Stress-induced changes in the aged-rat adrenal cortex. Histological and histomorphometric study

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Background: Stress exposure exerts direct effects on the morphology and functionality of the adrenal cortex. In addition, ageing effects growth, differentiation, apoptosis and cellularity of the cortex. The missing data is the combined effect of stress and ageing on the adrenal cortex. The aim of this study was to demonstrate the structural changes in the adrenal cortex following the exposure to stress in the adult and aged albino rats.

Materials and methods: Forty rats were divided into groups I and II (adult and senile). Each group was further subdivided into subgroups a and b (control and stressed). Light and electron microscopic studies were done. Area per cent of collagen fibres (Masson’s trichrome-stained sections), number of proliferating cells (optical density immunoreactivity in the Ki67 stained sections) and thickness of the three adrenal zones were also measured.

Results: Lamellar separation of the capsule with subcapsular spindle cell hyperplasia and areas of ghost cells were observed in zona glomerulosa (ZG) and zona fasciculata (ZF) in group I-b. Separation and indentation of the capsule with its lamellar separation were observed in group II-a with the existence of multiple scattered degenerative foci in ZF and zona reticularis (ZR). Similar and aggressive was the architectural pattern of ZF in group II-b with the presence of areas of homogenous degeneration. The nuclei of ZG had marginated chromatin in group I-b and were pyknotic with deformed irregular outlines in group II-b. Multiple lysosomes and vacuolar degeneration mitochondria were also seen in group I-b. The nuclei of ZF were irregular with condensed marginated heterochromatin in group I-b, irregular with scattered chromatin in group II-a and indented with areas of chromatin destruction in group II-b. Mitochondria with disrupted cristae and cristolysis were also detected in group I-b. Numerous lipofuscin granules and dilated smooth endoplasmic reticulum were revealed in group II-b. The mean collagen fibre area per cent and the mean number of the proliferating cells in group II-b were significantly higher by 39% and 23%. The thickness of ZG decreased significantly by 20% in group I-b. Contrary, the thickness of both ZF and ZR increased significantly by 10% in group I-b.

Conclusions: Histological alterations occurred in the adrenal cortex in response to stress, especially when coupled with the advance of age. This was accompanied by increase in the area per cent of collagen fibres and increase in the mean number of the proliferating cells in the adrenal cortex. (Folia Morphol 2018; 77, 4: 629–641)

Key words: stress, ageing, adrenal cortex
INTRODUCTION

Stress is a state of disturbed homeostasis that allows the living organisms to adapt to environmental pressure. Stress responses are mainly mediated by the activation of the hypothalamic-pituitary-adrenal axis [22]. The adrenal gland is the primary peripheral endocrine gland in the stress response [16]. It is composed of two functionally discrete organs, the cortex which synthesizes steroid hormones that mediate body homeostasis and chronic stress responses and the medulla which produces catecholamines that facilitate acute stress [27]. Stress exposure can exert direct effects on the morphology as well as the functionality of the adrenal gland [1].

Ageing of the human adrenal cortex presents several problems such as changes in growth, differentiation; apoptosis and cellular senescence affect the properties of this tissue over the lifespan [13]. The different layers of the adrenal cortex undergo changes with ageing, including a reduction in the thickness of the zona reticularis (ZR) and a relative increase of the thickness of the outer cortical zones.

With the advance of age, the cortex begins to show microscopic nodular changes [3]. Spindle or fusiform cell proliferation beneath the adrenal capsule is a common finding in mice with the advance of age [9].

The missing data is the combined effect of stress and ageing on the adrenal gland of each rat of all groups. They were kept in 10% formol saline, processed for paraffin sections of 5–7 µm thickness and mounted on glass slides for histopathological examination. The sections were stained with haematoxylin and eosin (H&E) stain to study the histological changes and stained with Masson’s trichrome stain to demonstrate the collagen fibres.

