In agreement with several reports, incorporation of exogenous PUFAs had no adverse effects on cell division (Bossie and Martin, 1989; Stukey *et al.*, 1989; Watson and Rose, 1980). However, PUFA-enriched *S. cerevisiae* were demonstrated to be markedly more susceptible towards Cd\(^{2+}\)-induced plasma membrane permeabilisation and whole-cell toxicity. Furthermore, Avery *et al.* (1996) had also demonstrated the elevated sensitivity of linoleate-enriched *S. cerevisiae* to plasma membrane permeabilisation and whole-cell toxicity upon exposure to the redox-active metal copper. While several reports have previously demonstrated heavy metal-induced perturbation of plasma membrane integrity, with associated cytosolic-solute leakage, e.g. K\(^+\) efflux, in *S. cerevisiae* (Aßmann *et al.*, 1996; Ohsumi *et al.*, 1988; De Rome and Gadd, 1987; Kessels *et al.*, 1985), this was the first report indicating increased susceptibility with increasing degrees of plasma membrane fatty acid unsaturation.

The mechanism underlying the increased metal sensitivity of PUFA-enriched cells was not elucidated. However, one general cause for loss of membrane integrity in biological systems, in response to a variety of stimuli, can arise through oxy-radical
mediated lipid peroxidation (Halliwell and Gutteridge, 1989; Mehlhorn, 1986). The accumulation of lipid peroxidation products, such as lipid hydroperoxides, within the hydrophobic core of plasma membranes can result in disturbances in the arrangement of phospholipid moieties, an increase in the membrane phase transition temperature, and ultimate impairment of membrane function (manifested as K⁺ loss) (Weckx and Clijsters, 1996; Van Ginkel and Sevanian, 1994). Indeed, the oxidative degradation of PUFAs was supported by the lower cellular fatty acid unsaturation indices evident during exponential growth in the presence of Cd(NO₃)₂ (Chapter 2). Depletion of PUFAs has been employed as a useful marker of lipid peroxidation (De Vos et al., 1993). Several studies have indicated that the susceptibility of fatty acids to lipid peroxidation increases with the degree of fatty acyl chain unsaturation (Porter et al., 1995; Vossen et al., 1995; De Vos et al., 1993), supporting a possible link with the results of Chapter 2.

As stated earlier, through redox-cycling, redox-active metals such as copper, are known to generate reactive oxygen species (ROS) such as the hydroxyl radical (HO·) and the superoxide anion/perhydroxyl radical (O₂⁻/HO₂⁻), via Fenton and Haber-Weiss reactions (Dix and Aikens, 1993; Shinar et al., 1989). In addition, nonredox-active metals such as cadmium have the capacity to indirectly promote oxidative stress through depletion of free radical scavengers such as glutathione and protein-bound sulphydryl groups, and by displacement of redox-active metals from enzyme active sites, thereby stimulating Fe³⁺/Cu²⁺-initiated lipid peroxidation (Figueiredo-Pereira et al., 1998; Hassoun and Stohs, 1996; Verstraeten and Otieza, 1995; Halliwell and Gutteridge, 1989). The metal hyper-sensitivity of various mutants of S. cerevisiae, defective in components of the oxidative stress response, supports a role of oxygen free radicals in microbial metal
toxicity. For example, SOD1 mutants deficient in copper/zinc superoxide dismutase activity are hypersensitive to copper (Greco et al., 1990) [these observations may be complicated, however, by the involvement of SOD1 in copper buffering (Culotta et al., 1995)]. Furthermore, mutations in the YAP1 and YAP2 genes of S. cerevisiae (key transcriptional activators of oxidative stress response genes) confer hypersensitivity to cadmium (Wu et al., 1993). An important role for lipid peroxidation in the toxicity of heavy metal ions towards higher organisms has also been implicated from several recent studies (Stohs and Bagchi, 1995; Kalyanaraman et al., 1992). However, to date no reports have investigated the relationship between heavy metal toxicity, plasma membrane fatty acid unsaturation, and lipid peroxidation in microorganisms.

In the present study, heavy metal-induced lipid peroxidation is demonstrated in a microorganism for the first time. Furthermore the increased susceptibility of PUFA-enriched S. cerevisiae to cadmium- and copper-induced plasma membrane perturbation and toxicity is shown here to correlate with elevated levels of lipid peroxidation in these cells.
3.2. Materials and Methods

3.2.1. Organism and culture conditions. *Saccharomyces cerevisiae* S150-2B (MATa, leu2-3, -112, *ura3-52, trp1-289, his3-A1, Gal*⁺) (kindly provided by D. J. Jamieson, Heriot Watt University, Edinburgh, U.K.) was selected for this study because of its well characterised responses to oxidative stress (Stephen and Jamieson, 1996; Stephen *et al.*, 1995; Jamieson, 1992). S150-2B was routinely maintained on solid YEPD and YNB medium as described previously for NCYC 1383 (see Section 2.2.1.). For experimental purposes, S150-2B was grown as previously described for NCYC 1383 (see Section 2.2.1.).

3.2.2. Preparation of cell homogenates. Whole-cell homogenates were prepared as described previously for *S. cerevisiae* NCYC 1383 (see Section 2.2.2.).

3.2.3. Lipid extraction and fatty acid analysis. Lipids were extracted from whole-cell homogenates as described previously (see Section 2.2.5.).

3.2.4. Preparation of cell suspensions for metal toxicity experiments. Cells from the late-exponential phase (13 h) were harvested by centrifugation at 1,500 g for 5 min and washed twice with distilled deionised water. Washed cells were suspended to a density of approximately 5 x 10⁷ ml⁻¹ in 40 ml of 10 mM Mes buffer (pH 5.5), 1% (w/v) glucose, and incubated at room temperature with orbital shaking at 120 rev. min⁻¹. After 10 min equilibration, Cd(NO₃)₂ or Cu(NO₃)₂ was added to the desired concentration.
3.2.5. **Cell viability.** Cell viability was once again determined as the ability to produce colony-forming units on OGYE medium, as described previously (see Section 2.2.7.).

3.2.6. **Potassium efflux.** At specified intervals after the addition of Cd(NO₃)₂ or Cu(NO₃)₂ to cell suspensions, 0.5 ml aliquots were removed, microcentrifuged for 3 min, and the supernatant then diluted with 4 volumes of distilled deionised water. Extracellular K⁺ was measured using a Corning 410 flame photometer, with reference to appropriate standard KCl solutions.

3.2.7. **Evaluation of lipid peroxidation.** Lipid peroxidation was quantified by two methods: determination of thiobarbituric acid (TBA)-reactive substances (TBARS) and conjugated dienes. TBARS were determined using a method adapted from that described by Aust (1994). At specified intervals (up to 3 h), after the addition of Cd(NO₃)₂ or Cu(NO₃)₂, 0.5 ml samples of cell suspension were removed and added to 1 ml of TBA reagent [0.25 M HCl, 15% (w/v) trichloroacetic acid, 0.375% (w/v) TBA]. Addition of the reagent terminated lipid peroxidation and initiated the assay. Samples were heated for 15 min in a boiling water bath and, after cooling, centrifuged at 1,000 g for 5 min in order to remove any cell debris. Absorbances of samples were then measured at 535 nm, using a Shimadzu UV-120-02 spectrophotometer, against a reference solution comprising 1 ml TBA reagent with the sample replaced by an equal volume (0.5 ml) of distilled deionised water. The concentrations of TBARS in samples were calculated by reference to a standard curve prepared using 1,1,3,3-tetramethoxypropane (TMP).
For the determination of conjugated dienes (Glende and Recknagel, 1994), 10 ml samples were removed from cell suspensions and lipid extracts prepared from whole-cell homogenates as described above. After evaporation of chloroform under nitrogen, dried lipids were resuspended in 3 ml cyclohexane. UV spectra of lipid solutions were recorded between 215 and 280 nm using a Perkin-Elmer lambda-7 UV/Visible spectrophotometer. Conjugated dienes absorb light between approximately 215 nm and 255 nm, with a peak at 233 nm (Corongiu et al., 1994). Peak absorbances at 233 nm were normalized against absorbance values at 260 nm and relative conjugated diene content expressed as A233/A260.

3.2.8. Fluorescence depolarization. Plasma membrane lipid-order was determined by measuring steady-state fluorescence anisotropy in whole cells labeled with trimethylammonium diphenylhexatriene (TMA-DPH). The cationic probe TMA-DPH becomes primarily anchored at the plasma membrane of intact cells (Gille et al., 1993; Block, 1991). TMA-DPH was added to S. cerevisiae cell suspensions (5 x 10^7 cells ml^-1, see above) from a 1 mM stock solution, prepared in concentrated dimethylformamide, to give a final concentration of 1 μM. After 30 min equilibration, when measurements had stabilised, Cd(NO₃)₂ was added to cell suspensions. Triplicate 3 ml samples of suspension were removed at intervals, transferred to 10-mm-pathlength quartz cuvettes, and analysed using a Perkin Elmer LS5 fluorescence spectrometer with polarization accessory. TMA-DPH was excited with vertically polarized light at 360 nm, and the vertical and horizontal vectors of emitted light measured at 450 nm. Membrane order
was expressed as the order parameter $S$, which reflects the orderliness of membrane phospholipids.

$$S = (r/r_o)^{0.5} \quad \text{(Gille et al., 1993)}$$

where $r_o$ is the theoretical limiting anisotropy (0.395 for TMA-DPH) in the absence of rotational motion, and $r$ is the steady-state anisotropy measured in the membrane. The effects of Cd$^{2+}$ on membrane order were determined by measuring $S$ at intervals over 60 min. Light scatter, determined in the absence of TMA-DPH, was found to account for less than 5% of the total emitted light and therefore no corrections were employed (Avery et al., 1995a).
3.3. Results

In the previous studies (Avery et al., 1996; Chapter 2), it was demonstrated that *S. cerevisiae* NCYC 1383 readily incorporated exogenous PUFAs (linoleate and linolenate) from its growth medium to greater than 65% of total fatty acids. *Saccharomyces cerevisiae* S150-2B was selected for this study because of its well characterised responses to oxidative stress (Stephen and Jamieson, 1996; Stephen et al., 1995; Jamieson, 1992). *Saccharomyces cerevisiae* S150-2B was also observed to incorporate exogenous PUFAs; levels of incorporation were very similar to those described previously (results not shown). Increases in PUFA content, as a proportion of total fatty acids, were contemporaneous with decreases in the monounsaturated fatty acids palmitoleate (16:1) and oleate (18:1). Cellular fatty acid unsaturation indices (average number of double bonds per fatty acid) decreased in the order 18:3-supplemented cells (2.28) > 18:2-supplemented cells (1.48) > unsupplemented cells (0.82). The fatty acid compositions of plasma membrane-enriched fractions from *S. cerevisiae* match their corresponding whole-cell homogenates very closely (Avery et al., 1996; Chapter 2).

3.3.1. Effect of cadmium and copper on the viability of unsupplemented and PUFA-supplemented *S. cerevisiae* S150-2B. *Saccharomyces cerevisiae* S150-2B previously grown in PUFA-supplemented media was markedly more susceptible to Cd$^{2+}$ (Fig. 3.1) and Cu$^{2+}$ (Table 3.1) toxicity than cells grown in unsupplemented medium. For example,
**Fig. 3.1.** Effect of cadmium exposure on the viability of unsupplemented and PUFA-supplemented *S. cerevisiae* S150-2B. Cells were grown for 13 h in unsupplemented (○), linoleate-supplemented (●), or linolenate-supplemented (□) medium and then harvested, washed, and resuspended in 10 mM Mes (pH 5.5), 1% w/v glucose. Cd(NO₃)₂ was added to a final concentration of 50 (a) or 200 (b) μM. Typical results from one of at least two experiments are shown. Points represent means of three replicate samples from single experimental flasks. Standard errors of the mean (SEM) are smaller than the dimensions of the symbols.
Table 3.1. Effect of Cu(NO₃)₂ on viability of *S. cerevisiae* S150-2B cells previously grown in unsupplemented and PUFA-supplemented media

<table>
<thead>
<tr>
<th>[Cu(NO₃)₂] (µM)</th>
<th>% Viability of the indicated cell type(^a)</th>
<th>18:2-supplemented</th>
<th>18:3-supplemented</th>
</tr>
</thead>
<tbody>
<tr>
<td>5(^b)</td>
<td>88 ± 1</td>
<td>70 ± 5</td>
<td>63 ± 3</td>
</tr>
<tr>
<td>40(^c)</td>
<td>11.0 ± 1.1</td>
<td>1.4 ± 1.6</td>
<td>0</td>
</tr>
</tbody>
</table>

\(^a\) Values are means ± SEM derived from three replicate determinations

\(^b\) Determined after 60 min

\(^c\) Determined after 5 min
after 1 h incubation in the presence of 50 \( \mu \text{M} \) \( \text{Cd(NO}_3\text{)}_2 \), only 40% and 16% of cells previously grown in 18:2- and 18:3-supplemented media, respectively, were still viable (capable of colony formation) (Fig. 3.1a), whereas 55% viability was maintained in cells from unsupplemented medium. Similar trends were evident at 200 \( \mu \text{M} \) \( \text{Cd(NO}_3\text{)}_2 \), albeit over a shorter time-scale. No viable cells were detected after 2 h incubation in the presence of 200 \( \mu \text{M} \) \( \text{Cd}^{2+} \). This was also the case after 5 min exposure of 18:3-supplemented cells to 40 \( \mu \text{M} \) \( \text{Cu(NO}_3\text{)}_2 \), although some capacity for colony formation was retained by \( \text{Cu}^{2+} \)-exposed unsupplemented and 18:2-supplemented cells (Table 3.1). Incubation for 60 min in the presence of 5 \( \mu \text{M} \) \( \text{Cu(NO}_3\text{)}_2 \) had far less marked effects on cell viability, although here again percentage viability decreased in the order unsupplemented cells > 18:2-supplemented cells > 18:3-supplemented cells.

### 3.3.2. Cadmium- and copper-induced K\(^+\) efflux from unsupplemented and PUFA-supplemented *S. cerevisiae* S150-2B.

Release of intracellular K\(^+\) was used as an indicator of plasma membrane permeabilisation (Avery *et al.*, 1996; Chapter 2; Ohsumi *et al.*, 1988). Following heavy metal addition, K\(^+\) efflux was initially very rapid, but slowed considerably after 15 min (Fig. 3.2). Metal-induced K\(^+\) release from *S. cerevisiae* previously grown in PUFA-supplemented medium was markedly higher than that from cells grown in unsupplemented medium. For example, amounts of K\(^+\) released from 18:2- and 18:3-enriched cells were approximately equal at 5.6 nmol (10\(^6\) cells\(^{-1}\)) after 15 min exposure to 50 \( \mu \text{M} \) \( \text{Cd(NO}_3\text{)}_2 \), whereas that from cells grown in unsupplemented medium was only 3.1 nmol (10\(^6\) cells\(^{-1}\)) (Fig. 3.2a). K\(^+\) loss continued after 15 min, and amounts
Fig. 3.2. Cadmium-induced $K^+$ efflux in unsupplemented and PUFA-supplemented *S. cerevisiae* S150-2B cells. Cells were grown for 13 h in unsupplemented (○), linoleate-supplemented (●) and linolenate-supplemented (■) medium and then harvested, washed, and resuspended in 10 mM Mes (pH 5.5), 1% w/v glucose. Cd(NO$_3$)$_2$ was added to a final concentration of 50 (a) or 200 (b) μM. Typical results from one of at least two experiments are shown. Values for $K^+$ release are means from three replicate determinations. SEM are smaller than the dimensions of the symbols.
Table 3.2. Copper-induced K⁺ release from *S. cerevisiae* S150-2B cells grown previously in unsupplemented and PUFA-supplemented media

<table>
<thead>
<tr>
<th>[Cu(NO₃)₂]</th>
<th>Exposure time</th>
<th>K⁺ release from the indicated cell type*</th>
</tr>
</thead>
<tbody>
<tr>
<td>(µM)</td>
<td>(min)</td>
<td>Unsupplemented</td>
</tr>
<tr>
<td>5</td>
<td>30</td>
<td>0.55 ± 0.04</td>
</tr>
<tr>
<td></td>
<td>210</td>
<td>0.62 ± 0.03</td>
</tr>
<tr>
<td>40</td>
<td>30</td>
<td>3.4 ± 0.1</td>
</tr>
<tr>
<td></td>
<td>210</td>
<td>4.1 ± 0.1</td>
</tr>
</tbody>
</table>

* Nanomoles of K⁺ released per 10⁶ cells. Values are means ± SEM derived from three replicate determinations
released from 18:3-supplemented cells after 3 h were approximately 17% higher than those from 18:2-supplemented cells. K+ efflux was greater at 200 μM Cd(NO₃)₂ than at 50 μM Cd(NO₃)₂, and differences between 18:2- and 18:3-supplemented cells were more marked. Thus, amounts of K⁺ released after 1 h exposure to 200 μM Cd²⁺ were approximately 4.8, 7.6 and 8.4 nmol (10⁶ cells)⁻¹ from cells previously grown in unsupplemented, 18:2- and 18:3-supplemented media, respectively (Fig. 3.2b). Similar trends were apparent during copper exposure (Table 3.2). Amounts of K⁺ released from 18:2- and 18:3-enriched cells incubated in the presence of 5 μM Cu(NO₃)₂ were similar, but approximately 100% and 84% higher after 30 min and 210 min, respectively, than those from unsupplemented cells under the same conditions. At 40 μM Cu(NO₃)₂, a greater relative permeabilisation of the plasma membrane of 18:3-supplemented cells compared to 18:2-supplemented cells also became apparent (Table 3.2).

3.3.3. Cadmium- and copper-induced lipid peroxidation in unsupplemented and PUFA-supplemented S. cerevisiae S150-2B. Lipid peroxidation was evaluated as thiobarbituric acid-reactive substances (TBARS) production and conjugated diene formation. Two different analytical methods were used to assess lipid-localised oxidative stress as no single method adequately measures the range of reactions possible (Coudray et al., 1995). Furthermore, the TBA assay alone was not suitable for direct comparison of linolenate-enriched cells with the other cell types as malondialdehyde, to which the assay is particularly sensitive, is only formed upon the breakdown of linolenate, and not linoleate or monounsaturated fatty acids (Mihaljevic et al., 1996; Aust, 1994).
Fig. 3.3. Cadmium-induced TBARS production in unsupplemented and linoleate-supplemented *S. cerevisiae* S150-2B. Cells were grown for 13 h in unsupplemented (○) and linoleate-supplemented (●) medium and then harvested, washed, and resuspended in 10 mM Mes (pH 5.5), 1% w/v glucose. Cd(NO$_3$)$_2$ was added to a final concentration of 50 (a) or 200 (b) µM. Typical results from one of at least two experiments are shown. Values for TBARS are means of three replicate determinations from single experimental flasks, ± SEM where these values exceed the dimensions of the symbols.
TBARS [pmol (10^6 cells)^{-1}]

**Time (h)**

- **a.**
- **b.**
**Table 3.3.** Copper-induced TBARS production in *S. cerevisiae* S150-2B cells grown previously in unsupplemented and linoleate-supplemented media

<table>
<thead>
<tr>
<th>[Cu(NO₃)₂] (µM)</th>
<th>Exposure time (min)</th>
<th>TBARS produced by the indicated cell typea</th>
<th>Unsupplemented</th>
<th>18:2-supplemented</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>0</td>
<td>5.7 ± 1.1</td>
<td>7.3 ± 0.9</td>
<td></td>
</tr>
<tr>
<td></td>
<td>60</td>
<td>8.9 ± 1.8</td>
<td>12.4 ± 1.5</td>
<td></td>
</tr>
<tr>
<td></td>
<td>180</td>
<td>14.3 ± 0.4</td>
<td>14.1 ± 3.1</td>
<td></td>
</tr>
<tr>
<td>40</td>
<td>0</td>
<td>5.4 ± 1.7</td>
<td>7.0 ± 0.4</td>
<td></td>
</tr>
<tr>
<td></td>
<td>60</td>
<td>14.2 ± 0.2</td>
<td>17.5 ± 1.1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>180</td>
<td>17.2 ± 1.0</td>
<td>16.0 ± 0.7</td>
<td></td>
</tr>
</tbody>
</table>

a Picomoles of TBARS produced per $10^6$ cells. Values are means ± SEM derived from three replicate determinations.
Difficulties were encountered with the purification of plasma membranes from cells following metal exposure: these were related to altered membrane density and partitioning behaviour. Thus, the values for whole-cell lipid peroxidation (given) include contributions from organellar membranes. Because the plasma membrane is a primary target for metal toxicity (Abemann et al., 1996; Verstraeten and Oteiza, 1995; Ohsumi et al., 1988), and its fatty acid composition is similar to that of organellar membranes (Avery et al., 1996), it can be inferred that lipid peroxidation per unit plasma membrane was at least equal to (probably greater than) that of organellar membranes.

