Molecular regulation of copper excretion in the liver

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Cu is an essential nutrient that is required for a broad range of cellular and molecular processes. Mammals have efficient systems to control Cu homeostasis that operate at the level of controlling uptake, distribution, sequestration and excretion of Cu. The study of diseases associated with disturbed Cu homeostasis has greatly enhanced our understanding of the molecular mechanisms involved in Cu metabolism. In man the liver is responsible for excreting excess Cu from the body by means of biliary secretion. Wilson disease is a severe human disorder characterized by Cu accumulation in the liver as a result of a deficiency in biliary Cu secretion. This disorder is caused by mutations in the gene that encodes a Cu-transporting P-type ATPase (ATP7B). The MURR1 gene was identified recently, and it was hypothesized that this gene is also essential for biliary Cu excretion and is presumed to act downstream of ATP7B. MURR1 is mutated in canine Cu toxicosis, a disorder with phenotypic characteristics similar to those of Wilson disease. MURR1 encodes a protein that is of unknown function and is without detectable sequence homology to known proteins. MURR1 is readily detected in all tissues and cell types, suggesting that it may exhibit a pleiotropic function in different organs, which may or may not be exclusively linked to Cu homeostasis. The use of genetic, biochemical and genomic tools, as well as the development of appropriate models in organisms other than dog, will allow the elucidation of the molecular and cellular function of MURR1 in relation to hepatic Cu homeostasis and biliary Cu excretion.

Copper metabolism: Biliary excretion: MURR1

A balanced diet contains adequate amounts of protein, carbohydrates, fat, vitamins and minerals, including trace metals. Trace metals can be essential (e.g. Fe, Zn, Cu and Mn) or non-essential (e.g. Pb and Hg). Non-essential trace metals can be toxic to life, even in extremely small quantities. In contrast, the essential trace metals sustain important biological functions, although excess amounts can also be toxic. Homeostatic control is defined as the balance between essentiality and toxicity, and it is maintained by various mechanisms, including uptake, transport, storage and excretion of metals.

Cu is one of the essential trace metals for all organisms, including man. Cu can exist in two redox states, which renders it an essential component of the active site of a number of cupro-enzymes, e.g. the cytochrome C oxidase complex, a component of the mitochondrial electron transport chain that plays an essential role in cellular respiration. Cu also forms the catalytic component of Cu/Zn superoxide dismutase, which protects against free radicals.

Paradoxically, this same property can also make Cu toxic when present in excess, because it is thought to lead to the formation of reactive oxygen species through Fenton chemistry. Free Cu ions can associate with adventitious sites, thereby impairing normal cellular function.

Recent developments and novel prospects in the understanding of the homeostatic control of Cu will be addressed. The recent exciting discovery of MURR1 as a Cu-homeostasis gene and the role of MURR1 in the regulation of biliary Cu excretion will be used as a recurrent theme to outline new genetic, genomic, proteomic and cell biological approaches to further our understanding of the regulation of Cu metabolism in general.

Copper

To maintain its essential functions, it is important that there is a sufficient daily intake of Cu in the diet. Cu is
found in a wide variety of foods, including cereals, meat, fruits and vegetables. Drinking water can also be a major source of Cu intake, especially when Cu water pipes are corroded. The acceptable range of oral intake of Cu is between 1.3 and 13 mg Cu/d for adults, and between 0.6 and 1 mg Cu/d for children. A schematic representation of the regulation of human Cu metabolism is presented in Fig. 1.

Approximately 15% of the Cu taken up from the diet is transported to various tissues, while the remaining 85% of the Cu is excreted. Under normal physiological conditions about 98% of the Cu excretion is via the bile and only 2% via the urine, indicating that the liver is the organ in which whole-body Cu homeostasis is achieved by regulation of biliary excretion. Dietary Cu is absorbed across the mucosal membrane of the upper intestine and transported to the rest of the body. Cu enters the bloodstream, where it binds to proteins or amino acids; thereafter Cu leaves the blood and enters the liver and kidneys. The liver takes up the majority of the Cu present in the circulation. Cu may enter the hepatocyte by means of Cu transporter 1 (CTR1), after which it is distributed intracellularly into one of three pathways by the Cu chaperones cytochrome C oxidase assembly factor 17 (COX17), ATOX1 and Cu chaperone for superoxide dismutase 1 (CCS1) for incorporation into cytochrome C oxidase, Cu-transporting P-type ATPase (ATP7B) and Cu/Zn superoxide dismutase respectively (Fig. 2). The cellular function of the Cu chaperone ATOX1 will be discussed in more detail later (see p. 33). Since there is virtually no free Cu available in the hepatocyte, any excess Cu is bound to metallothioneins, which act as scavengers. Our knowledge of the general mechanisms of cellular Cu homeostasis forms the foundation of our understanding of the role of the liver in whole-body Cu homeostasis. The characterization of Cu metabolism in the yeast Saccharomyces cerevisiae and other model organisms has contributed much to our knowledge of this cellular process (Askwith & Kaplan, 1998; De Freitas et al. 2003).