MATERIALS AND METHODS

Animals

Forty Wister male albino rats were divided into two groups. Each group was further subdivided into two subgroups. The rats were sacrificed after 14 days. The study was approved by the Ethics Committee, Faculty of Medicine, Cairo University (431/2017). All animal procedures followed the ethical guidelines of experimental animals. We followed the guidelines of the ethical standards of the National Institutes of Health guide for the care and use of Laboratory Animals (NIH Publications No. 8023, revised 1978).

Experimental design

Experimental design is shown in Table 1. Rats in subgroup a were housed separately in plastic restrainers (6 cm wide × 12 cm long) for 2 h daily for 14 days [24].

Light microscopic examination

The specimens were obtained from the adrenal gland of each rat of all groups. They were kept in 10% formol saline, processed for paraffin sections of 5–7 µm thickness and mounted on glass slides for histopathological examination. The sections were stained with haematoxylin and eosin (H&E) stain to study the histological changes and stained with Masson’s trichrome stain to demonstrate the collagen fibres.

Immunohistochemical staining

The Ki67 antigen (a cell division marker) was used to detect cell proliferation in the adrenal cortex. Immunohistochemistry for Ki67 antigen detects cells that are in late G1, S, G2, and M but not G0 phases of the cell cycle [5].

The technique was done according to the manufacturer’s instructions. The glands were mounted on polylysine slides. Endogenous peroxidase activity was blocked by immersing sections in 3% hydrogen peroxide in methanol for 30 min in the dark at room temperature. The sections were then rehydrated and washed in phosphate buffered saline (PBS; 2–5 min). To unmask antigenic sites, sections were microwaved at full power for 4–5 min in 0.01 M sodium citrate buffer, pH 6.0, and then cooled and washed in PBS (5 min). The sections were treated with blocking serum (20% normal rabbit serum in 1× Tris-buffered saline [TBS]), pH 7.6) for 30 min. Then, the sections were treated with the primary anti Ki67 anti-

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<th>Table 1. Experimental design</th>
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<td><strong>Group</strong></td>
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<td>Group I (adult group)</td>
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<td>Group II (senile group)</td>
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| Subgroup a (control group) | 10 |
| Subgroup b (stressed group) | 10 |

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body (Novocastra Lab Ltd.; NCL-Ki67-MM1) for 2 h. Biotinylated secondary antibody (rabbit anti-mouse IgG biotin, Dako E0354) was applied for 30 min. Then, the sections were treated with avidin-biotin-horseradish peroxidase solution (AB Complex/HRP kit; Dako K0355) for 30 min at room temperature. Finally, the sections were incubated for 3 min with 3,3'-diaminobenzidine solution (DAB isopac, Sigma D9015), comprising 25 μL DAB (2 mg/mL) + 5.9 mL 0.2 MTBS + 1 μL 30% H2O2. After lightly counterstaining with haematoxylin (Vector, H3401), slides were rinsed, dehydrated through graded ethanol, cleared in xylene and mounted under coverslips with DPX mounting medium. The cells were considered positive for the Ki67 antigen when the nuclei were stained with a brownish colour [6].

**Electron microscopic examination**

Specimens from the adrenal cortex were fixed in 4% glutaraldehyde then washed in phosphate buffer and post-fixed in 1% osmium tetroxide. Fixation was followed by dehydration and embedding in epoxy resins. Semithin sections (1 μm) were stained with toluidine blue. Ultrathin sections (50–60 nm) were stained with uranyl acetate and lead citrate. These sections were examined and photographed using a Joel, 100 CX II transmission electron microscope.

**Histomorphometric studies**

Quantitative data were obtained using “Leica Qwin 500 C” image analyser computer system Ltd. (Cambridge, England). The following parameters were measured in different fields with fixed field area 4 × 104 μm²: area per cent of collagen fibres (Masson’s trichrome-stained sections), the number of proliferating cells (optical density immunoreactivity in the Ki67 stained sections) and the thickness of the three adrenal zones.

