The effect of warfarin on skeleton development of fetuses’ rats
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ABSTRACT
The pregnancy period is characterized by the critical time of the pregnant mother’s exposure to external impacts, which are most severe during the so-called embryonic period. The study aimed toward study the effects of warfarin in the developing rat fetus. Three-month adult virgin female Wistar albino rats (Rattus norvegicus) were used. They are Left to adapt for one week and then isolated with males by two females with one male for mating determined the zero-gestation day. Pregnant females were arbitrarily divided into three equals with eight rats for each. The first group is control kept without any treatment The second and third groups take 0.1 or 0.05 mg/kg B.t of warfarin respectively, daily from (zero - 15th) day of gestation. The Collection of the blood samples from heart of pregnant rats for biochemical study via using Kit of ELISA technique to estimation the concentration of Osteocalcin and matrix Gla protein. The pregnant animals were sacrificed at (16th) day of gestation The fetuses’ specimens were collected for histological processing. Major anomalies were observed on incomplete ossification of the hind leg and wavy ribs by using double stained Alizarine red and Alcian blue. The histological examination of the vertebral column in the group of treated fetuses reveals degeneration at the vertebrae, damage to the intervertebral tissue, hemorrhage and necrosis. They are a significant decrease in the Matrix Gla protein and Osteocalcin in serum of pregnant rats.

Keywords: Warfarin, osteocalcin, MGP, skeletal abnormalities

INTRODUCTION
The anticoagulant rodenticides are a substance that prevent the vitamin K-dependent steps in the coagulation cascade in the liver, like warfarin, causing hemorrhage and death by interfering with normal blood clotting, the first-generation materials are not sufficiently toxic to reason death after a single nourishing while the Second products have a better affinity to binding the places in the vertebrate, liver and consequently show greater accumulation and perseverance in the body 1. The contact to the low levels of anticoagulants may pathological or behavioral effects, in most cases the animals appeared weak, have a movement with a slow reaction and more susceptible to the infection or accidents, the second-generation anticoagulants have a higher potency than first-generation anticoagulants is related to accumulation and slow elimination from the body after absorption, the anticoagulants accumulate in the liver which is the target organ for their action2. The warfarin is the most widely used as oral anticoagulant (OAC) in worldwide for the prevention of thromboembolic events in the great risk patients a heart attack3.
Previous studies revealed that the patient who attempted a suicide while taking a high dose of warfarin was successfully treated with blood transfusion and vitamin K injection without any complications. Currently, 0.5–1.5% of the world's population receives the medication on an annual basis 4. The Osteocalcin is the most abundant non-collagenous protein that is found in the bone and is comprised of 49 amino acids (AA) 5. Hauschka discovered Osteocalcin in bone, it has a highly conserved sequence among the vertebrates and binding with calcium to form hydroxyapatite crystals6. The matrix Gla protein (MGP) consist of 84 amino acid residues, it belongs part of the vitamin K-dependent Glα protein family, it’s a small, highly conserved protein and produced predominantly by chondrocytes and vascular smooth muscle cells. The matrix Gla protein was recognized the first time in 1983 in bovine bone through Price and co-workers. it was the second Glα protein isolated from bone and demonstrated strongly associated with the collagenous bone matrix7. The osteocalcin and matrix Glα protein biomarkers can give us a guide to examination pathological change which effect on the bone’s development 8. The study has given a new insight of the effect of warfarin on the level of osteocalcin and MGP and in serum of pregnant mothers that treated by warfarin during a gestation and effect of fetuses’ development.

MATERIALS AND METHODS
Three-month adult virgin female Wistar albino rats (Rattus norvegicus) were used. The average weighing is about (275 ± 25) g they were bred in the college's animal house Veterinary Medicine Universities of Basrah. These females were scanned periodically by the vaginal smear technique, to ensure that they were always in regular estrous. They are Left to adapt for one week and then isolated with males by two females with one male for mating determined the zero-gestation day 9. Pregnant females were arbitrarily divided into three equals with eight rats for each. The first group is control kept without any treatment. The second and third groups take 0.1 or 0.05 mg/kg B. of warfarin respectively, daily from (zero - 15th) day of gestation (organogenesis period). Depending on the half-lethal dose of warfarin (58 mg/kg) 10. The two different doses of Warfarin (low and high): were made. The pregnant animals were sacrificed at (16th) day of gestation under general anesthesia of a mixture of Xylazine (90 mg/kg) and Ketamine (10 mg/kg) 11. The Collection of the blood sample from heart of pregnant rats were placed in a covered gel tube then, the gel tube centrifuged at 3,000 rpm for 10 minutes 12. The serum directly transferred into Eppendorf tube till it was used in a biochemical study via using Kit of ELISA technique to estimation the concentration of Osteocalcin and matrix Glα protein (Depending on the manufacturer instructions, USA Al-Shkairate institution). The fetuses’ specimens were directly place in neutral buffered formalin fixative (10%) for (24-48) hrs. for histological processing. The tissues section was stained with eosin and hematoxylin13. The double staining technique proposed by Kimmel and Trammel, the skin of fetuses was removed and fetuses’ specimens were directly put for at least 24 hr. in the acid staining solution (pH 5.2.8) at room temperature and then dehydrated in 96% of ethanol for at least 6 hrs. Soaking of soft tissues was performed by putting specimens in the basic staining solution for 30 hrs. at room temperature, whereas renewing the solution at least three times. Clearing and hardening was performed by placing specimens in the cleaning solution for at least 8 hr. keeping of double-stained fetuses was performed in a 1:1 ethanol 70% and glycerin mixture14.