**Copper excretion from hepatocytes**

To allow efficient elimination of toxic compounds via biliary excretion, hepatocytes have developed mechanisms that generally apply to many different substances. Hepatocytes are polarized into an apical (canalicular) domain and a basolateral (sinusoidal) domain. The canalicular membrane is partly regulated at the transcriptional level (Trauner & Boyer, 2003). It has recently become clear that some of these ABC transporters reside in intracellular compartments and are delivered to the canalicular domain following increased physiological demand (Kipp & Arias, 2002). A putative role for the lysosomal compartment in biliary secretion has long been suspected, but currently remains unclear (Gross et al. 1989; Nakano et al. 1995).

Cu excretion from the hepatocyte is achieved in part by mechanisms that meet these general principles, but there are also some important notable differences. The current working model of the cellular machinery mediating Cu acquisition, redistribution and excretion in the hepatocyte is presented in Fig. 2 and described in more detail later (see p. 33).
Copper-transporting P-type ATPase ATP7B is a hepatocyte copper transporter

The transmembrane transport of Cu is mediated not by an ABC transporter but by ATP7B, a member of a highly-conserved family of heavy-metal-transporting P-type ATPases (Bull et al. 1993; Petrukhin et al. 1993; Tanzi et al. 1993; Yamaguchi et al. 1993). ATP7B is expressed predominantly in the liver and is structurally and functionally homologous with ATP7A, the gene mutated in the Cu-deficiency disorder Menkes disease (Fig. 1; Vulpe et al. 1993; Payne & Gitlin, 1998). ATP7B is mutated in Wilson disease, and the hepatic accumulation of Cu observed in patients with this autosomal recessive disorder suggests that ATP7B plays a central role in biliary Cu excretion, and that the biochemical function of ATP7B is in the transport of Cu across cellular membranes (Fig. 1). The ATP7B protein contains six tandemly-repeated heavy-metal-binding regions in its N-terminal domain. ATP7B is required for Cu insertion from the cytosol into the trans Golgi network, and most mutations associated with Wilson disease abolish this biochemical activity (Payne & Gitlin, 1998). Consistent with these observations, but in contrast to most other transporters in the liver cell, ATP7B is localized in the trans Golgi network under normal physiological conditions (Hung et al. 1997; Huster et al. 2003). Here the Cu is incorporated into apocaueruloplasmin, to form the redox-active holoenzyme caeruloplasmin.

In accordance with the observed induced intracellular relocation of hepatocyte transport proteins, ATP7B is also localized in different cellular compartments depending on the Cu status of the cell. When cellular Cu levels are raised this P-type ATPase is rapidly translocated from the trans Golgi network to a vesicular compartment diffusely localized within the cell (Hung et al. 1997). This Cu-dependent relocation is fast and does not require de novo protein synthesis. In addition, this phenomenon is reversible, since subsequent depletion of Cu results in the rapid return of ATP7B to the trans Golgi network. In polarized HepG2 hepatoma cells ATP7B redistributes to vesicular structures and to apical vacuoles reminiscent of bile canaliculi under high-Cu conditions (Roelofsen et al. 2000). Taken together, these data suggest that ATP7B is continuously cycling through different cellular compartments and that Cu availability regulates the efficiency and velocity of specific retrieval steps. The molecular mechanisms of Cu-dependent cycling of ATP7B are largely unknown, but some recent studies have suggested that Cu-dependent signal transduction pathways may be involved (Vanderwerf et al. 2001). It is possible that the unidentified ATP7B-containing vesicles represent some form of endosomal compartment, since excess Cu can be sequestered in lysosomes (Myers et al. 1993). Cu overload causes alterations in lysosomal morphology and increases lysosomal pH (Myers et al. 1993). Hepatocytes respond to a greater Cu load by sequestering excess Cu in more lysosomes that can empty their contents directly into the bile (Gross et al. 1989). These results provide evidence that exocytosis of lysosomal contents into biliary canaliculi is an important mechanism for biliary Cu excretion, at least during hepatic Cu overload (Gross et al. 1989; Myers et al. 1993).

