Manganese and Cobalt Concentrations in Hair and Nail of Some Kano Inhabitants

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ABSTRACT: Manganese and cobalt concentrations in hair and nail were determined by Flame Atomic Absorption Spectrophotometer (AAS). The mean manganese in hair and nail were $0.54 \pm 0.35\text{mg/g}$ and $0.68 \pm 0.30\text{mg/g}$ while the mean cobalt in hair and nail were $0.30 \pm 0.24\text{mg/g}$ and $0.46 \pm 0.37\text{mg/g}$ respectively. A progressive increase in cobalt concentrations in hair and nails with age indicated no significant difference in their means suggesting that cobalt in hair and nails originate from a common source. Comparing the mean manganese concentrations in hair with the nails a significant difference is indicated in the two tissues $(p < 0.05)$. Human hair and nails are hence recording filaments that can reflect metabolic changes of many elements over long periods of time and hence furnish a post nutritional event of some essential micro-elements.

Key words: Manganese, Cobalt, Hair, Nail, Determination, Pollution

INTRODUCTION

Hair is a site of excretion for essential, nonessential and potentially toxic elements (Ashraf et al., 1995). The amount of an element irreversibly incorporated into growing hair is proportional to the level of the element in other body tissues (Ajayi et al., 2001). Therefore, hair analysis provides an indirect screening test for physiological excess, deficiency or mal-distribution of elements in the body (Adeniyi and Aletor, 1999). Clinical researches indicated that hair levels of specific elements, particularly potentially toxic elements such as cadmium, mercury, lead and arsenic, are highly correlated with pathological disorders (Bache et al., 1991). For such elements, levels in hair may be more indicative of body stores than the levels in blood and urine. Scalp hair is vulnerable to external contamination by water, hair treatments and products (Kaspr et al., 1982; Sturaro et al., 1994; Bertazzo et al., 1996; Rosbong et al., 2003). The data that hair analysis does provide should be considered in conjunction with symptoms, diet analysis, occupation and lifestyle, physical examination and the results of other laboratory tests. Hair analysis may provide useful insights into the biochemical and hormonal condition of the body (DiPietro et al., 1989; Chattopadhyay et al., 1990; Contiera and Folin, 1994; Nnorom et al., 2005).

Trace metals are important, from nutritional and toxicological points of view. Small amounts of these essential elements are required for the maintenance of growth and normal health. Deficiencies occur when there are inadequate amounts to meet the metabolic needs. Clinical toxicity also occurs if over ingestion of some elements takes place (Baumgatner, 1993). It is present in all healthy tissues, its concentration from one species to the next is fairly constant, the amount of each element is maintained within its required limit, its withdrawal induces reproducibly the same physiological and/or structural abnormalities and their additions to the diet prevents or reverse the abnormalities (Tuormaa, 1995). Non – essential or toxic trace elements include, excess copper, lead, cadmium, mercury and aluminum acquired through environmental contamination (Kaspereck et al., 1982; Tuormaa, 1995; Rosbong et al., 2003).
The milk, urine, saliva and sweat measure the component that is absorbed but excreted. The blood measures the component absorbed and temporarily in circulation before excretion and/or storage (EPA, 1980). The hair, nails and teeth are tissues in which trace minerals are sequestered and/or stored and can be used to monitor the highest priority toxic trace metals (Barett, 1985). Manganese is an essential trace mineral in human nutrition (Keen et al., 1999). Manganese is the preferred metal co-factor for glycosyltransferases in the synthesis of glycoproteins and glycosaminoglycans (mucopolysaccharides). Manganese and iron scavenge for hydroxyl and superoxide radicals a component of the metalloenzyme manganese superoxide dismutase (MnSOD) found in the mitochondria and a constituent of the mitochondrial oxidant defense system. Symptoms associated with Mn deficiency include fatigue, lack of physical endurance, slow growth of fingernails and hair, impaired metabolism of bone and cartilage, dermatitis, weight loss, reduced fertility, increased allergic sensitivities and inflammation. Other deficiency signs include nausea, vomiting, change in hair color and neurologic sequel. Patients with end-stage liver disease accumulate manganese in their basal ganglia. Manganese plays a role in the hepatic encephalopathy in liver failure. It is eliminated primarily through the bile, and hepatic dysfunction leading to depressed manganese excretion (Krieger et al., 1995). Mine workers exposed to high concentrations of manganese dust develop “locuramanganica” or manganese madness with symptoms similar to Parkinson’s disease (Nagatomo et al., 1999). Cobalt is an essential nutrient part of vitamin B12. Excess exposure results in medium toxicity. Inhalation of high levels has effects on the lungs including asthma, pneumonia and wheezing (Petering, 1973). Realizing that both toxic metal excesses and trace mineral deficiencies are associated with all forms of reproductive failures, it has been advocated for the need of hair tissue analysis before conception (Barnes and Bradley, 1994; Bradley and Bennett, 1995). The aim of this work was to determine the levels of manganese and cobalt in the population group living in an urbanized and industrialized city of Kano-Nigeria. The dependence of the hair and fingernail contents on age and gender were examined.