Prior to the addition of metal, amounts of TBARS were generally approximately 25-55% larger in linoleate-enriched cells than in cells grown previously in unsupplemented medium (Fig. 3.3; Table 3.3). Marked increases in TBARS formation were evident following metal addition. Control experiments confirmed that no detectable rise in TBARS levels occurred during 3 h incubation in buffer in the absence of cadmium or copper (results not shown). Metal-induced increases were particularly large during the first minutes of metal exposure, although in certain cases TBARS formation continued up to 3 h (Fig. 3.3). The level of metal-induced TBARS production was considerably higher in S. cerevisiae previously grown in 18:2-supplemented medium than that in cells grown in unsupplemented medium (Fig. 3.3). For example, after 1 h exposure to 50 µM Cd(NO₃)₂, the level of TBARS in 18:2-enriched cells was approximately 33 pmol (10⁶ cells)⁻¹, whereas that from cells grown in unsupplemented medium was only 18 pmol (10⁶ cells)⁻¹ (Fig. 3.3a). The level of TBARS production was higher at 200 µM Cd(NO₃)₂ than at 50 µM Cd(NO₃)₂. Up to 2 h, levels of TBARS were between 1.6- and 2-fold greater in 18:2-supplemented cells than in unsupplemented cells at either Cd(NO₃)₂ concentration.
A reduction in cellular TBARS levels between 2 and 3 h incubation with cadmium was evident only in 18:2-supplemented cells, whereas lipid peroxidation continued to rise in unsupplemented cells during this period. Thus, after 3 h, the levels of TBARS measured in 18:2-supplemented cells were only approximately 1.2-fold higher than those of unsupplemented cells (Fig. 3.3). Similar trends were observed for copper exposure, with which TBARS production was also associated. Again, lipid peroxidation increased with metal concentration and, after 1 h, levels of Cu-induced TBARS were between 20% and 40% higher in 18:2-enriched cells than in unsupplemented cells. However, between 1 h and 3 h, TBARS levels in 18:2-supplemented cells either increased only slightly (5 µM Cu²⁺) or declined (40 µM Cu²⁺). In contrast, those of unsupplemented cells continued to increase and, at 3 h, exceeded TBARS levels evident in 18:2-enriched cells (Table 3.3).

Lipid peroxidation was also evaluated by measuring conjugated dienes, which are early intermediates in the lipid peroxidative chain (Glende and Recknagel, 1994). Relative levels of conjugated dienes were determined after 1 h exposure to Cd and Cu and expressed as A_{233}/A_{260}. At 50 µM Cd(NO₃)₂, A_{233}/A_{260} values for cells grown previously in unsupplemented and 18:2- and 18:3-supplemented media were 1.03, 1.12 and 1.18, respectively, indicating increased Cd-induced conjugated diene formation in cells displaying elevated fatty acid unsaturation indices (Fig. 3.4a). A similar trend was evident at 200 µM Cd(NO₃)₂, although the A_{233}/A_{260} values were higher in all cases at this increased cadmium concentration. Conjugated diene levels also increased with the unsaturation index in copper-exposed cells. A_{233}/A_{260} values after 1 h at 5 µM Cu(NO₃)₂, for cells grown previously in unsupplemented, 18:2- and 18:3-supplemented media were
Fig. 3.4. Cadmium- and copper-induced conjugated-diene formation in unsupplemented and PUFA-supplemented S. cerevisiae S150-2B. Cells were grown for 13 h in unsupplemented, linoleate- or linolenate-supplemented medium and then harvested, washed, and resuspended in 10 mM Mes (pH 5.5), 1% w/v glucose. (a) Cd\(^{2+}\) was added to a final concentration of 50 (☐) or 200 (■) µM. (b) Cu\(^{2+}\) was added to a final concentration of 5 (☐) or 20 (■) µM. Values are means for three individual lipid extracts ± SEM.
1.10, 1.21 and 1.41, respectively (Fig. 3.4b). Conjugated diene formation was greater at 20 μM than at 5 μM Cu(NO₃)₂, although again the A₂₃₃/A₂₆₀ values were highest for cells that had previously been grown in the presence of linoleate, and particularly linolenate. Interestingly, despite the lower copper concentrations employed, and in contrast to the TBARS results, copper-exposed cells generally displayed higher conjugated diene levels than cadmium-exposed cells. The elevated levels of lipid peroxidation evident (from both sets of data) in PUFA-enriched S. cerevisiae were correlated with the increased susceptibility of these cells to Cd²⁺ and Cu²⁺-induced plasma membrane permeabilisation and whole-cell toxicity.

3.3.4. Effect of cadmium on plasma membrane order of unsupplemented and PUFA-supplemented S. cerevisiae S150-2B. In order to assess metal-induced changes in plasma membrane physical properties, plasma membrane lipid order was monitored (by fluorescence depolarization with TMA-DPH-labelled cells) during the 50 min period following addition of 50 μM Cd(NO₃)₂ (Fig. 3.5). Prior to the addition of metal, S (membrane order parameter) values for cells previously grown in unsupplemented and 18:2- and 18:3-supplemented media were 0.74, 0.71 and 0.69, respectively, indicating decreased orderliness in plasma membranes enriched with PUFAs. After the addition of metal, decreases in plasma membrane order in all three types of cell were clearly evident, and these continued for at least 50 min. The Cd²⁺-induced change in S was slightly greater for cells previously grown in unsupplemented medium than for 18:2- and 18:3-enriched cells. However, after 50 min of incubation in the presence of Cd²⁺, values for S still decreased in the order unsupplemented cells > 18:2-supplemented cells > 18:3-
Fig. 3.5. Effect of cadmium on membrane order in unsupplemented and PUFA-supplemented *S. cerevisiae* S150-2B cells. Cells were grown for 13 h in unsupplemented (○), linoleate-supplemented (●), or linolenate-supplemented (□) medium and then harvested, washed, and resuspended in 10 mM Mes (pH 5.5), 1% w/v glucose. TMA-DPH was added to a final concentration of 1 μM, and 30 min later, Cd²⁺ was added to a final concentration of 50 μM. Typical results from one of at least two experiments are shown. Values for S are means for three replicate samples from single experimental flasks, ± SEM where these values exceed the dimensions of the symbols.
supplemented cells. The order parameter for plasma membranes of Cd\(^{2+}\)-exposed unsupplemented cells did not decrease below the values for PUFA-enriched cells determined prior to cadmium addition (Fig. 3.5).
3.4. Discussion

Metal-induced plasma membrane permeabilisation and whole-cell toxicity in *S. cerevisiae* S150-2B [a strain for which various components of the oxidative stress response have been well characterised (Stephen and Jamieson, 1996; Stephen et al., 1995; Jamieson, 1992)] were initially confirmed to be greater in cells enriched with the PUFAs linoleate and linolenate. Similar effects of plasma membrane fatty acid composition on copper (Avery et al., 1996) toxicity, albeit more marked, have also been observed in *S. cerevisiae*NCYC 1383. Hence, this effect is clearly not strain-specific.

The results also suggest that the ability of *S. cerevisiae* to incorporate exogenous PUFAs with little effect on growth rate may be widespread; previous reports of this characteristic were restricted to *S. cerevisiae*NCYC 1383 (Avery et al., 1996; Bossie and Martin, 1989; Chapter 2). The consequent experimental advantages of *S. cerevisiae* for modelling membrane-dependent effects have been highlighted elsewhere (Avery et al., 1996).

As in previous reports (Avery et al., 1996; Belde et al., 1988; Chapter 2; Ohsumi et al., 1988), extensive metal-induced potassium release was associated with a loss of viability of *S. cerevisiae*. Whereas a stoichiometric relationship between physiological metal uptake and potassium release can occur at non-toxic metal concentrations (Gadd and Mowll, 1983), our previous studies have confirmed nonstoichiometry at the toxic metal concentrations used here (Avery et al., 1996; Chapter 2). Thus, potassium release can be attributed to plasma membrane permeabilisation and toxicity (Avery et al., 1996; Chapter 2; Ohsumi et al., 1988). As is usually the case among yeast and fungi (De Rome
and Gadd, 1987; Gadd and Mowll, 1983), copper was more toxic than cadmium for \textit{S. cerevisiae}. However, copper exposure resulted in less K\(^+\) release than that observed during cadmium exposure at the concentrations used. Thus, it can be inferred that effects other than membrane permeabilisation, e.g. direct interaction with nucleic acids or misincorporation into metallothioneins (Cervantes and Gutierrez-Corona, 1994), may make a more important contribution in copper toxicity than cadmium toxicity to loss of \textit{S. cerevisiae} viability. An important role of copper-induced membrane effects was nevertheless implicated by the observed influence of membrane fatty acid composition.

A role of lipid-targeted oxy-radical attack in the toxicity of heavy metals towards higher organisms has been inferred from observations of elevated lipid peroxidation in metal-exposed organisms (Kalyanaraman \textit{et al.}, 1992; Halliwell and Gutteridge, 1989). In this study, metal-induced lipid peroxidation in a microorganism was demonstrated for the first time.

The integration of different analytical methods for the measurement of lipid peroxidation was necessary (Coudray \textit{et al.}, 1995). Results from conjugated-diene and TBARS analyses suggested contrasting pathways of cadmium- and copper-induced lipid peroxidation. Unlike TBARS, conjugated dienes are primary products in the lipid peroxidative chain, arising through oxy-radical mediated abstraction of hydrogen atoms from methylene groups separating PUFA double-bonds (Glende and Recknagel, 1994). Therefore, the higher relative levels of conjugated dienes in Cu-exposed cells suggest that initiation of lipid peroxidation occurs at a greater rate in the presence of copper than cadmium in \textit{S. cerevisiae}; such a conclusion is consistent with the respective redox-activities and inactivities of these metals (Stohs and Bagchi, 1995). The mechanism of
lipid peroxidation initiation is unlikely to differ as the principal initiating species generated by both copper (direct catalysis of the Fenton reaction) and cadmium (indirect promotion of the Fenton reaction by glutathione depletion) is most likely the hydroxyl radical (Dix and Aikens, 1993). However, observed differences in initial TBARS levels, prior to metal addition, do suggest the possibility of additional initiation via pre-formed lipid hydroperoxides (Dix and Aikens, 1993).

While the TBARS assay detects products arising from the decomposition of lipid hydroperoxides, such as the decomposition products of oleate and linoleate hydroperoxides detected in this study, the assay is particularly sensitive to malondialdehyde (Aust, 1994). Malondialdehyde is formed only upon oxidation of PUFAs containing at least three methylene-separated double bonds (Mihaljevic et al., 1996). Thus, while the results demonstrate elevated levels of lipid peroxidation in linoleate-enriched cells, compared to unsupplemented cells, the total cellular amounts of lipid hydroperoxides for all cell types are most likely underestimated.

The formation of high levels of TBARS can be precluded by glutathione peroxidase activity during normal cellular metabolism. Glutathione peroxidase converts lipid hydroperoxides to their corresponding hydroxy fatty acids (Tran et al., 1993). However, glutathione is a principal cellular target or sequestration site of cadmium (Figueiredo-Pereira et al., 1998; Li et al., 1997; Lesuisse and Labbe, 1995). Thus, the higher relative levels of TBARS in Cd-exposed than in Cu-exposed cells may reflect glutathione depletion, and hence, a reduced capacity of the former to repair lipid peroxidation damage.
The convergence of TBARS levels in unsupplemented and linoleate-supplemented *S. cerevisiae*, after prolonged metal-exposure may reflect a greater induction of antioxidant activity in the latter cells; the oxidative stress response of *S. cerevisiae*, which includes the induction of glutathione, catalase, metallothionein, and glutathione peroxidase, increases with the extent of initial insult (Steels *et al.*, 1994; Jamieson, 1992).

Exposure to neither metal was associated with a decline in the degree of cellular fatty acid unsaturation within one hour (results not shown). However, longer term reductions in the proportion of unsaturated fatty acids were previously noted during growth of *S. cerevisiae*NCYC 1383 in cadmium-supplemented medium (Chapter 2). This supports the view that reduced polyunsaturated fatty acid content can be a late marker of lipid peroxidation (De Vos *et al.*, 1993).

The similarities between the short initiation time-scales (noting that TBARS are not immediate products of lipid peroxidation) and between the relative extents, of lipid peroxidation and K⁺ release observed here, were consistent with the former process being associated with a deterioration of membrane integrity (Corongiu *et al.*, 1994; Van Ginkel and Sevanian, 1994). Loss of membrane impermeability during lipid peroxidation may arise through covalent bond formation between adjacent acyl radicals, resulting in increased membrane rigidity, or through the incorporation of short-chain and/or polar oxidation products, resulting in decreased membrane order (Van Ginkel and Sevanian, 1994; Dix and Aikens, 1993). The small cadmium-induced reductions in plasma membrane order reported here were indicative of a predominance of the latter type of effect, and in agreement with the reported effects of cadmium on *Schizosaccharomyces*
pombe plasma membranes (Ab mann et al., 1996). Unlike K⁺ efflux, the effect of Cd²⁺ on plasma membrane order was apparently no more pronounced in PUFA-enriched cells than in unsupplemented cells. Thus, a threshold value for membrane order, below which K⁺ efflux occurred, was not discernible. It appears that values for the membrane order parameter (S) do not provide a clear linear index of the state of a membrane in relation to either its integrity or its content of lipid peroxidation products. Nevertheless, values for plasma membrane order determined after metal-exposure decreased in the same order (unsupplemented cells > linoleate-supplemented cells > linolenate-supplemented cells) as that predicted from metal toxicity and lipid peroxidation experiments. Note that this is also the order to be expected on the basis of differences in initial fatty acid composition, irrespective of the extent of lipid peroxidation (Avery et al., 1996). In addition to the rapid gross effects on membrane permeability observed at the high metal concentrations used here, smaller changes in membrane physical properties at lower non-toxic metal concentrations might still elicit changes in cellular activity; the function of many membrane-bound enzymes is known to be sensitive to the enzymes' lipid environment (Hazel and Williams, 1990).

The accentuated levels of lipid peroxidation and heavy metal-toxicity in linoleate- and linolenate-enriched S. cerevisiae were consistent with the known susceptibility of polyunsaturated fatty acids to oxidation (Porter et al., 1995; Dix and Aikens, 1993). The possibility that the results might also be partly attributable to differences in the physical properties of PUFA-enriched plasma membranes cannot be discounted (Verstraeten and Oteiza, 1995); in common with studies on higher organisms (Weckx and Clijsters, 1996; Vossen et al., 1995), the fatty acid unsaturation index of S. cerevisiae (1.5-fold higher for
linolenate- than linoleate-enriched cells) was not linearly related to susceptibility to lipid peroxidation. Sensitivity may relate more closely to initial plasma membrane order.

The potentially major role of membrane lipid peroxidation in heavy metal toxicity towards yeast implicated from the present results suggests that changes in a number of highly-variable cellular properties, including antioxidant status (e.g. the induction of antioxidant defences during respiratory adaptation (Steels et al., 1994; Jamieson et al., 1994)) and membrane fatty acid composition (e.g. changes occurring during environmental acclimation (Hazel and Williams, 1990)), could seriously alter the cells’ ability to cope with heavy metal stress. As well as being of relevance to the stress biochemistry of yeast, these findings may have significant implications for metal-microbe interactions in the natural environment.
CHAPTER 4

Factors determining the differential sensitivity to copper toxicity among heterogeneous populations of *Saccharomyces cerevisiae*
4.1. Introduction

In our previous chapters a major role of plasma membrane fatty acid composition on the susceptibility of \textit{S. cerevisiae} towards the toxic effects of both the redox-active metal copper and the nonredox-active metal cadmium was demonstrated. \textit{Saccharomyces cerevisiae} enriched with the exogenously-supplied polyunsaturated fatty acids (PUFAs) linoleate and linolenate were markedly more susceptible towards heavy metal-induced plasma membrane permeabilisation and whole-cell toxicity (Avery \textit{et al.}, 1996; Chapters 2 and 3). While differences in the metal-sensitivities of unsupplemented and PUFA-supplemented \textit{S. cerevisiae} were examined, the differential susceptibility to metal toxicity observed within heterogeneous yeast cell suspensions was not investigated. In this particular study the nature of the heterogeneity in susceptibility towards metal toxicity observed for exponential phase \textit{S. cerevisiae} suspensions was examined. Thus, the relationship between cell cycle stage-repartition and susceptibility towards metal toxicity was investigated.

Cellular growth is considered as successive phases, characterised by specific biochemical processes and called, from one division to the next, the cell cycle. The yeast cell cycle consists of a DNA replicating phase, S phase, a mitotic phase, M phase, and G\textsubscript{1} and G\textsubscript{2} gap phases. Proper progression through the cell cycle of \textit{S. cerevisiae} requires the periodical expression of regulatory proteins such as the CLN- and B-type cyclins (Cln1p-Cln3p and Clb1p-Clb6p), cyclin-dependent kinases, primarily Cdc28p (Lew \textit{et al.}, 1997), and checkpoint proteins such as Rad9p, Mec1p, and Rad53p (Li and Cai, 1997; Paulovich
et al., 1997; Weinert et al., 1994). Cdc28p levels do not fluctuate during the yeast cell cycle. However, the pattern of cyclin accumulation/degradation leads to overlapping waves of activity of distinct cyclin/Cdc28p complexes, which in turn regulate gene transcription during the cell cycle (Lew et al., 1997).

Studies on cell cycle-specific gene expression are usually carried out using synchronous cultures, or on a relatively small number of cells, by time-lapse studies. Cell cycle-synchrony studies in yeast are performed using a variety of techniques, e.g. α-factor-induced G1 phase arrest, hydroxyurea-induced G1/S phase arrest, and nocodazole-induced G2/M phase arrest (Heichman and Roberts, 1996), sucrose density gradients, or through the use of temperature-sensitive cdc mutants, e.g. temperature-sensitive cdc7 (Siede et al., 1994; Kitada et al., 1992) and cdc15 mutants (Cocker et al., 1996). However, such methods tend to perturb the dynamic state of the cells (Avery et al., 1995b; Lloyd, 1987). Flow cytometry can be used to circumvent cell synchronisation in many cases, and has become the method of choice for cell cycle analysis (Kalejta et al., 1997; Avery et al., 1995b; Jayat and Ratinaud, 1993; Lloyd, 1993). Indeed, Porro et al. (1995) have recently applied flow cytometry to track the dynamics of cell cycle progression of new-born S. cerevisiae using the cell wall- and protein-specific fluorescent dyes, conjugated ConA-FITC and TRITC, respectively. Typically, monoparametric analyses, using cellular DNA content alone, are used to determine the approximate cell cycle stage (Jayat and Ratinaud, 1993). However, by simultaneously monitoring cell volume (proportional to forward-angle light scatter (FSC)) and cellular DNA content, one can more accurately estimate cell repartition in the various cell cycle stages (Lopezamoros et al., 1995; Jayat and Ratinaud, 1993). Therefore, by appropriate FSC-
histogram gate adjustment it is possible to analyse distinct cell cycle phase sub-
populations (of differing cell volumes) within a heterogeneous population (Avery et al., 1995b). Thus, flow cytometry permits distinction of a variety of types of sub-populations within heterogeneous microbial suspensions.

The phenotypic heterogeneity observed within microbial cultures is most likely a consequence of differential gene transcription, such as that regulated through cell cycle-dependent oscillations of cyclin/Cdc28p kinase activity (Lew et al., 1997; Heichmann and Roberts, 1996). The green fluorescent protein (GFP) of the jellyfish Aequorea victoria has been extensively used for both localisation and gene expression studies in S. cerevisiae, including microtubule visualisation, tracking of the kinetics of spindle pole body formation, and determination of expression levels of GAL1 and URA3 (Anderson et al., 1996; Niedanthal et al., 1996). The longer wavelength peak (470 nm) of GFPs bimodal absorption spectrum allows excitation with an argon ion laser (488 nm excitation) used in fluorescence-activated cell sorting (FACS) systems (Cormack et al., 1996; Ropp et al., 1995). Construction of GFP-promoter fusion vectors, and subsequent transformation of yeast allows the expression of the gene of interest to be determined in vivo using GFP as a reporter fluorophore, by flow cytometry. Therefore, using flow cytometry it is possible to analyse distinct cell cycle phase sub-populations for GFP fluorescence, and thus monitor cell cycle-changes in gene expression.

In this present study the differential susceptibility to metal toxicity in exponential phase S. cerevisiae cultures was investigated. Specifically, flow cytometry was used to examine whether repartition in the various cell cycle stages was associated with differential metal sensitivity. Furthermore, in this context, cell cycle stage-dependent
phenotypic alterations in yeast heavy metal resistance determinants, such as altered fatty acid composition, and differential levels of expression of metal homeostasis and oxidative stress response genes were examined.
4.2. Materials and Methods

4.2.1. Organisms and culture conditions. Yeast strains used for cell cycle studies were *S. cerevisiae* NCYC 1383 (*MATα, his3-D1, leu2-3, 112, ura3-52, trp1-289*) and DY150 (*MATα, ura3-52, leu2-3, 112, trp1-1, ade2-1, his3-11, can1-100*) (kindly provided by A. M. Avery, Emory University, Atlanta, GA). Maintenance of yeast strains, and preparation of strains for experimental purposes was as previously described (see Section 2.2.1.).

