Effects of Aroclor 1254 and vitamin E on arginase activity in adult, pregnant rats and their offsprings

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Summary: This study examined the effect of Aroclor 1254 on arginase activity and protective role of vitamin E among adult rats, pregnant rats and their offsprings. Totally 90 Sprague-Dawley rats were divided into three main groups as 30 pregnant, 30 adult and 30 offsprings (pups of the pregnant rats) based on the treatments administered during the pregnancy period (20 days) with subcutaneous daily injections of Aroclor 1254 (2 mg/kg/day) alone or associated with vitamin E (50 mg/kg/day) or normal saline in controls. Female rats and their offsprings (10 offsprings per groups) were analyzed in terms of the arginase activity in liver. A statistically significant increase was determined in liver arginase activity of adult rats exposed to Aroclor 1254 (p<0.05). Vitamin E administered simultaneously with Aroclor enabled liver arginase activity of adult rats to approach statistically normal values. A statistically insignificant increase was found in the arginase activity among pregnant rats and offsprings. While vitamin E administered simultaneously with Aroclor was ineffective in pregnant rats, it caused a statistically significant increase in offsprings (p<0.05). We recommend that addition of vitamin E can prevent the increase in the liver arginase activity caused by Aroclor 1254.

Keywords: Arginase, Aroclor 1254, offspring, pregnant, vitamin E.

Introduction

Polychlorinated biphenyls (PCBs) are halogenated aromatic hydrocarbons that were produced for industrial purposes and are now known as persistent environmental pollutants (1, 4). Their common usages are industrial fluids, flame-retardants, diluents, and fluids for capacitors and transformers (15). They have a tendency of bio-accumulation both in the environment and living organisms because of their lipophilic features and chemical stability (28, 32). It has been reported that PCBs have various adverse effects on human health including carcinogenicity, endocrine disruption, neural, immune, developmental and reproductive toxicity (17, 27, 31).

Since Arginase (L-arginine amidinohydrolase, EC 3.5.3.1) catalyses the hydrolysis of L-arginine to L-ornithine and urea in the final cytosolic step of the urea cycle, it exists mainly in the liver. This reaction provides the principal route in order to dispose nitrogenous waste from protein catabolism (26). The liver is the main target organ for PCBs in the body as they are detoxified here (22). PCBs are known to cause hepatotoxicity (17, 18, 23). Serum arginase activity increases in benign and malignant liver diseases (7).

Some studies have proved that there are some relationships between vitamin E and arginase. Park and Tappel (25) reported that rats fed with a vitamin supplemented diet had a lower liver arginase activity compared to those fed with a diet not containing vitamin E. Additionally, administration of dietary vitamin E caused a significant decrease in the liver arginase activity increased due to high doses of prednisolone in rats (12).

Aroclor 1254 ve vitamin E’nin yetişkin, gebe rat ve yavrularında arginaz aktivitesi üzerine etkileri

Özet: Bu çalışmada yetişkin, gebe rat ve yavrularında arginaz aktivitesi üzerine Aroclor 1254’in etkisi ve vitamin E’nin koruyucu rolü araştırıldı. Toplam 90 Sprague-Dawley rat 30 yetişkin, 30 gebe ve 30 yavru olmak üzere üç ana gruba ayrıldı. Gebelik süresince (20 gün) subkutan olarak Aroclor 1254 (2 mg/kg/gün) tek başına veya vitamin E (50 mg/kg/gün) ile birlikte ve kontrol grubuna serum fizyolojik uygulandı. Dişi ratlar ve onların yavrularının (her gruptan 10 yavru) karaciğerinde arginaz aktivitesi analiz edildi. Aroclor 1254 ile birlikte vitamin E uygulanmış yetişkin ratların arginaz aktivitesi istatistiksel olarak normal değerlerle yaklaştı. Gebe ratlar ve yavruların arginaz aktiviteleri istatistiksel olarak anlamsız bir artış tespit edildi. Aroclor 1254 ile birlikte vitamin E uygulanması gebe ratlarda etkili olmasa da; yavru ratlarda anlamlı bir artışa (p<0.05) sebep olmuştur. Vitamin E ilavesinin, Aroclor 1254’ün sebep olduğu karaciğer arginaz aktivitesindeki artış önlenebileceğini önermektediz.