**Statistical analysis**

Statistical analysis was performed using statistical package for the social sciences (SPSS) version 21.0 (IBM Corporation, Somers, NY, USA) statistical software. The data were expressed as means ± standard deviation (SD). Statistical evaluation was done using one-way analysis of variance (ANOVA) followed by Bonferroni pairwise comparisons. Significance was considered when p-value was ≤ 0.05. The percentage of increase or decrease (difference) of all study parameters were calculated per the following formula:

\[
\text{Percentage of difference} = \frac{\text{Mean difference value between two groups}}{\text{Value of the compared group}} \times 100
\]

**RESULTS**

**Histological section observations**

Zona glomerulosa (ZG) cells were arranged in clusters, zona fasciculata (ZF) cells were arranged in straight columns and ZR cells were arranged as anastomosing cords in group I-a (Fig. 1a). Lamellar separations of the capsule with areas of ghost cells were observed in ZG and ZF in group I-b (Fig. 1b, c). Separation and indentation of the capsule with its lamellar separation were observed in group II-a. In addition, the arrangement of ZG cells was disturbed with the existence of multiple scattered degenerative foci in ZF and ZR (Fig. 1d–f). Capsular separation with subcapsular spindle cell hyperplasia and disruption of the regular architectural pattern of ZF were seen in group II-b with the presence of areas of homogenous degeneration (Fig. 1g, h).

Spindle cell hyperplasia and vacuolated cell-areas were detected in ZR in the group I-b (Fig. 2a, b). Scattered brown granules, wide sinusoids, and areas of degeneration in ZR were observed in group II-a (Fig. 2c, d). The disturbed arrangement of the ZR cells with engorged central adrenal veins and congested sinusoids were seen in group II-b (Fig. 2e, f).

**Semithin sections findings**

Thickening with lamellar separation of the capsule with irregular shape and arrangement of its fibroblasts were observed in group II-b. ZG cells exhibited lipid droplet-loaded cytoplasm in the groups I-a, I-b, II-a. Their nuclei exhibited different shapes and sizes in group I-b, darkly stained in group II-a, pyknotic in group II-b (Fig. 3). The nuclei of ZF were rounded vesicular in groups I-a, II-a, while they were irregular in shape and darkly hyperchromatic in group II-b (Fig. 4).

**Masson’s trichrome-stained sections and immunohistochemical study**

The capsule was regular with parallel arranged collagen fibres in group I-a, while it was disrupted with thickened fibrous septa in group I-b. Lamellar separation of the capsule with irregularly dispersed collagen fibres was seen in group II-a. The capsule
Figure 1. Sections of the ZG and ZF (H&E): a. Capsule (C), ZG cells arranged in clusters, ZF cells arranged in straight columns and ZR cells arranged as anastomosing cords in a rat group I-a (×200). b. Capsular separation (arrowhead), multiple pleomorphic nuclei (N) and areas of ghost cells (dotted arrows) in ZF in a rat group I-b (×400). c. Lamellar separation of the capsule (dotted arrow), thin overlying capsule (arrowhead) and subcapsular spindle cell hyperplasia (arrow). Note an area of ghost cells (circle) in ZG in a rat group I-b (×400). d. Multiple scattered degenerative foci in the ZF (long arrow) and ZR (short arrows) in a rat group II-a (×100). e. Capsular separation (dotted arrow) with the disturbed arrangement of ZG cells (circle). Note a capsular artery (A) and vein (V) in a rat group II-a (×200). f. Capsule with lamellar separation (arrow) and indentation (dotted arrow). Note an area of degenerated ghost cells (circle) in a rat group II-a (×400). g. Capsular separation (arrows) with disruption of the regular architectural pattern (circle) of ZF in a rat group II-b (×150). h. Capsule (C) with subcapsular spindle cell hyperplasia (circle). Note: an area of homogenous degeneration (arrow) in a rat group II-b (×400); ZG — zona glomerulosa; ZF — zona fasciculata; ZR — zona reticularis.
Figure 2. Sections of the zona reticularis (ZR) and adjoining medulla (M) (H&E); a. Spindle cell hyperplasia (arrows) in a rat group I-b (×400). b. An area of vacuolated cells (arrows) in a rat group I-b (×400). c. Scattered brown granules (arrows) and wide sinusoids (arrowheads) in a rat group II-a (×400). d. An area of degeneration (arrows) in a rat group II-a (×150). e. A disturbed arrangement (circle) of the ZR cells and an engorged central adrenal vein (arrow) in a rat group II-b (×150). f. A congested sinusoids (arrows) in a rat group II-b (×400).

