ABSTRACT

OBJECTIVE: To study neuronal damage in brains of first- and second-generation pups, born to hypothyroid Wistar rats.

METHODS: Ten female adult Wistar rats, randomized into two equal groups- propylthiouracil (Group-P) and control (Group-C) were allowed to conceive. Group-P was given oral propylthiouracil throughout gestational period and weaning until 22nd day. Nine offspring from group-C (C/FC) and group-P (P/FC) were sacrificed on 22nd day of life to collect blood and brain samples. The dams of group-P (group-PP) were allowed to conceive and continued with propylthiouracil treatment throughout gestation and weaning. Their nine offsprings (PP/FC) were sacrificed on 22nd day of life. Serum levels of triiodothyronine (T₃), thyroxine (T₄) and thyroid stimulating hormone (TSH) were measured. Brain weight, apoptotic cell count (ACC) of Purkinje cells of cerebellar cortex and Pyramidal cells of hippocampus were documented.

RESULTS: Mean TSH levels were 21±1.3ng/dl, 23.3±3.5ng/dl and 10±2.1ng/dl in P/FC, PP/FC and C/FC respectively (p=0.003). Mean T₃, was 31.7±1.2ng/dl, 30.3±1.3ng/dl and 36.3±0.9ng/dl in P/FC, PP/FC and C/FC respectively (p=0.030). Mean brain weights was 1.21±0.21 mg, 1.20±0.41 mg and 1.42±0.01 mg group in P/FC, PP/FC and C/FC respectively (p>0.05). The normal Pyramidal and Purkinje cell count was low in P/FC and PP/FC groups compared to C/FC group (p<0.05). The ACC of Pyramidal and Purkinje cells was high in P/FC and PP/FC groups compared to C/FC group (p<0.05).

CONCLUSION: Maternal hypothyroidism adversely affected the morphology of Pyramidal and Purkinje cells by enhancing apoptosis, which increased further in second generation pups.

KEY WORDS: Thyroid Hormones (MeSH); Hypothyroidism (MeSH); Neurons (MeSH); Pregnancy (MeSH); Apoptosis (MeSH); Rats, Wistar (MeSH); Triiodothyronine (MeSH); Thyroxine (MeSH); Purkinje Cells (MeSH); Pyramidal Cells (MeSH).

INTRODUCTION

The thyroid gland synthesizes hormones named thyroxine (T₄) and triiodothyronine (T₃), which regulate growth and functions of almost all the vital organs and structures. Various experimental studies have unanimously proved that the main organ of the body directly targeted by thyroid hormones (THs) is the brain. These hormones are indispensable for the development and functioning of the brain throughout life. Disturbances in the functions of the thyroid gland are one of the most commonly encountered endocrine disorders. Hypothyroidism is the prevailing form of thyroid disease and it is manifested in the form of memory and learning impairment, depression, psychosis, slow motor activity, sleepiness, decline in intellectual ability and rarely coma. Since the thyroid hormones greatly affect the maturation of neurons concerned with learning and memory e.g., the neurons located in hippocampus and frontal cortex, the deficiency of these hormones during the neuronal maturation causes mental retardation, which may be permanent.

During development of brain, THs act by binding to nuclear receptors dispersed extensively in the fetal brain which are present even before the fetal ability to synthesize them. In humans, THs deficiency during brain development can lead to alterations in the structure and functions of cerebral cortex, hippocampus and cerebellar cortex. If hypothyroidism manifests during fetal or early neonatal period, it hinders the process of differentiation and connectivity of neurons. Usually THs deficiency during fetal or early postnatal life results in brain damage, which may be due to abnormal neuronal proliferation and migration. In humans, THs deficiency during the first four weeks after birth leads to cretinism, which manifests as mental retardation and neurological insufficiencies.

Lack of THs leads to extensive apoptosis during neurogenesis. Even subclinical hypothyroidism during early pregnancy can be deleterious to the development of fetal brain. Both the magnitude and the duration of apoptosis is increased in congenital hypothyroidism by suppression of the anti-apoptotic gene Bcl-2 and
upregulation of the pro-apoptotic gene Bax.

Rat pups are born with a comparatively undeveloped brain. The effects of THs on the development of brain in the rat occur in the early neonatal period. The first 2-3 weeks of life is defined as the time period during which THs are required for normal brain development in rodents. This period generally parallels the last trimester in humans. In both humans and rodents, if the hormone replacement therapy is delayed, it will cause structural and functional impairment of neurons permanently.

