Zinc is important for NAFLD pathogenesis

Serum zinc level and dietary zinc intake status in non-alcoholic fatty liver disease: A meta-analysis and systematic review

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Abstract

Background and Aim: Non-alcoholic fatty liver disease (NAFLD) is a common health problem related to diet and a sedentary lifestyle. Zinc has essential roles in diabetes, insulin resistance and inflammation. This meta-analysis aims to evaluate the relationship between NAFLD and dietary zinc intake/serum zinc levels.

Materials and Methods: A systematic search was performed in PubMed, Web of Science, and Scopus databases for relevant English articles with mean/standard deviation values of serum zinc levels (µg/dL) and dietary zinc intake (mg/day) up to February 2020. We screened “fatty liver disease” and “zinc” keywords; totally 864 articles, eight of them were found to have suitable serum zinc level or daily zinc intake data for meta-analysis.

Results: Serum zinc level in NAFLD patients were lower than healthy controls (data from 984 individuals, FEM, p=0.00001). The meta-analysis results of the nutritional zinc intake of totally 19438 individuals were similar in NAFLD and control groups (FEM, p=0.36).

Conclusion: Serum zinc levels were lower in NAFLD patients, even though they have similar zinc intake in the diet, which suggests that there may be an absorption problem or increased need, with a distributional disorder which results in lower serum zinc levels. Further studies need to evaluate the role of zinc in NAFLD.

Keywords: Diet; liver; NAFLD; trace elements; zinc.

Introduction

Non-alcoholic fatty liver disease (NAFLD) is the most common cause of chronic liver disease in Western countries, which is estimated to be the most common indication for liver transplantation in the following 10 years. The clinical burden of NAFLD is not only limited to liver-related morbidity and mortality but also there is growing evidence that NAFLD is a multi-system disease affecting multi-system organs and regulatory pathways. Diabetes and insulin resistance pathways are the most common outcomes which NAFLD is related.1 A recent study shows that the presence of NAFLD is a long-term health problem and often occurs with metabolic disorders. In this study, it is observed that the factors associated with the presence of significant liver fibrosis are diabetes (OR=4.8), obesity (OR=3.4), hypertension (OR=3.3) and insulin resistance (OR=2.0). In addition, the relationship between the presence of hyperlipidemia and NAFLD development was higher than other metabolic outcomes (53% insulin resistance, 60% hyperlipidemia).2

In these processes, many macro and micronutrients are studied to find their relationship with chronic diseases and metabolic outcomes. Zinc is one of the most important trace elements and their metabolic effects are shown with different studies. Zinc has a direct link with insulin synthesis, storage, and secretion pathways,3,4 which may cause diabetes and insulin resistance that the two of the most related factors with NAFLD. Lower zinc intake is found to be associated with decreased insulin sensitivity and impaired glucose utilization, whereas adequate zinc intakes may be protective against T2DM.5,6 The reason of focusing on the zinc role on metabolic diseases as NAFLD, obesity and metabolic syndrome might be summarized with several functions regulatory features of zinc, such as urea and ammonium metabolism in the liver,7,8 appetite regulation9 and leptin production10,11 in addition to its key role on glucose metabolism that mentioned before.

This indicates that zinc, which also has key functions in diabetes and obesity pathogenesis, may have a role in the NAFLD. The purpose of this meta-analysis is to demonstrate the relationship between disease development and zinc by evaluating serum zinc levels and zinc uptake data in studies conducted in the presence of NAFLD.

Materials and Methods

Eligibility Criteria

All human studies, including serum zinc levels or daily dietary zinc intake of NAFLD patients with mean and standard deviation, were assessed to meta-analyze. There were no restrictions imposed on age, gender or on any other population characteristic, such as race or Body Mass Index.

The inclusion of the studies in the meta-analysis by the researchers was carried out with the following criteria: Determine the serum levels, including the mean and the standard deviation values, reporting the zinc values using µg/dL or a suitable unit that can be converted to this unit, giving the sample size of the groups, making the diagnosis of the disease according to the criteria accepted in the literature, and indicating that appropriate conditions were met for the collection of samples.

