New Roles for Copper Metabolism in Cell Proliferation, Signaling, and Disease*

Over the past 2 decades, the molecular mechanisms by which cells acquire, distribute, and utilize copper have been under intense investigation. Significant progress has been made in the identification of genes encoding copper homeostasis proteins and in fundamental aspects of their structure, function, and mechanisms of action in copper balance. A number of more comprehensive reviews of the field with respect to the genetics, structure, function, and physiology of copper metabolism have recently appeared elsewhere (1–4). Here, we review general mechanisms for eukaryotic copper metabolism at the cellular level in the context of recent discoveries in the field, identifying potential new functions for copper and copper metabolism proteins in cell signaling, gene expression, tumor cell metastasis, and resistance to anti-neoplastic drugs.

Copper as a Catalytic and Structural Cofactor in Biology

Copper is a redox-active metal that is predominantly used by organisms living in oxygen-rich environments and that fluctuates between the oxidized (Cu²⁺) and reduced (Cu⁺) states (5). With these changes in redox state, copper can coordinate to a range of ligands that include carboxylate oxygen, imidazole nitrogen, cysteine thiolate, and methionine thioether groups and engages in cation-π interactions (6–8). Cu⁺ plays a well established structural role in proteins such as the fungal copper-activated metalloregulatory transcription factors (9). For example, Ace1 in Saccharomyces cerevisiae acquires profound conformational changes upon the cooperative binding of a tetra-Cu⁺ cluster, activating sequence-specific DNA binding (10).

Many enzymes harness the changes in bound copper oxidation state, in the presence of oxygen, to catalyze redox chemistry for a wide range of chemical transformations that are central to biology (Table 1). Significant phenotypes result from a diminution in these enzymatic activities (Table 1). Although copper-binding proteins and enzymes have been characterized for over a century, it is likely that a number of copper-dependent proteins and their functions have yet to be identified. The diversity of potential copper ligands along a polypeptide primary structure, in addition to the proper juxtaposition of bona fide ligands in the correct three-dimensional orientation in proteins, makes prediction of the entire copper proteome a challenging area for future studies (5).

Copper Homeostasis in Eukaryotic Cells

Over the past few decades, critical progress has been made in the identification of genes encoding proteins that function in copper uptake, intracellular distribution, and efflux and in the regulation of the copper homeostasis machinery (1–4). Many of the structural, functional, and regulatory details have been strikingly conserved from microbes to humans. Fig. 1 shows a model for copper homeostasis in a typical mammalian cell. Ctr1, an integral membrane protein, functions as a major copper importer at the plasma membrane. Genetic, biochemical, and structural studies support a model in which Ctr1 homodimerizes to form a central region of low electron density through which copper may traverse the plasma membrane (11–13). The genetic requirement for a metalloreductase in yeast for Ctr1-mediated high affinity copper uptake, in addition to other data (1, 14), supports the transport of Cu⁺ rather than Cu²⁺. Conserved methionine residues are present in the Ctr1 extracellular domain as Met-X-Met or Met-X₂-Met and, although not essential for activity, are needed for high affinity Cu⁺ import. Additionally, a conserved Met motif, Met-X₂-Met, present in the Ctr1 second transmembrane domain, is essential for Cu⁺ import (13). These and other experimental results are consistent with thioether-Cu⁺ coordination to Ctr1 at one or more steps in the import process.

Copper that is imported by Ctr1, or as yet unidentified, low affinity, Ctr1-independent activities (15), is delivered to intracellular proteins and compartments by the action of copper chaperones (1, 2, 16). CCS (copper chaperone for Cu,Zn-superoxide dismutase) delivers copper to Cu,Zn-SOD² via direct interactions and a complex mechanism that has been recently reviewed (17). Atox1 (Atx1 in bakers’ yeast, S. cerevisiae) is a small soluble protein that binds a single Cu⁺ atom in a specific but kinetically labile manner. As shown in Fig. 1, Atox1 directly interacts with repeated cytosolic metal-binding domains of the Cu⁺-transporting ATPases ATP7A and ATP7B to facilitate its delivery across the secretory compartment or, under conditions of high copper, across the basolateral membrane of cells.