There remains a gap in knowledge regarding the effect of low TH levels on functions of motor neurons. It has been documented that hypothyroidism causes impairment of brain function, but its direct effects on the process of apoptosis involving motor neurons has not been defined. Considering the critical part played by THs on brain development, the current study investigates the outcome of hypothyroidism in the neonatal rat brains and how it leads to decline in motor functions.

METHODS

This experiment was conducted in the Department of Anatomy, University of Health Sciences, Lahore, from September to December of 2018. This study was approved by the Ethical Committee of University of Health Sciences, Lahore (UHS) and was conducted according to the guidelines for the sacrifice of experimental animals, as laid down in American Veterinary Medical Association (AVMA). After procuring 10 adult female Wistar albino rats from animal house, each animal was weighed and thoroughly evaluated by gross inspection. All the apparently healthy rats were of 12-16 weeks of age with a weight range of 190 – 210 g. They were acclimatized in their allotted cages for one week before the commencement of the study. The rats were maintained in the animal house under controlled environment i.e. well-ventilated room at a temperature of 23±2°C, humidity 55±5% and light and dark cycles of 12 hours each (light on at 8:00 and off at 20:00 hours). They were fed on standard rat diet and water ad libitum.

After a week of acclimatization, the treatment of both the groups was started one week before conception. The appearance of the vaginal plug was considered as the first day of pregnancy. As the period of gestation is around 22-23 days in albino rats, and the experiment ended on 22nd postnatal day in the 2nd generation pups, the total time duration of this experiment was between 15 – 16 weeks.

A week before mating, 10 adult female rats were divided into 2 equal groups. This division was carried out randomly and each rat was tail marked for identification. Five adult females, serving as control group C, consisted of dams in a euthyroid state. Propylthiouracil (PTU) was given orally in a dose of 15mg/kg/day by mixing it in rat chow on daily basis to group P rats. The treatment was continued throughout gestation and weaning. Nine offspring from both the groups were chosen for further study and sacrificed on 22nd postnatal day, after receiving PTU indirectly throughout gestation and lactation. None of the mothers were sacrificed in this experiment.

The dams of P group were continued administration of PTU beyond 22nd day, and once again allowed to conceive around 30th day after delivery. This time they were renamed as PP group. The treatment was continued throughout next pregnancy till the end of 22nd day postnatally. Nine offspring, labelled as PP/FC group, were sacrificed respectively on 22nd postnatal day, after receiving PTU indirectly throughout gestation and lactation.

After euthanization, blood samples were immediately collected, from the cardiac region by inserting the needle under the sternum slightly to the left, to measure serum levels of T3, T4, and TSH using rat specific enzyme linked immunosorbent assay (ELISA) kits manufactured by Elabscience®, Texas, USA, according to manufacturer instructions.

Now the scalp was retracted to reveal the skull and an incision made in the bone along the midline. The brains were then extracted from the skull with the help of a spatula and observed for any gross lesion. There was no significant morphological change seen in any of the brains of the pups belonging to both first and second generation.

For histological examination, brains were fixed in 10% formalin, immediately after sacrifice, for 14-16 hours, dehydrated in ascending grades of alcohol, for 1 hour each and then cleared in xylene. Tissues were then embedded in paraffin wax (56 - 58°C melting point) and 3 μm thick sections were processed further for immune-histochemical (IHC) staining, by using monoclonal anti-Bax antibody as the primary anti-body and Horseradish Peroxidase (HRP) as the secondary antibody. The slides were later counter-stained with Hematoxylin and rinsed in tap water, dehydrated in ascending series of alcohol, cleared in xylene and mounted in DPX to be visualized under the light microscope. The parameters used to label the cells as apoptotic were size of the cell body, fragmentation of the nucleus, extent of Bax staining of the cell nucleus or cytoplasm. Neurons having a deeply stained cytoplasm and

### TABLE I: MEAN SERUM THYROID STIMULATING HORMONE (TSH), TRIIODOTHYRONINE (T3) AND THYROXINE (T4) LEVELS IN 22 DAYS OLD PUPS

| Groups (n=9) | TSH (μg/dL) (Mean±SD) | T3 (μg/dL) (Mean±SD) | T4 (μg/dL) (Mean±SD) |
|----------------|------------------------|-----------------------|----------------------|
| C/FC            | 10 ± 2.1               | 39.7 ± 0.5            | 36.3 ± 0.9           |
| P/FC            | 21 ± 3.7               | 39.3 ± 0.4            | 31.7 ± 1.2           |
| PP/FC           | 23.3 ± 3.5             | 39.3 ± 0.4            | 30.3 ± 1.3           |
| R2              | 68.18%                 | 27.49%                | 50.86%               |
| P value         | 0.003                  | 0.130                 | 0.030                |