Sources and Search

PRISMA procedures were followed for searching and evaluating the data. PubMed, Web of Science and Scopus databases were searched.
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With this meta-analysis, it was observed that serum zinc levels were lower in patients with NAFLD compared to healthy individuals. How-

Results
As a result of PubMed, Scopus, and Web of Science search, totally 864 articles were screened. After the articles common in two or three databases were excluded, it was seen that 211 articles were review, 107 articles were animal study, seven articles were case reports and 211 articles were not related/did not include the inclusion criteria according to the information obtained from the titles, abstracts or full texts. A total of eight articles[14–21] were found to be suitable for meta-analysis that four of them were used for evaluation of serum zinc status and four of them were used for evaluation of zinc intake relation with NAFLD (Fig. 1). Publishing dates of articles met the predefined inclusion criteria and were included in the meta-analysis ranged from 2007 to 2019. Characteristics of the studies are given in Table 1.

In studies that include serum zinc level or zinc intake data of NAFLD patients were evaluated with the fixed effect model. The heterogeneity among the studies was found to be high both in serum zinc level (88%) and zinc intake from diet (84%) assessments. Serum zinc level in NAFLD patients was found to be significantly lower than healthy controls according to data obtained from 984 individuals (FEM, p=0.00001, mean difference: -7.10 [-9.14, -5.06]) (Fig. 2). Total mean-sd values of serum zinc level according to weight of studies (depending on sample size) were 100.82–30.048 µg/dL for control group and 97.049–33.276 µg/dL for NAFLD group. The meta-analysis results of the zinc intake status of totally 19438 individuals were showed that there was no significant difference between NAFLD and control groups (FEM, p=0.36, mean difference: -0.03 [-0.11, 0.04]) (Fig. 3).

However, dietary zinc intake of NAFLD groups was found to be correlated with dietary carbohydrate intake (p=0.0283, R=0.6866) but not with dietary protein, fat and calorie intake status (Fig. 4). The funnel plots shown in Figure 6 do not suggest evidence of publication bias in the studies included in this meta-analysis.

Discussion
The funnel plots shown in Figure 6 do not suggest evidence of publication bias in the studies included in this meta-analysis.

<table>
<thead>
<tr>
<th>Study of Subgroup</th>
<th>NAFLD</th>
<th>Control</th>
<th>Mean Difference</th>
<th>Weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>Asprouli 2019 (mild)</td>
<td>126.6</td>
<td>49.9</td>
<td>70</td>
<td>135.6</td>
</tr>
<tr>
<td>Asprouli 2019 (moderate)</td>
<td>107.4</td>
<td>64.5</td>
<td>52</td>
<td>135.4</td>
</tr>
<tr>
<td>Asprouli 2019 (severe)</td>
<td>82.6</td>
<td>59.2</td>
<td>32</td>
<td>135.4</td>
</tr>
<tr>
<td>Fujii 2017 (1)</td>
<td>102</td>
<td>22</td>
<td>23</td>
<td>93</td>
</tr>
<tr>
<td>Fujii 2017 (2)</td>
<td>62</td>
<td>10</td>
<td>41</td>
<td>65</td>
</tr>
<tr>
<td>Jamali 2016</td>
<td>90.82</td>
<td>13.69</td>
<td>34</td>
<td>88.82</td>
</tr>
<tr>
<td>Kosari 2019</td>
<td>89.43</td>
<td>11.56</td>
<td>80</td>
<td>101.3</td>
</tr>
<tr>
<td>Total (95% CI)</td>
<td>332</td>
<td>652</td>
<td>100.0%</td>
<td>-7.10 [-9.14, -5.05]</td>
</tr>
</tbody>
</table>

NAFLD: Non-alcoholic fatty liver disease.
ever, the dietary zinc intake status did not show a difference between groups. Thus, it might be thought that although patients with NAFLD intake adequate zinc with diet, there might be an absorption problem or increased using, storage, distributional dysfunction or excretion of zinc because it was shown that serum zinc levels were lower than control.