In addition to delivery of copper to cytosolic proteins and to the secretory compartment, copper must be targeted to mitochondria, where cytochrome oxidase uses copper for oxidative phosphorylation. Genetic studies in microbes and the mapping of mutated genes that are responsible for defects in cytochrome oxidase assembly have identified many proteins involved in cytochrome oxidase copper metallation (2). These include Cox17 in the mitochondrial intermembrane space; Cox11 in the mitochondrial inner membrane; and Sco1 and Sco2, structurally similar proteins that play a more proximal role in copper delivery to cytochrome oxidase. Defects in Sco1 and Sco2 cause...
**A New Role for Ctr1 in Cisplatin Accumulation**

Although Cu\(^+\) is the only known physiological substrate for Ctr1, recent studies strongly suggest that Ctr1 in yeast and in mammals facilitates uptake of the metal-based anti-neoplastic drug cisplatin (19, 20). Cisplatin is an effective chemotherapeutic agent due to its ability to intercalate in DNA and thereby preferentially inhibit the growth of rapidly proliferating cells (21). Transposon mutagenesis studies in yeast identified Ctr1 as a loss-of-function mutation rendering these mutants highly resistant to cisplatin toxicity (19). Furthermore, *Ctr1*\(^{+/+}\), *Ctr1*\(^{+/−}\), or *Ctr1*\(^{−/−}\) mouse embryonic fibroblasts exhibit *Ctr1* gene dose-dependent acquisition of cisplatin, with *Ctr1*\(^{−/−}\) cells exhibiting elevated resistance to cisplatin compared with wild-type cells (19). Moreover, cell lines overexpressing human Ctr1 accumulate elevated cisplatin compared with controls cells (20). How does a dedicated Cu\(^+\) transporter facilitate the uptake of the structurally unrelated catastrophic defects in cytochrome oxidase assembly and result in severe human disease (18).

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### TABLE 1

| Enzyme/Protein | Function | Consequence of Deficiency |
|----------------|----------|---------------------------|
| Cu,Zn Superoxide Dismutase | Superoxide disproportionation; signaling | Oxidative stress; hepatocellular carcinoma; neurodegeneration |
| Cytochrome Oxidase | Mitochondrial oxidative phosphorylation; ATP production | Respiratory deficiency; cardiomyopathy; lethality |
| Tyrosinase/Laccase | Melanin synthesis; virulence and pathogenicity in fungi; innate immunity | Pigmentation Defects/Albinism; reduced fungal virulence |
| Pepidylglycine ε-aminomutase | Peptide amidation/maturaton | Heart rate defects; endocrine dysfunction; lethality |
| Dopamine β-Hydroxylase | Noradrenaline synthesis | Hypoglycemia; hypotension |
| Ceruloplasmin | Ferroxidase; Fe loading onto transferrin | Aceroeroulasminemia: progressive anemia, neurodegeneration, diabetes |
| Lysyl Oxidase | Covalent crosslinking of collagens and elastin | Anemia, impaired Fe absorption by peripheral tissues |
| Lysyl Oxidase-like Proteins | Oxidation of small transcription factor resulting in E-cadherin silencing and promoting EMT | Aortic aneurisms, cardiovascular dysfunction |
| Coagulation Factors V and VIII | Blood Clotting | Altered cell-cell contacts |
| Nitrous Oxide Reductase | Catalysis of reduction of NO and O2 in denitrification pathway of bacteria | Respiratory deficiency, imbalance in nitrogen cycle |
| Ethylene Receptor | Ethylene signal transduction | Plant senescence, fruit ripening, growth |
| XIAP | Inhibitor of apoptosis | Copper deficiency in mouse knock-out |
| Ace1 | *S. cerevisiae* transcription factor active under conditions of high intracellular copper | Inability to grow on high copper media |
| Copper Amine Oxidase | Deamination of primary amines to aldehyde | Impaired immune responses (in AOC3 knock-out mouse); reduced fat deposition in obese mice |