Group C/FC: control; Group P/FC: first generation born from PTU treated group; Group PP/FC: second generation born from PTU treated group.
Mean brain weights was 1.21 ± 0.21 mg, 1.20 ± 0.41 mg in P/FC and PP/FC groups respectively as compared to 1.42±0.01 mg in C/FC group (p >0.05). Light microscopic examination of IHC stained slides showed disturbed movement of Pyramidal cells in CA3 area of hippocampus in both P/FC (Figure 1 C) and PP/FC (Figure 1 D) groups, with apoptotic changes observed in their nuclei when compared to C/FC group (Figure 1 B). In case of Purkinje cells of cerebellar cortex, viable neurons observed in C/FC group had lightly stained spherical nuclei (Figure 2 B), whereas apoptotic neurons in P/FC and PP/FC groups showed shrunken and fragmented nuclei (Figure 2 C and D).

The count of normal Pyramidal cells in CA3 region of Hippocampus was 39.7±1.24 in P/FC and (38.0±2.62) in PP/FC groups as compared to 44.7±2.49 in C/FC group (P <0.05). One-way ANOVA was used for calculating the significance of the number of Pyramidal and Purkinje cells within groups.

RESULTS

The animals were examined daily to evaluate their health and growth. Pups belonging to group C/FC were active and healthy and there was no morbidity or mortality recorded among them. But the pups belonging to groups P/FC and PP/FC had a 30-50% mortality recorded. Moreover, they remained listless with decrease in appetite. Mean TSH levels were 21±3 g/dl, 23.3±3.5 g/dl and 10±2.1 g/dl in P/FC, PP/FC and C/FC respectively (p 0.003). Mean T, was 31.7±1.2 g/dl, 30.3±1.3 g/dl and 36.3±0.9 g/dl in P/FC, PP/FC and C/FC respectively (p 0.030) [Table I].

Figure 1: Immunohistochemical Bax stained slides from CA3 area of hippocampus in 22 days old pups. The square in slide A indicates the CA3 area and the slides B, C and D are the magnified CA3 areas of respective groups. B: control C/FC group, where the arrow points to a healthy nucleus. C: First generation PTU treated group P/FC where the arrow points to a Bax stained pyknotic nucleus. D: Second generation PTU treated group PP/FC, where apart from disturbed arrangement of pyramidal cells, the arrow points to a pyknotic Bax stained nucleus. The damage in D is significantly intense compared to B and C (40x magnification).

lightly stained spherical nucleus without any signs of Bax stain inside the nucleus were termed viable. Neurons with a shrunken or fragmented nucleus with visible Bax staining were termed apoptotic. IHC stained slides were used for counting both the normal and apoptotic Pyramidal cells in CA3 area of hippocampus and Purkinje cells in cerebellar cortex. Counting of the cells was done in all squares of the grid, excluding the lower and left border. The cells were counted randomly in each field chosen at the magnification of 100X and the mean was calculated using SPSS version 21.

One-way analysis of variance (ANOVA) was used for group comparisons. P-value ≤ 0.05 was considered as statistically significant. Mean ± SD was calculated for numeric variables (i.e. brain weight of pups, serum levels of T₄, T₃, TSH, cell count of Purkinje and Pyramidal cells) and one-way ANOVA was used for calculating the significance of the number of Pyramidal and Purkinje cells within groups.
the contrary, the count of apoptotic Pyramidal cells was 2.7±0.94 in P/FC group and 3.7±0.36 in PP/FC as compared to 2.0±0.81 in C/FC group (P < 0.05) [Figure 3].

Similar results were seen in Purkinje cells of cerebellar cortex, where the count normal cells was 10.3±1.24 and 9.2±1.36 in P/FC and PP/FC groups as compared to 12.2 ± 1.63 in C/FC group (p <0.05). On the contrary, the mean count of apoptotic Purkinje cells was 2.7±0.94 in first generation (P/FC group) and 3.7±0.36 in second generation (PP/FC group) propylthiouracil groups as compared to 1.3 ± 0.47 in control group (C/FC group) (P < 0.05) [Figure 4].

**DISCUSSION**

Metabolic inefficiency created due to THs deficiency during intrauterine life alters neuronal function in the developing rat brain, but how it leads to decline in the functions of neurons later is not well understood. THs play an essential role in many processes that lead to brain development and maturation and their deficiency during intrauterine life can have deleterious consequences on the brain of the developing fetus. THs have a strong influence on the movement of cells in the hippocampus and cerebellum and even their minor deficiencies are associated with migration defects. For instance, it was observed in a study that maternal hypothyroidism between embryonic days 12 to 15 significantly misplaced cells in the neocortex and hippocampus of the offspring when analyzed at 40 days of age, and the effects of maternal hypothyroidism on the anatomy of the cortex and hippocampus were prominent. Likewise, in the present study, the movement of Pyramidal cells during neurogenesis was disturbed in CA3 region of hippocampus of P/FC group (Figure 1C), when compared with control group. However, when group P/FC was compared to group PP/FC, it was observed that the migration of cells was further affected (Figure 1D). This led to the conclusion that the longer the mother is exposed to hypothyroid state, the more serious consequences occur in the newborn and lactating pups.

