Double Staining in Rats (Mini Review)

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ABSTRACT

The double skeleton staining method is among the methods which are particularly used on rats. In the literature, studies take place titled as the examination of the effects on fetus bones by giving various substances on pregnant rats. Two different types of ossification are described in the embryonic period called as intramembranous ossification and endochondral ossification. Different staining methods are used to determine this ossification. Studies in which binary skeletal staining have been used are formed in two groups as teratogenic studies and developmental studies.

Keywords: Double Staining; Ossification; Rat; Bone Mineral Density; Fetus

Introduction

Double skeleton staining method is among the methods which is used in the studies of literature. Generally, examination of the possible effect on fetus bones in pregnant rats is included in the studies which have been carried out [1]. The process of staining the bones and cartilage, which form the skeleton, with different stains in different colors or in different shades of the same colour is called double skeleton staining. The studies in which double skeleton staining have been used are collected in two groups as teratogenic studies and developmental studies. In teratogenic studies, the effects of various chemical substances or physical factors on the skeletal system and the fetus of experimental animals preferred in teratogenic studies are examined in the skeleton system by double skeletal staining at different stages of their development [2]. In developmental studies, double skeleton staining is performed on human embryos and fetus obtained as a result of abortus and the relation between bone and cartilage areas in different bones and age of fetus, ossification in other bones, etc.is examined [3,4]. Two different types of ossification, called intramembranous ossification and endochondral ossification, are defined in the embryonic period [1,5]. Intramembranous ossification: This type of ossification usually observed in flat bones. In areas where intramembranous ossification will occur; mesenchyme intensifies and begins to vascularize. There is no regular structure in the first formed bone. Bone, which are organized in time, gain a structure of lamellae. The lamellae structure formed around the blood vessels, while creating the Havers systems, the osteoblasts around the developing bone structures constitutes compact bone layers [6,7].

Endochondral Ossification

In this type of ossification, a cartilage model is formed primarily in the area where the bone will form. Then this cartilage model is eliminated by replacing the bone tissue. With this type of ossification, long and short bones are formed. In both types of ossification, the first bone tissue is the primary or immature bone tissue. Then the Primary bone tissue is replaced by secondary bone tissue [8]. Skeletal development in rats: At the end of the 9th day of embryonic life or in the beginning of the 10th day in rats, somit development starts. At the end of the 11th day there 25, on the 13th day 45 and on the 16th day 65 somit appear. From the 16th day onwards, the first ossification centers occur. The first ossification centers are seen in the clavicula, ribs and some bones of the skull. On the 16th day, the spaces of the hip, knee, shoulder and elbow joints begin to become clear and the vessels anlage fuse. On the 17th day, the ossification is disseminated to the extremity buds and other bones of the skull while the spaces of the wrist and ankle joints become evident. On the 18th day the head grows rapidly and the palate fuses. On the 18th day the head grows rapidly and the palate fuses. On the 19th and 20th days the spaces of the finger joints of the anterior and posterior extremities are seen. Birth occurs on the 22nd day in rats [7-10]. In a new born rat, the trunk and diaphysis of the long
bones are largely ossified, but the epiphyses are still in the cartilage structure. Most secondary ossification centers continue to exist in the days following birth [11-13]. Rat skeletal system: The skeletal system are examined by dividing into two parts as skeleton axiale and skeleton appendicular [14,15]. 1. Skeleton axiale: Columna vertebralis, which forms the body axis, consists of vertebralis, rib cage and skull bones. Skull bones: consist of a total of 28 bones; some of them are single while some of them are paired [10-14]. Spine: The rat spine consists of 7 cervical, 13 thoracic, 6 lumbal, 4 sacral and caudal (tail) vertebrae ranging from 25 to 30 [16,17].

Method

Double Staining Process

In order to stain the skeletal system of fetus with double skeleton staining method; all fetus is kept in 70% ethyl alcohol for 4-7 days and the water is provided to be absorbed. Following this process, they are kept in pure acetone for 1-3 days and they are purified from fats. Then the skin is removed. During all of these procedures, the number of experimental animals should be kept high in case some fetus has limb ruptures and bone tissue injuries. Damaged subjects are not stained. By peeling off the skin and removing the internal organs and the eyes for staining, the staining process of the skeleton of the fetus and the pups is carried out in accordance with the protocol described below in Table 1. Preparation of Stain Solution: Stain solution is obtained by combining the solutions prepared separately with specific methods.