MATERIALS & METHODS

Both hair and fingernails were sampled in the year 2004-2007 from (350) for hair and (300) for fingernails subjects living in urban population group in Kano-Nigeria. Samples were collected from subjects in the age range of 1-55years. Nail samples were collected in polyethylene containers. Hair samples were cut at the root of the occipital area of each subject. Surface contamination and grease were removed by washing the samples in detergent and distilled water after which they were kept in an alcohol–ether mixture for 45mins and dried at 60°C for 72hr (Nowak, 1998).

About 0.50g of each sample was digested in 10cm³ concentrated HNO₃ and the resulting solution was evaporated to dryness and redissolved in 0.1M nitric acid (Mehra and Juneja, 2005; Erzen and Kragelj, 2003; Seidel et al., 2001; Hinwood et al., 2003). Trace metal concentrations were determined by Flame Atomic Absorption on a Buck Model 210 VGP Spectrophotometer attached to IBM personal computer. The result of the absorbance of each sample was the average of ten sequential readings. Background light absorption and scattering were compensated for either by deuterium hollow cathode lamp or by tungsten/halogen lamp. Distilled water was digested as blank using the same procedure previously described (Ayodele and Abubakar, 1998; Ayodele and Abubakar, 2001). All statistical computations either were on the PC 486 66MHZ microcomputer using the integrated statistical package for windows from Umstat Ltd. (London) or dedicated micro instructions for the Excel spread sheets from Microsoft. The approach enabled the advantages of the various computational and graphical facilities of both types of software’s to be used with the ability to read different file formats. The analyses of variance (ANOVA) were carried out according to described procedures (O’Mahony, 1986).

RESULTS & DISCUSSION

The concentration of Mn and Co in hair and nails varied among individuals, thus large number of samples from a population was analyzed and the results treated statistically for meaningful that
correlation. The trace metal concentrations in hair and nails were determined using atomic absorption spectroscopic method. The age of the donors, sex and occupation were noted where necessary. The frequency distribution patterns for the elements in hair and nails vary widely among individuals, thus large number of samples from a population were treated statistically for meaningful correction. The Mn and Co metals in these samples, their mean and coefficient of variation are employed in assessing their levels. The frequency distribution pattern for the age of donors (years) is as shown in Fig. 1. The distribution is multimodal and is skewed towards high frequency of low age with a mean age of 27.51 ± 16.50 years. The frequency distribution pattern for manganese in hair is shown in Fig. 2. The distribution is multimodal and is skewed towards high frequency of low concentration with a mean and standard deviation of 0.54 ± 0.35mg/g while the frequency distribution pattern for manganese in nails (Fig. 3) is multimodal and is skewed towards high frequency of low concentration with a mean and standard deviation of 0.68 ± 0.30µg/g. Though the patterns are similar in both hair and nails, Pearson comparison has shown no correlation between the manganese content in hair and nails (p<0.05) (Table 1). Similarly, the analysis of variance (ANOVA) revealed that the mean concentration of manganese in hair is not significantly different from in the nails at p>0.05 (Table 2). The levels obtained in this study are in agreement with mean intake of manganese worldwide which ranges from 0.52 to 10.8mg (Ferguson et al., 1983; Foo et al., 1993). Manganese concentration in hair and nails with respect to age is as shown in Fig. 4. Manganese levels in both hair and nails decreased with age, but the decrease was pronounced in hair, indicating that manganese may be playing some physiological functions. The frequency distribution pattern for cobalt in hair is as shown in Fig. 5. The distribution is bimodal and is skewed towards high frequency of low concentration with a mean and standard deviation of 0.30 ± 0.24mg/g. The frequency distribution pattern for cobalt in nails is as shown in Fig. 6. The distribution is multimodal and is skewed towards high frequency of low concentration with a mean and standard deviation of 0.460 ± 0.37mg/g. A strong correlation exist between the cobalt content in hair and in nails (p<0.01) but the ANOVA (Table 4) indicated no significance difference between the mean cobalt concentration in hair and in nails at p>0.05. Fig. 7 represents cobalt concentration in hair and nails with respect to age. The pattern illustrates similar trends indicating the common source of the metal in them. Cobalt may not be age dependent because the concentration of 0.40mg/g was observed in all age groups. Though cobalt in hair increased with age, the
pattern appeared linear when compared with that of nails. These data are in agreement with values earlier reported (Weber et al., 1990; Hashem and Otham, 2001). From the levels of cobalt obtained, it is reasonable to believe that manganese and cobalt are playing some physiological roles in the respiratory organs where it plays special functions in man.