4.2.2. Preparation of cell homogenates. Cells were harvested by centrifugation at 1,500 g for 5 min and washed twice with distilled deionised water at 4 ºC. Cells were disrupted by shaking with 0.5 mm-diameter glass beads (Sigma, MO) using a Biospec Products Mini-Beadbeater™ (or Mini-Beadbeater-8™ for multiple samples) homogeniser for 4 x 30 s pulses. Samples were placed on ice between pulses.

4.2.3. Lipid extraction and fatty acid analysis. Lipids were extracted from whole-cell homogenates, and fatty acid methyl esters generated as described previously (see Section 2.2.5). In this study, FAMEs were separated using a Stabilwax-DA 30 m capillary column and analysed using a Perkin Elmer Autosystem gas chromatograph. Fatty acids were again identified by comparison with authentic standards.
4.2.4. Preparation of cell suspensions for metal toxicity experiments. Cells from the late-
exponential phase (15 h) were harvested by centrifugation at 1,500 g for 5 min and
washed twice with distilled deionised water. Washed cells were suspended to a density
of approximately $5 \times 10^7$ ml$^{-1}$ in 40 ml of 10 mM Mes buffer (pH 5.5), 1% (w/v) glucose,
and incubated at room temperature with orbital shaking at 120 rev. min$^{-1}$. After 10 min
equilibration, Cu(NO$_3$)$_2$ was added to the desired concentration.

4.2.5. Cell viability. For the flow cytometric assessment of viability, cells were stained with
propidium iodide (PI) (Sigma, MO), and fluorescein diacetate (FDA) (Molecular Probes
Inc., OR). PI is a phenanthridinium intercalator which is taken up by dead cells because
of loss of membrane impermeability (Kalejta et al., 1997; Lopezamoros et al., 1995).
Once bound to intracellular nucleic acids PI-fluorescence is enhanced 20- to 30-fold
(Lopezamoros et al., 1995). PI has an absorption maximum of 536 nm and an emission
maximum of 617 nm (Haugland, 1996). FDA is taken up by live cells and converted to
its fluorescent derivative fluorescein by non-specific intracellular esterases (Breeuwer,
1995). Fluorescein has an absorption maximum of 494 nm and emission maximum of
518 nm (Haugland, 1996).

Aliquots (1.5 ml) from control and Cu$^{2+}$-treated suspensions were taken at regular
intervals, washed twice with 10 mM Mes buffer (pH 5.5), 1% (w/v) glucose, and
resuspended in 1 ml phosphate-buffered saline (PBS) containing 10 $\mu$g ml$^{-1}$ FDA and 25
$\mu$g ml$^{-1}$ PI. Samples were incubated at 4 °C for 15 min prior to analysis. Stained cells
were visualised by fluorescence microscopy using an Olympus BH2 fluorescence
microscope and analysed by flow cytometry (see below).
4.2.6. **Measurement of in vivo molecular oxidation.** Direct evidence of in vivo intracellular molecular oxidation was assessed using the oxidant-sensitive probe 2',7'-dichlorodihydrofluorescein diacetate (H₂DCFDA) (Molecular Probes Inc., OR) (Davidson et al., 1996; Haugland, 1996). Nonionic H₂DCFDA traverses cell membranes and is enzymatically hydrolysed to the non-fluorescent compound H₂DCF by non-specific intracellular esterases (LeBel et al., 1992). H₂DCF reacts with intracellular reactive oxygen species, particularly hydrogen peroxide, and is oxidised to the highly fluorescent derivative 2',7'-dichlorofluorescein (DCF) (Davidson et al., 1996). DCF has an absorption maximum of 510 nm and emission maximum of 532 nm (Haugland, 1996).

Aliquots (1.5 ml) were taken at regular time intervals, washed with 1 ml of 10 mM Mes buffer (pH 5.5), 1% (w/v) glucose, and resuspended in 1 ml PBS containing 10 μl of 5 mM H₂DCFDA. Samples were incubated for 15 min at room temperature, washed twice in ice-cold distilled deionised water and resuspended in 1 ml PBS. Samples were kept at 4 °C until flow cytometric analysis (see below).

4.2.7. **Cellular DNA staining.** Cellular DNA was stained with PI according to the method of Butler et al. (1991), with the following modifications. At specified intervals after the addition of Cu(NO₃)₂ to cell suspensions (see above), 1.5 ml aliquots were removed, washed with 1 ml of 10 mM Mes buffer (pH 5.5), 1% (w/v) glucose, and immediately fixed in 70% (v/v) ethanol for 30 min at 4 °C. Samples were washed once with 1 ml 50 mM sodium citrate and resuspended in 1 ml of the same solution containing 1 mg ml⁻¹
boiled RNase A and 20 μg ml⁻¹ PI. Samples were then incubated at 37 °C for 3 h in the dark and stored at 4 °C until analysis.

4.2.8. Flow cytometry. All samples were analysed using a Becton Dickinson FACSCalibur™ instrument, equipped with a 15 mW, 488 nm argon ion laser. Prior to sample data acquisition FACSCComp software was run. FACSCComp software uses CaliBRITE unlabeled, fluorescein isothiocyanate (FITC)- and phycoerythrin (PE)-labeled beads to check both the instrument sensitivity and alignment, and the bead-positive signal and noise separation for forward-angle light scatter (FSC), side-angle light scatter (SSC) and fluorescence (FL1 and FL2). Fluorescein and 2′,7′-dichlorofluorescein fluorescence were monitored in channel FL1 (lower absorption maximum), while PI fluorescence was monitored in channel FL2. Fluorescence compensation was routinely adjusted to minimise overlapping emission spectra. The detection threshold was set in the FSC channel at a level just below the lowest yeast cell signal (threshold_{FSC} ~ 18). For sample data acquisition in non-sorting applications, cells were diluted in FACSFlow™ sheath fluid to a density of approximately 1 x 10⁷ cells ml⁻¹. Samples were aspirated into a stream of 0.22 μm pre-filtered FACSFlow™ sheath fluid and passed across the argon ion laser. Samples were analysed on high flow rate (60 μl min⁻¹). Typically, 10,000 cells were analysed per sample. For sorting applications, 0.22 μm pre-filtered PBS was used as sheath fluid, and samples were sorted at low flow rate (12 μl min⁻¹). Gated populations were physically separated (sorted) into sterile 50 ml Falcon tubes, previously coated with 4% (w/v) bovine serum albumin (BSA) in PBS for 1 h at 4 °C, to minimise adhesion of cells to the tube surface. Sorted populations were immediately placed on ice,
then centrifuged at 1,500 g for 5 min at 4 °C and resuspended in 0.5 ml PBS for analysis. Data acquisition and analysis were performed using CELLQuest software integrated with a Macintosh Quadra 650 computer with Apple 7.1 operating system software.

4.2.9. Bacterial genetic manipulations. *E. coli* DH5α competent cells (GIBCOBRL, MD) were routinely transformed with plasmid DNA according to standard transformation protocol (Ausubel *et al.*, 1998), with the following modifications. Competent cells (50 μl) were transferred to pre-chilled microcentrifuge tubes and placed on ice. Plasmid DNA (<1 μg) was gently mixed with the competent cells and incubated on ice for 30 min. The cells were heat-shocked at 42 °C for 45-90 sec. Cells were placed on ice for 2 min. Room temperature S.O.C. medium (900 μl), comprising 1.2% (w/v) tryptone, 2.4% (w/v) yeast extract, 0.4% (v/v) glycerol, 0.23% (w/v) KH₂PO₄, 1.25% (w/v) K₂HPO₄ (GIBCOBRL, MD), was added and the mixture incubated at 37 °C for 2 h with shaking at 225 rev. min⁻¹. Aliquots were spread-plated onto LB medium containing 50 μg ml⁻¹ ampicillin (Sigma, MO) and incubated at 37 °C overnight. Transformation efficiencies were determined using pUC19 DNA (0.1 μg ml⁻¹) as control plasmid. Plasmids were isolated from bacterial strains using the standard alkaline lysis mini-prep procedure (Ausubel *et al.*, 1998). Restriction enzyme digestions were performed according to standard protocol (Ausubel *et al.*, 1998). DNA fragments were typically separated by agarose gel electrophoresis performed on 1% (w/v) agarose (GIBCOBRL, MD) mini-gels, using Pharmacia Biotech gel electrophoresis apparatus (GNA-100 or GNA-200) coupled to a Pharmacia Biotech electrophoresis power supply EPS-200. Agarose gels were run.
for 45 min at 75 V and 60 mA. Hi-Lo™ DNA marker (Minnesota Molecular, MN) was used to track DNA bands.

PCR amplification was performed in 100 µl reaction volumes containing 1 x PCR reaction buffer (Boehringer Mannheim, IN), 1 µg ml⁻¹ plasmid DNA or 10 µg ml⁻¹ chromosomal DNA as template, 1 µM of each primer, 200 µM each of 2'-deoxythymidine-5'-triphosphate (dTTP), 2'-deoxyadenosine-5'-triphosphate (dATP), 2'-deoxyguanosine-5'-triphosphate (dGTP) and 2'-deoxycytidine-5'-triphosphate (dCTP), 100 U ml⁻¹ of Taq DNA polymerase (Boehringer Mannheim, IN), and nuclease-free sterile, distilled deionised water. PCR was performed in a Perkin Elmer Cetus DNA thermal cycler. For all reactions, strand denaturation was carried out at 94 °C for 3 min, and was followed by five preliminary cycles of 94 °C for 60 s, 40 °C for 60 s and 72 °C for 90 s, and 25 cycles of 94 °C for 60 s, 45 °C for 60 s and 72 °C for 90 s. PCR products were purified using phenol/chloroform/isoamyl alcohol (25:24:1 v/v/v) and ethanol precipitated prior to gel electrophoresis and/or restriction enzyme analysis.

4.2.10. Yeast genetic manipulations. Yeast cells were transformed with plasmid DNA using the Frozen-EZ™ yeast transformation kit (Zymo Research Inc., CA).

Both plasmid and chromosomal DNA were isolated from yeast according to standard protocols (Ausubel et al., 1998), with the following modifications. Cell pellets were resuspended in 400 µl breaking buffer (2% (v/v) triton X-100, 1% (w/v) SDS, 100 mM NaCl, 10 mM Tris.Cl (pH 8.0), 1 mM EDTA) and 400 µl phenol/chloroform/isoamyl alcohol (25:24:1 v/v/v). The cell suspension was transferred to a 2 ml screw-cap microcentrifuge vial filled one-half to two-thirds full with 0.5 mm diameter glass beads
Yeast cells were disrupted using a Biospec Products Mini-Beadbeater™ (or Mini-Beadbeater –8™ for multiple samples) homogeniser for 4 x 30 s pulses. Yeast suspensions were placed on ice between pulses. For yeast plasmid purification, the aqueous phase was extracted and 2 μl used to transform *E. coli* DH5α cells (see above). Chromosomal DNA was precipitated with 95% ethanol and washed once with 70% ethanol. The pellet was dried under vacuum and resuspended in 100 μl sterile, distilled deionised water.

4.2.11. Plasmid construction. The *CUPl::GFP* transcriptional promoter gene fusion plasmid, pYEX-GFP*CUPl* (kindly provided by A. M. Avery, Emory University, Atlanta, GA) had been constructed by ligation of the 0.72 kb green fluorescent protein (GFP) coding sequence of *Aequorea victoria* to the large EcoRI-XhoI fragment of pYEX 4T-1 (Clontech, CA), downstream of the Cu²⁺-inducible *CUPl* promoter.

The *SODI::GFP* gene fusion plasmid, pYEX-GFP*SDPl* was constructed by first amplifying a 0.6 kb *SODI* promoter sequence, from −600 to +5 (relative to the ATG start codon). The 0.6 kb fragment was amplified from genomic DNA using the following oligonucleotide primers in a PCR reaction:

\[
\begin{align*}
5'\text{-GTGGGTTCAGAATACTACCAATGGTGC-3'} \\
\text{(XbaI)} \\
5'\text{-ACTGCGACTGCGAATTCCATTATAAATTAA-3'} \\
\text{(EcoRI)}
\end{align*}
\]

The amplified fragment was then digested with XbaI and EcoRI, and ligated to the large XbaI-EcoRI fragment of pYEX-GFP*CUPl*, according to the following ligation protocol.
Following digestion with *XbaI* and *EcoRI*, the *SOD1* promoter fragment was purified with phenol/chloroform/isoamyl alcohol (25:24:1 v/v/v) and ethanol precipitated. 

*XbaI/EcoRI*-digested pYEX-GFP<sub>CUP1</sub> was ethanol precipitated and calf intestinal-alkaline phosphatase- (Promega, WI) treated to prevent re-circularisation. Phosphatase-treated vector DNA was subsequently purified with phenol/chloroform/isoamyl alcohol (25:24:1 v/v/v) and ethanol precipitated prior to the ligation reaction. The ligation reaction was performed in a 10 µl reaction volume containing 1 x T4 DNA ligase buffer (Promega, WI), 300 U ml<sup>−1</sup> T4 DNA ligase (Promega, WI), approximately 17 µg ml<sup>−1</sup> vector DNA and 7 µg ml<sup>−1</sup> insert DNA (1:5 molar ratio of vector:insert DNA ends) and nuclease-free sterile, distilled deionised water. The ligation reaction was performed at 16 °C overnight. 

*E. coli* DH5α competent cells were transformed with 5 µl of the ligation reaction mixture, as described above. Insertion of the 0.6 kb *SOD1* promoter fragment created a transcriptional fusion with the *GFP* gene.
4.3. Results

4.3.1. Relationship between percentage maximum cell volume, cellular DNA content and cell cycle stage. In order to establish a direct correlation between cell volume and cell cycle stage, the cellular DNA content of ten different fractions, distinguished by cell volume, of an exponential phase *S. cerevisiae* NCYC 1383 suspension was monitored using flow cytometry. *Saccharomyces cerevisiae* NCYC 1383 was selected for this study and subsequent toxicity experiments because of its well characterised interactions with heavy metals (Avery *et al.*, 1996; Chapter 2). Cell volume increases as cells progress through the cell cycle (Lord and Wheals, 1980). Forward-angle light scatter (FSC) histograms correspond to cell volume distributions in heterogeneous yeast cell suspensions (Lloyd, 1992). Cell volume is directly proportional to FSC signal. Hence, by simultaneously monitoring cellular FSC signal and PI fluorescence intensity (FL2) (PI fluorescence is proportional to cellular DNA content), it was hoped to be able to demonstrate a correlation between cell volume and cell cycle stage. An FSC histogram was plotted, which demonstrated the normal distribution of cell volume, and divided into ten regions (R2-R11), each representing 10% of the total population (10,000 cells region\(^1\)) (Fig. 4.1). Individual FL2 histograms were subsequently plotted for each of the 10% regions (Fig. 4.2c-I). An FSC-SSC dot plot of the total heterogeneous yeast cell population is shown in Figure 4.2a. The cellular DNA content of the total population is displayed in Figure 4.2b. Two distinctly resolved peaks can be observed, which
Fig. 4.1. Forward-angle light scatter (FSC) histogram of *S. cerevisiae* NCYC 1383. The FSC histogram representing the total cell population has been gated into 10 x 10% regions (R2-R11). The FSC histogram contains data collected from at least 100,000 cells, and each region represents approximately 10,000 cells.
correspond to cells comprising 1C and 2C DNA contents. The number of cells consisting of 1C (G1 phase) and 2C (G2/M phase) DNA were approximately equal (Fig. 4.2b). The cellular DNA contents of regions R2 through R11, from Figure 4.1 are shown in Figures 4.2c-l. Cells in the 0-10% maximum cell volume fraction (R2) consist exclusively of 1C DNA, and thus correspond to cells in G1 phase (Fig. 4.2c). Cells at 10-20% of maximum cell volume (R3) consist largely of 1C DNA, however a small 2C DNA peak, corresponding to 1% of the total number of cells, was observed. It can be inferred that cells approaching 20% maximum cell volume have reached the critical threshold size necessary to traverse START and enter S phase (S phase cells have a cellular DNA content between 1 and 2C) (Lew et al., 1997). Figures 4.2e, f and g depict progression through S phase and emergence of the 2C DNA peak. Cells at 40-50% of maximum cell volume (R6) have a larger 2C DNA peak, corresponding to 53% of the total number of cells, and thus consist predominantly of G2/M phase cells (Fig. 4.2g). Cells 90-100% maximum cell volume consist almost exclusively of 2C DNA and thus most likely correspond to cells approaching the M/G1 phase boundary (Fig. 4.2l). The results clearly demonstrate the relationship between cell volume and cellular DNA content, and thus validate the use of FSC measurements (cell volume) for cell cycle analysis.

4.3.2. Relationship between cell volume (cell cycle stage) and susceptibility to copper toxicity. Here we sought to determine if sensitivity to copper toxicity was influenced by cell cycle stage. Cells previously grown in unsupplemented medium were exposed to 100 μM Cu2+ for 15 min. Samples were stained with FDA and PI and analysed by flow
Fig. 4.2. Relationship between % maximum cell volume and cellular DNA content (cell cycle stage) in *S. cerevisiae* NCYC 1383. Figure 4.2a represents a forward-angle light scatter (FSC) versus side-angle light scatter (SSC) dot plot of the exponential phase population. A region gate (R1) was applied to separate cells from cell debris. Figure 4.2b represents the propidium iodide fluorescence (FL2) histogram of the total population (R1). Figure 4.2c-l represent FL2 histograms for regions 2 to 11, derived from the FSC histogram (Fig. 4.1). Each histogram contains data collected from approximately 10,000 cells.
cytometry. At least 100,000 cells were analysed per sample. An FSC histogram was plotted and divided into fifty 2% regions (defined as gates) (see Fig. 4.1). An FL1 versus FL2 dot plot was subsequently prepared for each of the fifty 2% regions to determine the number of viable cells within each region (see Fig. 4.4a). FL1 versus FL2 region statistics were then used to construct a plot of % viability against % maximum cell volume (Fig. 4.3). The largest cells (98-100% of maximum cell volume) were the most resistant to copper, whereby after 15 min exposure to 100 μM Cu²⁺, approximately 69% of these cells remained viable. Interestingly, the smallest cells (0-2% of maximum cell volume) were also relatively resistant to copper, whereby after 15 min exposure to 100 μM Cu²⁺, approximately 57% of these cells remained viable. However, as % of maximum cell volume increased beyond 0-2%, Cu²⁺-sensitivity increased and was maximal at 38-40% of maximum cell volume, whereby only 32% of cells remained viable after Cu²⁺-exposure (Fig. 4.3). With increased cell volume beyond this point, % viability increased to a maximum with the largest cells. These results indicate differing susceptibilities of cells at different stages of the cell cycle towards copper toxicity.

4.3.3. Cellular DNA content of *S. cerevisiae* NCYC 1383 exposed to a toxic concentration of copper. To further characterise the relationship between cell cycle stage and sensitivity towards copper toxicity, *S. cerevisiae* was exposed to a toxic concentration of copper, and the cellular DNA content of sensitive and resistant sub-populations was compared. Cells previously grown in unsupplemented YEPD medium were exposed to 100 μM Cu²⁺ for 15 min. Samples were stained with FDA and PI and analysed by flow cytometry. An FL1 versus FL2 dot plot was created to distinguish between live and dead
**Fig. 4.3.** Relationship between cell volume (cell cycle stage) and susceptibility to copper toxicity in *S. cerevisiae*NCYC 1383. Cells were exposed to 100 μM Cu(NO₃)₂ for 15 min, and viability was determined by staining with FDA (FL1) and PI (FL2). An FSC histogram of the total population was gated into 50 x 2% regions. The number of viable cells of each region was determined from gate statistics attained from an FL1 versus FL2 dot plot. Data was collected from at least 100,000 cells, and each region represents approximately 2,000 cells.
sub-populations (Fig. 4.4a). Dead and live populations were gated R1 and R2, respectively. Both regions were sorted (physically separated into different Falcon tubes), fixed, and cellular DNA was stained with PI. The cellular DNA content of the copper-sensitive sub-population (R1) is shown in Figure 4.4b. Of the copper-sensitive sub-population, approximately 53% of cells comprised 2C DNA, as indicated by the slightly greater 2C DNA peak (Fig. 4.4b). The cellular DNA content of copper-resistant cells (R2) is shown in Figure 4.4c. The copper-resistant sub-population consisted predominantly of G1 phase cells, whereby approximately 72% of cells comprised 1C DNA. These results indicate that cells at the cell cycle stage corresponding to a 1C DNA content (G1 phase) are considerably less sensitive to the toxic effects of copper. It was also evident that the copper-resistant sub-population also consisted of a small proportion of 2C DNA cells (approximately 28% of the total number of cells) (Fig. 4.4c). It could be inferred from Figure 4.3 that this 2C DNA peak corresponds to cells of maximum cell volume. These results further demonstrate the differing sensitivities of cells of different cell cycle stage to the toxic effects of copper.