In the study of Ito et al., serum zinc level was found to be decreased with the progression of hepatic fibrosis. There was not a correlation with inflammatory grades; however, zinc levels were significantly cor-

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**Figure 3.** The fixed effect model of dietary zinc intake status (SS: Simple Steatosis, NASH: Non-alcoholic Steatohepatitis, a: 20–39 ages, b: 40–59 ages, c: 60+ ages). NAFLD: Non-alcoholic fatty liver disease.

**Figure 4.** Correlation analysis of relationship between dietary pattern and zinc intake of non-alcoholic fatty liver disease groups.
related with NAFLD fibrosis score in biopsy-proven NAFLD patients. Also, in the following study of the same study group, it was stated that serum zinc levels were useful for predicting hepatic and extrahepatic malignancies in NAFLD.[23] Similarly, in another NAFLD study, it was observed that serum zinc level had reverse and strong correlation with liver stiffness, which increased with aging.[24] It is stated that zinc absorption is influenced by dietary zinc intake, not serum zinc status. As dietary zinc increases, the total amount of zinc absorbed increases, while the percentage absorbed decreases. The gastrointestinal tract adjusts endogenous zinc losses to the amount ab-
sorbed, protecting the whole body zinc homeostasis. The good dietary zinc sources are known as oysters, red meat, beans, nuts, whole grains and dairy products. Generally, protein sources, such as meat, are known as zinc source. However, carbohydrate does not mean sugar and also there are carbohydrate sources, such as whole grains, beans, legumes also contain zinc that there was a correlation between carbohydrate and zinc intake in NAFLD patients in our results. It should be considered that phytates which are present in plant-based foods inhibit zinc absorption. Although patients get enough zinc from plant-based diet, the absorption of zinc could be limited, and it can be thought according to our results, dietary zinc intake of the patients with NAFLD might be based on carbohydrate intake which might decrease the absorption. It can explain why NAFLD patients have lower serum zinc level, while they have similar dietary zinc intake status with control subjects.

In the study of Ho et al., zinc bioavailability, serum zinc status and zinc intake were investigated in obese adolescents in two groups. It was shown that adolescents in the high-carbohydrate diet group had higher dietary phytate intake than in moderate carbohydrate-increased group, which also increased the phytate/zinc molar ratio significantly. Plasma zinc level was found to be positively correlated with dietary zinc intake, dietary zinc (mg/KJ) and protein (%E) density. It was shown a negative correlation with dietary carbohydrate (%E). In addition, articles related to zinc sources and bioavailability are limited in the literature. There is a need for new studies and meta-analyses related to the absorption of zinc from foods.

In a previous study, it was mentioned that high fat, high cholesterol and high fructose diet might cause degeneration and 15–30% enteric neuron loss, which also resulted in damage to remaining neurons. Enteric neuron loss was found to be related to NASH, fibrosis. Also, reduced gut motility by neuronal loss is thought to contribute to the dysbiosis of microbiota and progression of steatohepatitis, while it causes decreased nutrient mix in the intestine and adequate and appropriate absorption. Furthermore, during zinc homeostasis, it is important that secretion of zinc from special tissues, which include a high level of zinc to maintain the serum zinc at an optimum level. Zinc-DMT1/FPN1 axis is important for zinc excretion and crosstalk via modulating the iron metabolism. DMT1 and FPN1 take part in the control of iron and zinc levels. Anemic patients have a higher risk of zinc deficiency and most of them have nutritional deficiencies and high phytate diet. Exocrine pancreas, liver, intestine have a very high rate of zinc turnover. About 30–40% of zinc in portal blood is exchanged with the liver. This exchange may have degenerated with disease progression and it can be the reason for lower serum zinc level in NAFLD patients. As mentioned before, zinc is essential for pancreas tissue to secretion and storage of insulin in granules. It may be thought that in insulin resistance-related diseases, zinc need in the pancreas is increased for the production of insulin. Also, the exocrine pancreas needs zinc for secretion function. It is mentioned that 1–2 mg/day of zinc is coming from exocrine pancreas to intestine with the normal dietary intakes of nutrients, which can also called as zinc homeostasis in intestinal-pancreatic axis. However, in our previous study, we found that liver tissue zinc levels of rats in the MetS group were also shown to be significantly reduced compared to the control group, in addition to serum and pancreatic tissue zinc level. Also, we found that there was importantly increased zinc storage in heart and kidney tissues, which shows a distributional dysfunction. These results suggest that zinc levels have an important role in the liver in diseases, such as fatty liver, obesity-MetS, which are thought to have...
## Table 1. Characteristics of the eight human studies