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**FIGURE 1. Mammalian intracellular copper homeostasis.** Copper is reduced by an as yet unidentified reductase and imported into cells by the high affinity copper transporter Ctr1. Once inside the cell, copper is escorted to destinations via the action of copper chaperones. CCS incorporates copper into the cytosolic protein Cu,Zn-SOD. Atox1 delivers copper to the secretory compartment; the protein possesses an N-terminal copper-binding motif (CXXC) that directly interacts with the N-terminal copper-binding domain of ATP7A and ATP7B (MTCXXC), facilitating copper delivery to ATP7A and ATP7B. ATP7A and ATP7B are P-type copper-transporting proteins that transport copper into the lumen of the Golgi, where the metal can then be incorporated into copper-dependent proteins such as the secreted form of LOX. The P-type ATPases also export copper out of the cell by translocating to the plasma membrane when intracellular copper levels are high. Copper is escorted to the mitochondria by an as yet uncharacterized ligand. Once inside the intermembrane space, copper is handed off to Cox17 and then passed onto either Sco1, which transfers copper to the Cox2 subunit of cytochrome oxidase, or Cox11, which transfers copper to the Cox1 subunit of cytochrome oxidase (CCO). TGN, trans-Golgi network; ER, endoplasmic reticulum.
metal complex cisplatin? Recent mutagenesis experiments and FRET studies between yeast Ctrl subunits suggest distinct mechanisms for Cu⁺ import compared with cisplatin uptake (22). Lee and co-workers (22) demonstrated that copper enhances the FRET signal of functional Ctrl1-enhanced cyan fluorescent protein and Ctrl1-enhanced yellow fluorescent protein monomers coexpressed in yeast cells in a manner that is dependent on copper transport-competent Ctrl1, suggesting that copper-induced conformational changes occur in concert with Cu⁺ movement through the plasma membrane. In contrast, although the Ctrl1 FRET pairs were capable of stimulating cisplatin uptake, cisplatin neither induced FRET nor abrogated the copper-induced FRET signal. Moreover, a Ctrl1 mutant lacking the Met-rich ectodomain was competent for Cu⁺ uptake but completely defective in cisplatin accumulation, further underscoring differences between Cu⁺ uptake and cisplatin accumulation. Similar mutations in human Ctrl1 extracellular Met-X-Met motifs abrogated cisplatin-stimulated Ctrl1 multimerization (23), suggesting that cisplatin may interact with these methionine thioether groups. Taken together, these studies suggest that Ctrl1 may function similarly to a transporter for Cu⁺ acquisition but may enhance cisplatin uptake in a methionine-rich, ectodomain-dependent, receptor-mediated endocytic mechanism. Although there are conflicting reports on whether, like copper, cisplatin enhances Ctrl1 endocytosis (19, 22–24), this question must be resolved to understand how Ctrl1 facilitates cisplatin accumulation in a manner distinct from that of copper.

**Ctrl1 in Growth Factor Signaling**

The generation of a mouse Ctrl1 knock-out illustrated the essentiality for Ctrl1 in metazoan development (25, 26). Ctrl1⁻/⁻ mutant embryos showed growth retardation at embryonic day 7.5, a stage in development when gastrulation begins, with embryonic lethality by embryonic day 12.5. Homozygous mutants histologically demonstrated poorly developed neural ectoderm and mesoderm and a defect in neural tube closure. Although the precise mechanisms underlying the mutant phenotypes have not been elucidated, it is possible that defects in multiple copper-dependent proteins culminate in catastrophic developmental arrest. However, recent work in *Xenopus* may provide an alternative explanation for the requirement for Ctrl1 in normal growth and development (27). *Xenopus Ctrl1 (Xctr1)* was identified as part of a complex consisting of Laloo and SNT-1, proteins required for FGF-mediated mesoderm induction (27). Misexpression of Xctr1 promoted mesodermal and neural ectodermal differentiation, whereas, conversely, loss of Xctr1 through morpholino injection inhibited these processes. Knockdown of Xctr1 also had an effect on pathway activation dynamics as evidenced by partial inhibition of SNT-1 phosphorylation by Laloo and strong inhibition of FGF-dependent ERK phosphorylation. Interestingly, knockdown of Xctr1 at later stages of development only modestly inhibited induction of mesoderm markers, suggesting that Xctr1 is required either before or during gastrulation for mesoderm development, a time window that parallels the developmental arrest of Ctrl1⁻/⁻ mutant embryos.

