The ingestion of considerable amounts of water or food contaminated with nickel can be very toxic. The present work was conducted aiming to evaluate the effects of nickel exposures on ascending colon of adult Wistar male rats at histometric level. We used 12 animals that were divided in a control (ingested uncontaminated water) and a nickel-contaminated (i.e., 25 mg de nickel/L of water) groups. Nickel chloride was offered in declorinated water and the experiment had a 56 days exposure period. A portion of the ascending colon was removed of the animals and subjected to histological labelling processes using blue toluidin (for general histometric description), Alcian Blue (AB, for acid mucins) and periodic acid-Schiff (PAS) technique (for neutral mucins). The potential differences between groups were desgined by applying the Whitney test and $t$ test ($p < 0.05$). The crypts were smaller for the nickel-contaminated group, even though these organism exhibited broader and higher crypts. Nickel-contaminated animals exhibited a smaller amount of calyceform cells with AB and PAS positive reactions as well as a less mucus quantities when compared with nickel-uncontaminated animals. Such reductions on the amount of calyceform cells with AB and PAS positive reactions may be related with the shallower crypts, which possibly reduced the synthesis and secretion of mucins, compromissing the functional aspects (e.g., lubrification and intestinal mucosa protection) of the nickel-contaminated large intestines. Interestingly, the wider and higher crypts and higher epithelium collumn on the nickel-contaminated animals may represent a relevant trade-off for the intestinal mucosa protection.

**Keywords:** nickel contamination, toxicology, hystology, morphology.
INTRODUCTION

Nickel is a heavy metal naturally found in the earth’s crust at about 0.009%, existing as soluble and insoluble compounds in soils and waters (Bertini et al., 2001). However, the environmental nickel concentration has increased due to the anthropogenic actions (mining and industrial activities) that are involved in the production of clocks, mobile phones, coins and cardiovascular stents (Schmidt and Goebeler, 2011).

The daily nickel requirements by human beings is estimated to be between 5 and 50μg (EMEA, 2008), which has leading it to be considered as ‘potential essential’ for animals, including humans, in 1970’s (Nielsen and Ollerich, 1974). Interestingly, such categorization was based on some experiments using model animals but exclude the fact that no nickel-containing enzymes were found on such investigations. The potential reasons for this nickel need are still controversial but may be related to the use of nickel by the mammals’ microbiota (Monachese et al., 2012). For instance, Gerbino et al. (2012) demonstrated that maximum tolerated nickel (Ni²⁺) concentrations without reductions on the biomass of Lactobacillus kefir CIDC3A 8348 and JCM 5818 were of 5mM and 2.5 mM, respectively, which led to the conclusion that Ni²⁺ is an essential metal in vestigial concentrations and toxic to bacteria when in higher concentrations. Thus, when nickel is ingested in higher amounts, it becomes toxic to live organisms (Gonzalez, 2016).

The high toxicity levels of heavy metals to organisms and their easy way of entering and accumulating along the trophic chains reinforce the relevance of investigating these metal concentrations in aquatic environments (Rocha and Azevedo, 2015). The humans daily ingest circa of 170μg of nickel through food and drink approximately 20μg of nickel. Such nickel-containing food and water ingestion have to be considered as it has been demonstrated that starving individuals absorb much more nickel on their gastrointestinal tract (Ahmad and Ashraf, 2011). The nickel absorption may vary from 12% up to 33% in starving organisms, and from 0.7 up to 6% when mixed with food (Sunderman et al., 1989; Haber et al., 2000). In investigations conducted with animals, it was demonstrated that nickel absorption by ingestion also varies with the nickel solubility, being higher for nickel soluble compounds, 9.8%, 11% and 34% for sulphate, chloride and nitrate ions, respectively; and lower for sulphite salts and green oxides: 0.47 and 0.01%, respectively (Haber et al., 2000).

Pharmacocynetics in human beings demonstrated that nickel is absorbed through lungs, intestinal tracts and skin. In the...
respiratory tract, the nickel can be removed by mucociliary transport being delivered to gastrointestinal tract, where its absorption is not high. The absorption level is influenced by metal particles’ size, chemical structure, density and electric charge as well as by accumulation place in the organism (Clancy and Costa, 2012).