was thickened with lamellar separation and thickened disrupted collagen bundles in group II-b. Dilated congested blood capillaries with extravasation of blood and exudation were also seen in group II-b (Figs. 5, 6).

The immunoreaction of Ki67 was nuclear. It was absent in groups I-a, II-a, while it was positive in groups I-b, II-b (Fig. 7).

Ultrastructural remarks

The nuclei of ZG were euchromatic with prominent nucleoli in groups I-a and II-a, had margined chromatin in group I-b and were pyknotic with deformed irregular outlines and disrupted nuclear envelopes in group II-b. Multiple lysosomes and vacuolar degeneration mitochondria were also seen in group I-b (Fig. 8).

The nuclei of ZF were regular and rounded in group I-a, irregular with condensed marginated heterochromatin in group I-b, irregular with scattered chromatin in group II-a and indented with areas of chromatin destruction in group II-b. Mitochondria with disrupted cristae and discontinuity of their envelopes and mitochondria with cristolysis and loss
Figure 3. Semithin sections of the zona glomerulosa (ZG) and zona fasciculata (ZF) (Toluidine blue ×1000); a. Capsular connective tissue fibres (dotted arrows), fibroblasts (white arrows) and ZG cells (thick black arrows) with the cytoplasm loaded with lipid droplets (arrowheads) in a rat group I-a. Note a blood capillary (thin black arrow). b. Large swollen disrupted arrangement fibroblasts (arrowheads). The nuclei of the ZG cells exhibited different shapes and sizes (arrows), while the cytoplasm was loaded with lipid droplets (dotted arrows) in a rat group I-b. c. A distended capillary (arrow) with folded endothelium (dotted arrows) in a rat group II-a. d. Capsular separation (long arrow) and a markedly dilated blood vessel (short arrow). The ZG shows abundant lipid droplets (circles) and darkly stained nuclei (arrowheads) in a rat group II-a. e. A lamellar separation of the capsule (arrows) and ZG with pyknotic nuclei (dotted arrow) in a rat group II-b. f. A marked thickening of the capsule with irregular shape and arrangement of its fibroblasts (arrows) and collagen fibres (dotted arrow) in a rat group II-b.

Figure 4. Semithin sections of the zona fasciculata (ZF) and zona reticularis (ZR) (Toluidine blue ×1000); a. ZF with rounded vesicular nuclei (arrowheads) and lipid droplets loaded cytoplasm (arrows) in a rat group I-a. b. ZF with rounded nuclei (arrowheads) and prominent nucleoli. Note nuclei with double nucleoli (arrows) in a rat group I-b. c. ZF with vesicular regular shaped nuclei (thick arrows) and lipid droplets loaded cytoplasm (thin arrows) in a rat group II-a. Note pyknotic nuclei (dotted arrows). d. ZF with irregular shape nuclei (long arrows), darkly stained hyperchromatic nuclei (short arrows) in a rat group II-b. Note lipid droplets (circles).
Figure 5. Collagen fibres in zona glomerulosa (ZG) and zona fasciculata (ZF) (Masson’s trichrome); a. A regular capsule (C), trabeculae of collagen fibres (dotted arrow) and parallel arranged collagen fibres (arrows) perpendicular to the surface in a rat group I-a. Note blood sinusoids (arrowheads) (×200). b. A thickened fibrous septum (arrow) and a disrupted area in the capsule (dotted arrow) in a rat group I-b (×200). c. Lamellar separation (dotted arrows) of the capsule (C) and irregularly dispersed collagen fibres (arrows) in a rat group II-a (×400). d. Accessory adrenocortical tissue (arrow) in a rat group II-a (×100). e. Thickened-capsule (C) with lamellar separation of its collagen bundles (dotted arrows), thickened collagen bundles (white arrowheads), irregular collagen fibres (black arrowheads) and areas of exudate (arrows) in a rat group II-b (×400). f. A thickened fibrous septum (arrows) and a thickened partially separated capsule (dotted arrow) in a rat group II-b (×200).