4.3.4. Fatty acid composition of cells at different cell cycle stage. The increased susceptibility of cells displaying higher degrees of membrane fatty acid unsaturation to heavy metal toxicity had previously been demonstrated (Avery et al., 1996; Chapters 2 and 3). Here we sought to ascertain whether the above differences in the sensitivities of cells at different stages of the cell cycle towards copper toxicity could be attributed to cell cycle-induced changes in fatty acid composition. *Saccharomyces cerevisiae* NCYC 1383
Fig. 4.4. Cellular DNA content of live and dead *S. cerevisiae* NCYC 1383 after exposure to copper. Cells were exposed to 100 μM Cu(NO₃)₂ for 15 min, and viability was determined by staining with FDA (FL1) and PI (FL2). Figure 4.4a represents an FL1 versus FL2 dot plot indicating dead (R1) and live (R2) sub-populations. R1 and R2 were sorted, the cellular DNA of each sub-population was stained with PI, and FL2 histograms were plotted (Fig. 4.4b,c). Data was collected from at least 10,000 cells.
previously grown in unsupplemented YEPD medium was analysed by flow cytometry. An FSC histogram was plotted and divided into four 25% regions (R1-R4), representing cells at different stages of the cell cycle (see Fig. 4.1). Each region was physically sorted, cells were resuspended in PBS, fatty acids were extracted, and FAMEs were analysed by gas-liquid chromatography as described previously (see Section 4.2.3). The cellular fatty acid compositions of regions 1, 2, 3 and 4 are displayed in Table 4.1. Levels of the monounsaturated fatty acids, 16:1 and 18:1 were similar for each region, only varying between approximately 21 and 26%, and 22 and 26% of total cellular fatty acids, respectively (Table 4.1). Thus, the unsaturation indices (average number of double bonds per fatty acid) were found to be similar, ranging between 0.44 and 0.49, for each of the sorted regions. The results indicate that the cellular fatty acid composition of *S. cerevisiae* remains relatively constant throughout the cell cycle, and the small differences that are evident appear too small to significantly influence Cu$^{2+}$-sensitivity (Chapter 3).

4.3.5. Cellular DNA content of *S. cerevisiae* NCYC 1383 previously stained with the oxidative stress probe 2',7'-dichlorodihydrofluorescein diacetate (H$_2$DCFDA). The oxidant status of cells at different stages of the cell cycle was next examined, to determine if repartition in the various cell cycle stages was associated with differences in endogenous levels of oxy-radicals. *Saccharomyces cerevisiae* NCYC 1383 was stained with the oxidant-sensitive probe H$_2$DCFDA and examined by flow cytometry. An FL1 histogram was plotted and on the basis of FL1 intensity, three sub-populations of low (R1), medium (R2) and high (R3) cellular oxidant status were gated and sorted for
Table 4.1. Cellular fatty acid composition of *S. cerevisiae* NCYC 1383 at different stages of the cell cycle

<table>
<thead>
<tr>
<th>Fatty acid</th>
<th>Region</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>R1</td>
</tr>
<tr>
<td>16:0</td>
<td>32.4±0.6</td>
</tr>
<tr>
<td>16:1</td>
<td>23.0±0.5</td>
</tr>
<tr>
<td>18:0</td>
<td>19.9±0.4</td>
</tr>
<tr>
<td>18:1</td>
<td>24.7±0.8</td>
</tr>
<tr>
<td>18:2</td>
<td>tr</td>
</tr>
<tr>
<td>U.I.</td>
<td>0.48</td>
</tr>
</tbody>
</table>

Fatty acids are abbreviated with the first figure representing the number of carbon atoms in the fatty acyl chain and the second representing the number of double bonds. U.I. refers to the unsaturation index (average number of double bonds per fatty acid). Values for percentage fatty acid composition are means from two replicate determinations ± SD. (tr. < 0.1).
Fig. 4.5. Cellular DNA content of *S. cerevisiae*NCYC 1383 previously stained with the oxidant-sensitive probe 2',7'-dichlorodihydrofluorescein diacetate (H$_2$DCFDA) (FL1).

Figure 4.5a represents an FL1 histogram gated on regions R1, R2 and R3 of low, medium and high FL1 intensity, respectively. R1, R2 and R3 were sorted, the cellular DNA of each sub-population was stained with PI, and FL2 histograms were plotted (Fig. 4.5b,c,d). Data was collected from at least 10,000 cells.
subsequent determination of the cells' DNA content (Fig. 4.5a). R1 (low oxidant status) comprised predominantly of 1C DNA cells (64% of the total number of cells), with a small proportion of 2C DNA cells (Fig. 4.5b). R2 (medium oxidant status) consisted of a greater proportion of 2C DNA cells (61% of the total number of cells) (Fig. 4.5c). Finally, R3 (high oxidant status) consisted predominantly of 2C DNA cells (69% of total number of cells), with a small proportion of 1C DNA cells (Fig. 4.5d). These results suggest that within a heterogeneous yeast population the generation of oxy-radicals, by normal metabolic processes, occurs to a greater extent at later cell cycle stages.

4.3.6. Influence of initial oxidant status on the susceptibility of *S. cerevisiae* NCYC 1383 towards copper toxicity. Here we sought to determine if differences in the levels of endogenously generated oxy-radicals of different cell cycle stage sub-populations could be correlated with differential susceptibility towards copper toxicity. Cells were stained with H2DCFDA, and three different sub-populations were sorted as described above (see Fig. 4.5a). Cells were harvested and resuspended to a density of approximately 2 x 10⁵ cells ml⁻¹ in PBS, and exposed to 10 μM Cu²⁺ for 10 min. Samples were subsequently stained with PI and re-examined by flow cytometry to determine the number of viable cells of each sub-population. The % viability of regions 1, 2 and 3 of unsupplemented, 18:2- and 18:3-supplemented cultures after 10 min exposure to copper is shown in Figure 4.6. It is evident from the graph that cells of region 3 (high initial oxidant status) are the most susceptible to copper toxicity. Hence, with cells previously grown in unsupplemented medium, after 10 min exposure to 10 μM Cu²⁺ 72% of cells of region 3
**Fig. 4.6.** Influence of initial oxidant status on the susceptibility of *S. cerevisiae* NCYC 1383 to copper toxicity. Cells were stained with the oxidant-sensitive probe H$_2$DCFDA (FL1) and on the basis of initial oxidant status three different sub-populations of low (R1), medium (R2) and high (R3) FL1 intensity were sorted (see Fig. 4.5a). Each sub-population was subsequently exposed to 10 μM Cu(NO$_3$)$_2$ for 10 min and the number of viable cells of each sub-population was determined by staining with FDA and PI. Figure 4.6 shows the % viability of regions 1, 2 and 3 of unsupplemented (○), linoleate- (●) and linolenate- (□) supplemented cells after 10 min exposure to 10 μM Cu(NO$_3$)$_2$. 
remained viable compared with 93% of cells of region 1. However, this effect was greatly accentuated with cells previously grown in 18:3-supplemented medium. Here only 17% of cells of region 3 remained viable after 10 min exposure to 10 \( \mu \text{M} \) Cu\(^{2+} \), compared with 76% of cells of region 1. The results clearly demonstrate that cells with high initial oxidant status are predisposed to killing in the presence of toxic concentrations of copper.

4.3.7. *CUP1* expression in *S. cerevisiae* DY150 exposed to copper. The differential sensitivity of different cell cycle stage fractions towards copper toxicity had previously been demonstrated. Here we sought to determine whether the cell cycle stage-dependency of susceptibility towards copper toxicity could be attributed to heterogeneous basal levels of *CUP1* metallothionein expression. While ideally promoter fusion experiments would also have been performed with *S. cerevisiae* NCYC 1383, this strain was particularly difficult to transform with plasmid DNA. Thus, the yeast strain *S. cerevisiae* DY150 (Clontech, CA) was selected for molecular biological experimentation because of its ease of transformation relative to NCYC 1383. DY150 was transformed with pYEX-GFP\(_{CUP1}\) (Fig. 4.7a), an episomal plasmid containing the *GFP* coding sequence under the control of the *CUP1* promoter. Thus, *CUP1* promoter activity could be monitored by GFP fluorescence (FL1) using flow cytometry. Transformed DY150 was exposed to 10 \( \mu \text{M} \) Cu(NO\(_3\))\(_2\) for 10 min to preclude *CUP1* induction, cells were stained with PI (FL2) and analysed by flow cytometry. Exposure to 10 \( \mu \text{M} \) Cu(NO\(_3\))\(_2\) for 10 min resulted in approximately 64% toxicity as demonstrated by the large proportion of
**Fig. 4.7.** The plasmid pYEX-GFP_{CUP1} (a) was constructed by ligation of the 0.72 kb green fluorescent protein (GFP) coding sequence of *Aequorea victoria* to the large EcoRI-XhoI fragment of pYEX 4T-1 (Clontech, CA). The plasmid pYEX-GFP_{SOD1} (b) was constructed by ligating a 0.6 kb *SOD1* promoter sequence, from −600 to +5 (relative to the ATG start codon), to the large XbaI-EcoRI fragment of pYEX-GFP_{CUP1}. 
Fig. 4.8. *CUP1* expression in *S. cerevisiae* DY150 exposed to copper. Cells were exposed to 10 μM Cu(NO₃)₂ for 10 min and stained with PI (FL2). Figure 4.8a represents an FL1 (GFP) versus FL2 dot plot indicating dead (R1) and live (R2) sub-populations. Figure 4.8b shows the FL1 histogram of dead cells (R1) while figure 4.8c shows the FL1 histogram of live cells (R2). Data was collected from at least 10,000 cells.
PI-stained cells (Fig. 4.8a). The GFP fluorescence signal of dead cells (R1) is shown in Figure 4.8b, while that of live cells (R2) is shown in Figure 4.8c. The GFP fluorescence signal for both sub-populations demonstrated considerable heterogeneity, indicative of fluctuations in episomal plasmid copy number during the cell cycle (Mason, 1991).

However, there were no significant differences in basal levels of \textit{CUP1} promoter activity, as determined by GFP fluorescence, between live and dead sub-populations. Therefore, the results indicate that differences in basal levels of \textit{CUP1} expression do not account for differential sensitivity towards copper toxicity. One would expect to observe increased live cell-GFP fluorescence for a gene whose differential expression was associated with increased Cu\textsuperscript{2+}-resistance.

Several reports have indicated that \textit{CUP1} is an inducible component of the \textit{S. cerevisiae} oxidative stress response system (Jensen \textit{et al.}, 1996; Liu and Thiele, 1996). Here we also sought to confirm Cu\textsuperscript{2+}-induced \textit{CUP1} transcription. DY150 transformed with pYEX-GFP\textit{CUP1} was exposed to 10 \textmu{}M Cu(NO\textsubscript{3})\textsubscript{2} for up to 1 h, cells were stained with PI and analysed by flow cytometry. Exposure to 10 \textmu{}M Cu(NO\textsubscript{3})\textsubscript{2} for 1 h resulted in approximately 70\% killing, as demonstrated by the high proportion of PI-stained cells (R1) (Fig. 4.9a). GFP fluorescence in the absence of Cu\textsuperscript{2+} (t = 0) and after 1 h exposure to 10 \textmu{}M Cu\textsuperscript{2+} (t = 1 h) is displayed in Figure 4.9b. Thus, exposure to 10 \textmu{}M Cu(NO\textsubscript{3})\textsubscript{2} for 1 h results in both an increase and decrease in GFP fluorescence intensity as demonstrated by the lengthening of the GFP peak (Fig. 4.9b). GFP fluorescence after 1 h exposure to Cu\textsuperscript{2+}, gated on R1 and R2 is shown in Figure 4.9c. As can be seen from the histogram, live cells (R2) account for the high GFP fluorescence signal, while dead cells (R1) predominantly account for the reduction in GFP fluorescence signal. The results
Fig. 4.9. CUP1 induction in *S. cerevisiae* DY150 exposed to copper. Cells were exposed to 10 μM Cu(NO₃)₂ for 1 h and stained with PI (FL2). Figure 4.9a represents an FL1 (GFP) versus FL2 dot plot indicating dead (R1) and live (R2) sub-populations. Figure 4.9b shows the FL1 histogram of cells prior to Cu²⁺ addition (t = 0), and after 1 h exposure to Cu²⁺ (t = 1 h). Figure 4.9c shows the FL1 histogram of both dead (R1) and live (R2) cells, after 1 h exposure to Cu²⁺. Data was collected from at least 10,000 cells.
demonstrate induction of the CUP1 promoter in S. cerevisiae in the presence of 10 μM Cu²⁺. Furthermore, the results demonstrate that live cells have higher levels of CUP1 activity, as evidenced by their higher GFP signal. It is most likely that the decrease in GFP fluorescence observed for the dead cells is a consequence of membrane permeabilisation. GFP is normally found in the cytoplasm and leaks upon prolonged incubation with membrane permeabilising-agents, such as copper (Kalejta et al., 1997).

4.3.8. SOD1-expression in S. cerevisiae DY150 exposed to copper. The yeast strain DY150 was also transformed with pYEX-GFP SOD1 (Fig. 4.7b), containing the GFP coding sequence under the control of the SOD1 promoter, in order to determine whether differences in basal levels of SOD1 expression could account for the differential susceptibility of different cell cycle stage fractions to copper toxicity. Transformed DY150 was exposed to an approximately 50% cytotoxic dose of Cu(NO₃)₂ for 10 min to preclude SOD1 induction. Cells were stained with PI and analysed by flow cytometry (Fig. 4.10a). Thus, within a heterogeneous population SOD1 appears to be constitutively expressed to varying degrees (Fig. 4.10a). The GFP fluorescence signal of dead cells (R1) is shown in Figure 4.10b, while that of live cells (R2) is shown in Figure 4.10c. Two distinct GFP peaks were evident for the live cells; a sub-population with high levels of GFP fluorescence (39% of the total number of cells), and a smaller sub-population with no appreciable levels of GFP fluorescence (10% of the total number of cells). It is most likely that the smaller sub-population corresponds to plasmid-free cells, as yeast episomal plasmids are highly unstable during growth on non-selective medium (Mason, 1991). Indeed, also using flow cytometry, Hjortsu et al. (1985) reported approximately
12% plasmid-free yeast cells during exponential batch growth. In addition, yeast cells not expressing GFP have a low level of autofluorescence when irradiated with the 488 nm argon ion laser (Niedenthal et al., 1996). While a statistically significant difference between the GFP fluorescence signal of live and dead sub-populations could not be established, the results are not as outright as those of the CUP1 experiments (Fig. 4.10b,c). The results are complicated by the fact that the emission spectra of PI and GFP overlap (an inherent disadvantage of the simultaneous use of red and green fluorophores for flow cytometric analysis), and thus PI-fluorescence is also detected by the FL1 detector (Haugland, 1996). Therefore, it remains to be elucidated whether heterogeneous basal levels of SOD1 activity play a significant role in the cell cycle stage-dependency of susceptibility towards copper toxicity.

We also sought to determine if SOD1 promoter expression could be induced upon prolonged exposure to either Cu\(^{2+}\) or the specific superoxide anion generating compound menadione. Transformed DY150 was exposed to 10 μM Cu\(^{2+}\) or 2 mM menadione for 1 h, cells were stained with PI and analysed by flow cytometry (Fig. 4.11). An FL1 versus FL2 dot plot of transformed DY150 prior to toxin exposure is shown in Figure 4.11a. Again two populations are evident; a smaller sub-population with no appreciable levels of GFP fluorescence (R1) (11% of the total number of cells) and a larger sub-population displaying appreciable levels of GFP fluorescence (R2). Again, it is most likely that the smaller sub-population corresponds to plasmid-free cells (see above). Figure 4.11b and c represent FL1 versus FL2 dot plots of transformed DY150 after 1 h exposure to 10 μM Cu\(^{2+}\) and 2 mM menadione, respectively. Exposure to 10 μM Cu\(^{2+}\) and 2 mM menadione for 1 h resulted in approximately 65% (R3) and 56% (R4) killing, respectively (Fig.
4.10b,c). However, exposure to either toxin did not appear to result in \textit{SOD1} induction (Fig. 4.11b,c).
Fig. 4.10. SOD1 expression in S. cerevisiae DY150 exposed to copper. Cells were exposed to 10 μM Cu(NO₃)₂ for 10 min and stained with PI (FL2). Figure 4.10a represents an FL1 versus FL2 dot plot indicating dead (R1) and live (R2) sub-populations. Figure 4.10b shows the FL1 histogram of dead cells (R1) while figure 4.10c shows the FL1 histogram of live cells (R2). Data was collected from at least 10,000 cells.
Fig. 4.11. Exposure of *S. cerevisiae* DY150, transformed with pYEX-GFP<sub>SOD1</sub>, to copper and menadione. Cells were exposed to 10 µM Cu(NO<sub>3</sub>)<sub>2</sub> and 2 mM menadione for 1 h, and stained with PI (FL2). Figure 4.11a represents the FL1 (GFP) versus FL2 dot plot of DY150 prior to toxin exposure. Figure 4.11b represents an FL1 versus FL2 dot plot after 1 h exposure to copper while figure 4.11c depicts 1 h exposure to menadione.
In this study we describe the use of flow cytometry for the analysis of specific cell cycle stage populations within heterogeneous populations of *S. cerevisiae*. More specifically we have examined the cell cycle stage-dependency of susceptibility towards copper toxicity in *S. cerevisiae*. Studies on specific cell cycle stage populations are normally carried out following cell synchronisation procedures (Porro *et al.*, 1995; Lloyd, 1987). However, conventional methods employed to induce synchronous populations often perturb cell population dynamics (Avery *et al.*, 1995b; Porro *et al.*, 1995; Lloyd, 1987). Thus, results obtained using conventional cell synchronisation methods often may not accurately reflect the normal physiological state of cells. Using flow cytometry, we have simultaneously monitored forward-angle light scatter (FSC), which is proportional to cell volume, and propidium iodide fluorescence, which is proportional to cellular DNA content. When cells reach a critical threshold size, and if environmental conditions are favourable, cells traverse START, DNA replication is initiated and cellular DNA content increases from 1C DNA during G1 phase, to between 1C and 2C DNA during S phase, to 2C DNA during G2/M phase, whereby cells undergo mitosis to produce haploid progeny (Lew *et al.*, 1997). By simultaneously monitoring both parameters we thus validated the use of FSC measurements for the analysis of specific cell cycle stage populations. Furthermore, by appropriate 'gate' setting on FSC histograms it was possible to physically separate and analyse cells differentiated by volume, i.e. cell populations at different stages of the cell cycle. A novel flow cytometric method for monitoring the
promoter activity of key oxidative stress response proteins of specific sub-populations, using the green fluorescent protein of *Aequorea victoria* was also described. Cell cycle-analysis using flow cytometry is relatively non-perturbing and facilitates rapid analysis of distinct cell cycle stage populations, and thus circumvents many of the problems encountered with conventional cell synchronisation methods.

Using the above methodology, we initially sought to examine the differential susceptibility to copper toxicity observed within a heterogeneous population exposed to a semi-lethal Cu²⁺ concentration. A distinct relationship between cell cycle stage and susceptibility to copper toxicity was observed, whereby cell populations consisting predominantly of G₂/M phase cells, excluding those approaching maximum cell volume, were the most susceptible to the toxic effects of copper. In chapters 2 and 3 we demonstrated that the sensitivity of *S. cerevisiae* towards both the redox-active metal copper and the non-redox-active metal cadmium is greatly influenced by cellular fatty acid composition. However, the elevated copper-sensitivity of cell fractions consisting predominantly of G₂/M phase cells could not be attributed to changes in cellular fatty acid composition, as cellular fatty acid composition did not vary significantly throughout the cell cycle.

The largest volume cell fraction, consisting exclusively of 2C DNA cells, i.e. budded cells about to undergo mitosis (M/G₁ phase boundary), were found to be the most resistant to copper. The smallest volume cell fractions, consisting almost exclusively of 1C DNA cells (G₁ phase cells) were also found to be relatively resistant towards copper toxicity. However, as cells traversed START and progressed through S phase, sensitivity towards copper toxicity increased to a maximum at a point corresponding to equal
proportions of $G_1$ and $G_2/M$ phase cells. Cellular DNA staining of live and dead cells following exposure to an approximately 50% cytotoxic dose of copper revealed a predominance of $G_1$ phase cells in the live sub-population, with a small proportion of 2C DNA cells. Live 2C DNA cells corresponded to cells at the M/$G_1$ phase boundary.

Staining with the oxidant-sensitive probe 2',7'-dichlorodihydrofluorescein diacetate (H$_2$DCFDA), followed by cellular DNA staining revealed greater intracellular basal levels of reactive oxygen species in cell fractions consisting predominantly of $G_2/M$ phase cells. ROS are generated endogenously in *S. cerevisiae* through mitochondrial respiration, steroidogenesis, and the $\beta$-oxidation of high-molecular weight fatty acids (Rapoport *et al*., 1998; Davidge *et al*., 1995). Copper toxicity experiments confirmed that the high oxidant status of cell fractions consisting predominantly of $G_2/M$ cells correlated with their increased susceptibility towards copper toxicity. Numerous studies have examined the relationship between metal ion homeostasis and toxicity, and oxy-radical metabolism in yeast (Lapinskas *et al*., 1995; Liu and Culotta, 1994; Greco *et al*., 1990). Furthermore, in chapter 3 we demonstrated oxy-radical-mediated lipid peroxidation as a major means of heavy metal toxicity in *S. cerevisiae*. Thus, the results presented here are again suggestive of a major role of oxy-radical-mediated cellular damage during heavy metal exposure in yeast.

