Dietary copper is an essential trace element with roles in both functional and structural aspects of the cardiovascular system. In particular, the vascular response to inflammatory stimuli is known to be significantly augmented in copper-deficient rats. The current study was designed to quantify the extent of injury-induced neointimal proliferation and stenosis in rats fed diets either adequate or deficient in copper. Male, weanling Sprague-Dawley rats were fed purified diets that were either adequate (CuA; 5.6 µg Cu/g) or deficient (CuD; 0.3 µg Cu/g) in copper for 4 weeks. Balloon injury was induced in the left external carotid arteries. Fourteen days after injury, histomorphometric analysis of cross-sections from carotid arteries showed increased neointimal formation in the CuD group compared with the CuA controls (neointima/media ratio: 4.55 ± 0.93 vs 1.45 ± 0.2, respectively). These results correspond with data indicating that the activity of Cu/Zn-superoxide dismutase (SOD) is depressed in rats fed this CuD diet. Because superoxide anion and redox status are known to play a key role in the extent of neointimal formation in response to injury, we propose that the exaggerated neointimal proliferation seen in the CuD group is the result of the diminished Cu/Zn-SOD activity.

Materials and Methods

Animals and Diet. This project was approved by the University of Louisville Animal Care and Use Committee. Twelve male weanling Sprague-Dawley rats were purchased from Charles River Breeding Laboratories (Wilmington, MA). On arrival, rats were housed individually in stainless steel cages in a temperature- and humidity-controlled room with a 12:12-hr light:dark cycle. The rats were given free access to distilled water and to one of two purified diets for 4 weeks. The basal diet (16) was a casein-sucrose-cornstarch-based diet (no. TD 84469; Teklad Test Diets, Madison, WI) containing all known essential vitamins and minerals except for copper and iron. The copper-adequate (CuA) diet consisted of the basal diet (940 g/kg of total diet) with safflower oil (50 g/kg) and a copper-iron mineral mix that provided 0.22 g of ferric citrate (16% Fe) and 24 mg of CuSO4·5H2O per kilogram of diet. The copper-deficient (CuD) diet was the same except for replacement of copper with cornstarch in the mineral mix. Diet analysis by atomic absorption spectrophotometry (model 503; Perkin Elmer, Norwalk, CT) indicated that the CuA diet contained 5.6 mg copper/kg diet and the CuD diet contained 0.33 mg copper/kg diet. Parallel assays of National Institute of Standards and Technology (NIST; Gaithersburg, MD) reference samples (citrus leaves, no. 1572) yielded values within the specified range, which validated our copper assays.
Arterial Injury Model. Animals were anesthetized by intraperitoneal injection of a ketamine (37.5 mg/kg) and xylazine (5 mg/kg) mixture. The level of anesthesia was verified before and during surgery by evaluating vibrissa movement, tail tonus, and breathing rate in response to handling. If the animal did not attain an adequate level of anesthesia in the first 7 min following administration, it was returned to the breeding colony for use in later experiments. This protocol ensured that animals remained under complete anesthesia for at least 40 min.

The left common and external carotid arteries were exposed and a balloon catheter (PE50) was passed through the external carotid artery and advanced into the aorta, inflated with saline, then pulled back to distend the common carotid artery. This procedure was repeated twice, after which the external carotid artery was ligated, and the incision was sutured.

Histomorphometric Analysis. Fourteen days after balloon injury, the rats were anesthetized and transcardially perfused with phosphate-buffered saline (PBS) followed by 4% paraformaldehyde at a fixed perfusion pressure of 100 mmHg. The right and left common carotid arteries were removed, and the adipose and connective tissue in the adventitia were excised. The vessels were then divided into three equal segments and placed for 24 hr in 4% paraformaldehyde for further fixation.

The arterial segments were embedded in paraffin, cut on a rotary microtome (5 μm), and stained with hematoxylin and eosin (H&E). Cross-sectional rings stained with H&E were used to determine the extent of neointima formation. Analysis was performed using an Axiosvert microscope (Zeiss, Jena, Germany) attached to an imaging system analyzer Image-1/Metamorph (Universal Imaging Corp., Westchester, PA) at ×100 magnification. Microscopic images of cross-sectional rings were digitally transferred into a computerized image analyzer and calibrated for area measurement. Two independent operators using a double-blind protocol performed the area measurements by contouring the luminal, luminal plus neointimal, and total arterial area between uninjured carotid arteries from CuA and CuD groups. Luminal and medial area values of uninjured arteries were (mean ± SEM, in mm²): CuA, 8.81 ± 0.77; CuD, 7.66 ± 0.49 for lumen, and CuA, 0.74 ± 0.32; CuD, 0.98 ± 0.27 for media. In contrast, the histomorphometric analysis of balloon-injured carotid arteries from CuD animals revealed a dramatic increase in neointima formation when compared with CuA animals (Table I), resulting in most cases in almost complete obstruction of the arterial lumen (Fig. 1).

Cross-sectional segments from injured carotid arteries were further analyzed under higher magnification ×400. The CuA group had concentric intimal hyperplasia consisting of totipotential mesenchymal cells and an intact intimal surface (Fig. 1). The CuD group had lumen occluding intimal proliferation with recanalization. There was also early collagen deposition with significant loss of inner medial fibers that were apparently incorporated into the neointima formation of injured CuD animals (Fig. 1). Figure 1 also demonstrates dramatic retraction of the medial wall in the CuD group.