Figure 6. Collagen fibres in zona fasciculata (ZF), zona reticularis (ZR), and adjoining medulla (M) (Masson’s trichrome ×200); a. Thickened collagen bundles (arrow) in a rat group I-b. b. A network of collagen bundles (arrowheads) and numerous blood capillaries (arrows) in a rat group II-a. c. A thickened disrupted collagen bundles (circle) dilated congested blood capillaries (long arrows) with extravasation of blood (short arrows) in a rat group II-b.
of their envelopes were also detected in group I-b (Fig. 9).

The absence of nuclear affection was perceived in the ZR cells of the different groups. Lipofuscin granules and dilated smooth endoplasmic reticulum were also observed in group II-b (Fig. 10).

**Morphometric measurements (Tables 2–4)**

The mean collagen fibre area per cent of group II-b was 39%, significantly higher than that of group II-a. Using the optical density of the Ki67, the mean number of group II-b proliferating cells was significantly higher by 23% than that of group I-b.

The thickness of ZG decreased significantly by 20% in group I-b as compared to that of group I-a. Contrary, the thickness of both ZF and ZR increased significantly by 10% in group I-b as compared to their respective of group I-a.

**DISCUSSION**

Stress is a state of endangered homeostasis. The effect of stress on the adrenal cortex was evaluated as the adrenal gland is the chief stress responsive organ involved in the overall homeostasis. Various types of experimental stressors have been styled such as restraint, cold/heat, water immersion, water deprivation, noise, crowding and electric shock [10]. Among all these methods, restraint is the method that attains the least pain, suffering or distress to the animal.
Figure 8. Electron photomicrograph of the zona glomerulosa; a. A nucleus (N) containing euchromatin (Eu), clumps of peripheral heterochromatin (Ht), prominent nucleolus (nu) and multiple elongated mitochondria (M) with tubulolamellar cristae in a rat group I-a (TEM ×20000). b. A nucleus (N) with clogged marginated deeply stained chromatin distorting the nuclear contour. The cytoplasm contains multiple lysosomes (arrows), mitochondria (M) with tubulolamellar cristae and mitochondria with vacuolar degeneration (thick arrow) in a rat group I-b (TEM ×20,000). c. A nucleus (N) with distinct nucleolus (nu), heterochromatin (Ht) and undulation of its nuclear envelope (black arrow) in a rat group II-a. Note lipid droplets (L) (TEM ×10,000). d. A pyknotic nucleus (N) with a deformed irregular outline (dotted arrow), dense clumps of clogged chromatin associated with disrupted nuclear envelope (thick arrow) in a rat group II-b. Note mitochondria (M), desmosomes (white arrows) and bundles of collagen fibres (black arrow) (TEM ×15,000).

The present study was confined to male rats since the adrenal gland of female rats undergo several changes such as the appearance and regression of the X-zone [9] and the response to stress is a sex-specific due to a gender difference in the hormonal receptors [2].

Areas of subcapsular spindle cell hyperplasia were detected in the adult and senile stressed groups (groups I-b, II-b). These cellular clusters were located initially in the ZG and then extend centripetally between the underlying ZF cell cords. Several explanations had been suggested: a mast cells role [15], sex hormones association [20] or misspecifications of subcapsular mesenchymal stem cells that do not differentiate to enter the steroidogenic lineage [28].

Ghost cell areas, ghost nuclei, multiple pleomorphic and pyknotic nuclei existed in ZG of the adult stressed group. The nuclei of ZG cells are extremely sensitive to stress [16]. Areas of vacuolated cells at the junctional zone between ZR and medulla existed in this group. Stress induces pronounced changes at the interphase between ZR and medulla where connective tissue cords loosened, cell count decreased and cell-to-cell spaces extended [18]. These vacuolated areas might denote an early stage of cystic degeneration [21].