This study provides a potential explanation for the developmental abnormalities seen in mouse Ctrl1⁻/⁻ mutant embryos and forges a new intersection between copper homeostasis and a cell signaling pathway. However, the role of Xctr1 in mesoderm development is suggested to be independent of its copper-transporting activity, as misexpression of Xctr1 carrying mutations in conserved residues required for copper transport gave the equivalent phenotype as misexpression of wild-type Xctr1 (27). This unanticipated copper transport-independent role for Ctrl1 in FGF signaling, morphogenesis, and differentiation must be further explored, as other reports using copper chelators suggest that copper levels modulate hematopoietic stem cell differentiation (28, 29).

**Copper-dependent Metabolic Changes in Cancer Cells**

Recent exciting work has implicated copper-handling and copper-utilizing proteins in controlling the striking metabolic changes that have long been known to occur in cancer cells. Otto Warburg (30) discovered that respiratory capacity is down-regulated in many cancer cell types with a concomitant dependence on glycolysis for cellular energy generation. Although this phenomenon has been known for over 75 years, the cellular regulatory mechanisms governing this metabolic switch are still not well understood. Matoba et al. (31) reasoned that this metabolic reengineering may result from regulatory changes common to many cancer cell types. Because mutations in the gene encoding the p53 sequence-specific DNA-binding transcription factor are among the most common genetic changes found in a broad range of cancer cells, they explored whether cancer cell lines harboring p53 mutations exhibit changes in their dependence on respiration versus glycolysis. Indeed, human colon cancer cells with p53-inactivating mutations also showed significant reductions in oxygen consumption while generating increased levels of lactate, a by-product of glycolysis, metabolic characteristics reflecting a switch from oxidative phosphorylation to glycolysis typical of the Warburg effect.

As the SCO2 gene was previously identified as a potential p53 target, the expression of SCO2 was explored as a function of p53 gene dosage (31, 32). Although SCO2 mRNA and protein levels were considerably reduced in p53⁺/⁺ and p53⁻/⁻ mouse livers compared with wild-type livers, SCO1 mRNA levels were unchanged. Furthermore, the p53-dependent stimulation of SCO2 expression was dependent on a canonical p53-binding site in the SCO2 promoter, establishing a direct functional link between p53 and SCO2 expression levels. Taken together, these studies identified a link between the p53 status of cancer cells and metabolic reprogramming toward glycolytic energy generation first observed by Warburg. With the essential role for copper in cytochrome oxidase function, it would be interesting to ascertain if p53-dependent changes in the expression of other factors that function in copper acquisition might also play a role in the Warburg effect.

**Copper in Signaling and Tumor Cell Metastasis**

Cell migration and tissue invasion from the primary tumor site require a loss of integrity of target tissue cell-cell junctions. The epithelial-mesenchymal transition is a critical step in the initiation of cell invasion, as it results in compromised cell-cell...
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Copper Deficiency: Who Goes First?

Important insights into the hierarchy of copper metalla- tion in vertebrates have recently been gained through a small molecule chemical screen in zebrafish (42). A search was conducted for molecules that interfered with copper metabolism, specifically giving rise to copper-deficient phenotypes as evidenced by lack of pigmentation. Zebrafish lateral line pigmentation is mediated by tyrosinase, a copper-dependent enzyme required for melanin synthesis (43). The most severe copper-deficient phenotype, obtained by treating embryos with the copper chelator neocuproine, resulted in a deformed notochord, impaired neurogenesis, inappropriate cartilage maturation, poor vascular development, and hematopoietic defects. Furthermore, it was found that varying copper chelator concentrations would generate a continuum of phenotypes that could be correlated with the severity of copper deficiency within the embryo, from a lack of pigmentation but with a normal notochord to a lack of pigmentation and abnormal notochord development. Continued exposure of embryos to neocuproine resulted in a lack of pigmentation, notochord defects, and neurogenesis defects. These results suggest that, at early stages of copper deficiency, pigmentation processes mediated by tyrosinase would be among the first to be lost, freeing up limited copper reservoirs for more essential developmental processes. As the developing embryo encounters increasingly stringent access to copper, mechanisms appear to be in place that prioritize copper utilization to meet physiological demands.