Statistical analysis
Data were given as mean ± standard error of the mean. The litter was regarded as the experimental unit of comparison for all group, effects of warfarin on fetal growth and effected MGP and Osteocalcin in serum of pregnant rats were determined by one way ANOVA.

RESULTS
Double Staining (Whole mount)
The effect of warfarin on skeleton development of fetuses’ rats

**FIG 1:** photomicrography of two rat fetuses (16th day) of gestation, (B) is control, (A) fetus obtained from rat treated orally with high dose of warfarin0.1/kg b. wt. during the organogenesis period showing incomplete ossification of hind limb and wavy ribs.

**FIGURE 2:** fetus obtained from a rat treated orally with warfarin 0.01mg/kg b. wt. during the organogenesis period showing the areas without ossification and no stained cartilage

**Microscopic study of histopathological changes of fetus**

**FIG 3:** photomicrography of vertebra column of fetus 16th days age of control group shows normal vertebral structure (back arrow), normal intervertebral tissue (blue arrow) H&E 125X arrow) H&E 125X
The effect of warfarin on skeleton development of fetuses’ rats

Estimation of biomarker Osteocalcin in the serum of pregnant rats treated with warfarin

TABLE 1: The mean value of Osteocalcin at 16 days of gestation.

<table>
<thead>
<tr>
<th>Name of group</th>
<th>Mean ± S.D of Osteocalcin (16 day)/(pg/ml) N=8</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>8.64± 0.71</td>
</tr>
<tr>
<td>Low Dose(0.05mg/kg)</td>
<td>7.35± 0.45</td>
</tr>
<tr>
<td>High Dose(0.1mg/kg)</td>
<td>7.60± 0.33</td>
</tr>
</tbody>
</table>

The small letter (a, b) refers to significant value at level of P≤0.05.
TABLE 2: The mean value of Matrix Gla protein at 16 days of gestation.

<table>
<thead>
<tr>
<th>Name of group</th>
<th>Mean± S.D of MGP (16 day)/(pg/ml) n=8</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>1.07a ± 0.12</td>
</tr>
<tr>
<td>Low Dose (0.05mg/kg)</td>
<td>0.40b ± 0.40</td>
</tr>
<tr>
<td>High Dose (0.1mg/kg)</td>
<td>0.42b ± 0.42</td>
</tr>
</tbody>
</table>

The small letter (a) refers to significant value at level of P≤0.05

DISCUSSION
This study demonstrated the congenital malformations and teratogenicity of the warfarin in the fetuses obtained from a mother treated orally with warfarin through the organogenesis period show large incomplete ossification of the skull, large opened fontanel, sacral and coccygeal vertebrae, metatarsal bones, coccygeal and sacral vertebrae, incomplete ossification of hind limb and wavy ribs these results agree with Finkelstein et al.15. These changes occur because of warfarin may be impaired formation synthesis of proteins crucial for bone and cartilage in fetal which result warfarin embryopathy16. A number of fetuses show areas without ossification and no stained cartilage, while other show is as stained with blue indicate cartilage formation. The results transverse section in the skull of rat fetuses, obtained from treated group of warfarin 0.1mg/kg B. Wt. through the organogenesis period time noticed no calcification happen and incomplete development of bone and the Importance of Half-life described the congenital anomalies after exposure to the warfarin in human include congenital anomalies in bones and hemorrhage complications, these changes occur due to warfarin may be impaired the synthesis of proteins important for bones and cartilage in the fetal. The vertebral column of the fetus on the (16 day) of gestation in control group noticed normal vertebral structure and normal intervertebral tissue, Whereas the sections of vertebral column of fetus which treated with the warfarin revealed hemorrhage and moderate degeneration at the intervertebral tissue, as well as, column of fetus 16 day of gestation treated with high dose of warfarin appears a marked Necrosis and degeneration at the vertebrae, these similar to Li et al 18. The changes in vertebra column of fetus may be because of the role of warfarin to inhibition vitamin K regeneration by inhibiting the necessary enzyme epoxide reductase 19. There is a significant decrease in the Matrix Gla protein in the serum of the pregnant rats in both treated group with low doses of warfarin and high doses of warfarin in both (16th) day gestation at level (p≤ 0.05) compared with the control group, our study agrees with Elango et al 20.by which the warfarin reduces gamma-carboxylation of MGP, which may be act as a central role in abnormal development of fetuses. There is a significant decrease in the osteocalcin at the level (p≤ 0.05) in the serum of treated group with low dose of warfarin and the groups of the pregnant rats treated with High dose of warfarin compared with the control group, our study agrees with Sugiyama et al 21. This study shows the effect of Warfarin on fracture risk when the patient was giving a high warfarin dose. The effect of warfarin may block peripheral vitamin K activity, so inhibiting the γ-carboxylation of osteocalcin. The endogenous change of vitamin K1 to menaquinone-4 may be contribute to the direct effects on osteocalcin level 22. There are new visions into the pathogenesis of the fetal anomalies related with the prenatal warfarin exposure, warfarin is vitamin K antagonists prevent the recycling of vitamin K in the cell, bones, cartilages, and the developing CNS and vitamin K – dependent proteins have been recognized 23. Animal studies established an effect of warfarin on the developing of bone 24,25.

CONCLUSION
The results suggest the role of Warfarin on the inhibition of MGP and osteocalcin in blood and tissue.
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