Haloperidol-Induced Neuronal Damage in Guinea Pig Hippocampus: A Microscopic Study

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Summary

Haloperidol, a neuroleptic drug, is still discussed if it is neurotoxic. In the present study, neurotoxic effects of haloperidol on guinea pig hippocampus were examined. Sixteen guinea pigs were randomly divided into two groups of 8 animals each. The animals of treatment group were given intraperitoneally (i.p.) haloperidol diluted in saline (dose of 5 mg/kg daily) for 6 weeks. Control animals received only the same volume of saline i.p. Hippocampi from both groups were examined by light and electron microscope. Numerous dark neurons with a heterochromatin nucleus and dense cytoplasm were observed in the hippocampi of the treatment group by light microscope and ultrastructural levels. Also some neurons have increased vacuolar contents and lysosomes. These findings suggest that haloperidol leads to prominent damage of hippocampal neurons.

Key words: Haloperidol, Hippocampus, Guinea Pig, Light Microscopy, Electron Microscopy

INTRODUCTION

Antipsychotic agents in the central nervous system have been considered to play an important role in modulation of mental diseases. Various clinical(10,14) and experimental(3) studies have been carried
out the effects of different neuroleptic drugs which are considered to affect via neurotransmitter systems\(^{(8)}\). It has been continuing the discussions about both effect mechanisms\(^{(21)}\) but discussions about clinical results\(^{(35)}\) of these drugs are continuing up to now.

Haloperidol (HAL) that is known as a dopamine receptor antagonist and sigma receptor–active neuroleptic\(^{(2)}\) is a potent typical antipsychotic to be frequently used in-patients with schizophrenia to medication the behavioral disturbances\(^{(30)}\). About its mechanism, reported that hypersensitivity of dopamine receptors\(^{(4)}\) or a decrease in the activity of glutamic acid decarboxylase, GABA-synthesizing enzyme, following long-term administration of antipsychotic drugs may contribute to the drug-induced antipsycotic's extrapyramidal effects\(^{(6)}\).

Although it has been obtained a lot of useful information from different type studies such as structural\(^{(12)}\) ultra structural\(^{(34)}\) quantitative\(^{(25)}\) and clinical\(^{(16)}\) etc. Numerous previous studies are present about effects of haloperidol\(^{(23,28)}\) and only two of them are ultrastructural hippocampal studies\(^{(17,32)}\). And introduced immediately after learning, haloperidol shows a tendency towards deterioration in the phase of memory consolidation\(^{(29)}\).

The hippocampal region is an essential component of learning and memory processes\(^{(13)}\) and has been shown to be damage by a number of toxicants\(^{(15,24,36)}\) some of which are neuroleptic agents\(^{(1-7)}\).

In the present study, we attempted to examine possible neuronal changes in guinea pig hippocampus following haloperidol treatment by comparing with those of the control group by light and electron microscopy.

**MATERIAL AND METHODS**

**Experimental design and applying the drug.**

The Ethical Committee of Ataturk University Hospital approved this study. Sixteen adult male guinea pigs were used in this study. All animals were maintained under standardized conditions of light and temperature, with tap water.

The experimental animals were divided into 2 groups (8 pigs per group). All the animals were treated simultaneously according to the following conditions; I)It given haloperidol 5 mg/kg/day for first group (n:8) calling that is chronic administration; ii) It given saline at equal volume (n:8) calling that is the control group, respectively. Experimental drug or saline treated animals were injected i.p daily for 6 weeks.

**Perfusion and fixation**

At the end of applying, all animals were anaesthetized via a short inhalation of ether, and then were fixed. Initially 0.9% saline (30 ml) solution was given intracardially followed by a mixture of 2% paraformaldehyde+2% glutaraldehyde (150 ml) in 0.1 M phosphate buffer, pH 7.4 at room temperature, for approximately 30 min.

Brains were removed and stored in the same fixative overnight at 4 °C. On the following day, the hippocampus in each brain was dissected out as described previously\(^{(11,31)}\). The hippocampal samples belonging to the each animal were embedded separately.