While elevated endogenous levels of ROS correlated with the increased sensitivity of cell fractions consisting predominantly of $G_2/M$ phase cells towards copper toxicity, this relationship did not hold true for cells approaching maximum cell volume. The late-exponential phase asynchronous yeast cultures used during this study consisted of a small proportion of older stationary phase cells (initial stationary phase inoculum).
Entry to stationary phase is known to be associated with increased resistance towards oxidative stress (Jamieson et al., 1994; Jamieson, 1992). The oxidative stress response genes CTT1, CTA1, CCP1, SOD1, and SOD2 are all repressed by glucose (Krems et al., 1995). Following glucose exhaustion and respiratory adaptation during yeast batch growth, de-repression of antioxidant genes occurs, thus contributing to the increased oxidative stress-resistance of stationary phase yeast (Steels et al., 1994). De-repression of antioxidant genes facilitates tolerance to the greater levels of ROS that are generated during respiratory growth. Thus, despite the greatest endogenous levels of ROS being evident in cells approaching maximum cell volume (oldest cells), greater resistance towards the toxic effects of copper could be attributed partly to the presence of elevated levels of oxidative stress response proteins, prior to copper exposure.

Oxy-radical induced DNA damage has been implicated as a major means of copper toxicity (Flowers et al., 1997; Lloyd et al., 1997). The hydroxyl radical is a key culprit in DNA damage, generating several base lesions in double-stranded DNA including thymine glycol, 8-oxoguanine and formamido-pyrimidine as well as other base oxidation products (Brennan et al., 1994; Hutchinson, 1985). As stated previously, G1 phase cells (1C DNA), were also found to be relatively resistant towards the toxic effects of copper. Early G1 phase cells have not yet reached the critical threshold cell size necessary to traverse START and enter S phase (Lew et al., 1997). Maintenance of genome integrity during early G1 phase, and during periods of quiescence is facilitated by chromatin condensation and basal transcription repression (Kadonaga, 1998). Previous studies have demonstrated that the physical distance between the compact-aggregates of ‘condensed’ or ‘native’ chromatin is greater than the effective damaging range of HO'
radicals (Nygren *et al.*, 1995). Furthermore, the water concentration of condensed chromatin aggregates has been shown to be too low to facilitate HO' radical-mediated DNA damage (Nygren *et al.*, 1995). Therefore, the elevated resistance of cell fractions consisting predominantly of G1 phase cells could possibly be attributed in part to condensed chromatin assembly (Kadonaga, 1998; Demple and Harrison, 1994; Ljungman and Hanawalt, 1992). Furthermore, chromatin condensation occurs during late G2 phase prior to M phase, to facilitate packaging of newly synthesised DNA into chromatin, thus affording greater protection of cells approaching maximum cell volume towards oxy-radical-mediated DNA damage (Kadonaga, 1998).

To further characterise the nature of the differential susceptibility of cells towards copper toxicity, we also investigated whether cell cycle-dependent transcription of critical heavy metal resistance genes could be correlated with differential Cu$^{2+}$-sensitivity. Thus, we specifically examined levels of *CUP1* and *SOD1* (key Cu$^{2+}$-resistance genes) promoter activity in live and dead cells, following exposure to a toxic concentration of copper, using GFP-promoter fusion plasmids. Despite the heterogeneity in the GFP fluorescence signal of live and dead sub-populations (most likely a function of variation in plasmid copy number during the cell cycle), it was apparent that basal levels of *CUP1* promoter activity did not differ significantly between live and dead sub-populations. Thus, differences in susceptibility towards copper toxicity could not be attributed to differences in *CUP1* activity at the time of Cu$^{2+}$ addition (it was confirmed that *CUP1* promoter activity did not change during the short time of Cu$^{2+}$ exposure). Increases in *CUP1* promoter activity were only evident after approximately 1 h exposure to Cu$^{2+}$, conducive with several reports indicating that *CUP1* is an inducible component
of the *S. cerevisiae* oxidative stress response system (Jensen *et al.*, 1996; Liu and Thiele, 1996). While results from the SOD1::GFP promoter fusion plasmid experiments also failed to reveal significant differences in basal levels of SOD1 promoter activity between live and dead sub-populations, our findings were inconclusive. The increased heterogeneity in the GFP fluorescence signal of pYEX-GFP<sub>SOD1</sub>-transformed DY150 (compared to pYEX-GFP<sub>CUP1</sub>-transformed cells) appears to indicate that SOD1 is constitutively expressed to varying degrees in different cell fractions. Indeed, changes in SOD1 activity during the cell cycle have been demonstrated in both vascular endothelial and smooth muscle cells (Kong *et al.*, 1993). Furthermore, the likelihood that the FL1 signal of the dead sub-population includes contributions from PI fluorescence could possibly explain the lack of detectable differences between the GFP fluorescence of live and dead sub-populations (John Daly, Dana Farber Cancer Institute Core Cytometry facility, personal communication). Thus, it remains to be clearly elucidated whether differences in basal levels of SOD1 activity play a role in the differential sensitivity of cells to copper toxicity.

Consistent with the findings of Jamieson *et al.* (1994), exposure to the specific superoxide anion-generating compound menadione for prolonged periods failed to induce SOD1 gene promoter expression, indicating that SOD1 is not an inducible part of the superoxide adaptive stress response. In addition, SOD1 was not induced in the presence of copper. The promoter region of SOD1 has previously been demonstrated to contain a single ACE1 metalloregulatory trans-acting factor, which mediates transcriptional activation of SOD1 in the presence of copper (Pena *et al.*, 1998; Culotta *et al.*, 1995; Gralla *et al.*, 1991). However, there are several possible explanations for the lack of
inducibility of our SOD1::GFP construct. The SOD1 promoter fragment used in our construct may not contain all the regulatory factors necessary for induction. Also, it is possible that the late exponential/early stationary phase, aerobic cultures used for toxicity experiments already have maximally induced levels of SOD1. Furthermore, ACEI-dependent stimulation of the CUP1 promoter has been demonstrated to be up to 50-fold greater than that of SOD1 (CUP1 contains four ACE1 binding sites) (Culotta et al., 1995; Gralla et al., 1991). Thus, our flow cytometric method for analysis of SOD1 promoter-stimulated GFP fluorescence may not be of sufficient sensitivity to detect such low levels of Cu^{2+}-induced, ACE1-dependent SOD1 stimulation.

The use of episomal plasmids in this study prevented the direct comparison of CUP1 and SOD1 promoter activity of different cell cycle stage fractions, as plasmid copy number increases proportionately with cell volume, and cell volume increases as cells progress through the cell cycle (Mason, 1991; Lord and Wheals, 1980). The problems associated with the use of episomal plasmids encountered during this study, i.e. variations in plasmid copy number throughout the cell cycle, and inherent genetic instability, can be overcome through the use of integration plasmids (Wach et al., 1997; Niedenthal et al., 1996). Indeed, the specific targeted integration of the mutant GFP, GFPS65T (exhibiting elevated FACS-activated fluorescence), reporter module to the 3'-end of several yeast genes has recently been described (Wach et al., 1997).

Thus, we have examined the differential susceptibility to copper toxicity in heterogeneous S. cerevisiae cultures. Sensitivity towards copper toxicity was demonstrated to be cell cycle-stage dependent, whereby G2/M phase cells were the most susceptible to the toxic effects of copper. The increased sensitivity of G2/M phase cells
was correlated with increased endogenous levels of ROS in these cells. While differences in basal levels of \textit{CUP I} activity between Cu\textsuperscript{2+}-resistant and Cu\textsuperscript{2+}-sensitive sub-populations were not evident, further experimentation is necessary to determine whether constitutive differences in \textit{SOD I} expression, in association with many other factors, plays an important role in the cell cycle stage-dependency of susceptibility to copper toxicity.
CHAPTER 5

The effects of heavy metals on DNA replication and the cellular DNA content of Saccharomyces cerevisiae
Heavy metals such as nickel, chromium, cadmium, and possibly cobalt, are carcinogenic to humans (Kasprzak, 1995; Snow, 1994). It is postulated that the mechanisms of their carcinogenic activity include metal-mediated pro-mutagenic oxidative damage to DNA and nuclear proteins, and decreasing the fidelity of DNA replication and repair processes (Kasprzak, 1995). For example, human DNA ligase I, the major form of the enzyme in replicative cells, is inhibited by Zn$^{2+}$ and Cd$^{2+}$ ions (Yang et al., 1996), while Cr$^{3+}$-induced DNA-DNA cross-links cause DNA polymerase I and T7 DNA polymerase arrest (Xu et al., 1996; Bridgewater et al., 1994). Furthermore, as stated previously, redox-active metals are capable of generating reactive oxygen species (ROS), such as the superoxide anion ($O_2^-$) and the hydroxyl radical (OH), through redox cycling activity, while nonredox-active metals indirectly promote oxidative stress by depletion of free radical scavengers such as glutathione and protein-bound sulhydryl groups, e.g. Cd$^{2+}$ coordination by glutathione (Figueiredo-Pereira et al., 1998; Li et al., 1997; Hassoun and Stohs, 1996; Stohs and Bagchi, 1995). The interaction of ROS with DNA may result in the generation of a wide variety of lesions including apurinic/apyrimidinic (AP) sites and both single- and double-strand breaks with 3'-deoxyribose fragments (Demple and Harrison, 1994; Ramotar et al., 1991). Double-strand breaks in particular have been implicated as the causative lesions of genome rearrangements such as deletions, duplications, and translocations, which have been associated with the onset of carcinogenesis (Brennan and Schiestl, 1998; Manivasakam and Schiestl, 1998; Brennan
et al., 1994). The hydroxyl radical has been implicated as the key causative agent of DNA damage, generating several base lesions in double-stranded DNA including thymine glycol, hypoxanthine, 8-oxoguanine and formamido-pyrimidine (Hutchinson, 1985). A role of the hydroxyl radical in gross chromosomal rearrangements has also been implicated during hydrogen peroxide-induced intracellular and intercellular recombination events in S. cerevisiae (Brennan et al., 1994). The class II AP endonucleases/3' esterase, encoded by the APNI gene, is the major cellular enzyme of S. cerevisiae responsible for initiating the direct repair of ROS-induced DNA lesions, such as AP sites and 3'-fragmented strands (Ramotar et al., 1991). In addition, genes belonging to the RAD52 epistasis group repair double-strand breaks by homologous recombination (Guzder et al., 1994; Klein, 1988). While numerous reports have examined the nature of the direct deleterious effects of ROS on DNA, and the physiological effects of mutations in key DNA damage repair genes, e.g. apnl and rad52 mutants (Xiao and Chow, 1998; Kramer et al., 1994), no studies have sought to examine the effects of PUFA-enrichment and heavy metals on cellular DNA and DNA replication in vivo.

In order to ensure the production of two viable progeny after mitosis, cell cycle events such as DNA replication and mitosis must be both temporally and spatially coordinated (Paulovich et al., 1997; Weinert et al., 1994). The prevailing theory of cell cycle control in S. cerevisiae involves two central coordinating mechanisms; the cell-cycle clock based on the serine/threonine cyclin-dependent kinases, particularly Cdc28p, and the checkpoint controls. Checkpoint controls ensure that cells maintain genome integrity (e.g. remain euploid) despite the low but continuous levels of DNA damage that
occur during normal cell growth, or during exposure to exogenous agents such as heavy metals (Weinert et al., 1994). For example, the S-M phase checkpoint which inhibits mitosis in response to DNA damage is dependent on the RAD53 and MEC1 genes; mec1-1 and rad53 mutants replicate rapidly in the presence of DNA damage (Paulovich et al., 1997; Weinert and Hartwell, 1988). Therefore, cells can be actively prevented from continuing in the cell cycle due to checkpoints that sense ongoing or aberrant DNA replication (Heichman and Roberts, 1996; Weinert et al., 1994).

Chromosomal DNA replication in S. cerevisiae is initiated at bidirectional sequence-specific origins (Oris) during S phase (Dalton and Whitbread, 1995). Oris are bound by the pre-replicative (pre-RC) protein complex from late M to early S phase, and by the post-replicative complex (post-RC) from late S to early M phase (Cocker et al., 1996; Lei et al., 1996; Tye, 1994). The initiation of DNA replication requires the minichromosome maintenance (MCM) family of polypeptides, which include Mcm2p, Mcm3p, Cdc54p/Mcm4p, Cdc46p/Mcm5p, Mcm6p and Cdc47p/Mcm7p (Dalton and Hopwood, 1997; Young and Tye, 1997; Lei et al., 1996). The six MCM proteins carry a highly conserved motif (the MCM box), which resembles a conserved domain associated with transcription and replication factors with known or assumed DNA-dependent ATPase activity (Lei et al., 1996). The role of MCM proteins as direct regulators of the initiation of DNA replication has been suggested by a number of lines of evidence; they are necessary for the maintenance of autonomously replicating sequence (ARS)-containing plasmids, mutants defective in MCM gene products have diminished usage of Oris (Young et al., 1997), and members of the MCM gene family interact with components of the origin recognition complex (ORC) (Dalton and Hopwood, 1997).
Therefore, the MCM proteins may play an important role in controlling origin activity through direct recruitment into pre-RCs. The cyclin-dependent kinase (CDK) Cdc6p is necessary for origin firing and maintenance of pre-RCs in *S. cerevisiae* (Cocker *et al.*, 1996). Without Cdc6p, genomic footprints of pre-RCs closely resemble those of post-RCs suggesting that the ORC is also a component of the pre-RC (Cocker *et al.*, 1996).

Cdc6p-dependent pre-RCs assemble at Oris at the end of M phase. Assembly of pre-RCs at Oris during late M phase is consistent with the cell cycle-regulated nuclear localisation of several of the MCM proteins, e.g. Cdc46p and Cdc47p (Dalton and Whitbread, 1995; Tye, 1994). The activation of the pre-RC results in origin firing and initiation of DNA replication. This activation step is known to require the activity of the CDKs Cdc7p and Cdc28p in association with any one of the B-type cyclins, Clb1p-Clb6p (Cocker *et al.*, 1996). Thus, it has been proposed that a key replication initiation protein of the pre-RC is phosphorylated by the Cdc7p- or Cdc28p-Clb1p-6p kinases, signalling its usage and targeting it for subsequent Cdc16p-Cdc23p-Cdc27p-dependent proteolysis (Heichman and Roberts, 1996). Temperature-sensitive *cdc6* and *cdc27* mutants over-replicate their DNA despite having continuously elevated mitotic Cdc28p-Clb2p and Cdc28p-Clp5p activity (Heichman and Roberts, 1996). DNA over-replication in these mutants involved all chromosomes and did not require passage through M phase or START. Cdc6p, Mcm2p and Mcm3p have been implicated as potential key phosphorylation target proteins of the pre-RC complex. Both Mcm2p and Mcm3p are chromatin-associated from late M phase through *G1* phase and dissociate from chromatin as cells progress through *S* phase (Young and Tye, 1997). Furthermore, a key role of Cdc6p has been suggested by recent studies with *S. pombe* whereby 10- to 20-fold overexpression of the
Cdc6p homolog, Cdc18p, induced over-replication of cellular DNA; de novo protein synthesis was not required for this effect (Stern and Nurse, 1996). Overexpression of the CDK inhibitor rum1p of *S. pombe* also leads to sequential, uninterrupted S phases (Moreno and Nurse, 1994; Correa-Bordes and Nurse, 1995). Stern and Nurse (1996) have proposed that a very low CDK activity towards the end of M phase enables dephosphorylated initiator protein-associated pre-RCs to assemble at Oris in *S. pombe*. A subsequent increase in CDK activity to a moderate level leads to phosphorylation of the initiator protein, origin firing and initiation of DNA replication. Phosphorylation of the initiator protein(s) simultaneously prevents the assembly of further pre-RCs, as phosphorylated initiator protein(s) cannot bind to Oris. Thus, according to the proposed model, DNA replication is limited to once per cell cycle, with each round of DNA replication requiring a change in CDK activity.

The purpose of this study was to examine the effects of heavy metal-exposure on DNA replication and the cellular DNA content in unsupplemented and PUFA-supplemented *S. cerevisiae*. Heavy metal-induced DNA over-replication in PUFA-enriched *S. cerevisiae* is demonstrated for the first time, whereby exposure of PUFA-enriched cells, particularly 18:3-enriched cells to redox-active metals caused repeated rounds of DNA replication, accumulation of up to 8C DNA, and the uncoupling of DNA synthesis from cell cycle progression.
5.2. Materials and Methods

5.2.1. Organisms and culture conditions. Yeast strains used for DNA replication experiments were *S. cerevisiae* NCYC 1383 (MATα, his3-Δ1, leu2-3, leu2-112, ura3-52, trp1-289) and RJD621 (MATa, can1-100, leu2-3, -112, his3-11, -15, trp1-1, ura3-1, ade2-1, cdc28::CDC28-HA::TRP1) (a gift from Ray DeShaises, Caltech, CA). RJD621 contains an HA-tagged *CDC28* gene in a W303 strain background. Maintenance of yeast strains, and preparation of strains for experimental purposes was as previously described (see Section 2.2.1.).

5.2.2. Preparation of cell suspensions for toxicity experiments. The preparation of cell suspensions for toxicity experiments was as described previously (see Section 4.2.4.). For the hydroxyurea experiments, cultures were incubated with 0.2 M hydroxyurea (Sigma, MO) for two hours prior to the addition of toxin. Hydroxyurea prevents DNA replication by inhibiting ribonucleotide reductase, thereby depleting cells of dNTP precursors (Weinert et al., 1994).

5.2.3. Cellular DNA staining. Cellular nucleic acids were stained with PI and examined using flow cytometry as described previously (see Sections 4.2.7. and 4.2.8.).

5.2.4. Diphenylamine determination of DNA concentration. To confirm results from flow cytometry experiments, cellular DNA concentrations were determined using the
diphenylamine (DPA) assay (Burton, 1968), as modified by Sasaki (1992), with the following additional modifications. At specified time points after the addition of toxin, 20 ml samples (~1 x 10^9 cells) from control and toxin-treated cells were harvested by centrifugation at 1,500 g for 5 min, and washed twice with cold distilled deionised water. Cells were washed with 4 ml of 75% (v/v) ethanol and allowed to dry overnight at room temperature. Pellets were resuspended in 1.25 ml of 1.5 N perchloric acid and heated at 70°C for 30 min. Samples were allowed to cool, 0.75 ml of DPA solution (4% (w/v) diphenylamine, 1.5% (v/v) sulphuric acid, 0.01% (v/v) paraldehyde) was added, and the mixture was incubated at 30°C for 18 h. Samples were allowed to cool and 1.1 ml amyl acetate was added. Absorbances of samples were measured at 595 nm, using a Pharmacia Biotech Ultraspec 2000 UV/Visible spectrophotometer, against a reference solution comprising 1.1 ml amyl acetate, 0.75 ml DPA solution, and 1.25 ml 0.2 N perchloric acid. The concentrations of DNA in samples were calculated by reference to a standard curve prepared using bovine DNA (Promega, WI).