The role of ATOX1 in hepatic copper excretion

An important protein in the Cu excretion pathway mediated by ATP7B is the small cytosolic Cu chaperone ATOX1. ATOX1 is a protein of sixty-eight amino acid residues with a similar structural fold to each of the six metal-binding sites in ATP7B (Klomp et al. 1997; Wernimont et al. 2000). ATOX1 itself can bind Cu and it undergoes a transient Cu-dependent association with ATP7B (Hamza et al. 1999; Larin et al. 1999). This Cu-dependent association between ATP7B and ATOX1 seems essential for efficient Cu transport, since mutations in ATP7B that abolish this interaction are associated with Wilson disease (Hamza et al. 1999). Consistent with this finding, mice with a deletion of the murine Atox1 locus have severe abnormalities in cellular Cu metabolism (Hamza et al. 2001). Based on work in yeast, it has been postulated that ATOX1 binds and detoxifies cytosolic Cu and presents the metal to ATP7B for subsequent transmembrane transport (Pufahl et al. 1997). A second hypothesis for the biological function of ATOX1 in the cell has been put forward by Lutsenko and coworkers (Walker et al. 2002), who showed that ATOX1 stimulates the catalytic activity of ATP7B. Most recently, elegant experiments by Hamza et al. (2003) have revealed that Cu-dependent subcellular trafficking of ATP7A is markedly impaired in cells isolated from Atox1–/– mice. They proposed that ATOX1 is essential for the regulation of Cu-dependent trafficking of ATP7A, and presumably also that of ATP7B. Future research should be aimed at the exciting possibility of unifying these three hypotheses into a single coherent working model for the molecular role of ATOX1 in the regulation of ATP7B-mediated cellular Cu excretion.

MURR1, a new copper-metabolism gene?

Solving the aetiology of disorders of Cu homeostasis other than Menkes disease and Wilson disease, both in man and animals, may further increase our knowledge of the regulation of Cu homeostasis. Some years ago the authors became interested in a form of Cu toxicosis in dogs that has long been considered an excellent model for studying hepatic Cu overload and developing therapeutic approaches in man. Canine Cu toxicosis (CT) is an autosomal recessive disorder with a high frequency in Bedlington terriers (Hardy et al. 1975). Dietary intake of Cu in these animals is normal, but biliary Cu excretion is markedly reduced compared with control animals (Fig. 1; Su et al. 1982). Animals that are affected have Cu concentrations of >1000 μg/g liver dry weight. The excess Cu is stored in the lysosomes of hepatocytes, resulting in liver cirrhosis, and unless treated the dogs die at between 2 and 6 years of age. The Cu-chelation therapy used in Wilson disease is effective in Bedlington terriers. The disorder has a clear idiopathic component, since reduced penetrance of CT has also been put forward by Lutsenko and coworkers (Walker et al. 2002), who showed that ATOX1 stimulates the catalytic activity of ATP7B. Most recently, elegant experiments by Hamza et al. (2003) have revealed that Cu-dependent subcellular trafficking of ATP7A is markedly impaired in cells isolated from Atox1–/– mice. They proposed that ATOX1 is essential for the regulation of Cu-dependent trafficking of ATP7A, and presumably also that of ATP7B. Future research should be aimed at the exciting possibility of unifying these three hypotheses into a single coherent working model for the molecular role of ATOX1 in the regulation of ATP7B-mediated cellular Cu excretion.
of CT and those of the classic Wilson disease: the absence of neurological manifestations; the normal caeruloplasmin concentrations in plasma. Cu toxicosis resembles Indian childhood cirrhosis (Tanner et al. 1979) and endemic Tyrolean infantile cirrhosis (Müller et al. 1996), two fatal human hepatic Cu overload disorders of unknown etiology (Fig. 1). A breakthrough in the molecular characterization of CT was the localization of the gene defect to canine chromosome 10, region q26 (van de Sluis et al. 1999). Based on the map location of the CT locus, mutations in the canine ATP7B and ATOX1 genes could be excluded as the cause of the disorder because these genes mapped to canine chromosomes 22q11 and 4q24–q31 respectively (Dagenais et al. 1999; Nanji & Cox, 1999; van de Sluis et al. 2001).