Preparation of solutions:

a) Solution 1: It is obtained by stirring and melting 300 mg of Alcian Blue constantly in 100ml of 70% ethyl alcohol.

b) Solution 2: It is obtained by stirring and melting 100 mg Alizarin Red-S constantly in 100ml of 95% ethyl alcohol.

c) Solution 3: It is obtained by adding the second solution gradually into the first solution which has been prepared beforehand. After complete mixing of these two solutions, 100 ml of Glacial Acetic Acid is added to this mixture.

d) Solution 4: It is prepared by adding 1700 ml of 70% ethyl alcohol into the prepared third solution.

Table 1: Double Skeletal System Staining Method.

| Technical Stages | Solutions | Time          |
|------------------|-----------|---------------|
| Fixation         | 70% ethyl alcohol | 4-7 days     |
| Degreasing       | Pure acetone           | 1-3 days     |
| Double Staining  | preparing the double staining solution | 7 days | 38-40°C incubated |
|                   | 1st solution: 300mg Alcian Blue + 100ml 70% ethyl alcohol |  |  |
|                   | 2nd solution: 100mg Alizarin Red S + 100ml 95% ethyl alcohol |  |  |
|                   | 3rd solution: 1st solution + 2nd solution + 100ml Glacial acetic acid |  |  |
|                   | 4th solution: Prepared by adding 1700 ml of 70% ethyl alcohol to the first three solutions. |  |  |
| Transparency      | 1) 1% KOH | 1-3 days | 5-7 days | 5-7 days |
|                   | 2) 1% KOH (80 ml) +50% glycerin (20 ml) |  |  |
|                   | 3) 1% KOH (50 ml) +50% glycerin (50 ml) |  |  |
|                   | 4) 1% KOH (20 ml) +80% glycerin (80 ml) |  |  |
| Safekeeping       | 100% Pure acetone |  |  |  |

Staining Process

The tissues of the fetus and pups are provided to be stained in the stain solution prepared according to the method described above, by being kept in the oven at 38-40°C for 7 days. At the end of this period, the fetus is washed under running tap water for 2 hours to have them reacted with water. Then, the process of transparency proceeds. Transparency phase: It consists of 4 stages.

a) Stage 1: The stained fetus is soaked in 1% KOH for 1 day.

b) Step 2: For 5 days, they are soaked in the mixture of 80 ml of 1% KOH and 20 ml of 20% glycerin.

c) Step 3: For 5 days, they are soaked in the mixture of 50 ml of 1% KOH and 50 ml of 50% glycerin.

d) Step 4: They are soaked in 20cc 1% KOH and 80 ml of 80% glycerin solution in for 5-7 days.

After these stages, the bone cartilage ratio is calculated in Alcian Blue, and Alizarin Red S-stained fetus bones (Figure 1).
Discussion

In many studies conducted in the literature, different stains have been used in double staining methods and the aim is to provide different colours of bone and cartilage. Teratogenic studies are usually available on pregnant Rats Yilmaz 2018; Güneş 2011; Tizabi 2000. In their studies, the pregnant rats have been given different doses of nicotine. In their study, they reported the variability of bone and cartilage ratios in fetus by using double skeleton staining method [1,18,19]. Yılmaz (2018) investigated the protective effects of melatonin against the effects of nicotine on bones in their study; and they used double staining method in their study. They found that the percentage and rate of ossification was low in the upper and lower extremity bones of the group [1]. Kurtoglu (2007) investigated the negative effects of nicotine on bone development. In their study they used double staining method and they administered 3 mg/kg nicotine given i.p during pregnancy and lactation [20]. Hu (2018) evaluated the osteoclastogenesis and endochondral ossification of long bones of fetuses in pregnant rats exposed to nicotine they used double staining skeletal method [21]. In their study, Tomaszewski et al, reported with the double staining method that they had common abnormalities related to the most deteriorated mineralization of bone tissue in fetus of pregnant rats which have been given caffeine [22].

Conclusion

One of the methods used in many studies is the double skeleton staining method. When literature is examined, many publications in this area especially in determining teratogenic effect can be seen. In many of our studies, we have used the double skeleton staining method. We believe that this study will be instructive by contributing the literature in terms of specifying the method more dearly.

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