5.2.5. Yeast subcellular fractionation. Nuclear membrane-enriched fractions were isolated according to Ausubel et al. (1998) and Young and Tye (1997), with the following modifications. After 6 h incubation in the presence of Cu(NO₃)₂, cells were harvested by centrifugation at 1,500 g for 5 min and washed twice with 10 mM Mes buffer (pH 5.5), 1% (w/v) glucose. The approximate wet weight (in grams) of cells was measured, cells were resuspended in approximately 3 vol ice-cold distilled deionised water, and immediately centrifuged at 1,500 g for 5 min. Cells were resuspended in 1 vol spheroplasting buffer (50 mM Tris.Cl (pH 7.5), 10 mM MgCl₂, 1 M sorbitol, 30 mM
DTI) and incubated at room temperature for 15 min. Cells were harvested by centrifugation at 1,500 g for 5 min and resuspended in 1 vol spheroplasting buffer containing 1 mM DTT, protease inhibitors (0.1 μg ml⁻¹ chymostatin, 2 μg ml⁻¹ aprotinin, 1 μg ml⁻¹ pepstatin A, 1.1 μg ml⁻¹ phosphoramidon, 5 μg ml⁻¹ E-64, 0.5 μg ml⁻¹ leupeptin, 2.5 μg ml⁻¹ antipain, 0.1 mM benzamidine, 0.1 mM sodium metabisulphite), and phosphatase inhibitors (5 mM sodium pyrophosphate, 0.1 mM sodium metavanadate, 50 mM sodium fluoride). Cell walls were digested with 20 U ml⁻¹ recombinant lytic enzyme (ICN Biochemicals, CA) and 20 U ml⁻¹ lytic enzyme 100T (ICN Biochemicals, CA) at 30 °C for 2 h with rotary shaking at 50 rev. min⁻¹ (approximately 1 ml of cell suspension in sterile 15 ml centrifuge tubes). Spheroplast formation was followed by examining water-induced spheroplast lysis by light microscopy. Approximately 80-90% spheroplast yield was usually observed after 2 h incubation. Spheroplasts were centrifuged at 1,500 g for 10 min, washed twice in 2 vol ice-cold spheroplast buffer, and resuspended in 0.5 vol nuclei buffer (1 M sorbitol, 20 mM PIPES (pH 6.3), 0.5 mM CaCl₂, 1 mM DTT, 0.1 mM EDTA, 1 mM PMSF, with protease and phosphatase inhibitors). Spheroplasts were lysed by pipetting drop by drop into approximately 20 vol ice-cold Ficoll buffer (18% (w/v) Ficoll 400 (Pharmacia Biotech, NJ), 20 mM PIPES (pH 6.3), 0.5 mM CaCl₂, 1 mM DTT, 0.1 mM EDTA, 1 mM PMSF, with protease and phosphatase inhibitors) at 4 °C. Ficoll lyases spheroplasts but does not lyse nuclei (Lohr, 1988). Cell debris was removed by centrifugation at 3,000 g for 20 min at 4 °C, using a Sorvall RC-5B refrigerated superspeed centrifuge and SS-34 rotor. The resulting supernatant was centrifuged at 20,000 g for 30 min, yielding a total nuclear pellet. The supernatant was collected and stored at 4 °C (cytosolic fraction, C). The nuclear pellet was resuspended in 100 μl of
low salt extraction buffer (10 mM HEPES (pH 7.5), 1 mM EDTA, 0.5 M NaCl), transferred to a microcentrifuge tube and microcentrifuged at 14,500 rev. min$^{-1}$ for 20 min. The low salt nucleosolic fraction (NSL) supernatant was collected and stored at 4 °C. The pellet was resuspended in 100 µl of high salt extraction buffer (10 mM HEPES (pH 7.5), 1 mM EDTA, 2.0 M NaCl) and microcentrifuged at 14,500 rev. min$^{-1}$ for 20 min. The high-salt nucleosolic fraction (NSH) supernatant was collected and stored at 4 °C. The nuclear pellet was resuspended in 100 µl of low salt extraction buffer and incubated with four hundred units of deoxyribonuclease I (DNase I) (Boehringer Mannheim, IN) at 37 °C for 2 h. The debris was pelleted at 14,500 rev. min$^{-1}$ for 20 min and the nuclear pellet fraction (NP) removed and stored at 4 °C.

5.2.6. Lipid extraction and fatty acid analysis. Lipids were extracted from nuclear membrane fractions as described previously (see Section 2.2.5). Fatty acid methyl ester separations and analysis were performed as described in Section 4.2.3.

5.2.7. DAPI staining of yeast nuclei. The purity of isolated yeast nuclei was assessed by staining with the DNA-binding fluorochrome 4',6-diamidine-2'-phenylindole dihydrochloride (DAPI) (Molecular Probes Inc., OR), using a modified method of M'Millan and Tatchell (1994). At specified intervals after the addition of toxin to cell suspensions, one and a half ml aliquots were removed, washed with 1 ml of 10 mM Mes buffer (pH 5.5), 1% (w/v) glucose, and immediately fixed in 70% (v/v) ethanol for 30 min at 4 °C. Samples were washed twice with distilled deionised water and resuspended in 1 ml of PBS containing 1 µg ml$^{-1}$ DAPI for 20 min. DAPI has an absorption
maximum of 358 nm and emission maximum of 461 nm (Haugland, 1996). Samples were washed twice with distilled deionised water, resuspended in 1 ml of 50% (v/v) glycerol, and stored at 4 °C until analysis. DAPI-stained nuclei were visualised by video fluorescence microscopy, using an Olympus BH-2 fluorescence microscope and Javelin video camera.

5.2.8. **Protein determination.** Protein concentrations of yeast subcellular fractions were determined using the Micro-Lowry assay, with the following modifications. Samples (100 μl) were added to microcentrifuge tubes and made up to 1 ml with sterile, distilled deionised water. Protein was precipitated by adding 100 μl of 0.15% (w/v) sodium deoxycholate and 100 μl of 72% (w/v) trichloroacetic acid (TCA). Samples were microcentrifuged at high speed for 15 min, washed twice with 100% (v/v) ethanol, and the final pellet was dried under vacuum. The pellet was resuspended in 800 μl of CTC reagent (0.1% (w/v) CuSO₄, 0.2% (w/v) C₄H₄Na₂O₆.2H₂O, 10% (w/v) Na₂CO₃) and 200 μl of 17% (v/v) Folin-Ciocalteau reagent (Sigma, MO), and incubated at room temperature for 30 min. Absorbances of samples were measured at 750 nm using a Pharmacia Biotech Ultraspec 2000 UV/Visible spectrophotometer. Protein concentrations were calculated by reference to a standard curve prepared using bovine serum albumin (BSA) (Sigma, MO).

5.2.9. **SDS-PAGE and immunoblotting.** SDS-PAGE was performed according to standard protocol (Ausubel *et al.*, 1998). In preparation for SDS-PAGE, proteins were precipitated by mixing with an equal volume of 20% (w/v) TCA, and microcentrifuged at
full speed for 10 min. Protein pellets were washed twice with ice-cold 100% (v/v) ethanol, resuspended in sample loading buffer (50 mM Tris.Cl (pH 6.8), 100 mM DTT, 2% (w/v) SDS, 10% (v/v) glycerol, 0.1% (w/v) bromophenol blue), and denatured by boiling for 1-2 min. Protein samples were stored at 4 °C until analysis. Proteins were separated using the standard Laemmli system with homogenous 7.5% SDS-PAGE gels as follows. Separating gels (12 ml) comprised 2.4 ml of 1.875 M Tris (pH 8.8), 1.8 ml of 50% (w/v) acrylamide solution (24:1 (w/w) acrylamide:bis-acrylamide) (Bio-Rad Laboratories, CA), 120 µl of 10% (w/v) SDS, and 7.68 ml of sterile, distilled deionised water. Immediately before pouring, 12 mg of ammonium persulphate, and 6 µl N,N,N’,N’-tetra-methyl-ethylenediamine (TEMED) (Bio-Rad Laboratories, CA) were added to facilitate polymerisation. The separating gel was layered with saturated n-butanol, to prevent the formation of internal air bubbles. Upon polymerisation, the overlay was decanted and the stacking gel was applied. The stacking gel (3 ml) comprised 0.3 ml of 1.25 M Tris (pH 6.8), 240 µl of 50% (w/v) acrylamide solution, 30 µl of 10% (w/v) SDS, and 2.43 ml of sterile, distilled deionised water. Immediately before pouring, 3 mg of ammonium persulphate and 3.5 µl TEMED were added. Proteins were separated at 20 mA for 1.5-2 h, in Laemmli buffer (0.025 M Tris, 0.192 M glycine, 0.1% (w/v) SDS), using the Bio-Rad Mini-PROTEAN II electrophoresis cell. Gels were stained for 1 h with gentle rotation in 0.25% (w/v) Coomassie blue (Bio-Rad Laboratories, CA), 50% (v/v) methanol, 10% (v/v) glacial acetic acid. Gels were destained by incubating in primary destaining solution (30% (v/v) methanol, 10% (v/v) glacial acetic acid) for 1 h, and in secondary destaining solution (16.5% (v/v) methanol, 5% (v/v) glacial acetic acid) (with frequent changes) overnight. Immunoblotting was
performed using standard techniques (Ausubel et al., 1998). Separated proteins were transferred to 0.45 μm nitrocellulose membranes (Bio-Rad Laboratories, CA) at 150 mA overnight in transfer buffer (39 mM glycine, 48 mM Tris, 20% (v/v) methanol, pH 8.3) at 4 °C, again using the Bio-Rad Mini-PROTEAN II electrophoresis cell. Nitrocellulose membranes were washed in 5 x Tris-buffered saline (TBS) (0.25 M Tris (pH 7.4), 1 M NaCl), and incubated in 15 ml blocking buffer, comprising 1.5 ml of 10 x TBS, 750 μl of 10% (w/v) gelatin (EIA grade, 60 bloom) (Bio-Rad Laboratories, CA), 12.75 ml of sterile, distilled deionised water for 1 h. Nitrocellulose membranes were again washed with 5 x TBS, and incubated in antibody-binding buffer, of the same composition as blocking buffer, with antibody. The primary antibodies used for immunoblotting were 1:500 dilutions of affinity purified rabbit anti-Mcm3 (a gift from Rolf Knippers, Universite Konstanz; Hu et al., 1993), and 5 μg ml⁻¹ mouse anti-HA (12CA5) (Boehringer Mannheim, IN). Nitrocellulose membranes were then incubated in secondary antibody-binding buffer for 1 h. The secondary antibodies used were 1:5000 dilutions of alkaline phosphatase-conjugated anti-rabbit IgG (Fc) (Promega, WI) and alkaline phosphatase-conjugated anti-mouse IgG (H + L) (Promega, WI). Alkaline phosphatase was detected using the BCIP/NBT colour system (Promega, WI), as follows. Nitrocellulose membranes were incubated in 15 ml substrate buffer, comprising 1.5 ml of 1 M Tris.Cl (pH 9.5), 500 μl of 3 M NaCl, 75 μl of 1 M MgCl₂, 99 μl of 5-bromo-4-chloro-3-indolyl phosphate (BCIP), 49.5 μl of nitro blue tetrazolium (NBT), and 12.8 ml of sterile, distilled deionised water, and incubated with gentle rotation until the appearance of bands.
5.3. Results

5.3.1. Effect of Cu(NO₃)₂ on the cellular DNA content of unsupplemented and PUFA-supplemented *S. cerevisiae* NCYC 1383. The increased susceptibility of PUFA-enriched *S. cerevisiae* to both cadmium- and copper-induced membrane permeabilisation and whole-cell toxicity had previously been demonstrated (Chapters 2 and 3). The increased susceptibility of PUFA-enriched cells towards metal toxicity was attributed to greater levels of ROS-mediated lipid peroxidation in these cells. We therefore sought to investigate the effects of PUFA-enrichment on heavy metal-induced ROS-mediated DNA damage. Thus, we investigated the effects of Cu(NO₃)₂ on the cellular DNA content of *S. cerevisiae* previously grown in unsupplemented, 18:2- and 18:3-supplemented media, using flow cytometry. Late-exponential phase cells were exposed to Cu(NO₃)₂ in buffer for up to 20 h. Samples were taken at regular time intervals, fixed, and cellular DNA was stained with PI. Linear-scale histograms of FL2 against counts were plotted. The cellular DNA content of cells grown previously in unsupplemented medium, was examined at specified intervals during 20 h exposure to 100 μM Cu(NO₃)₂ (Fig. 5.1). Exposure of unsupplemented cells to 100 μM Cu(NO₃)₂ for up to 20 h had no marked effects on cellular DNA content, as evidenced by the presence of clear and resolved 1C and 2C DNA peaks (1C corresponds to an arbitrary FL2 intensity unit of ~ 120, 2C ~ 240) (Fig. 5.1f). In addition, exposure of unsupplemented cells to copper did not appear to result in checkpoint control-induced cell cycle arrest, as numbers of 1C and 2C DNA cells remained relatively constant throughout the 20 h exposure to copper. The cellular
**Fig. 5.1.** Effect of 100 μM Cu(NO₃)₂ on the cellular DNA content of *S. cerevisiae* NCYC 1383 previously grown in unsupplemented medium. The FL2 histograms depict the cellular DNA content of cells before copper addition (a), and after 15 min (b), 30 min (c), 4.5 h (d), 6 h (e) and 20 h (f) exposure to 100 μM Cu(NO₃)₂. Each histogram contains data collected from at least 10,000 cells.
Fig. 5.2. Effect of 100 μM Cu(NO₃)₂ on the cellular DNA content of *S. cerevisiae* NCYC 1383 previously grown in linoleate-supplemented medium. The FL2 histograms depict the cellular DNA content of cells before copper addition (a), and after 15 min (b), 30 min (c), 4.5 h (d), 6 h (e) and 20 h (f) exposure to 100 μM Cu(NO₃)₂. Each histogram contains data collected from at least 10,000 cells.
**Fig. 5.3.** Effect of 100 µM Cu(NO₃)₂ on the cellular DNA content of *S. cerevisiae* NCYC 1383 previously grown in linolenate-supplemented medium. The FL2 histograms depict the cellular DNA content of cells before copper addition (a), and after 15 min (b), 30 min (c), 4.5 h (d), 6 h (e) and 20 h (f) exposure to 100 µM Cu(NO₃)₂. Each histogram contains data collected from at least 10,000 cells.
DNA content of cells previously grown in 18:2-supplemented medium, was also examined during 20 h exposure to 100 μM Cu(NO₃)₂ (Fig. 5.2). During the initial period of copper exposure 1C and 2C peak intensities were reduced, and peak broadening was evident (Fig. 5.2c,d,e). Furthermore, after 20 h exposure over-replication of cellular DNA was evident, whereby both 1C and 2C DNA peaks were severely diminished, and DNA levels upwards of 4C (FL2 > 480) were evident (Fig. 5.2f). The cellular DNA content of cells previously grown in 18:3-supplemented medium was also examined during 20 h exposure to 100 μM Cu(NO₃)₂ (Fig. 5.3). Here the effect of Cu²⁺ on DNA content was considerably more dramatic than in 18:2-supplemented cells; 1C and 2C peak intensities were significantly diminished after only 15 min exposure (Fig. 5.3b). Furthermore, after only 4.5 h cellular DNA levels greater than 4C and up to 8C (FL2 ~ 960) were evident (Fig. 5.3d). The results demonstrate continued DNA synthesis in PUFA-enriched S. cerevisiae exposed to toxic concentrations of Cu(NO₃)₂.

5.3.2. Influence of hydroxyurea on Cu(NO₃)₂-induced over-replication in 18:3-enriched S. cerevisiae NCYC 1383. In order to confirm that the flow cytometric DNA profiles of Cu²⁺-exposed PUFA-enriched S. cerevisiae genuinely represented continued DNA synthesis, cells were incubated in the presence of hydroxyurea for two hours prior to the addition of Cu(NO₃)₂, and cellular DNA content was again examined by flow cytometry. Hydroxyurea prevents DNA replication by inhibiting ribonucleotide reductase, thereby depleting cells of dNTP precursors (Weinert et al., 1994). The cellular DNA content of 18:3-enriched cells exposed to 100 μM Cu(NO₃)₂ for up to 20 h, after pre-incubation in the presence of 0.2 M hydroxyurea, is shown (Fig. 5.4). Exposure to Cu(NO₃)₂, after
Fig. 5.4. Influence of hydroxyurea on Cu(NO₃)₂-induced over-replication in linolenate-enriched S. cerevisiae NCYC 1383. Cells were incubated in the presence of 0.2 M hydroxyurea for 2 h prior to the addition of Cu(NO₃)₂. The FL2 histograms depict the cellular DNA content of linolenate-enriched cells before copper addition (a), and after 15 min (b), 30 min (c), 4.5 h (d), 6 h (e) and 20 h (f) exposure to 100 μM Cu(NO₃)₂. Each histogram contains data collected from at least 10,000 cells.
incubation with hydroxyurea, resulted in a reduction in both 1C and 2C DNA peak intensities, and peak broadening (Fig 5.4b-f). However, there is little evidence of continued DNA synthesis, even after 20 h exposure. The results demonstrate that pre-incubation of 18:3-enriched *S. cerevisiae* with the DNA synthesis inhibitor hydroxyurea prevents Cu(NO₃)₂-induced over-replication of cellular DNA. Thus, the results confirm continued DNA replication in 18:3-enriched *S. cerevisiae* upon prolonged exposure to toxic concentrations of Cu(NO₃)₂.

5.3.3. Determination of cellular DNA concentrations. Here we sought to further confirm DNA over-replication in PUFA-enriched *S. cerevisiae* upon exposure to Cu(NO₃)₂, using an alternative assay for cellular DNA content. Therefore, we measured the DNA content of unsupplemented and PUFA-supplemented *S. cerevisiae* incubated in the presence and absence of Cu(NO₃)₂ for 6 h, using the diphenylamine assay (Burton, 1968) (Table 5.1). Exposure to 100 μM Cu(NO₃)₂ for 6 h had only a small effect on the average amount of DNA per unsupplemented cell, resulting in a reduction of approximately 16.3%. However, marked increases in the average amount of DNA per cell were observed for PUFA-enriched cells. Here, the average amount of DNA per PUFA-enriched cell increased by approximately 38.5% and 76.9%, in 18:2- and 18:3-enriched cells, respectively, after 6 h exposure to 100 μM Cu(NO₃)₂ (Table 5.1). It was also evident that the average amount of DNA per PUFA-enriched cell was approximately 40% lower than that of unsupplemented cells, previously incubated in the absence of Cu(NO₃)₂. These results are most likely a consequence of variations in DNA extraction efficiencies, due to inherent differences in physical membrane properties, between unsupplemented and
Table 5.1. Cellular DNA concentrations of unsupplemented and PUFA-supplemented *S. cerevisiae* NCYC 1383 incubated in buffer, in the presence and absence of 100 μM Cu(NO₃)₂ for 6 h

<table>
<thead>
<tr>
<th>Medium</th>
<th>[DNA] (fg cell⁻¹)</th>
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<tbody>
<tr>
<td></td>
<td>no metal</td>
</tr>
<tr>
<td>unsupplemented</td>
<td>20.8 ± 2.6</td>
</tr>
<tr>
<td>18:2-supplemented</td>
<td>13.0 ± 0.3</td>
</tr>
<tr>
<td>18:3-supplemented</td>
<td>12.1 ± 1.2</td>
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Values for [DNA] are means from two replicate determinations ± SEM.
PUFA-supplemented cells (Alterthum and Rose, 1973). Nevertheless, the results strongly support flow cytometric data indicating DNA over-replication in PUFA-supplemented *S. cerevisiae* exposed to Cu(NO$_3$)$_2$.

5.3.4. Effects of non-toxic concentrations of Cu(NO$_3$)$_2$ on the cellular DNA content of unsupplemented and PUFA-supplemented *S. cerevisiae* NCYC 1383. Here we investigated whether Cu(NO$_3$)$_2$-induced DNA over-replication in PUFA-enriched cells would also occur at lower, non-toxic concentrations of Cu(NO$_3$)$_2$. Cells grown previously in unsupplemented and PUFA-supplemented medium were exposed to 1 and 10 μM Cu(NO$_3$)$_2$ for up to 9 h. Exposure of 18:3-supplemented cells to 1 and 10 μM Cu(NO$_3$)$_2$ for 1 h resulted in approximately 7% and 28% lethality, respectively (results not shown). The cellular DNA content of cells previously grown in 18:3-supplemented medium, exposed to 10 μM Cu(NO$_3$)$_2$ is shown (Fig. 5.5). After 2 h exposure, both 1C and 2C DNA peak intensities were much reduced (similar detrimental effects were observed after only 15 min exposure of 18:3-enriched cells to 100 μM Cu(NO$_3$)$_2$ (see Fig. 5.3b)) (Fig. 5.5a). Furthermore, after 9 h exposure both 1C and 2C DNA peaks were severely diminished and DNA levels greater than 4C were evident (Fig. 5.5d). Over-replication of cellular DNA was not evident for unsupplemented or PUFA-supplemented cells exposed to 1 μM Cu(NO$_3$)$_2$, or unsupplemented and 18:2-supplemented cells exposed to 10 μM Cu(NO$_3$)$_2$ (results not shown). The results indeed demonstrate DNA over-replication in 18:3-enriched *S. cerevisiae* exposed to non-toxic concentrations of Cu(NO$_3$)$_2$. Furthermore, the results indicate a threshold concentration of Cu(NO$_3$)$_2$ below which over-replication of cellular DNA does not occur.
**Fig. 5.5.** Effect of 10 μM Cu(NO₃)₂ on the cellular DNA content of linolenate-enriched *S. cerevisiae* NCYC 1383. The FL2 histograms depict the cellular DNA content of cells after 2 h (a), 4.5 h (b), 6 h (c), and 9 h (d) exposure to 10 μM Cu(NO₃)₂. Each histogram contains data collected from at least 10,000 cells.
5.3.5. Effects of the free radical scavengers mannitol and DMSO on Cu(NO$_3$)$_2$-induced over-replication in PUFA-supplemented *S. cerevisiae*NCYC 1383. Redox-active metals such as copper directly promote oxidative stress through the generation of reactive oxygen species such as the hydroxyl radical (OH$^-$) (Flowers *et al.*, 1997; Lloyd *et al.*, 1997). As the hydroxyl radical has been implicated as a key causative agent of DNA damage in *S. cerevisiae*, we examined whether Cu$^{2+}$-induced DNA over-replication in 18:3-enriched cells could be inhibited by the hydroxyl radical scavengers dimethyl sulfoxide (DMSO) and mannitol. Cells previously grown in unsupplemented, 18:2- and 18:3-supplemented medium were exposed to 20 μM Cu(NO$_3$)$_2$ for up to 22 h, in the presence of 0.2 M mannitol and 0.4 M DMSO. Incubation with DMSO and mannitol during Cu$^{2+}$-exposure had little protective effect against over-replication of cellular DNA in PUFA-enriched cells (results not shown). Repeated rounds of DNA replication, and accumulation of cellular DNA levels of up to 8C DNA were again evident, despite the presence of both DMSO and mannitol (see Fig. 5.3). The results suggest that the hydroxyl radical is most likely not directly involved in Cu$^{2+}$-induced DNA over-replication in PUFA-enriched cells.