The availability of large Bedlington terrier pedigrees segregating CT has made it possible to embark on a positional cloning strategy to identify the gene mutated in CT (an outline of positional cloning is depicted in Fig. 3; van de Sluis et al. 1999, 2000).

More recently, it was shown that this disorder is associated with a deletion of exon 2 of the MURR1 gene, at the cDNA level (van de Sluis et al. 2002). At the genomic level the mutation in Bedlington terriers spans at least 13 kb and encompasses exon 2 of the MURR1 gene. Hence, this rearrangement is predicted to cause an in-frame deletion of the MURR1 transcript, presumably leading to the formation of a shortened MURR1 protein product of ninety-four amino acid residues instead of the predicted 188 residues (Fig. 3). It was predicted that this short protein product would be unstable and therefore lead to a complete loss of MURR1 function. Consistent with this hypothesis, it has been established experimentally that the deletion is indeed associated with a complete absence of detectable MURR1 protein from the livers of affected Bedlington terriers, whereas the protein is expressed as a 23kDa single-chain polypeptide in the livers of unaffected dogs (Klomp et al. 2003b). Importantly, mutations in MURR1 are not associated with Indian childhood cirrhosis and endemic Tyrolean infantile cirrhosis in man (Müller et al. 2003). Taken together, these data strongly suggest that MURR1 plays a rate-limiting role in Cu excretion from hepatocytes during Cu overload, at least in the dog. Experiments using Murr1 knockout mice and knock-down of MURR1 by RNA interference are underway to extend these observations to other species. Another exciting possibility includes the structural characterization of MURR1. Much of the current knowledge of the biochemistry of intracellular Cu homeostasis has come from x-ray crystallography and NMR structural analysis of proteins involved in Cu metabolism. It has now become essential to determine the crystal structure of MURR1.

Clearly, MURR1 deficiency provides a unique opportunity to study the cellular biology of hepatic Cu excretion. MURR1 encodes a protein that is currently of unknown function and has no detectable sequence homology to known proteins. MURR1 mRNA is expressed abundantly in liver tissue, but is also readily detected in other tissues. Using a specific polyclonal antibody raised against MURR1, ubiquitous Murr1 protein expression was recently identified in murine organs and human cell lines. From these data it has been postulated that MURR1 could perform a pleiotropic function in different organs, not exclusively linked to Cu homeostasis. Several lines of indirect evidence suggest that MURR1 may function in the ATOX1–ATP7B-mediated pathway of Cu excretion, probably downstream of ATP7B. MURR1-deficient Bedlington terriers have normal plasma holocaeruloplasmin activity, but a markedly elevated lysosomal Cu content and a strongly reduced biliary Cu excretion. This finding suggests that Cu transport by ATP7B into the Golgi apparatus is normal, and indicates that MURR1 deficiency does not lead to a complete inhibition of the Cu-transporting activity of ATP7B. Rather, MURR1 deficiency may lead to an inefficient degranulation of lysosomal contents into bile during Cu overload (Fig. 2). Consistent with the lysosomal Cu accumulation in affected Bedlington terriers, it was observed that exogenous and endogenous MURR1 is partially localized to a vesicular compartment, reminiscent of the endosomal–lysosomal system, in several cell lines (Klomp et al. 2003b). These preliminary findings will have to be further corroborated both in cultured cells and in liver tissue.

Is there a link between copper import and export?