**Microscopy for histopathological examination at ultrastructural level**

The hippocampal samples were processed; post fixed in 1% osmium tetroxide for 1 h, dehydrated through a graded alcohol series and embedded in Epon resin for ultrathin sections. Ultra thin sections with a gold interference colour were obtained using an ultramicrotome equipped with a diamond knife and were mounted on grids coated with formvar support film. From each block, approximately 30 serial sections were collected on ten grids. These grids containing sections were taken from three different regions of the blocks, which are called S1, S2 and S3. S1, S2 and S3 were
the proximal, medial and distal level of the embedded tissue in the block, respectively. The section were stained with uranyl acetate and lead citrate and viewed using a JEM-100SX electron microscope. Then It was taken electron micrographs of these specimen at X5000 magnification to examine the morphologic changes of neuron histopathologically. **Microscopy for histopathological examination at structural level** After the hippocampal samples to be received for applying of stereological method and histopathological examination were blocked as above, semi-thin sections (1 µm) were cut serially with a Nova LKB Bromma ultratome, and stained with toluidine blue. Also histopathological photographs were obtained under a light microscope with camera attachment (Olympus BX 51, JAPAN). **RESULTS** **Histopathological results at ultrastructural level** In the control animals, the cell body of neurons was usually large, dilated region of the cell that contain a large, euchromatic nucleus with one or two prominent nucleoli. The perikaryon also had masses of free ribozomes, profiles of granular endoplasmic rediculum (GER), lysosomes, numerous mitochondria, and a large perinuclear Golgi organelle (Figure 1a). It observed some difference according to control group in first group calling that is acute administration. Most of neurons were clearly determined smaller nuclei compared to that of control. In second and third groups calling that is chronic administration, It was established increasing of amount of ultrastructural changes. It was seen the dark neuron with the dens nucleus (Figure 1b), increasing the lipofuscin granules, abundant lysosomes (Figure 2c), especially third groups of which are. In addition, perinuclear edema and large vacuolar structures appeared clearly in perikaryon (Figure 2a) and the distance between the inner and outer nuclear membrane was enlarged (Figure 2b), and the inner nucleus membrane fragmented in some areas (Figure 2b). Finally, distorted myelin sheaths were encountered in the second and third groups (Figure 2d).

**Figure 1:** Electron micrographs of the hippocampus in guinea pigs (a) Neuronal cell nucleus (black arrow), illustrating apparently normal ultrastructural morphology. (b) Electron micrographs of neuronal degeneration; two neuron in the guinea pig hippocampus that is given high dose haloperidol (3 mg/kg), exhibiting the increased condensation and margination of nuclear chromatin (arrow head) and one nucleus with normal shape (black arrow).
Histopathological results at structural level

The histopathological findings in light microscope correlated well with the findings in electron microscope (Figure 3). When it is compared the morphological changes of hippocampal neuron between control and first, second and third group. It determined some alternation toward necrosis such as swelling and clumping in nuclear chromatin and also observed pathologiacal changes about nuclei of neurons such as condensation of chromatin and shrinkage of the nucleus (Pyknosis), and fragmentation of the nucleus (Kayorrhexis) (Figure 4).

It was seen more dramatic changes in second and third groups calling that is chronic administration (Figure 3c,d). The amount of necrotic neurons increased both second and third group. It observed that the nucleolus is disappeared at most of neurons. At the same time, the nucleus displaced eccentrically and the cell body enlarged, causing conversion of its counter from concave to convex for especially second and third groups.

Figure 2: Electron micrographs of neuronal degeneration in the hippocampus of guinea pigs that is applied high dose, illustrating apparently abnormal ultrastructural morphology. which is contained (a) large vacuole (v), (b) swollen mitochondria (m), fragmentation (black arrow) and dilatation (white arrow) of nuclear envelope, (c) increased density of lysosomes (ly) and finally (d) degenerating myelin (M) profile
Figure 3: Light micrographs of the hippocampus in guinea pigs (a) Neuronal cell nucleus (arrow) in control group, illustrating apparently normal ultrastructural morphology. (b) In first group, gradually to be started increasing amount of degenerated neurons (arrowhead) (c) It seem like the first group arrangement neurons in the second group. Although there is both normal and necrotic neuron, It was in favor of image of normal neuron (arrow) that is dominated the most of interesting area (d) In the third group, It was attracted attention increasing the number of necrotic neurons (arrowhead). Nucleus and cell body affected neurons are shrunken and lack morphologic detail.

Figure 4: Light micrograph of neuron one of which is affected (arrow), other has normal (arrowhead) structural shape apparently. It was observed clumping and condensation of chromatin and shrinkage of the nucleus and also nucleus was displaced peripherally, shrunken and darkly staining (asterisk). Internal details of ischemic neuron are obscured. Finally Cell counters are conversed from concave to convex.
DISCUSSION

Various clinical observations regarding the actions of neuroleptic drugs, which are one group of antipsychotic agents, have led to conjectures that these drugs may induce structural alterations in the brain. It has been purposed a lot of idea about the morphological effects in neuronal level of antipsychotic drugs.

