Effects of Diclofenac Sodium on the Rat Liver in Postnatal Period

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Abstract: Diclofenac Sodium (DS) is a nonsteroidal anti-inflammatory drug used in painful conditions, trauma, dysmenorrheoa, rheumatoid arthritis, degenerative joint disease and ankylosing spondylitis. There is the paucity of the research investigating DS effect on the rat liver in the fetal period. The present study aimed to investigate the possible postnatal effects of DS on the liver tissues of the offspring of rats. DS was administrated to pregnant rats for 15 days from the 5th-20th day of pregnancy. After mating day, pregnant female rats were separated into the control and DS treated groups. DS (1 mg kg⁻¹ daily) was injected intraperitoneally to the drug-treated group and physiological saline (1 mL kg⁻¹) to the control group. Liver tissues were collected from pups at the 20th week and paraffin sections were dyed with hematoxylen-eosin. DS caused proliferation of bile ducts, enlargement of perportal area, dilatation of sinusoids and the central vein, parenchymal degeneration and mononuclear cell infiltration. Significant changes between the control and the treated groups were found (p<0.05). However, no significant difference was observed between the sexes in both groups. This study using DS with low dose did not show parenchymal cell death, as a different finding from the previous studies. However, these observations need to be confirmed by further studies using different doses of DS during pregnancy.

Key words: Diclofenac sodium, pregnancy, rat, liver, fetus

INTRODUCTION

Nonsteroidal Anti-Inflammatory Drugs (NSAIDs) have been widely used for many years in palliation of pain and decreasing inflammation and fever throughout the world (Simon, 1994). Diclofenac (sodium-[o-[2,6-dichlorophenyl]-amino[phenyl]-acetate) (DS) is an NSAID and the treatment choice for alleviation of fever and inflammation in rheumatoid arthritis, degenerative joint disease and ankylosing spondylitis. Nonsteroidal anti-inflammatory drugs are also used in painful conditions, trauma, dysmenorrheoa, dental medicine and in the treatment of pain resulting from minor surgery (Brogden et al., 1980; Machiko et al., 2001; Power et al., 2007).

DS inhibits Cyclo-Oxygenase (COX) and hence, the Prostaglandin (PG) production (Vane, 1971), which has the potential to cause negative maternal and fetal effects during pregnancy. Maternal effects include prolongation of pregnancy and labour. DS also causes the ductus arteriosus constriction, renal dysfunction and hemostatic abnormalities in the fetus and neonates. DS passes through the placental barrier and shows its effect on fetus (Ostensen, 1998). NSAIDs also affect central nervous development and compromise the neural activity, such as the impairment of sciatic nerve morphology (Canaan et al., 2008). Diclofenac is mainly metabolized in the liver (Castel et al., 1997) and causes liver damage of the mammalian and avian species (Reckly et al., 2006). DS is also used widely in the women of child-bearing age for the treatment of common gynecological problems. On the other hand, the toxicity of diclofenac has been confirmed on rat embryos (Chan et al., 2001). There is the paucity of the work investigating DS effect on the rat liver in the fetal period. In the pups of the rats treated with DS during pregnancy, 4 weeks after birth, significant morphological changes were observed in the liver (Gökçimen et al., 2001). The aim of the present study was to examine the postnatal hepatic effects of DS administered during pregnancy.

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MATERIALS AND METHODS

This study was approved by the Animal Use Ethics Commission from Yuzuncu Yil University and all procedures were performed according to the Animal Experimentation Ethics Committee.

Animals and experimental procedures: In the present study, male and female Wistar rats (150-200 g) were used. Animals were grouped into the control and diclofenac sodium group. Each group was separately left to mating in the same day. After observing the vaginal plug produced by male vesicular and coagulating gland secretion in the next day, the animals were accepted as pregnant rats (Ragheb et al., 2007). Pregnant animals were kept in a standard plastic cage on sawdust bedding in an air-conditioned room (20°C), under a 12/12 h light/dark cycle and fed ad libitum.

After mating day, pregnant rats from the control group (n= 20) received physiologic saline (1 mL kg⁻¹) and the rats of the treatment group (n= 20) were injected with DS (1 mg kg⁻¹) for 15 days from the 5-20th day of pregnancy. From each group, 12 (6 male, 6 female) offspring were chosen randomly after delivery and housed for 20 weeks. On the 21st day after the birth, under anesthesia, intracardiac perfusion was performed with 0.9% saline and 10% formalin from a total of 24 offspring. Tissue samples were processed by graded alcohol and xylene and then, embedded in paraffin blocks. Tissue sections (4 μm) were taken and stained with hematoxylin and eosin for histological examination. The preparations were examined with a light microscope (Zeiss axioskop 40). Observations concerning bile duct proliferation, mononuclear cell infiltration, enlargement in the periporal area, sinusoidal dilatation, vena centralis dilatation and parenchymal degeneration were evaluated by the researchers of this study and a pathology expert. The evaluation of the changes mentioned above was made according to the method described by Gökmen et al. (2001). Briefly, normal appearance (+) mild = 0-25% degeneration (+++) moderate = 25-50% degeneration (+++) severe = 50-75% degeneration (++++) very severe = >75% degeneration (Table 1).

Table 1: Hepatic changes found in the control and the treated groups (DS).