Cu import in yeast is mediated by the homologous Cu transporters Ctrl1–3 (Puig et al. 2002). The human orthologous genes, hCTR1 and hCTR2, show limited sequence homology with the yeast Ctr1 gene (Zhou & Gitschier, 1997). The hCTR1 gene encodes a protein of 190 amino acids, which is glycosylated, forms oligomers and is ubiquitously expressed (Klomp et al. 2002, 2003a; Eisses & Kaplan, 2002). CTR1 proteins span the membrane multiple times, permitting formation of a channel, which is consistent with its proposed role as a high-affinity Cu permease (Puig et al. 2002; Eisses & Kaplan, 2002; Klomp et al. 2003a). The subcellular localization of CTR1 differs markedly between different cell types, ranging from a predominant intracellular vesicular perinuclear compartment to a location on the plasma membrane. The subpool of CTR1 present on the plasma membrane undergoes rapid internalization and degradation when the extracellular Cu concentration is elevated (Petris et al. 2003). It is tempting to speculate that the Cu-dependent internalization of CTR1 is regulated by mechanisms similar to those for the metal-dependent redistribution of ATP7A and ATP7B, thereby providing a mechanistic link between the regulation of Cu import and export. Homozygous null mice at the Ctrl1 locus exhibit profound delays in growth and development and die at embryonic day 7.5, presumably as a consequence of Cu deficiency (Kuo et al. 2001; Lee et al. 2001). These studies demonstrated a crucial rate-limiting role for CTR1 in Cu acquisition and embryonic development in mammals and support the hypothesis that cellular Cu homeostasis is regulated at the level of both Cu import and Cu excretion. It would be interesting to determine the exact role of CTR1 in dietary Cu uptake and investigate whether putative mild variations in CTR1 structure and/or regulation of plasma membrane expression are associated with differences in acquisition of Cu in the population.
Molecular regulation of trace element metabolism

Genetic markers, STS, ETS

Database search BAC

Radiation hybrid mapping

Physical map, transcript map

BAC Marker FH2523 CF10B18 CF10B17 C04107

CT-region (±500 kb)

Sequence analysis MURR1 in affected CT dogs

RNA level

Protein level

Normal CT

469 bp; expected length

187 bp; deletion fragment

CT-Affected dog Normal dog

188 amino acid protein; ubiquitously expressed

Ninety-four amino acid protein; loss of function mutation

Fig. 3. Schematic overview of positional cloning strategy employed in cloning the MURR1 gene. Genetic mapping was used to map the disease gene onto the dog genome (Yuzbasiyan-Gurkan et al. 1997). The map location was determined by comparative mapping (van de Sluis et al. 1999). Radiation hybrid mapping was used to construct a physical map (van de Sluis et al. 2000). A bacterial artificial chromosome (BAC) contig was constructed and new polymorphic markers were isolated from the BAC clones to narrow down the canine copper toxicosis (CT) gene region further by genetic mapping. Haplotype sharing revealed a region of approximately 500 kb to be shared by all affected animals. This region contained seventeen putative transcripts, four of which were subjected to mutation analysis by sequencing. The MURR1 gene was mutated in affected Bedlington terriers, which was associated with a short RNA product (van de Sluis et al. 2002). The mutation results in a loss of function, since no protein can be detected in the livers of affected dogs (Klomp et al. 2003).

STS, sequence tagged site; ETS, expression tagged site.
**Functional characterization of MURR1**

The current challenge is to enhance the understanding of MURR1 function in relation to hepatic Cu homeostasis. In addition to the cell biology approach to the dissection of MURR1 function described earlier, the availability of whole genome sequences and high-throughput genomic and proteomic technologies will offer new opportunities to address MURR1 function and to identify the cellular and molecular pathways in which MURR1 acts. Such a multidisciplinary approach will not only allow us to investigate phenotype-related processes, but will also provide tools to unravel the molecular aspects of the normal adaptive response of cells, tissues and organisms to dietary Cu (Fig. 4).

**Comparative genomics**

The complete genome sequences of more than eighty-five bacteria, five different yeast strains and eight eukaryotes have been determined. By the end of 2003 whole genome sequences of many additional eukaryotes will be available. Genome analysis and sequence comparisons will contribute to our understanding of gene structure, gene function and gene evolution.

Comparing the gene sequence of MURR1 with the fully-sequenced genomes available to date showed that orthologues of MURR1 are present in a number of vertebrates, including man, but they could not be detected in lower eukaryotes and prokaryotes (van de Sluis et al. 2002). This finding is in contrast to the results for other proteins of Cu metabolism, which are clearly well conserved in lower eukaryotes. The presence of the MURR1 proteins may be related to the presence of bile and its participation in the excretion of toxic compounds, which is a unique feature of vertebrate organisms. Much of our current insight into the homeostatic control of Cu metabolism is in fact based on studies of Cu homeostasis in yeast (Askwith & Kaplan, 1998; De Freitas et al. 2003).