Table 1. Parametric Correlation Coefficients for Manganese in Hair and Nails

<table>
<thead>
<tr>
<th></th>
<th>Hair</th>
<th>Nails</th>
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</thead>
<tbody>
<tr>
<td>Hair Pearson correlation</td>
<td>1</td>
<td>91</td>
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<tr>
<td>Sig. (2-tailed)</td>
<td>.</td>
<td>501</td>
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<tr>
<td>N</td>
<td>350</td>
<td>300</td>
</tr>
<tr>
<td>Nails Pearson Correlation</td>
<td>091</td>
<td>1</td>
</tr>
<tr>
<td>Sig. (2-tailed)</td>
<td>507</td>
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<td>N</td>
<td>300</td>
<td>300</td>
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</table>

** Correlation is significant at the 0.01 level
Table 2. Analysis of variance for Manganese in Hair and Nails

<table>
<thead>
<tr>
<th>Source of variation</th>
<th>SS</th>
<th>Df</th>
<th>MS</th>
<th>F</th>
<th>P-value</th>
<th>F crit</th>
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<tbody>
<tr>
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<td>0.57456818</td>
<td>6.38530834</td>
<td>0.0136633</td>
<td>3.9290115</td>
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<tr>
<td>Within Group</td>
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<td>108</td>
<td>0.09141448</td>
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<td></td>
</tr>
<tr>
<td>Total</td>
<td>10.44733</td>
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Table 3. Parametric Correlation Coefficients for Cobalt in Hair and Nails

<table>
<thead>
<tr>
<th></th>
<th>Hair Pearson correlation</th>
<th>Hair Sig. (2-tailed)</th>
<th>Nails Pearson Correlation</th>
<th>Nails Sig. (2-tailed)</th>
<th>Sig. (2-tailed)</th>
<th>N</th>
<th>Sig. (2-tailed)</th>
<th>Sig. (2-tailed)</th>
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</thead>
<tbody>
<tr>
<td>Hair Pearson correlation</td>
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<td>.</td>
<td>703</td>
<td>.</td>
<td>000</td>
<td>.</td>
<td>350</td>
<td>300</td>
</tr>
<tr>
<td>Sig. (2-tailed)</td>
<td>.</td>
<td>.</td>
<td>000</td>
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<tr>
<td>N</td>
<td>350</td>
<td>300</td>
<td>1</td>
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</tbody>
</table>

** Correlation is significant at the 0.01 level

Table 4. Analysis of variance for Cobalt in Hair and Nails

<table>
<thead>
<tr>
<th>Source of variation</th>
<th>SS</th>
<th>Df</th>
<th>MS</th>
<th>F</th>
<th>P-value</th>
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<td>3.9290115</td>
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<tr>
<td>Within Group</td>
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<td>108</td>
<td>0.0963329</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>11.1992</td>
<td>109</td>
<td></td>
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</tbody>
</table>

Fig. 4. Manganese Concentration (mg/g) in hair and Nails with respect to age
Fig. 5. Frequency Distribution Pattern for Cobalt in Hair

Fig. 6. Frequency Distribution Pattern for Cobalt in Nails

Fig. 7. Cobalt concentration (mg/g) in Hair and Nails with respect to age
CONCLUSION

Human exposure to toxic trace elements has been the focus of increasing attention among researchers, formulators and managers of health and nutrition policies due to its damages to health. The levels Mn and Co in hair and nails vary and may be affected by various factors (Siedel et al., 2001). Age was observed to be a factor influencing their levels. The values of two elements recorded revealed sex dependence. Hair colour, nutritional status, geographic, racial/ethnic and ecological can have a significant impact on the levels of these elements in hair and nails (Sandra and Silva, 2002; Chojnacka and Gorecki, 2006a&b); but no correlation with any of these factors was observed, since our samples were collected from the same geographical location. The high trace element levels in hair and nails makes analysis easy; slow metabolic turnover rate of hair; and its being a reliable status of body without daily variation. The collection of samples were simple and non traumatic. Hair and nails may be regarded as complementary to body fluids in biological monitoring. Comparing hair and nails as points of excretion the latter appear superior to the former. The former enables monitoring of elements accumulated over a time span up to several months. They are easily sampled, handled and transported, and less prone to post – sampling contamination because of higher elemental concentration. Therefore human hair and nails are recording filaments that reflect metabolic changes of many elements over long periods of time and hence furnish post nutritional event as dietary levels of some essential micro – elements have been reported corresponding to hair concentrations of the elements (Maugh, 1978).

REFERENCES

Mn & Co Concentration in Hair and Nail