Copper Chelation and Cancer Chemotherapy

Copper has been proposed to be essential for angiogenesis, the formation of new blood vessels that provide a delivery route for nutrients, growth factors, and other signaling agents important for tumour growth and survival. Although a number of copper-dependent roles in angiogenesis have been proposed (44), the entire range of functions important for efficient angiogenesis that require copper has not been elucidated. This is an important challenge, as TTM, a potent Cu²⁺ chelator, has been reported to be of therapeutic value in the treatment of several types of cancers as an anti-angiogenesis and anti-cancer molecule. Recently, ATN-224, an orally available choline salt derivative of TTM, has been suggested to preferentially target Cu,Zn-SOD in tumor and endothelial cells, with the implication that SOD1 plays a stimulatory role in growth factor signaling (45, 46). SOD1 protects cells from oxidative stress and catalyzes the disproportionation of superoxide to hydrogen peroxide, molecules that have been established to play signaling roles in biology (47).

In an initial report, TTM was shown to potently inhibit Cu,Zn-SOD activity, thereby increasing superoxide anion accumulation and inducing programmed cell death in multiple myeloma cells (45). In parallel with these features, the phosphorylation of ERK, which has critical functions downstream of Ras in cell proliferation, was also inhibited (45). Recently, ATN-224 was reported to inhibit growth factor-stimulated phosphorylation of the EGF and insulin-like growth factor receptors in parallel with an increase in superoxide and a decrease in hydrogen peroxide accumulation (46). This decrease in hydrogen peroxide was proposed to reduce the inactivation of the protein-tyrosine phosphatase PTP1B, thereby inhibiting growth factor-stimulated receptor phosphorylation and attenuating downstream activation of proliferation pathways involving ERK. This is a potentially important mechanism for the action of copper chelators, but many questions remain. It is unclear why ATN-224 is effective for inhibiting SOD1 activity in cell culture in the nanomolar range but exhibits potent anti-proliferative effects at micromolar concentrations. Moreover, RNA interference-mediated knockdown of SOD1 did not exhibit anti-proliferative effects, and Sod1 knock-out mice did not show any obvious growth defects (48). Although Cu,Zn-SOD may be an important target for the anti-angiogenesis and anti-cancer activity of clinically relevant copper chelators, it is not currently clear whether other copper-dependent activities might play a role. A thorough understanding of the anti-proliferative mechanism of copper chelators will require a comprehensive identification of the copper proteome and a deeper understanding of the downstream effects of copper chelation.
understanding of the mechanisms of copper homeostasis and their regulation.

Future Directions

To date, genetic studies in model systems and biochemical, cell biology, and structural studies have identified many proteins involved in eukaryotic copper homeostasis and provided insights into their structures and mechanisms of action. It is likely that there are other functions and mechanisms to be discovered for known copper homeostasis proteins, and new components of this machinery will be discovered and integrated into our current understanding of cellular copper balance. Many questions have yet to be tackled in the field of copper metabolism, from new, exciting, and perhaps serendipitous perspectives. How does copper make its way from the site of import at the plasma membrane, or efflux from intracellular storage vesicles, to the copper chaperones and other intracellular ligands? What is the entire constellation of copper-binding proteins in a mammalian cell? How do the components of the copper homeostasis machinery communicate, and what other signals, including chemical, hormonal, and environmental, trigger changes in copper homeostasis? How do multicellular organisms sense copper deficiency in peripheral tissues and transmit this signal to mobilize copper stores? The field of copper homeostasis is currently similar to a high school graduate going off to college, armed with some basic information; we will see how things develop with additional outside influence.

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