Comparative genomics can also be a powerful tool for determining which parts of a gene harbour important functions, as these parts are usually subject to strong selection and are, therefore, expected to show the highest level of sequence conservation. Furthermore, a comparison of the non-coding regions surrounding a gene may also identify highly-conserved regions, which generally indicate important regulatory elements such as promoters and enhancers (Muller et al. 2002).

**Gene expression profiling**

Microarrays are being used to measure the levels of expression of thousands of genes simultaneously. Commercial microarrays (‘chips’) with a large number of genes are currently available for different species. There are a number of different technologies and platforms on the market, but the general principles underlying each of them are similar. In brief, equal amounts of two fluorescently-labelled RNA preparations are simultaneously hybridized onto a chip. Transcripts with differential expression between the two samples will show only a signal from one fluorescent dye (Duggan et al. 1999).

![Diagram](https://www.cambridge.org/core)

**Fig. 4. Overview of the multidisciplinary approach taken to understand MURR1 gene function and to identify the molecular pathways in which MURR1 is involved. A gene-driven approach using genomic information for identifying genes (DNA and RNA) and gene products (protein) involved in certain cellular processes can be performed. This approach will also be extremely useful in nutrigenomics research, when studying the adaptive response to copper in relation to genetic background. Alternatively, a phenotype-driven approach analyses phenotypes from human pathologies or animal mutants. This approach will initially unravel the primary genes involved in the disease process and subsequently the molecular pathway(s) in which such a gene operates.**

This technology is potentially an extremely powerful tool for assigning gene function and deciphering regulatory networks. To identify the possible cellular and molecular mechanisms in which MURR1 plays an essential role, cells with a mutant MURR1 can be profiled under different Cu conditions. Mutant MURR1 cells may be obtained from *in vivo* models (e.g. affected dogs or knock-out mice), or from *in vitro* models (e.g. knock-out or knock-down cells). The current disadvantage of the natural MURR1 mutant, the Bedlington terrier, is the lack of canine-specific microarrays.

Microarray studies can also be a powerful tool for providing information on the normal adaptive cellular
response to Cu exposure and the regulatory mechanism involved. A recent study in yeast grown under changing Cu conditions (Gross et al. 2000) indicated the presence of a limited set of yeast genes that are differentially expressed under Cu-excess or Cu-replete conditions. A preliminary time-course experiment (H van Bakel and C Wijmenga, unpublished results), in which both the acute and long-term responses in either Cu-excess or Cu-starvation conditions were measured in yeast, revealed seven new genes that had not previously been implicated in Cu metabolism. Although most genes involved in Cu transport in yeast have human homologues, the situation is not the same for the Cu-induced transcription factors Ace1 and Mac1. It has been suggested that in vertebrates Cu itself is not involved in regulating the expression of genes (Linder, 2001). However, a pilot experiment in human cultured fibroblasts showed that increasing Cu levels promotes a rapidly-induced expression of a number of genes, including metallothioneins (P Muller and C Wijmenga, unpublished results). These results have two major implications. First, they suggest that Cu-dependent regulation of transcription does indeed exist in mammalian cells. Second, they indicate that in addition to the regulation of Cu import and Cu export from the cells Cu may also directly regulate intracellular sequestration of the metal. It is not yet clear whether Cu is associated with transcription factors that bind to promoters of Cu-regulated genes. The identification of Cu-responsive genes will allow their promoter regions to be analysed and will promote the search for common elements that may be responsible for Cu-mediated regulation of expression.

More recently, it has also become evident that microarray technology may become an important tool in toxicology (Hamadeh et al. 2002; Waring & Halbert, 2002; Vrana et al. 2003). In the future it should be possible to compare the patterns of gene expression associated with novel chemicals with databases that contain patterns of expression for known chemicals and toxic compounds, including transition metals. Moreover, microarray studies could also be instrumental in determining the toxic dose of Cu from dose–response relationships. For the latter it is evident that there should be some knowledge of the normal adaptive response to Cu in individual organs or cell types.