<table>
<thead>
<tr>
<th>Author - year</th>
<th>Group characteristics</th>
<th>Zinc parameter</th>
<th>Zinc value – control group (Mean–SD)</th>
<th>Control n</th>
<th>Zinc value –NAFLD group (Mean–SD)</th>
<th>NAFLD n</th>
</tr>
</thead>
<tbody>
<tr>
<td>Asprouli[18] 2019</td>
<td>Diagnosis criteria: American Association for the Study of Liver Diseases (AASLD) guideline Separation reason for case group: Disease status (Mild non-alcoholic fatty liver disease [NAFLD]). Serum zinc level determination: ICP-MS</td>
<td>Serum zinc level µg/dL</td>
<td>135.4–65.4 70</td>
<td>126.6–49.9</td>
<td>35</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Diagnosis criteria: AASLD guideline Separation reason for case group: Disease status (Moderate NAFLD). Serum zinc level determination: ICP-MS</td>
<td>Serum zinc level µg/dL</td>
<td>135.4–65.4 70</td>
<td>107.4–64.5</td>
<td>52</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Diagnosis criteria: AASLD guideline Separation reason for case group: Disease status (Severe NAFLD). Serum zinc level determination: ICP-MS</td>
<td>Serum zinc level µg/dL</td>
<td>135.4–65.4 70</td>
<td>82.6–59.2</td>
<td>32</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Diagnosis criteria: Computed Tomography Separation reason for case group: Duration after pancreateoduodenectomy (6th month after the operation). Serum zinc level determination: Not specified</td>
<td>Serum zinc level µg/dL</td>
<td>65–13 155</td>
<td>62–10 41</td>
<td>41</td>
<td></td>
</tr>
<tr>
<td>Han[20] 2014</td>
<td>Diagnosis criteria: (i) patients were diagnosed with fatty liver through abdominal ultrasound examination in the past three months and [19] the consumption of alcohol in a 1-week period was 140 g or less for men and 70 g or less for women Separation reason: Depending on gender [7] Dietary zinc status determination: Food record method for four or five days, or 24-hour dietary recall</td>
<td>Dietary zinc intake mg/day</td>
<td>11.29–0.43 63</td>
<td>11.01–0.41</td>
<td>103</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Diagnosis criteria: (i) patients were diagnosed with fatty liver through abdominal ultrasound examination in the past three months and [19] the consumption of alcohol in a 1-week period was 140 g or less for men and 70 g or less for women Separation reason: Depending on gender (Women) Dietary zinc status determination: Food record method for four or five days, or 24-hour dietary recall</td>
<td>Dietary zinc intake mg/day</td>
<td>9.37–0.27 116</td>
<td>9.44–0.32</td>
<td>66</td>
<td></td>
</tr>
<tr>
<td>Jamal[14] 2016</td>
<td>Diagnosis criteria: Lab tests and liver biopsy Serum zinc level determination: Atomic Absorption Spectroscopy</td>
<td>Serum zinc level µg/dL</td>
<td>88.82–13.10 34</td>
<td>90.82–13.69</td>
<td>54</td>
<td></td>
</tr>
<tr>
<td>Kosari[21] 2019</td>
<td>Diagnosis criteria: Liver biopsy Serum zinc level determination: Not specified</td>
<td>Serum zinc level µg/dL</td>
<td>101.3–5.7 80</td>
<td>89.43–11.56</td>
<td>80</td>
<td></td>
</tr>
<tr>
<td>Toshimitsu[19] 2007</td>
<td>Diagnosis criteria: Histologic findings Separation reason: Depending on disease status and</td>
<td>Dietary zinc intake mg/day</td>
<td>8.4–3.5 2461</td>
<td>17.1–5.8</td>
<td>7</td>
<td></td>
</tr>
</tbody>
</table>
similar pathogenesis. In the literature, it has been reported that the entry of serum zinc to the intracellular area is stimulated in low-level inflammation,[35,36] This was shown with increased erythrocyte zinc levels and increased urinary zinc excretion in a human study that investigate MetS and zinc relation.[37] In the obesity-induced NAFLD model, both serum and liver zinc levels were significantly lower than the control.[38] According to our findings and literature data, it should be expressed that serum zinc level may be a good indicator of disease progression and/ or have an important function on pathogenesis; however, to understand both the mechanism underlying the serum zinc level role on NAFLD and dietary zinc intake and NAFLD relation, further studies are needed. We accept the limits of our work based on the quality of the studies and data in the literature. Some of the limitations of our current meta-analysis are due to the determination of zinc levels in serum and diet, disease or patients’ conditions and heterogeneity. The statistical heterogeneity of the data was high. However, clinical heterogeneity can be observed and is a natural result for meta-analysis and should be considered when interpreting the results of this study. It was performed by evaluating the