Discussion

Smooth muscle cell proliferation is a common event featured in several cardiovascular diseases. From the development of the atheromatous plaque to accelerated restenosis

<table>
<thead>
<tr>
<th>Group</th>
<th>n</th>
<th>Body Weight (g)</th>
<th>Liver Cu (mg/g dry wt)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Arrival Before injury</td>
<td>After injury</td>
</tr>
<tr>
<td>CuA</td>
<td>6</td>
<td>35 ± 5</td>
<td>221 ± 31</td>
</tr>
<tr>
<td>CuD</td>
<td>6</td>
<td>35 ± 5</td>
<td>210 ± 17</td>
</tr>
</tbody>
</table>

Values are mean ± SEM.  
* P < 0.05 by Student’s t test.
after angioplasty, smooth muscle cell plasticity has been associated with some interesting functions. It is able to regress into its undifferentiated embryological form and from there become a highly specialized collagen producer (21). Evidence of this change in the vascular smooth muscle is clearly seen in the vascular response of the rat carotid artery to disruption of the endothelium by balloon injury (Fig. 1). In particular, the response is accentuated in the CuD group where both the intimal hyperplasia consisting of the totipotential mesenchymal cells and the luminal deposition of collagen are significantly greater in the CuD group only 2 weeks after vascular injury (Fig. 1).

The most likely mechanism for this exaggerated stenosis in the CuD group is related to the redox imbalance, which occurs during the repair reaction and which has been shown to be a major contributor to restenosis following angioplasty (22–26). Azevedo et al. (26) have reported that the timing of the oxidative stress corresponds with the early inflammatory and proliferative phase of vascular repair.

There are several copper-dependent antioxidant enzymes in the cardiovascular system (for review, see Ref. 27). Of these, the activity of Cu/Zn-SOD is implicated by studies showing a role for superoxide in the early vascular smooth muscle mitogenesis associated with restenosis (28, 29). We have previously shown that Cu/Zn-SOD activity is significantly reduced in the wall of the aorta from rats fed the CuD diet used in the current study (30). Further, we have reported that the concentration of plasma peroxynitrite is increased in CuD rats, indicating an increased interaction between superoxide anion and nitric oxide in vivo (4). Together, these results demonstrate a decrease of Cu/Zn-SOD activity and consequent increase in superoxide in CuD rats.

**Table II. Histomorphometric Analysis of Injured Common Carotid Arteries**

<table>
<thead>
<tr>
<th>Group</th>
<th>n</th>
<th>Lumen (mm²)</th>
<th>Media (mm²)</th>
<th>Neointima (mm²)</th>
<th>Total area (mm²)</th>
<th>Neo/Media</th>
</tr>
</thead>
<tbody>
<tr>
<td>CuA</td>
<td>6</td>
<td>1.62 ± 0.14</td>
<td>1.18 ± 0.06</td>
<td>1.62 ± 0.18</td>
<td>3.98 ± 0.16</td>
<td>1.45 ± 0.22</td>
</tr>
<tr>
<td>CuD</td>
<td>6</td>
<td>0.99 ± 0.30ᵃ</td>
<td>0.44 ± 0.07ᵃ</td>
<td>2.03 ± 0.35</td>
<td>2.91 ± 0.29ᵃ</td>
<td>4.55 ± 0.93ᵃ</td>
</tr>
</tbody>
</table>

Values are mean ± SEM. Lumen, luminal area; Media, medial area; Total Area, total arterial area; Neo/Media, ratio of neointima area/media area; n, number of animals.

ᵃ P < 0.05 by Student’s t test.

Figure 1. Photomicrographs of representative cross-sections of uninjured and balloon injured carotid arteries from rats fed CuA or CuD diets. (A) Uninjured CuA. (B) Uninjured CuD. (C and E) Injured CuA at ×25 and ×400. (D and F) Injured CuD at ×25 and ×400. M, media; NI, neointima.
Therefore, the exaggerated degree of restenosis seen in the CuD rats (Fig. 1) correlates with the inactivation of Cu/Zn-SOD by dietary copper restriction. This conclusion follows the assertion of Azevedo et al. (26) that oxidative stress may be a “dose-dependent” sensor of the degree of injury.

Generation of the reactive oxygen species following vascular injury occurs predominantly in the vascular smooth muscle cells (VSMC) or adventitial fibroblasts rather than in the endothelium (31, 32). Leukocyte-induced oxidative stress is also a component of the early inflammatory response to the balloon-induced injury (33). However, the respiratory burst is significantly diminished in CuD neutrophils (34). Therefore, VSMC or fibroblasts are probably the cell type involved with the balloon injury-induced oxidative stress in the CuD group.

A second possible mechanism for the augmented response seen in the CuD group involves the copper-dependent enzyme, lysyl oxidase. Lysyl oxidase is an amine oxidase in the vascular wall that initiates the covalent crosslinking of collagen and elastin. The enzyme is also potently chemotactic for VSMCs and is critical for repair of injuries in the cardiovascular system (35). However, this lysyl oxidase-dependent chemotaxis is abolished by inhibition of the enzyme’s active site (35). Dietary copper deficiency is also known to inhibit lysyl oxidase activity in the rat (36). Therefore, it seems unlikely that lysyl oxidase-mediated chemotaxis of VSMCs is involved in the augmented neointimal proliferation and stenosis seen in the CuD group of the current study.

In conclusion, neointimal formation and stenosis are exaggerated following balloon-induced injury in carotid arteries of CuD rats. The likely mechanism is an increased superoxide-induced VSMC proliferation in the presence of reduced Cu/Zn-SOD activity. The current study, combined with previous data demonstrating significantly larger atherosclerotic lesions in aortas of CuD and copper-marginal rats (37), suggest an important role for copper nutrition in the vascular response to injury or atherosclerosis.

The authors acknowledge the technical assistance of Sharon Young, Gwen Dahlen, and Pete Leary.

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