In the large intestine occurs the absorption of water and inorganic salts, leading to the formation of fecal bolus. To achieve that, the colicular cells present microvils in the apical surface and \( \text{Na}^+ / \text{K}^+ \)-ATPases in the basolateral membranes. The caliciform cell are in great number and the mucus contribute to the fecal bolus compactuation, facilitating the its sliding and lubricating the epithelial surface (Gartner and Hiatt, 2017; Lowe and Anderson, 2015). In addition to lubrication, which turns the fecal bolus progression an easier task, the colons’ mucus barrier promote the chemical protection in the internal intestinal lumen against potential aggression against digestive enzymes, antigens and toxins (Hoebler et al., 2006; Gaudier et al., 2009). The mucus still present bactericidal properties, regulating the bacterial strains that are in direct contact with the epithelial surface, dificultating these organisms enter into the internal environment (Finnie et al., 1995; Corfield et al., 2000; Hoebler et al., 2006; Gaudier et al., 2009).

Thus, this research was conducted aiming to evaluate the nickel effects on the ascending colon of adult Wistar male rats at hystometric level and trying to correlate the potential alterations with the functional aspects of digestive tract.

MATERIAL & METHODS

Animals, treatment and material collection. The animals were handled and euthanized according to the Animal Research Ethical Principles established by the Colégio Brasileiro de Experimentação Animal (COBEA, i.e., Brazilian College of Animal Experimentation - Protocol 001/11). Twelve adult Wistar rats of the same age (90 days), weighing between 276 and 302g, from the Central Biotery of the Center for Biological and Health Sciences at Universidade Federal de Viçosa (UFV, Viçosa, Minas Gerais, Brasil) were used. The animals were kept in the biotery of the Health and Nutrition Department at UFV throughout the experimental period (56 days). The ambient temperature was 22 ± 1°C and the photoperiod 12:12h (light:dark). The animals were divided into two groups (of six individuals each), individually housed in cages (18 x 34 x 41cm) and fed daily (at 9:00 a.m.) with 30g of pelleted commercial food (Labcil, SOCIL, São Paulo, São Paulo, Brasil). The control group received free water and whereas the nickel-treated animals received 25mg of nickel chloride/L of tap drinking water over the experimental period. We monitored the average daily intake of 30mL of water per animal, which corresponds to a nickel consumption of 0.75mg/day. The animals were weighed every two weeks in order to evaluate the weight gain.

After the experimental period, the animals were anesthetized using 30mg/Kg of the commercial combination of ketamine and xylazine (Xylazin and Cetamin, Syntec, Santana de Parnaíba, São Paulo, Brasil) and euthanized by prolongating anesthesia. After that, the animals were weighed, and a cranial portion fragment of the ascending colon was removed of each animal. The fragments were then fixed in Karnovsky solution for 24 h and then left in 70% alcohol.
Hystometric evaluations of ascending colon region. The fixed ascending colon fragments were dehydrated in a growing ethyl series (alcohol 80, 90, 95% and absolute I) and included in historesin. Semi-serial sections of 3µm thickness were obtained using a rotating microtome (model RM2245, Leica, Lincolnshire, Buffalo Grove, United States). The sections were stained with toluidine blue (in 1% sodium borate) for tissue hystomorphological description and histometric analysis, with periodic acid-Schiff (PAS) and Alcian Blue (pH 2.5) techniques for quantifications of neutral and acidic, respectively, mucin-producing cells and the stored mucus in these cells.

Observations and photographic recordings of the sections were performed under an optical microscope with an attached digital camera (model BX53, Olympus, Miami, Florida, United States) and photomicrographs were obtained using the 10x objectives for the intestinal crypt’s depth and width (at midpoint) and for height apical epithelium height. Objectives of 20x were used for photomicrographs used at the quantification of other caliciform cells and PAS- and AB-positive mucus-producing cells. Five fragments of each animal were randomly selected, and photomicrographs were obtained of five fragment fields per each fragment, totaling 25 photomicrographs per animal and 150 photomicrographs per animal group (control and nickel-contaminated animals).

To obtain the histometric data, the photomicrographs obtained were digitalized and analyzed using the Image J software (Media Cybernetcs, Inc, Madison, Wisconsin, United States). From the histological images, we conducted five measurements for each parameter (depth and width of intestinal crypts and height of the apical epithelium), obtaining 125 measurements/animal/ parameter. The quantification of caliciform cells and PAS- and AB-positive mucus-producing cells per µm² (photographic field area) were obtained by counting all stained cells in the field.

Statiscal analyses. The normality of Kolmogorov-Smirnov and homoscedasticity of Cochran tests were conducted using the Prisma (Graphpad software, San Diego, California, United States) software. Once these assumptions were achieved, a t-test ($P < 0.05$) was conducted for comparing the values obtained for nickel-unexposed (control) and nickel-treated animals. When the normality and homoscedasticity were not achieved, it was conducted such comparison using the Mann-Whitney nonparametric test ($P < 0.05$).