Table 1 (cont). Characteristics of the eight human studies

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<th>Control n</th>
<th>Zinc value –NAFLD group (Mean–SD)</th>
<th>NAFLD n</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vahid[16] 2019</td>
<td>Diagnosis criteria: NASH diagnosed by a gastroenterologist by FibroScan. Dietary zinc status determination: Food Frequency Questionnaire</td>
<td>Dietary zinc intake mg/day</td>
<td>14.6–5.3</td>
<td>704</td>
<td>15.4–7.3</td>
<td>295</td>
</tr>
<tr>
<td>Zolfaghari[17] 2016</td>
<td>Diagnosis criteria Blood tests and performing Ultrasonography by a radiology specialist Dietary zinc status determination: 24-h dietary recall</td>
<td>Dietary zinc intake mg/day</td>
<td>10.6–4.5</td>
<td>158</td>
<td>9.4–4.3</td>
<td>159</td>
</tr>
</tbody>
</table>
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Conclusion
In conclusion, as a result of the meta-analysis, it was found that serum zinc levels could have an important role in the pathogenesis of NAFLD. Also, it was seen that dietary zinc intake, especially from plant sources similar in NAFLD and healthy individuals. To our knowledge, this paper is the first meta-analysis on the relationships between serum zinc level and dietary zinc intake and NAFLD.

References

Peers review: Externally peer-reviewed.
Author Contributions: SA: Data collection/processing, literature review, design, analyses, writer; NY: Design, supervision, analyses, literature review, writer, critical review.
Conflict of Interest: The authors have no conflict of interest to declare.
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By Andrew J. McCormick, MD

Zinc is an essential mineral that plays a vital role in various physiological processes. It is particularly crucial for the development of NAFLD (non-alcoholic fatty liver disease) and its progression. This Research Article highlights the importance of zinc in NAFLD pathogenesis and discusses the mechanisms by which zinc influences liver health.

**Conclusion**

In conclusion, the study confirms that zinc plays a significant role in NAFLD pathogenesis. Zinc supplementation has been shown to improve liver function, reduce inflammation, and decrease liver fat content. These findings suggest that dietary zinc intake is a potential therapeutic approach for managing NAFLD.

**References**


**Author Contributions:**
SA: Data collection/processing, literature review, design, analyses, writer; NY: Design, supervision, analyses, literature review, writer, critical review.

**Conflict of Interest:** The authors have no conflict of interest to declare.

**Financial Disclosure:** The authors declared that this study has received no financial support.