Anahtar sözcükler: Arginaz, Aroclor 1254, gebe, vitamin E, yavru.
PCBs transfer to foetus and to newborn by means of placenta and milk, respectively (2, 21, 30). Aroclor 1254 is also known to be a dangerous environmental pollutant for offsprings (5, 6). There is no study investigating the effect of Aroclor 1254 on liver arginase activity. Therefore, the purpose of this study was to examine the effect of Aroclor 1254 on arginase activity and protective role of vitamin E among adult rats, pregnant rats, and their offsprings.

Materials and Methods

Experimental design: Totally 90 female Sprague-Dawley rats having a weight between 150-180 g were used in the study. The rats were kept in cages (n=5 per cage) at standard temperature (21 ± 1°C) in a 12:12 light/dark cycle. Once the sexual cycle periods and/or pregnancy were determined according to the vaginal smear method, rats were randomly divided into groups based on chemical treatments and physiological status. Groups 1 and 4 served as nonpregnant (n=10) and pregnant (n=15) controls and were administered with normal saline subcutaneously for 20 days, respectively. While in groups 2 (nonpregnant females, n=10) and 5 (pregnant females, n=15), only Aroclor 1254 was subcutaneously administered to females at 2 mg/kg/day for 20 days (since the first day of pregnancy in pregnant rats); in groups 3 (nonpregnant females, n=10) and 6 (pregnant females, n=15), vitamin E (50 mg/kg/day) were administered with the Aroclor 1254. The last groups (groups 7, 8 and 9) corresponded to offsprings (10 in each group) from the control pregnant females, from females only administered with Aroclor 1254, and from females administered with Aroclor 1254 and vitamin E, respectively. On the day 20, liver tissue of 10 rats from each group was extracted during deep ether anaesthesia. Liver tissue samples were stored at -80°C until analysis.

Arguments activity was defined as µmol of the product formed per min at 4 ºC. The supernatants were used in order to carry out assay of the arginase activities.

Statistical analysis: Measurement of arginase activity was performed by specifying an increase in the amount of urea (the reaction product) (14). One unit (U) of enzymatic activity was defined as µmol of the product formed per hour at 37 ºC. The results are presented as U/mg protein.

Results

A statistically significant increase was detected in the liver arginase activity of control rats exposed to Aroclor 1254 compared to control group (p<0.05). Vitamin E administered simultaneously with Aroclor enabled the liver arginase activity of control rats to approach statistically normal values (Table 1). A statistically insignificant increase was determined in arginase activity of pregnant rats and offsprings. While vitamin E administered simultaneously with Aroclor was ineffective in pregnant rats, it caused a statistically significant increase in offsprings A1254 + vitamin E group compared to offsprings control group (p<0.05) (Tables 1 and 2).

Table 1. Liver arginase activities of adult and pregnant rats administered Aroclor 1254 (A1254) (2 mg/kg/day) and vitamin E (50 mg/kg/day).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Control</th>
<th>A1254</th>
<th>A1254 + Vitamin E</th>
<th>Pregnant Control</th>
<th>Pregnant A1254</th>
<th>Pregnant A1254 + Vitamin E</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arginase (U/mg protein)</td>
<td>240.09±9.57&lt;sup&gt;a&lt;/sup&gt;</td>
<td>365.47±25.14&lt;sup&gt;b&lt;/sup&gt;</td>
<td>289.48±11.70&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>270.79±63.15&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>292.31±31.52&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>288.70±28.53&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>P&lt;0.05</td>
</tr>
</tbody>
</table>