A significant decrease in the thickness of the cortical zones was observed in the adult stressed group (group I-b) which may be related to shifting in hormonal stress responsiveness [7].

Increase thickness, indentation with lamellar separation were constant findings of the capsule of the senile control group (group II-a). In addition, replacement of the capsular cellular elements with collagenous tissue was also detected; such replacement was suggested as a mechanism responsible for the age-dependent capsular changes [14].

Disturbance of the arrangement the ZG was noticeable in both senile groups (groups II-a, II-b). The
imperfect repair of damage with the advance of age combined with compensatory overgrowth can lead to more and more abnormal tissue architecture [12]. In addition, increased vacuolation of the ZG cells reflected a decrease in the secretory activity of ZG cells [25]. The three cortical zones normally have some degree of vacuolation; these vacuoles contain neutral lipid and cholesterol. The increased cortical vacuolation can be due to age-dependent drop in cholesterol biosynthesis or metabolism leading to accumulation of cholesterol and steroids [25].

Areas of degeneration with a cytoplasmic heterogeneity of the ZF cells was seen in group II-a which reflects different cell sensitivity to destructive factors and an increase in the subpopulation of newly formed adrenocortical cells [17]. Another notable finding in this group was the predominance of lipofuscin-like inclusions seen in the cytoplasm of the ZR cells. Lipofuscin is a biomarker of ageing and its progressive accumulation is one of the most prominent changes with ageing [14]. Deposition of lipofuscin pigment is most probably due to decreased metabolic activities in the cells [14]. It accumulates gradually in lysosomes of post-mitotic tissue and it is not removed by degradation or exocytosis [4].

Accessory adrenocortical tissue was also encountered in group II-a. This tissue is composed of normal cortical tissue, in which ZG and ZF are notable while the medullary tissue is not normally associated [11]. The accessory cortical tissues undertake the same ageing changes as the main body of the adrenal [11].

The senile stressed group (group II-b) exhibited marked disruption of the normal architecture of three adrenocortical zones which may be related to the effect of elevated corticosterone level [8]. This disturbance was associated with different vascular changes in the form of patches of exudate in ZF, congestion of the capillaries in ZR and marked...
Figure 10. Electron photomicrograph of the zona reticularis; a. A euchromatic nucleus (N) with a nuclear envelope (arrowhead), a profile of smooth endoplasmic reticulum (SER) (S), lipid droplets (L) and a spindle-shaped fibroblast (arrow) in a rat group I-a (TEM ×10,000). b. A rounded euchromatic nucleus (N) possessing double nucleoli (nu) and mitochondria (M) with numerous tubular cristae in a rat group I-b (TEM ×15,000). c. A euchromatic nucleus (N), SER (S), multiple mitochondria (M) and few lipid droplets (L) in a rat group II-a. Note an elongated endothelial nucleus (black arrow) (TEM ×10,000). d. A euchromatic nucleus (N) with a distinct nucleolus (nu), mitochondria (M), dilated SER (S), lysosomes (white arrows), and Golgi body (black arrow) in a rat group II-b (TEM ×10,000). e. Few lipid droplets (L), numerous lipofuscin granules (white arrows), dilated SER (black arrow) and free ribosomes (arrowheads) in a rat group II-b (TEM ×20,000).