Animal experiments have even indicated that haloperidol can damage cholinergic pathways\(^{(30)}\). Haloperidol may initiate a number of harmful effects on central nervous system neurons, including damage to cholinergic pathways on effect that could be especially deleterious to that experiencing memory dysfunction\(^{(30)}\).

There are a lot of morphological studies about useful effect of neuroleptic agents such as neuroprotective and neurogenesis at microscopic level\(^{(35,5)}\) but it has been mentioned by only some researcher that treatment with these agents have some side effects such neurodegeneration or neuronal cell loss\(^{(1,20)}\) in some area of brain, for examples hippocampus, striatum, medial prefrontal cortex.

Our study was limited to only one time point for treatment (42 days), which was selected to correspond to most published studies on typical neuroleptic. In guinea pigs, a 42-day, chronic treatment period corresponds to about 6 years of treatment in patients\(^{(35)}\). Therefore, our results correspond to chronic haloperidol treatment in humans. Because it is unknown whether these effects of haloperidol are dependent dosage or not\(^{(37,22)}\). We aimed to evaluate effects of haloperidol on hippocampal neurons at different dosages.

In the low dose, Dawirs and colleagues have reported that the low dose treatment with haloperidol increases cell proliferation on hippocampus\(^{(5)}\) but we could not encounter any mitotic figures at ultrastructural level. In addition, we could not find out any information by stereological approach about more less neuronal height, to be thought an important evidence of mitosis in low dose treatment group. Briefly, there was not any abnormal morphological appearance neither structural and ultrastructural investigation nor morphometrical analysis.

It has been reported that the addition of haloperidol to the cell culture medium reduced HT22 cell survival dose-dependently\(^{(27)}\). This finding also was reported for other cell lines and for rat primary cells\(^{(33)}\). Another study, Post et al. showed that haloperidol treatment in rats was associated with increases in the expression of p53 and the ratio of pro-apoptotic (Bax) to anti-apoptotic (Bcl-2/Bcl-x(L)) proteins in the hippocampus and caudate putamen (CPu)\(^{(26)}\).

We showed a decrease of neuronal height, one of the most evident indicating cell deaths by using one of the stereological approaches in our previous study\(^{(31)}\). There was a significant decrease in mean neuronal height and we especially determined a significant difference statistically in neuronal height of high dose group, comparing with other two groups\(^{(31)}\).

Our findings at structural and ultrastructural level support to our stereological results. In toxic situations as known cytoplasmic swelling in neurons occurs owing to an impairment of the sodium pump mechanism at the cell membrane and they have proved that in cell swelling, the nuclear envelope, which is functionally continuous with the granular endoplasmic reticulum (GER), is also susceptible to osmotic damage resulting in membrane separation and cisternal dilatation\(^{(19,16)}\). We determined much important evidence of degenerations such as pyknosis in nucleus, the increases in cytoplasmic density and in the number of lysosomes and vacuolar structures, perinuclear edema (or cytoplasmic swelling), and distorted myelin sheaths.
show that high doses of haloperidol caused hippocampal neurodegeneration (figure). We also saw that the perinuclear cistern between the inner and outer nuclear membranes was expanded, and the inner nuclear membrane was fragmented in some areas. It was determined that dark neurons were increased in number from first group to third, and the clear neurons with euchromatic nucleus were less in third group than those in other groups as have been declared by some scientists that haloperidol causes necrotic cell death in hippocampus\(^2,18\)\(^{18}\). The hypothesis that atypical neuroleptics (olanzapine and risperidone) stimulate neurogenesis\(^{35}\) and the typical neuroleptic, haloperidol (0.4 mg/kg/day), does not stimulate neurogenesis\(^{9}\)\(^9\) resemble with our findings. Although it has been claimed by some authors that haloperidol increase neurogenesis in the hippocampus\(^5\)\(^5\).

It has been claimed a lot of findings about the morphological effects in neuronal level of antipsychotic drugs. The discrepancies in the results of these haloperidol studies might be due to differences in experimental designs, drug dosages, and the species studied.

In conclusion, our findings may support that high dose haloperidol is toxic to neurons. Especially, cognitive impairment may be a result of hippocampal neuron necrosis by high-dose chronic haloperidol treatment. In clinical settings, should be avoided from high dose neuroleptic treatment.

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