The hippocampus is extremely sensitive to the changes in the levels of THs during the first month of life in rats and TH receptors are present in the neurons of hippocampus. It is stated that hypothyroidism during the neonatal period in rats reduces the dentate granule cells and retards the growth of pyramidal cells in CA3 region of hippocampus. Since it was previously reported that the number of granule cells of the dentate gyrus is reduced in hypothyroid animals, therefore, in the present experiment, it was decided to extend this observation to the pyramidal cells of the

Figure 2: Features observed under immunohistochemical Bax stained slides of Purkinje neurons of cerebellar cortex. Slide A represents the 3 layers of the cerebellar cortex and the arrows point to the middle Purkinje cell layer (10x magnification). Slide B represents the control group C/FC and the arrow points to a healthy Purkinje cell (100x magnification). Slide C represents the magnified image of first generation PTU treated group P/FC (100x magnification), in which the arrow points to a damaged Purkinje cell with irregular outline. Slide D represents second generation PTU treated group PP/FC, where the arrow points to a shrunken pyknotic Bax stained nucleus of Purkinje cell (100x magnification).
NEURONAL DAMAGE IN BRAINS OF FIRST- AND SECOND-GENERATION PUPS BORN TO HYPOTHYROID WISTAR RATS

Cerebellar development is regulated by THs. The development of cerebellum is fundamentally during the first month of life in rodents. Morphological changes in the brain tissue due to deficiency of THs have been observed in rodents during the neonatal period, and these changes are very similar to the changes observed in humans under similar conditions. In case of Purkinje cells observed in the current study, viable neurons had lightly stained spherical nuclei (Figure 2B), whereas apoptotic neurons showed shrunken and fragmented nuclei, clearly visible in P/FC and PP/FC groups (Figure 2C and D). A significant decrease was seen in normal cell count of Purkinje cells in P/FC and PP/FC groups compared to control (Figure 4), and in case of apoptotic cells count, PTU treated groups P/FC and PP/FC, had a higher count as compared to control (Figure 4).

In the present study, there was an insignificant reduction in brain weight of hypothyroid pups as compared to the control. Similar to these findings, there was a study conducted by Schwartz et al., in 1997, in which no significant decrease was reported in the brains of 21 days old fetuses taken out from hypothyroid rats. In the second generation PP/FC group, the weights of hypothyroid brains were slightly reduced compared to the control. This slight reduction in brain weight indicates that when the mother is exposed to impaired levels of THs, be it for a brief period or a longer period, the fetal and neonatal brain is liable to suffer loss.

The thyroid gland of the fetus does not start THs synthesis until during middle of gestation in humans, and if the mother is hypothyroid before this time period, it has adverse consequences on the development of nervous system of the developing fetus. There was a significant increase in TSH levels in both first and second generation hypothyroid pups showed how due to maternal insufficiency of thyroid hormones, the pups could not compensate for the loss (Table I), as indicated by a rise in the serum levels of TSH and a significant decrease in the serum levels of T4. In the present study, with the results of all the parameters combined, including...
histological, morphometric, enzymatic analysis of T₃, T₄, and TSH, it is stated that maternal hypothyroidism during gestation and lactation can have serious consequences on the structure of motor neurons of neonate brain.

Further research is required for better understanding of the effect of maternal low TH levels on fetal brain development and function to devise optimum management strategies for this disorder.

LIMITATIONS OF STUDY

The effect of low maternal thyroid levels has been confined to only the motor neurons, specifically the Purkinje and Pyramidal cells of the neonate brain. Staining the sensory neurons and the interneurons too would have further provided strength to the study. Moreover, this study has only been conducted on 22 days old pups, as it was not possible to scoop the brains out as a whole from the skull of newborn pups.

CONCLUSION

The present study concludes that maternal hypothyroidism causes neuronal apoptosis of the neonates in TH deficient brain, which is further aggravated in the second generation pups, if left untreated. This was indicated by structural destruction and apoptosis in the pyramidal neurons of CA3 area in hippocampus along with Purkinje neurons of the cerebellum, more intense in second generation, when compared to the first generation pups.

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