5.3.6. Effects of Cd(NO$_3$)$_2$, CrK(SO$_4$)$_2$, H$_2$O$_2$, and menadione on the cellular DNA content of unsupplemented and PUFA-supplemented *S. cerevisiae*NCYC 1383. We sought to determine whether toxic concentrations of the nonredox-active metal ions Cd$^{2+}$ and Cr$^{3+}$, the oxidative toxin H$_2$O$_2$, or the superoxide anion-generating compound menadione could also induce DNA over-replication in PUFA-supplemented *S. cerevisiae*. Cells previously grown in unsupplemented, 18:2- and 18:3-supplemented medium were
exposed to either 200 μM Cd(NO₃)₂, 5 mM CrK(SO₄)₂, 5 mM H₂O₂ or 2 mM menadione for up to 20 h. Prolonged exposure to all four toxins had no evident adverse effects on the cellular DNA content of both supplemented or PUFA-supplemented cells, whereby both 1C and 2C DNA peaks were still evident and clearly resolved (results not shown).

5.3.7. Effects of reduced-glutathione on Cu²⁺-induced DNA damage and over-replication in linolenate-enriched S. cerevisiae NCYC 1383. The dithiol glutathione is one of the most important cellular antioxidants of S. cerevisiae (Stephen and Jamieson, 1996). Glutathione possesses a redox-active sulphydryl group which reacts with oxidants to produce oxidised glutathione (GSSG). Here we sought to determine if redox-cycling was essential for Cu²⁺-induced DNA over-replication in linolenate-enriched cells, by inhibiting redox-cycling in the presence of excess reduced-glutathione. Cells previously grown in 18:3-supplemented medium were exposed to 20 μM Cu(NO₃)₂ for up to 6 h, in the presence and absence of 0.1 M reduced-glutathione (Fig. 5.6). Again, after only 1 h exposure to 20 μM Cu(NO₃)₂, 1C and 2C DNA peaks intensities were diminished (Fig. 5.6a). Furthermore, after 6 h exposure to 20 μM Cu(NO₃)₂ DNA levels greater than 4C were evident (Fig. 5.6e). However, DNA over-replication was not evident after 6 h exposure of 18:3-enriched cells to 20 μM Cu(NO₃)₂ in the presence of 0.1 M reduced-glutathione, whereby both 1C and 2C DNA peaks remained distinctly resolved (Fig. 5.6f). The results indicate that Cu²⁺ redox-cycling activity is essential for DNA over-replication in 18:3-enriched cells.
Fig. 5.6. Influence of reduced-glutathione on Cu(NO₃)₂-induced DNA over-replication in linolenate-enriched S. cerevisiae NCYC 1383. Cells were exposed to 20 µM Cu(NO₃)₂ in the presence and absence of 0.1 M reduced-glutathione for up to 6 h. The FL2 histograms depict the cellular DNA content of cells after 1 h (a), 4.5 h (c) and 6 h (e) exposure to Cu(NO₃)₂, in the absence of reduced-glutathione, and 1 h (b), 4.5 h (d) and 6 h (f) exposure in the presence of reduced glutathione. Each histogram contains data from at least 10,000 cells.
5.3.8. **Effect of \( \text{CrO}_3 \) on the cellular DNA content of unsupplemented and PUFA-supplemented \( S. \text{cerevisiae} \) NCYC 1383.** \( \text{Cr}^{6+} \) is a highly redox-active metal ion, exemplified by the limited number of chromium (VI) compounds, which include \( \text{CrO}_3 \) and peroxy compounds, that exist (Mackay and Mackay, 1989). Hence, we investigated whether metal-induced over-replication of cellular DNA in PUFA-enriched cells could also be induced by \( \text{Cr}^{6+} \). Cells previously grown in 18:3-supplemented medium were exposed to 5 mM \( \text{CrO}_3 \) for up to 6 h. The cellular DNA content of 18:3-enriched \( S. \text{cerevisiae} \) exposed to \( \text{CrO}_3 \) is shown (Fig. 5.7). After only 15 min exposure to \( \text{CrO}_3 \) 1C and 2C DNA peak intensities were reduced (similar to that observed for 18:3-enriched cells exposed to 100 \( \mu \text{M} \) \( \text{Cu(NO}_3\text{)}_2 \) (see Fig. 5.3)) (Fig. 5.7b). After 2 h exposure to 5 mM \( \text{CrO}_3 \) the cellular DNA content of 18:3-enriched cells increased to greater than 4C (Fig. 5.7d). Thus, the onset of DNA over-replication appeared to occur more rapidly in the presence of 5 mM \( \text{Cr}^{6+} \), compared with 100 \( \mu \text{M} \) \( \text{Cu}^{2+} \) (see Fig. 5.3). Furthermore, upon continued exposure to \( \text{CrO}_3 \) DNA levels greater than 6C were evident (Figs. 5.7e,f). These results were very similar to those attained upon exposure of 18:3-enriched cells to 100 \( \mu \text{M} \) \( \text{Cu(NO}_3\text{)}_2 \) (see Fig. 5.3). These results confirm that exposure of 18:3-enriched \( S. \text{cerevisiae} \) to the redox-active metal \( \text{Cr}^{6+} \) also results in continued rounds of DNA synthesis.

5.3.9. **Fatty acid composition of nuclear membrane-enriched fractions.** We had previously demonstrated the incorporation of exogenous linoleate and linolenate into whole-cell and plasma membrane lipids (see Tables 2.1 and 2.4). As this particular study was primarily
Fig. 5.7. Effect of 5 mM CrO$_3$ on the cellular DNA content of linolenate-enriched *S. cerevisiae* NCYC 1383. The FL2 histograms depict the cellular DNA content of cells before chromium addition (a), and after 15 min (b), 30 min (c), 2 h (d), 4.5 h (e) and 6 h (f) exposure to 5 mM CrO$_3$. Each histogram contains data collected from at least 10,000 cells.
concerned with the effects of toxins on cellular DNA content and DNA replication, we sought to confirm incorporation of exogenous 18:2 and 18:3 into nuclear membrane lipids, and examine the fatty acid composition of nuclear membranes after exposure to Cu(NO₃)₂. In order to assess the purity of isolated nuclear membrane fractions, nuclei were stained with the DNA-binding fluorochrome DAPI (McMillan and Tatchell, 1994). The purity of nuclear membrane-enriched fractions from cells grown in unsupplemented, 18:2- and 18:3-supplemented media was confirmed using both light and fluorescence microscopy, and by observing characteristic nuclear electrophoretic profiles using SDS-PAGE, e.g. the presence of the 38 kDa nucleolar protein fibrillarin (Aris and Blobel, 1991) (results not shown). Isolated nuclear membrane fractions were thus considered of sufficient purity for lipid analysis.

The fatty acid compositions of nuclear membrane-enriched fractions from cells grown for 16 h in unsupplemented, 18:2- and 18:3-supplemented media (Table 5.2) were generally similar to those of their corresponding whole-cell homogenates (Table 2.1), and plasma membrane-enriched fractions determined previously (Table 2.4). However, incorporation of the exogenous PUFAs 18:2 and 18:3 into nuclear membrane lipids was approximately 48% lower and 33% lower, respectively, than those of the corresponding plasma membrane-enriched fractions (Table 5.2). Nevertheless, fatty acid unsaturation indices calculated for nuclear membrane-enriched fractions again decreased in the order 18:3-supplemented > 18:2-supplemented > unsupplemented (Table 5.2). While the results confirm incorporation of exogenous PUFAs into nuclear membrane lipids, genuine differences between levels of incorporation of PUFAs into nuclear membrane and plasma membrane lipids were apparent. However, several factors, such as
Table 5.2. Fatty acid composition of nuclear membrane-enriched fractions from *S. cerevisiae* NCYC 1383 previously grown in unsupplemented, linoleate- and linolenate-supplemented media

<table>
<thead>
<tr>
<th>Fatty acid</th>
<th>Medium</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Unsupplemented</td>
<td>18:2-supplemented</td>
<td>18:3-supplemented</td>
<td></td>
</tr>
<tr>
<td>16:0</td>
<td>24.5±1.9</td>
<td>20.2±1.7</td>
<td>18.8±2.2</td>
<td></td>
</tr>
<tr>
<td>16:1</td>
<td>44.2±4.4</td>
<td>11.8±0.2</td>
<td>4.6±0.3</td>
<td></td>
</tr>
<tr>
<td>18:0</td>
<td>7.2±0.1</td>
<td>14.9±0.9</td>
<td>24.7±7.6</td>
<td></td>
</tr>
<tr>
<td>18:1</td>
<td>24.1±2.4</td>
<td>16.2±0.3</td>
<td>8.4±2.3</td>
<td></td>
</tr>
<tr>
<td>18:2 tr.</td>
<td>tr.</td>
<td>36.9±0.6</td>
<td>tr.</td>
<td></td>
</tr>
<tr>
<td>18:3</td>
<td>n.d.</td>
<td>n.d.</td>
<td>43.6±2.0</td>
<td></td>
</tr>
<tr>
<td>U.I.</td>
<td>0.68</td>
<td>1.02</td>
<td>1.44</td>
<td></td>
</tr>
</tbody>
</table>

Values for percentage fatty acid composition are means from two replicate determinations ± standard deviation (SD). (n.d. = not detectable; tr. < 0.1). U.I. refers to the unsaturation index (average number of double bonds per fatty acid).
differences in subcellular fractionation techniques, FAME separation and analysis procedures, and experimental conditions, e.g. ambient temperature, and presence or absence of buffer, during nuclear membrane and plasma membrane isolation (see Sections 4.2 and 5.2), may have contributed to the discrepancies observed.

The fatty acid composition of nuclear membrane-enriched fractions from S. cerevisiae incubated in buffer, in the presence of 100 μM Cu(NO₃)₂ for 6 h showed significant differences to that of nuclear membrane-enriched fractions of cells incubated in the absence of Cu(NO₃)₂ (Table 5.3). Nuclear membrane enriched-fractions from cells incubated in the presence of Cu(NO₃)₂ generally displayed a lower content of unsaturated fatty acids, as a proportion of total fatty acids, than those from cells incubated in the absence of Cu(NO₃)₂. This was most evident as lower proportions of 18:2 and 18:3. Hence, after 6 h exposure of PUFA-enriched cells to Cu(NO₃)₂, the unsaturation indices of 18:2- and 18:3-supplemented nuclear membrane-enriched fractions were approximately 37% and 17% lower, respectively, than nuclear membrane-enriched fractions from cells incubated in the absence of Cu(NO₃)₂.

5.3.10. Determination of Mcm3p in nuclear fractions of S. cerevisiae NCYC 1383. Mcm3p is present in high abundance and at relatively constant levels throughout the cell cycle in actively proliferating cells (Young and Tye, 1997). Furthermore, while Mcm3p is present in both the cytoplasm and nucleus throughout the cell cycle, it becomes tightly associated with chromatin only from late M phase to early S phase (Young and Tye, 1997). Mcm3p is thus thought to be involved in the assembly of the pre-replication
Table 5.3. Fatty acid composition of nuclear membrane-enriched fractions from S. cerevisiae NCYC 1383 previously grown in unsupplemented, linoleate- and linolenate-supplemented media, exposed to 100 μM Cu(NO₃)₂ for 6 h

<table>
<thead>
<tr>
<th>Fatty acid</th>
<th>Medium</th>
<th>Unsupplemented</th>
<th>18:2-supplemented</th>
<th>18:3-supplemented</th>
</tr>
</thead>
<tbody>
<tr>
<td>16:0</td>
<td>25.1±4.4</td>
<td>36.1±0.6</td>
<td>28.3±0.2</td>
<td></td>
</tr>
<tr>
<td>16:1</td>
<td>38.4±1.9</td>
<td>7.1±1.3</td>
<td>4.2±1.0</td>
<td></td>
</tr>
<tr>
<td>18:0</td>
<td>10.5±1.5</td>
<td>22.2±0.4</td>
<td>21.4±0.6</td>
<td></td>
</tr>
<tr>
<td>18:1</td>
<td>26.0±4.0</td>
<td>12.2±0.1</td>
<td>11.9±0.8</td>
<td></td>
</tr>
<tr>
<td>18:2</td>
<td>tr.</td>
<td>22.4±1.2</td>
<td>tr.</td>
<td></td>
</tr>
<tr>
<td>18:3</td>
<td>n.d.</td>
<td>n.d.</td>
<td>34.5±2.2</td>
<td></td>
</tr>
<tr>
<td>U.I.</td>
<td>0.64</td>
<td>0.64</td>
<td>1.20</td>
<td></td>
</tr>
</tbody>
</table>

Footnote as for Table 5.2.
initiation complex that precedes DNA replication initiation in *S. cerevisiae* (Aparicio *et al.*, 1997).

Therefore, Mcm3p in nuclear membrane-enriched fractions from unsupplemented and PUFA-supplemented cells, incubated in the absence and presence of 100 μM Cu(NO₃)₂, was examined to determine differences, if any, in the nuclear localisation, i.e. chromatin-bound or nucleosolic, of Mcm3p in over-replicating cells. After 6 h incubation, cells were sequentially fractionated in to nucleosolic (NS) and nuclear pellet (NP) fractions. Equal volumes of the NS and NP fractions (containing variable amounts of protein) were separated using 7.5% SDS-PAGE and analysed on nitrocellulose Western blots (Fig. 5.8). Mcm3p was detected with anti-Mcm3 antibodies (Hu *et al.*, 1993). The Mcm3p antibody cross-reacted with numerous proteins (Fig 5.8). Several studies with Mcm antibodies have revealed similar results (Merchant *et al.*, 1997; Lei *et al.*, 1996; Chong *et al.*, 1995), suggestive of cross-reactivity with the remaining members of the MCM family (all proteins carry a highly conserved motif (the MCM box)), and possibly protein degradation (Lei *et al.*, 1996). Mcm3p (107 kDa) was detected in the nucleosolic fractions of unsupplemented (lane 1), 18:2- (lane 2) and 18:3- (lane 3) supplemented cells, and in the nuclear pellet fractions of unsupplemented (lane 7) and 18:2-supplemented (lane 8) cells, incubated without Cu(NO₃)₂ (Fig. 5.8b). Significantly, of the Cu²⁺-exposed cells, Mcm3p was only detected in the nucleosolic fraction of unsupplemented cells (lane 4).

5.3.11. Determination of Cdc28p in nuclear fractions of *S. cerevisiae* RJD621. Levels of the cyclin-dependent kinase Cdc28p do not fluctuate throughout the cell cycle. However,
Fig. 5.8. Detection of Mcm3p in nuclear fractions of *S. cerevisiae* NCYC 1383 incubated in the presence and absence of 100 μM Cu(NO₃)₂. After 6 h incubation nuclear fractions were purified, proteins were electrophoresed on 7.5% SDS-PAGE gels, western blotted onto nitrocellulose membranes, and probed with monoclonal anti-Mcm3p. Samples are abbreviated with the first letters, Y, L, and E, representing cells previously grown in unsupplemented, linoleate- and linolenate-supplemented medium, respectively. The letter M following Y, L, or E, corresponds to cells incubated in the presence of Cu(NO₃)₂. NS and NP represent nucleosolic and nuclear pellet fractions, respectively.
Mcm3p →

<table>
<thead>
<tr>
<th>YNS</th>
<th>LNS</th>
<th>ENS</th>
<th>YMNS</th>
<th>YNP</th>
<th>LNP</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

— 97.4 kDa
— 116 kDa
through association with CLN- and B-type cyclins Cdc28p kinase activity rises from undetectable levels in early G1 to maximal levels during nuclear division (Lew et al., 1997). Cdc28p, in association with any of the six B-type cyclins, is required for the activation of pre-replication initiation complexes, resulting in the firing of origins of replication (Lew et al., 1997). Cdc28p is thus central to the interdependence of DNA replication and mitosis under normal conditions. As these processes are evidently uncoupled in Cu²⁺-exposed PUFA-enriched cells, we investigated whether differences, if any, in levels or localisation of Cdc28p in DNA over-replicating cells could be detected.

The yeast strain S. cerevisiae RJD621 (cdc28::CDC28-HA) was used for this particular study as the epitope-tagged Cdc28p can be conveniently detected using anti-HA antibodies. Toxicity experiments confirmed even more accentuated DNA over-replication in 18:3-enriched RJD621 exposed to Cu(NO₃)₂ (Fig. 5.9). After only 2 h exposure to 100 μM Cu(NO₃)₂, DNA levels of up to 8C were detected (Fig. 5.9b). Thus, after 6 h incubation, cells were sequentially fractionated in to cytosolic (C), high-salt nucleosolic (NSH), low-salt nucleosolic (NSL) and nuclear pellet (NP) fractions. Equal volumes of all fractions (containing equal amounts of protein, verified using the Micro-Lowry assay) were separated using 7.5% SDS-PAGE, and analysed on nitrocellulose Western blots (Fig. 5.10). Cdc28p-HA was detected using anti-HA antibodies. Cdc28p was not detected in any fractions from cells previously incubated in the presence of Cu(NO₃)₂. Significantly, an approximately 90 kDa band, most likely representing Cdc28p (34 kDa) in association with Clb2p (56 kDa) (Surana et al., 1991), was detected in the high-salt nucleosolic fractions of unsupplemented (lane 7), 18:2- (lane 8) and 18:3- (lane 9) supplemented cells, incubated without Cu(NO₃)₂ (Fig. 5.10). Elevated levels of
Cdc28p-Clb2p kinase activity are observed in *S. cerevisiae* during late G2 phase prior to mitosis (Surana *et al.*, 1993).
Fig. 5.9. Effect of 100 μM Cu(NO₃)₂ on the cellular DNA content of linolenate-enriched S. cerevisiae RJD621. The FL2 histograms depict the cellular DNA content of cells before copper addition (a) and after 2 h (b), 4 h (c) and 6 h (d) exposure to 100 μM Cu(NO₃)₂. Each histogram contains data collected from at least 10,000 cells.
Fig. 5.10. Detection of Cdc28p in nuclear fractions of *S. cerevisiae* RJD621 incubated in the presence and absence of 100 μM Cu(NO₃)₂. After 6 h incubation nuclear fractions were purified, proteins were electrophoresed on 7.5% SDS-PAGE gels, western blotted onto nitrocellulose membranes, and probed with anti-HA. Samples are abbreviated with the first letters, Y, L, and E, representing cells previously grown in unsupplemented, linoleate- and linolenate-supplemented medium, respectively. NSH, NSL and NP represent high-salt nucleosolic, low-salt nucleosolic and nuclear pellet fractions, respectively.
Cdc28p-HA → YNSH LNSH ENSH

- 97.4 kDa
- 66.2 kDa
5.4. Discussion

In this particular study we have investigated the effects of heavy metals and oxidative toxins on the cellular DNA content of *S. cerevisiae* NCYC 1383. Lipid peroxidation as a major means of heavy metal toxicity in *S. cerevisiae* had previously been demonstrated in Chapter 3. Levels of the lipid peroxidation byproducts, thiobarbituric acid-reactive substances (TBARS) and conjugated dienes were found to be markedly elevated in cells exogenously-enriched with the polyunsaturated fatty acids (PUFAs) linoleate and linolenate, upon exposure to Cu\(^{2+}\) and Cd\(^{2+}\). Lipid peroxidation is initiated in the presence of heavy metals through the generation of reactive oxygen species (ROS), such as the hydroxyl radical (HO'), superoxide anion (O\(_2^-\)), singlet oxygen (O), and the carbon-centered radical (L'), through Fenton and Haber-Weiss reactions (Dix and Aikens, 1993). These ROS have also been implicated as the causative agents of a broad spectrum of DNA damage, including double-strand breaks, which are thought to induce genome rearrangements such as deletions, duplications, and translocations (Brennan and Schiestl, 1998; Manivasakam and Schiestl, 1998). Therefore, we sought to investigate whether enrichment of *S. cerevisiae* with the PUFAs linoleate and linolenate would also predispose cells to greater levels of heavy metal-induced, ROS-mediated DNA damage.

Significantly, DNA over-replication was demonstrated in PUFA-enriched cells upon exposure to the redox-active metal species Cr\(^{6+}\) and Cu\(^{2+}\). Exposure of PUFA-enriched *S. cerevisiae*, particularly linolenate-enriched cells, to both Cu\(^{2+}\) and Cr\(^{6+}\) resulted in continued DNA synthesis and accumulation of DNA levels of up to 8C, in
certain cases. Metal-induced continued DNA synthesis in linolenate-enriched cells was confirmed by complete inhibition of DNA over-replication in the presence of the ribonucleotide reductase inhibitor hydroxyurea, and by the diphenylamine assay (Weinert et al., 1994).