Table 2. Area per cent of collagen fibres

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<th>Groups</th>
<th>Mean ± SD</th>
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<tr>
<td>Group I-a</td>
<td>31.6 ± 4.7</td>
<td>0.09</td>
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<tr>
<td>Group I-b</td>
<td>38.7 ± 0.9</td>
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<tr>
<td>Group II-a</td>
<td>41.4 ± 3.7</td>
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<tr>
<td>Group II-b</td>
<td>57.8 ± 2.2</td>
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*Significant (p < 0.05); SD — standard deviation

Table 3. Optical density of Ki67 immunoreactivity in the stressed groups

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<th>Groups</th>
<th>Mean ± SD</th>
<th>P</th>
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<tr>
<td>Group I-b</td>
<td>67.46 ± 5.17</td>
<td>0.02*</td>
</tr>
<tr>
<td>Group II-b</td>
<td>83.05 ± 1.63</td>
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*Significant (p < 0.05); SD — standard deviation
engorgement of the central adrenal veins. The morphological changes under stress conditions adrenals developed against a background of haemodynamic changes that involved all layers of the adrenal cortex where the sinusoids and capillaries were dilated and plethoric [18].

Immunoreactivity to Ki67 was detected only in the stressed groups (groups I-b, II-b) as the cellular turnover in the adrenal cortex occurs only to accommodate the physiological needs or in response to experimental manipulations [23]. The pattern of distribution of immunoreactivity varied; in the adult stressed group, the reactivity was located in the subcapsular region, while in the senile stressed group it was confined to the junctional zone between ZG and ZF. The stem/progenitor cells exist in the adrenal capsule, subjacent cortex, and juxtamedullary region. These cells proliferate, differentiate, and repopulate in the cortex centripetally or centrifugally [23].

The ultrastructural remodelling of the adrenals of the stressed groups was mainly manifested as changes in the constituents involved in steroid hormone biosynthesis, namely, the lipid droplets, mitochondria, and smooth endoplasmic reticulum. The lipid droplets were seen packed together. These packed lipid droplets represent packets of steroid hormones awaiting release by exocytosis [8]. Other changes correlate with adrenal stimulation was the increasing number and size of mitochondria with more infolding of their inner membranes and their transformation to a more lamellar membrane pattern.

Another ultrastructural feature encountered in the senile stressed group was the degenerative changes of mitochondrial in the form of cristolysis, disruption of mitochondrial envelopes with the presence of intra-mitochondrial electron-dense inclusions [17]. The number of secondary lysosomes increased in this group which indicated autophagocytosis in the adrenocortical cells [19].

The senile stressed rats (group II-b) showed evident nuclear changes. The ZG cells exhibited nuclei with pyknosis, highly deformed irregular outlines and dense clumps of clogged chromatin. The ZF cells also showed highly indented bilobed nuclei with the undulation of the nuclear envelopes, electron-lucent spots and focal areas of chromatin destruction. These features of nuclear degeneration could reflect the superimposed effect of stress when coupled with ageing. The senile rats which were already exposed to the deleterious effect of ageing, showed marked deterioration upon stress exposure. Age prompted such effects through the affection of the hypothalamic-adrenocorticotropic-adrenal axis which may be confounded by multiple factors, including stress [26].

CONCLUSIONS

In conclusion, histological alterations occurred in the adrenal cortex in response to stress, especially when coupled with the advance of age. This was accompanied by proliferative changes.

REFERENCES


| Table 4. Thickness of the three adrenal zones |
| ZG   | Mean ± SD    | P   |
| Group I-a | 51.7 ± 6    | 0.02* |
| Group I-b | 41.3 ± 5.5  |      |
| Group II-a | 61.1 ± 9.4  |      |
| Group II-b | 57.1 ± 9.7  | 0.074 |

| ZF   | Mean ± SD    | P   |
| Group I-a | 173.0 ± 6.1  | 0.02* |
| Group I-b | 190.7 ± 5.5  |      |
| Group II-a | 167.3 ± 6.4  |      |
| Group II-b | 170.7 ± 4.2  | 0.130 |

| ZR   | Mean ± SD    | P   |
| Group I-a | 115.0 ± 5.0  |      |
| Group I-b | 127.3 ± 5.1  | 0.003* |
| Group II-a | 83 ± 4.3     |      |
| Group II-b | 85.2 ± 3.4   | 0.069 |

*Significant (p < 0.05); SD — standard deviation; ZG — zona glomerulosa; ZF — zona fasciculata; ZR — zona reticularis