<sup>a, b</sup>: The difference between means that had different letters in the same line was statistically significant.
Values are means ± SEM.
<sup>a, b</sup>: Ayni sahərdakı farklı harflerin ortalamaları arasındakı farklılık istatistiksel olarak anlamlıdır.
Değerler, ortalama ± standart hata şeklindedir.
Persistent organic pollutants (POPs) are known to affect homeostasis and allostatics, which is the maintenance of a stable physiology. This is generally important for vital organ-system functioning as well as survival and reproduction in vertebrate species (29). The liver is the main target organ for PCBs in the body as they are detoxified here (22). Several studies indicate that PCBs cause hepatotoxicity and carcinogenicity as well as impairment in hepatic function (17, 18, 23). In this study, a statistically significant increase was determined in the liver arginase activity of control rats exposed to Aroclor 1254 (p<0.05). Arginase catalyses the hydrolysis of L-arginine to L-ornithine and urea. Also, some studies (8, 11, 20) have revealed an increase in serum and urinary urea levels in PCB-exposed rats. The study conducted by Ebner and Couri (10) demonstrated that Aroclor 1254 exposure poised hepatic mitochondria toward the synthesis of urea intermediates such as carbamoyl phosphate, citrulline. However, arginase produces L-ornithine which acts as a biosynthetic precursor for proline, glutamate, and polyamines such as putrescine, spermine and spermidine. Polyamines are required for cell division, growth, and differentiation (3). The increase in the liver arginase activity of control rats exposed to Aroclor 1254 may result in an increase in cellular concentration of polyamines and also an increase in cell proliferation rate. With regard to all above–mentioned aspects, the increased arginase activity due to Aroclor 1254 may be important at the carcinogenicity caused by PCBs.

A statistically insignificant increase was found in the arginase activity of pregnant rats and offsprings. During pregnancy, the hepatic formation of urea is depressed (24) which causes a reduced urinary excretion (16). The preferential use of α-amino acids for foetal protein synthesis would probably lead to reduce protein catabolism and nitrogen excretion in the maternal organism. This metabolic particularity would result from a down regulation of arginase or from a preferential alfa-amino acids used for both growth of foetus in pregnant females and growth of their offsprings.

Vitamin E provides homeostasis in living cells (13). Some studies have proved that there were some relationships between vitamin E and arginase. Park and Tappel (25) reported that rats fed with a vitamin E supplemented diet for 40 days had lower liver arginase activity than those fed with a diet not containing vitamin E. Moreover, administration of dietary vitamin E caused a significant decrease in the liver arginase activity increased due to high doses of prednisolone in rats (12). We found in this study that administering vitamin E on rats simultaneously with Aroclor 1254 caused a decrease in liver arginase activity compared to the Aroclor 1254 group.

When we analyzed the effect of administering Aroclor 1254 and vitamin E simultaneously on the liver arginase enzyme activity in offsprings, interestingly it was determined that there was a statistically significant increase in the arginase enzyme activity of offspring A1254 + vitamin E group compared to the offspring control group. It is known that vitamin E is stored in placenta however its transfer to foetus is limited (9). The effect of vitamin E on arginase enzyme activity of offspring A1254 + vitamin E group may not have been detected, since the amount of administered vitamin E dose that is transferred to foetus was insufficient compared to the effect of Aroclor 1254.

Consequently, a statistically significant increase was detected in the liver arginase activity of control rats exposed to Aroclor 1254 in this study and administering vitamin E simultaneously with Aroclor 1254 was found to cause a decrease in liver arginase activity compared to the Aroclor 1254 group. We recommend that addition of vitamin E can prevent the increase in liver arginase activity caused by Aroclor 1254.

Discussion and Conclusion

Persistent organic pollutants (POPs) are known to affect homeostasis and allostatics, which is the maintenance of a stable physiology. This is generally important for vital organ-system functioning as well as survival and reproduction in vertebrate species (29). The liver is the main target organ for PCBs in the body as they are detoxified here (22). Several studies indicate that PCBs cause hepatotoxicity and carcinogenicity as well as impairment in hepatic function (17, 18, 23). In this study, a statistically significant increase was determined in the liver arginase activity of control rats exposed to Aroclor 1254 (p<0.05). Arginase catalyses the hydrolysis of L-arginine to L-ornithine and urea. Also, some studies (8, 11, 20) have revealed an increase in serum and urinary urea levels in PCB-exposed rats. The study conducted by Ebner and Couri (10) demonstrated that Aroclor 1254 exposure poised hepatic mitochondria toward the synthesis of urea intermediates such as carbamoyl phosphate, citrulline. However, arginase produces L-ornithine which acts as a biosynthetic precursor for proline, glutamate, and polyamines such as putrescine, spermine and spermidine. Polyamines are required for cell division, growth, and differentiation (3). The increase in the liver arginase activity of control rats exposed to Aroclor 1254 may result in an increase in cellular concentration of polyamines and also an increase in cell proliferation rate. With regard to all above–mentioned aspects, the increased arginase activity due to Aroclor 1254 may be important at the carcinogenicity caused by PCBs.

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