RESULTS AND DISCUSSION

Some researchers and toxicologists are focusing on the risk of environmental pollution by heavy metals, which are dangerous and cause effects related to animal studies, toxic effects due to oral exposure, with the possibility of slow development (Domshlak et al., 2005).

In the present study, the animals used with 25 mg/L NiCl₂ showed no significant changes ($P > 0.05$) in body weight, although there was a tendency to decrease in weight gain (Figure 1). Several studies have evaluated nickel toxicity by oral exposure at different chronic doses, and a significant reduction in body weight has been observed (Dieter et al., 1988; Obone et al., 1999; Cempel and Janicka, 2002; Gill et al., 2002). The toxic effects by chemicals in biological systems are determined by the concentration mechanism and the exposition time (Oga et al., 2008). Therefore, the dose used in this study may not have been sufficient to cause significant reduction in body weight of these animals.
However, the other hystometric evaluated parameters showed significant differences between the animal groups (Figures 2 and Figure 3). Here, it was showed a reduction in the number of caliciform cells and in the amount of PAS- (Figures 2A and B; 3C and D) and AB-positive mucus (Figure 2C and D; Figures 3E and F) in the large intestine of nickel-exposed Wistar male rats. The number of caliciform cells demonstrates the potential for mucus secretion and colonic gland activity (Glotzer et al., 1981). Changes in the composition and pattern of mucin secretion in the intestinal crypts have been observed in several inflammatory diseases that reach the colon, such as bacterial colitis, nonspecific ulcerative colitis, Crohn’s disease, exclusion colitis, adenomatous polyps and colorectal cancer (Filipe 1969; Finnie et al., 1995; Keli et al., 1997). In nonspecific colitis, the total production of acid and neutral mucins are reduced, being absent in some cases and in regions with severe inflammatory process (Filipe, 1969). Thus, a reduction in mucin secretion is seen as a detrimental effect in the intestinal protection and lubrication as well as in mucin-mediated processes such as cell-cell and cell-substrate interactions, wounding and immune response modulations (Kim and Gum, 1995).

Figure 1. Effect of nickel chloride exposure on body weight of adult male Wistar rats. Four weight measurements were performed at two week intervals.

There was no statistically significant difference. Data are expressed as mean ± standard error (SE). * P < 0.05 (t-test).

![Body weight graph](image)
Data are expressed as mean ± standard error (SE). * $P < 0.05$ (t-test).

**Figure 2.** Effects of nickel chloride exposure on the number of caliciform cell with PAS- (**A**) and AB- (**C**) positive reactions (**A** and **C**, respectively) on colon of adult Wistars rats. The intensity of PAS- and AB-positive mucin marking (**B** and **D**) were also presented.
The results obtained for control (i.e., nickel-unexposed) animals are presented in panels (A), (C) and (E) while the results obtained for nickel(25mg/L)-exposed animals are presented in panels (B), (D) and (F). The epithelium (ep) and intestinal crypts (c) are shown in the material stained with toluidine blue (A – B). Material stained with periodic acid-Schiff (PAS) (C – D) and Alcian Blue (AB) (E – F) techniques are also shown. The arrowheads indicated the neutral (C – D) and acid (E – F) mucing-producing cells.

**Figure 3.** Hystological microphotographies of colon of Wistar rats.
The decrease in crypt depth (Figures 3A and B; Figure 4A) reflects the reduction in the number of mucus producing cells. Previous studies have shown that animals that had their fecal transit decreased, also had a decrease in the number of mucous cells, which led to significant reduction in crypt depth (Mello et al., 2012). In the present investigation, it was also demonstrate an increase in crypt width (Figures 3A and B; Figure 4B) and epithelium height (Figure 3A and B; Figure 4C) in animals exposed to nickel chloride. Such changes can be understood as a compensatory mechanism for protection of intestinal mucosa, as the mucus barrier was subjected to relevant alterations (reduction in the number of caliciform cells).

![Figure 4](image_url)

**Figure 4.** Effect of nickel chloride exposures on the colon of adult Wistars rats. Crypt’s depth (**A**), and width (**B**). (**C**) Epithelium height. Data are expressed as mean ± standard error (SE). *P* < 0.05 (Mann-Whitney test).
CONCLUSIONS

The chronic exposure to nickel impacted the colong region of Wistar adult male rats. The smaller number of caliciform cells and lower PAS- and AB-positive mucus production is certainly related to reduced crypt depth in nickel-exposed animals, which may compromised the functional aspects of the large intestine (e.g., lubrication and protection of the intestinal mucosa). The larger values for crypt width and epithelium height may be a compensatory mechanism for protection of the intestinal mucosa as the mucus barrier was also altered.

Conflict of interest: The authors declare that there is no conflict of interest.

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