Proteomics

Recently, it was shown that the Cu chaperones exhibit a specific transient protein–protein interaction with their respective target proteins. These interactions are Cu dependent and necessary for efficient Cu transfer from the chaperone to the target proteins (Pufahl et al. 1997). These data support the existence of a protein–protein interaction network in which Cu is transferred from one protein to another depending on the requirement to synthesize copper-enzymes in different organelles, or alternatively to store or excrete the metal. The presence of this network provides an opportunity to identify novel proteins that play a role in Cu homeostasis simply by isolating ‘new’ proteins that interact with known proteins. During the last decade, novel high-throughput methods have been developed and successfully applied to the identification of protein–protein interaction networks. A comprehensive analysis of protein–protein interactions was carried out recently in the yeast S. cerevisiae (Ito et al. 2000; Uetz et al. 2000). The yeast–two hybrid method has been applied to the identification of MURR1 interacting proteins. In preliminary experiments a previously unknown protein that specifically interacts with MURR1 was isolated and tentatively named MURR1-interacting protein 1 (P de Bie, LW Klomp and C Wijmenga, unpublished results). The interaction of MURR1 with MURR1-interacting protein 1 has been confirmed by independent techniques. Although the function of MURR1-interacting protein 1 and the relevance of its interaction with MURR1 need to be further established, these preliminary data illustrate the relevance of this approach in characterizing functional pathways in Cu homeostasis, which may also lead to the identification of novel candidate genes for human Cu-overload disorders, including Indian childhood cirrhosis and endemic Tyrolean infantile cirrhosis.

Model systems

Yeast has been used widely as a model organism in the study of Cu metabolism (Askwth & Kaplan, 1998; De Freitas et al. 2003). Although the yeast system shows strong homology in Cu uptake and cellular distribution to the systems in higher vertebrates, including man, the excretion process is rather different. A mouse model with a MURR1 mutation similar to that seen in Bedlington terriers is currently being constructed by homologous recombination in mouse embryonic stem cells. These mice will serve different purposes. They will confirm the disease-causing phenotype, they will help in defining the MURR1 gene function and, finally, they may become useful for the study of gene–diet interactions. These mice will be subjected to the techniques described earlier; microarray analysis, for example, is expected to reveal multiple genes that act downstream of MURR1. It could become rather time-consuming and expensive to produce knock-out or transgenic mice for a number of these downstream targets in order to functionally characterize them. The availability of RNA interference has introduced a powerful tool for studying gene function in mammals by inducing post-transcriptional gene silencing, both in vitro and in vivo (Brummelkamp et al. 2002).

Future perspectives

It is becoming evident that most traits and disorders have a genetic basis. It is therefore tempting to speculate that an individual’s susceptibility to dietary Cu may also have a genetic background. Individual variation in genes involved in either the uptake or excretion of Cu might render an individual or animal more susceptible or resistant to Cu. The cases of Indian childhood cirrhosis in India and endemic Tyrolean infantile cirrhosis are clear examples. However, both the lack of appropriate molecular markers for the assessment of Cu-related genotypes, and the lack of biomarkers for the assessment of intracellular Cu levels have hampered studies carefully designed to address this problem. The new research field of nutrigenomics is the
application of high-throughput genomic tools to nutrition research (Müller & Kersten, 2003). This approach will increase our understanding of how nutrition influences metabolic pathways and homeostatic control, and how this regulation is disturbed under changing dietary conditions or diet-induced diseases in relation to the individual’s background.

Currently, ATP7B is the only Cu toxicity-related gene known to be present in man. Given that the frequency of Wilson disease is one in 30 000, approximately one in every eighty-seven individuals must carry at least one ATP7B mutation. It would be interesting to investigate whether there is a potential risk associated with being an ATP7B mutation carrier when exposed to conditions of high levels of exogenous Cu, e.g. through drinking water contaminated by domestic Cu piping. Large population-based studies will be required to establish ‘Cu-related cellular phenotypes’ by the application of a range of genomic, proteomic and metabolomic tools (Bochner, 2003; Freimer & Sabatti, 2003).

It is expected that the work on MURR1 and its pathway will reveal a group of genes that are directly or indirectly involved in modulating hepatocellular Cu levels. New technological advances will make it possible to compile extensive genetic profiles using single nucleotide polymorphisms that capture the genetic variation of these genes.

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References