In addition, flow cytometric cellular DNA profiles, in the early stages of Cr\textsuperscript{6+} and Cu\textsuperscript{2+} exposure, were suggestive of minor levels of DNA fragmentation. DNA fragmentation was manifested as decreased 1C and 2C DNA peak intensities and resolution, and a decrease in FL2 positive signal to noise ratio. The induction of hypoploidy and loss of G\textsubscript{1} peak resolution have been used as markers of DNA fragmentation, characteristic of apoptosis in tumorigenic cells (increased noise signal is also a characteristic of apoptotic-cell DNA histograms) (Takasu et al., 1998; Vernole et al., 1998). Also, while there was little evidence of DNA over-replication in cells pre-incubated with hydroxyurea, decreased 1C and 2C DNA peak intensities and resolution were still evident. The progression of DNA over-replication in the temperature-sensitive cdc16 and cdc27 mutants of S. cerevisiae, and cdc13 mutant of S. pombe, i.e. the absence of 1C and 2C DNA peak broadening and no decrease in the FL2 positive signal to noise ratio, lends further support for Cu\textsuperscript{2+}- and Cr\textsuperscript{6+}-induced DNA fragmentation, prior to DNA over-replication, observed in this study (Heichmann and Roberts, 1996; Hayles et al., 1994). It is plausible that DNA fragmentation might be a consequence of degradation of single strands from exposed 5' ends of Cu\textsuperscript{2+}- and Cr\textsuperscript{6+}-induced double-strand breaks during 'single-strand annealing' or 'one sided invasion' intrachromosomal recombination processes (Galli and Schiestl, 1998). However, further experimentation in the form of
TdT-mediated dUTP-biotin nick end-labelling (TUNEL) and pulsed-field electrophoresis are required to conclusively confirm DNA fragmentation.

Cu$^{2+}$-induced DNA over-replication was found to occur at a range of concentrations between 10 and 100 μM, indicating a threshold level of Cu$^{2+}$ ions below which induction does not occur. In addition, neither DNA over-replication or loss of peak resolution were evident in unsupplemented S. cerevisiae exposed to toxic concentrations of Cu$^{2+}$. This finding is consistent with earlier reports of hydroquinone (HQ) or benzoyl peroxide (BzPO) and Cu$^{2+}$-induced DNA-protein crosslinks (DPC) and DNA strand breaks, while Cu$^{2+}$ alone elicited no significant DNA damage (Altman et al., 1995; Akman et al., 1993; Li and Trush, 1993). Thus, it appears that Cu$^{2+}$ genotoxicity requires the presence of reducing agents such as HQ, BzPO or PUFAs.

DNA over-replication was also not observed after prolonged exposure of either unsupplemented or PUFA-supplemented S. cerevisiae to a highly toxic concentration of the nonredox-active metal Cd$^{2+}$. Exposure to the nonredox-active metals Cd$^{2+}$, Mn$^{2+}$ and Zn$^{2+}$, in the presence of HQ, has previously been shown to result in negligible deleterious effects to cellular DNA (Li and Trush, 1993). However, exposure to redox-active Cu$^{2+}$/HQ has been associated with the generation of extensive DNA lesions (Li and Trush, 1993). Thus, it is proposed that redox-cycling activity is essential for DNA over-replication and damage in metal-exposed linolenate-enriched cells. Significantly, in agreement with other reports (Chiu et al., 1995; Li and Trush, 1993; Rodriguez-Montelongo et al., 1993), Cu$^{2+}$-induced over-replication in linolenate-enriched cells was completely inhibited in the presence of the antioxidant reduced-glutathione. Reduced-
glutathione maintains a strong cellular reducing environment thereby preventing the
redox-cycling activity of Cu$^{2+}$ (Stephen and Jamieson, 1996; Li and Trush, 1993).

Hydrogen peroxide ($H_2O_2$), the hydroxyl radical (HO'), and the superoxide
anion/perhydroxyl radical ($O_2^{'-}H_2O_2$), have been implicated as the major ROS
responsible for eliciting deleterious effects to DNA (Keyer and Imlay, 1996; Brennan et
al., 1994). The perhydroxyl radical exists in dynamic equilibrium with the superoxide
anion in biological systems (Rego and Oliveira, 1995; Dix and Aikens, 1993). However,
Cu$^{2+}$-induced DNA over-replication was also evident in the presence of elevated levels of
the hydroxyl radical scavengers mannitol and DMSO. In addition, neither menadione or
$H_2O_2$ alone induced deleterious effects to DNA in any of the cell types examined. Thus,a
direct role of the ROS, HO', $O_2^{'-}$ or $H_2O_2$ in eliciting DNA over-replication in PUFA-
enriched cells exposed to Cu$^{2+}$ or Cr$^{6+}$ could not be established. Conducive with our
findings, the induction of DNA lesions in isolated nuclear chromatin by the Cu$^{2+}$- and
Fe$^{2+}$-EDTA/$H_2O_2$ systems has also been shown to occur in the presence of both mannitol
and DMSO (Chiu et al., 1995).

The reduction of Cu$^{2+}$ and Cr$^{6+}$ by endogenous intracellular reductants, such as
NADH or ascorbate, results in the generation of the ROS, $O_2^{'-}$, $H_2O_2$, and the reduced
forms of both metals, Cu$^+$ and Cr$^{3+}$ (Xu et al., 1996; Bridgewater et al., 1994; Rodriguez-
Montelongo et al., 1993). UV-visible spectroscopy and ESR have also indicated the
facilitation of reduction of Cu$^{2+}$ by NADH with the subsequent production of the highly
reactive carbon-centered radical (L') (Oikawa and Kawanishi, 1996). Furthermore, ROS-
mediated propagation of the lipid peroxidative chain in PUFA-enriched cells results in
the generation of the lipid hydroperoxides (LOOH) linoleate hydroperoxide and
linolenate hydroperoxide (Evans et al., 1998; Dix and Aikens, 1993). LOOHs are known to form delocalised lipid radicals which self-react to form dienoic dimers, or react with other LOOHs resulting in the generation of highly reactive lipid peroxyl radicals (LOO') (Evans et al., 1998). We therefore propose that DNA over-replication in PUFA-enriched cells, particularly linolenate-enriched cells, is mediated through Fenton-type interactions of LOOHs with Cu$^+$ or Cr$^{3+}$, resulting in the generation of Cu$^+$- and Cr$^{3+}$-peroxyl complexes (Chiu et al., 1995). Copper is also known to bind to non-histone proteins at the nuclear matrix attachment sites of DNA loops (Chiu et al., 1995). Thus, in addition to possible direct effects on nucleic acids, Cu$^{2+}$- and Cr$^{6+}$-induced DNA over-replication in linolenate enriched cells may be a consequence of Cu$^+$- and Cr$^{3+}$-peroxyl complex interactions with key DNA replication proteins.

Mcm3p and Cdc28p have been implicated as key DNA replication proteins in S. cerevisiae (Lew et al., 1997; Young et al., 1997). Mcm3p is constitutively present in the cytoplasm and nucleus throughout the cell cycle (Young et al., 1997; Young and Tye, 1997). Mcm3p remains chromatin-associated throughout G$_1$ phase and dissociates from chromatin, most likely as a result of Cdc28p-Clb2p or Cdc7p-Dbf4p phosphorylation, as cells enter S phase (Lew et al., 1997; Young et al., 1997; Young and Tye, 1997; Lei et al., 1996). Thus, Mcm3p is thought to form part of the pre-RCs that assemble at Ori from late M phase to early S phase, and thereby plays a significant role in restricting Ori activity to once per S phase (Dalton and Hopwood 1997; Young and Tye, 1997). Our results with unsupplemented and PUFA-supplemented cells, incubated in the absence of Cu(NO$_3$)$_2$, confirmed recent observations that two forms of Mcm3p exist in the nucleus of S. cerevisiae; a readily extractable form and a tightly chromatin-associated form, that
is released only upon treatment with DNase (Young and Tye, 1997). However, chromatin-associated Mcm3p was not detected in 18:3-enriched cells, previously incubated in the absence of Cu(NO₃)₂. This result may be a consequence of unequal protein loading; inherent differences in membrane physical properties of 18:3-enriched cells were associated with lower DNA and protein yields following nuclear fractionation. Of cells incubated in the presence of Cu²⁺, Mcm3p was only detected in the nucleosolic fraction of unsupplemented cells (DNA over-replication not evident). Thus, our preliminary results indicate that Mcm3p is not present in the nucleosolic or nuclear pellet fractions of over-replicating cells. Therefore, it could be postulated that interactions of Cu⁺-peroxyl complexes with Mcm3p prevent assembly of Mcm3p in to pre-RCs, resulting in continued firing of Oris for DNA replication.

Furthermore, Clb2p-associated Cdc28p (Surana et al., 1991), was only detected in the high salt nucleosolic fractions of unsupplemented and PUFA-supplemented S. cerevisiae, incubated in the absence of Cu(NO₃)₂. CLB2 transcription is activated during G₂ and M phases, and repressed during G₁ phase (Lew et al., 1997). In addition, S. cerevisiae has a requirement for Cdc28p-Clb2p kinase activity to enter M phase (Surana et al., 1993). Cdc28p was not detected in the nucleosolic or nuclear pellet fractions of unsupplemented or PUFA-supplemented cells previously incubated in the presence of Cu(NO₃)₂. Schizosaccharomyces pombe deleted for cdc13⁺, encoding a B-type cyclin, or overexpressing rum1p, a specific CDK inhibitor, undergoes continued DNA replication and sequential S phases in the absence of mitosis (Correa-Bordes and Nurse, 1995; Moreno and Nurse, 1994). Thus, it has been proposed that moderate levels of Cdc2p-Cdc13p kinase activity are required in G₂ phase to prevent additional rounds of DNA
replication in *S. pombe* (Stern and Nurse, 1996). Therefore, Cu\(^{2+}\) or Cr\(^{3+}\)-peroxyl complex interactions with Cdc28p-Clb2p in PUFA-enriched cells may result in diminished CDK activity, which is manifested as DNA over-replication. Furthermore, unlike the *cdc2\(^{2+}\)* gene of *S. pombe*, which is down-regulated in response to inhibition of DNA replication, mitotic CDK activity (Cdc28p-Clb2p) in *S. cerevisiae* remains elevated in cells arrested in S and G\(_2\)M phases by replication inhibition (Li and Cai, 1997). Our proposition is however complicated by the fact that elevated levels of Cdc28p-Clb2p kinase activity have previously been demonstrated during over-replication in the *S. cerevisiae cdc16* and *cdc27* temperature-sensitive mutants (Heichmann and Roberts, 1996).

We have also previously demonstrated Cu\(^{2+}\)-induced plasma membrane lipid peroxidation in PUFA-enriched cells (Chapter 3). Plasma membrane-lipid peroxidation was associated with a deterioration of membrane integrity and loss of membrane impermeability (Chapter 2; Van Ginkel and Sevanian, 1994). Nuclear membrane fatty acid compositional data of Cu\(^{2+}\)-exposed PUFA-enriched cells obtained in this study was also suggestive of lipid peroxidative membrane damage. Oxidative degradation of PUFAs was supported by the lower unsaturation indices of nuclear membranes of Cu\(^{2+}\)-exposed cells (De Vos *et al.*, 1993). Thus, in addition to possible direct interactions with Cu\(^{2+}\)-peroxyl complexes, loss of membrane impermeability may also account for the absence of Mcm3p and Cdc28p from nuclear membrane fractions of PUFA-enriched cell exposed to Cu\(^{2+}\).

As stated previously, repeated rounds of DNA replication have been observed for the *cdc16* and *cdc27* *S. cerevisiae* mutants (Heichmann and Roberts, 1996), the *cdc2* and
cdc13 S. pombe mutants, and upon overexpression of rum1+ and cdc18+ (homolog of S. cerevisiae cdc6p) in S. pombe (Stern and Nurse, 1996; Correa-Bordes and Nurse, 1995; Moreno and Nurse, 1994), indicating the critical function of these proteins in limiting DNA replication to once per cell cycle. Here we have demonstrated redox-active metal-induced DNA polyploidy in PUFA-enriched S. cerevisiae for the first time. We propose that DNA over-replication could be mediated through direct interactions of Cu2+ and Cr3+-peroxyl complexes with key pre-RC proteins (Cdc6p, Mcm3p or Mcm2p), critical cyclin-dependent kinases (Cdc28p-Clb2p or Cdc7p-Dbf4p), or by Cu2+- and Cr6+-induced gross lipid peroxidation-mediated nuclear membrane structural alterations, leading to loss of nuclear membrane impermeability. While further extensive experimental work is required to elucidate the exact mechanism of redox-active metal-induced DNA over-replication in PUFA-enriched S. cerevisiae, a major role of membrane fatty acid composition on heavy metal-induced genomic instability has been demonstrated.
CHAPTER 6

Conclusions
The effects of altered membrane fatty acid composition on the toxic interactions of heavy metals with the model organism *S. cerevisiae* were investigated. While other studies have investigated the effects of altered plasma membrane phospholipid unsaturation on the kinetics of solute accumulation in *S. cerevisiae* (Hoptroff et al., 1997; Keenan et al., 1982), this was the first study to examine the effects of such alterations on toxic interactions with heavy metals. *Saccharomyces cerevisiae* was selected for this study for several reasons; many aspects of the cellular organisation, function and structure of *S. cerevisiae* are common to those found in higher eukaryotes. Also, many regulatory mechanisms found in yeast, e.g. cell cycle-control, are conserved among larger, more complex eukaryotes. With the advent of the *Saccharomyces cerevisiae* genome database (Cherry et al., 1997), the development of powerful yeast genetic techniques, combined with its low DNA content and ease of culture, *S. cerevisiae* has been established as an excellent model eukaryotic system. Numerous studies have also proven *S. cerevisiae* highly amenable as a model organism for the specific examination of metal-microbe interactions (Aßmann et al., 1996; Ohsumi et al., 1988). In addition, studies of membrane biology in lower eukaryotes are most advanced for *S. cerevisiae* (Van der Rest et al., 1995; Van der Westhuisen et al., 1994). In agreement with several reports, we have demonstrated that *S. cerevisiae* is highly amenable to cultural manipulation of membrane composition, without adverse effects on growth rate (Avery et al., 1996; Parks et al., 1995; McDonough et al., 1992; Bossie and Martin, 1989). The consequent advantages of this system of membrane compositional manipulation, over conventional methods have been highlighted throughout this thesis. While the enrichment of *S. cerevisiae* with PUFAs elicited no discernible adverse phenotypic variations under non-
stressed conditions, susceptibility towards the toxic effects of heavy metals was greatly accentuated. Thus, cells that were previously enriched with the PUFAs linoleate or linolenate displayed elevated sensitivity towards both copper- and cadmium-induced plasma membrane permeabilisation and whole-cell toxicity. The implications of these findings for metal-microbe interactions in the environment are potentially very significant. The possibility that the large variability in microbial fatty acid composition, which can result from adaptive responses to changes in external physico-chemical parameters (Hazel and Williams, 1990; Suutari et al., 1990) as well as genetically-determined intrinsic differences, may partly dictate differing heavy metal sensitivities among microorganisms is potentially very important. Indeed, recent studies examining changes in microbial fatty acid profiles following metal addition to soil have suggested a greater impact on microorganisms rich in PUFAs (Pennanen et al., 1996). Thus, a passive role of modification of membrane fatty acid composition as an additional (in association with constitutive and inducible elements) determinant of heavy metal resistance is plausible.

Initial results implied a role of oxy-radical mediated effects in the toxicity of cadmium towards *S. cerevisiae*. Prolonged incubation of both unsupplemented and PUFA-supplemented *S. cerevisiae* in the presence of cadmium was associated with reduced cellular fatty acid unsaturation indices during the exponential growth phase. The depletion of unsaturated fatty acids has been used as a marker for lipid peroxidation (De Vos et al., 1993). As PUFAs are well known to be markedly susceptible to oxy-radical mediated lipid peroxidative degradation (Porter et al., 1995; Dix and Aikens, 1993), the increased susceptibility of PUFA-enriched cells to plasma membrane permeabilisation
and whole-cell toxicity was postulated to be a direct consequence of increased lipid peroxidation in these cells. Elevated levels of TBARS and conjugated dienes (lipid peroxidation decomposition products) confirmed accentuated oxidative degradation of cellular unsaturated fatty acids in metal-exposed PUFA-enriched cells. Thus, lipid peroxidation as a major means of heavy metal toxicity in microorganisms was demonstrated for the first time. The present results may prove useful in modelling lipid peroxidative processes associated with a multitude of pathological conditions in higher organisms. For example, the accumulation of fibrous plaques within the endothelial lining of blood vessels (culminating in atherosclerosis) is a direct consequence of low density lipoprotein (LDL) oxidation (Steinbrecher et al., 1990). More significantly, transition metal-induced (e.g. aluminium, iron, and mercury) lipid peroxidation has been implicated in the aetiology of the neurodegenerative diseases Parkinson's disease and Alzheimer's dementia (Bondy et al., 1998; Hock et al., 1998; Rajan et al., 1997).

The influence of cellular fatty acid composition on the interactions of heavy metals with cellular nucleic acids was also investigated. PUFA-enrichment was clearly associated with enhanced heavy metal-induced genotoxic effects. Gross genomic instability, manifested as continued DNA synthesis in the absence of cell division, was evident following exposure of PUFA-enriched cells, particularly linolenate-enriched cells, to the redox active metals Cu²⁺ and Cr⁶⁺. While ROS-induced DNA adducts and double-strand breaks have been associated with gross genome rearrangements, e.g. deletions, duplications, and translocations (Manivasakam and Schiestl, 1998; Brennan et al., 1994), this is the first report of heavy metal-induced polyploidy. It is postulated that interactions of Cu⁺ or Cr³⁺-peroxyl complexes with key DNA replication proteins or
cyclin-CDK complexes could be one reason for the loss of cell cycle checkpoint control, leading to DNA over-replication. Elucidation of the precise molecular mechanism underlying heavy metal-induced DNA over-replication in PUFA-enriched cells may serve highly useful for carcinogenesis studies, as greatly accelerated endogenous gene amplification is a feature of tumorigenic cells. The expression of viral oncoproteins in human fibroblasts results in the inactivation of the p53 and Rb tumor suppressor genes, leading to subunit alterations of cyclin-CDK complexes (Xiong et al., 1996; Schaefer et al., 1993). The consequences of such alterations include loss of cell cycle checkpoint control and cytogenetic aberrations (Xiong et al., 1996). Also, individuals with Li-Fraumeni Syndrome (LFS) carry congenital mutations in one of their p53 alleles, leading to accelerated gene amplification (polyploidy) and loss of cell cycle checkpoint control (Thea D. Tlsty, unpublished results).

We also examined the heterogeneity in copper sensitivity of genetically-homogeneous *S. cerevisiae* using flow cytometry. Flow cytometry is a powerful technique which enables the simultaneous, multiparametric analysis of single cells within heterogeneous populations (Jayat and Ratinaud, 1993; Lloyd, 1993). The use of flow cytometry for the analysis of specific cell cycle stage populations was validated. Using this methodology, a distinct relationship between cell cycle stage and sensitivity to copper toxicity was demonstrated, whereby cell populations consisting predominantly of 2C DNA cells (G2/M phase) were the most susceptible to copper toxicity. The fatty acid compositions of different cell cycle stage sub-populations remained relatively constant, thus eliminating a role of fatty acid compositional variations in the cell cycle stage-dependency of sensitivity to copper toxicity. However, using the oxidant-sensitive probe
H₂DCFDA, the increased sensitivity of G₂/M phase cells was correlated with elevated levels of endogenous reactive oxygen species in these cells. Our findings are consistent with several reports demonstrating the elevated toxin-sensitivity of S and G₂ phase cells (Nitiss and Wang, 1996; Galli and Schiestl, 1995). We also describe a novel flow cytometric technique for the analysis of the promoter activity of genes of interest in distinct sub-populations, using the green fluorescent protein of *Aequorea victoria*. While differences in basal levels of *CUP1* activity between live and dead sub-populations were not evident, our results with *SOD1* were inconclusive and further experimentation is required to determine if the large heterogeneity in *SOD1* activity can be correlated with the cell cycle stage-dependency of susceptibility to copper toxicity. Indeed, several studies with higher organisms have demonstrated changes in *SOD1* activity during the cell cycle (Oberley *et al.*, 1995; Kong *et al.*, 1993). The major biological importance of the SOD oxidative stress response proteins has been highlighted by their role in familial amyotrophic lateral sclerosis (FALS), a motor neuron degenerative disease (Wiedau-Pazos *et al.*, 1996).

Thus, the effects of altered membrane fatty acid composition on the toxic interactions of heavy metals with *Saccharomyces cerevisiae* have been examined. The impact of increased membrane fatty acid composition on susceptibility to heavy metal-induced reactive oxygen species-mediated cellular damage was marked. As well as being of relevance to the stress biochemistry of yeast, the findings may have significant implications for metal-microbe interactions in the natural environment, and for deciphering the mechanisms of loss of cell cycle control under stress conditions.
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