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Control</th>
<th>DS</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bile duct proliferation</td>
<td>-</td>
<td>++</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Enlargement of periporal area</td>
<td>-</td>
<td>++</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Sinusoidal dilatation</td>
<td>+</td>
<td>+++</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Vena centralis dilatation</td>
<td>+</td>
<td>+++</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Parenchymal degeneration</td>
<td>-</td>
<td>++</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Mononuclear cell infiltration</td>
<td>-</td>
<td>++</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Fibrosis</td>
<td>-</td>
<td>-</td>
<td>&gt;0.050</td>
</tr>
</tbody>
</table>

Histopathologic assessment of the experimental parameters were graded as follows: (-) showing no changes (+) (+++) indicating mild, moderate and severe changes, respectively. The number of specified parameters were counted under the light microscope (n = number of observations).

Statistical analysis: The data from the present study was compared for two groups and sexes by a statistician. Mann-Whitney U-test for comparison was used, converting the finding to numeric counterpart.

RESULTS

There was full agreement on yes/no choice among the observers. Significant changes between the control and the treated groups were found in terms of pathological changes in present study (p<0.05) (Fig. 1-3). However, no significant changes were observed between

Fig. 1: Histologic structure of the liver of a rat from the control group (HE ×20)

Fig. 2: a-c) Bile duct proliferation of portal area (+Φ), dilatation of vena centralis (+Φ) and sinusoidal dilatation (ΦΦΦΦΦΦ in liver of a rat from the DS group (HE ×40)
the sexes in both groups. The male and female rats were equally affected by DS in the treated group. With the little change, the same appearance was observed between the sexes in the controls. Briefly, no significant difference were observed between the sexes in both the control and DS-treated group for bile duct proliferation, mononuclear cell infiltration, enlargement in the portal area, sinusoidal dilatation, vena centralis dilatation and parenchymal degeneration.

Mild to moderate enlargement in the portal area and mononuclear cell infiltration (p<0.001), mild to severe changes in bile duct proliferation (p<0.001), moderate to severe changes in sinusoidal dilatation and vena centralis dilatation, (p<0.001) and mild changes in parenchymal degeneration were observed in both male and female rats treated with DS compared to the controls. Mild changes in sinusoidal dilatation, vena centralis dilatation and parenchymal degeneration from the control group were seen in both sexes (Table 1).

DISCUSSION

Diclofenac sodium as a Nonsteroidal Anti-inflammatory Drug (NSAIDs) is one of the safest agents of its kind for the treatment of a broad range of rheumatic conditions (Willkens, 1985). On the other hand, NSAIDs are contraindicated during pregnancy because of their teratogenic potential. Nonsteroidal anti-inflammatory drugs given during pregnancy prolong the pregnancy and labour, inhibiting cyclo-oxygenase (Ostensen, 1998). All Non-Steroidal Anti-Inflammatory Drugs (NSAIDs) are prostaglandin inhibitors, which explain their fotal toxicity (Beavo, 1992). These drugs are either avoided or given in a low dose in the first trimester of pregnancy.

The present study showed significant morphologic changes in the liver of the 20 weeks old offspring of the DS-administrated rats. In rats, diclofenac decreases hepatic ATP content and induces hepatocyte apoptosis (Boc et al., 1999). The adverse gastrointestinal effects and hepatic toxicity were also observed in pregnant rat administrated orally with non-selective COX inhibitors (Ersad et al., 2004). Gokcen et al. (2001) reported histopathologic changes including enlargement in the perportal area, pyknosis in the nucleus of hepatocytes and vascular degeneration in parenchymal cells of the 4 weeks old pups of DS-treated rats. Although, these findings are generally similar to those of this study, pyknosis was not observed in the present study in postnatal 20th week. The occurrence of the pyknosis in the liver of 4-weeks old offspring was attributed directly to the influence of DS metabolites (Gokcen et al., 2001). We are unable to give commentary for the absence of the pyknosis in DS-treated group. However, if the pyknosis has taken place, a possable improvement of the parenchymal cell might occur in the postnatal period between 4 and 20th week.

These findings clearly demonstrate that a 15-days maternal administration of DS during pregnancy has significant negative effects on the developing liver compared to control, such as mild to moderate regressive structural changes. This study also showed that gestation period was significantly prolonged in DS-treated rats, similar to Kokcu et al. (1992) and Gokcen et al. (2001). With light microscopy, no significant alterations were observed in this control group, 20-weeks-old rats. Because of the parenchymal degeneration, however, some of the hepatocytes have the appearance of clear swelling and fuzzy in the present study (Fig. 2 and 3).

DS appears to have some negative effect on both development and differentiation of liver cells. After partial resection or massive injury, unlike other organs, the mammalian liver can rapidly regenerate and regain its original size, structure and function (Chung et al., 2006).

The toxic effect of diclofenac on hepatocytes may be caused by drug-induced mitochondrial impairment, together with a futile consumption of NADH (Boc et al., 1999). Recent studies using rat liver mitochondria and freshly isolated rat hepatocytes showed that diclofenac decreased hepatic ATP content and induced hepatocyte apoptosis (Chung et al., 2006).

CONCLUSION

We found that DS treatment during pregnancy caused mild to moderate bile duct proliferation and mononuclear cell infiltration and enlargement in the perportal area in 20 weeks-old female and male rats. This study did not find pyknosis, which is a different finding from that of previous studies and suggests that DS does not cause a parenchymal cell death in low doses.
However, these observations should be confirmed by further studies using different doses of DS during pregnancy.

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REFERENCES


