Differential effect of high fat diet (HFD) on the cardiac muscle of adult and aged female mice and the possible protective role of artichoke treatment: Histomorphometric and ultrastructural study

Asmaa F. A. Dawood and Hemmat H.G. Hareedy

1 Department of Histology, Faculty of Medicine, Assiut University, Assiut, Egypt, Department of Biomedical Science, Faculty of Medicine, King Faisal University, Alhasa, Saudi Arabia, 2Department of Basic Medicine, Majma'ah University, Saudi Arabia.

ABSTRACT

Background: Coexistence of hyperlipidemia and aging is a highly serious condition that massively deteriorates cardiac function. Artichoke is a well-known antioxidant commonly used in long-lived countries diet. Recent researches showed its hypolipidemic and antiaging effects.

Objective: This work aimed to investigate the histopathological effects of high fat diet (HFD) on adult mice cardiac muscle, to address and compare the extent of myocardial damage in aged mice and to assess beneficial effects of artichoke in mitigating these effects.

Materials and Methods: A total of 42 adult (Group I) and aged (Group II) female mice (3 and 26 months old respectively) were divided into six equal subgroups (7 per each); Groups Ia and IIa: adult and aged control, Groups Ib and IIb: adult and aged HFD groups were fed diet enriched with 1.5% cholesterol and 10% lard oil for 6 weeks. Groups Ic and IIc: adult and aged artichoke treated group, were received HFD for 6 weeks as HFD groups concomitant with artichoke extract at a dose of 1.5 gm/kg once daily via gastric tube.

Results: Adult HFD laden cardiac myocytes showed loss of cross striations, inflammatory cell infiltrate, increase collagen content, myofiber focal degeneration, disruption of myofibrils, sarcomeres and T tubule system, lipid vacuolation, mitochondrial damage and nuclear pyknosis. These effects were the worst in aged HFD fed animals. Artichoke markedly improved these changes in both adult and aged mice.

Conclusion: combination of HFD and aging caused more extensive insults on cardiac myocytes than each factor alone. Artichoke is highly efficient in ameliorating these insults.

Key Words: HFD, fibrillolysis, cristolysis, sarcopenia, artichoke.

INTRODUCTION

The prevalence of obesity has increased at an alarming rate during the last two decades. It has almost reached to epidemic proportions and this appears to be more pronounced in women. Changes in food consumption, physical activity, urbanization, socioeconomic and demographic factors are being important factors that contribute to the increased prevalence of obesity[1].

Lipotoxic cardiac morbidity is one of the most serious consequences of obesity in human. In both human and animal models, lipotoxicity resulted in diverse myocardial dysfunction including left ventricular chamber dilation, cardiac hypertrophy, contractile dysfunction, atrial fibrillation and ventricular extrasystoles[2]. One of the most serious and fatal complications of morbid obesity is congestive heart failure which occurs even in absence of hypertension or underlying organic cardiac disease[3].

Aging is a natural process which is associated with gradual decline in biological functions, that directly affects lifestyle. Aging remains the single major risk factor for heart attacks, stroke, diabetes, cancers, and most chronic diseases[4]. With the current increase in life expectancy, the prevalence of obesity also raises steadily among elderly people. Both obesity and aging can lead to serious health problems mainly cardiac diseases that can lead to death. Aging is associated with an increase in abdominal obesity, a major contributor to insulin resistance and the metabolic syndrome[5]. Obesity in the elderly is thus a serious concern and comprehension of the related histopathological changes and the key underlying mechanisms become a necessary matter[6]. Actually, studies of histological and ultrastructural alterations of cardiac muscle in case of HFD
feeding in adult or when associated with aging are very limited. Moreover, little is known about the protective effect of concomitant treatment with antioxidants on aged HFD laden myocardium.

Antisteatotic drugs are commonly used in treatment plan of hyperlipidemia but recent researches have reported their adverse effect on human health. Thereby, nowadays there is increasing interest for the use of natural lipid-lowering agents that may delay or circumvent drug therapy\(^{[6]}\).

Artichoke (Cynarascolymus) is a plant frequently grown in Mediterranean countries. It is highly rich in natural active antioxidant ingredients and thus is commonly used as an herbal drug\(^{[7]}\). Recent investigations have suggested that artichoke extract has potential lipid-lowering, hepatoprotective and also antiaging effects on skin\(^{[8,9]}\). However, its potential role against cardiac lipotoxicity is not yet investigated.

Accordingly, the present study aimed to address the histopathological effects of HFD feeding on cardiac muscle of aged mice, comparing such impact with that on adult myocardium and to demonstrate the beneficial effects of artichoke in mitigating lipotoxic cardiac damage.

**MATERIALS AND METHODS**

**Chemicals:**

Cholesterol and Lard oil were purchased from Sigma Chemical (St. Louis, MO, USA). They were added to standard diet at a percentage of 1.5 and 10\% respectively. The high fat diet was freshly prepared every week and stored at 4C.

Artichoke was provided in the form of commercial capsules (MEPACO- EGYPT -Enshas El Raml-Sharkeiya) sold in the market as Cyncholine 100mg. A suspension was made by mixing the content of the capsules with distilled water (each 100 mg was dissolved in 5ml distilled water).

**Animals:**

Forty-two adult and aged female albino mice (21 for each group) were used in the current study. They were provided by animal house of Assiut University, Assiut, Egypt. Adult and old mice were 3 month-old (25-30gm) and 26-month old (40-50gm). All animals were kept in clean ventilated stainless-steel cages under controlled laboratory conditions. They were allowed free access to water and were under daily weight monitoring. All procedures on experimental animals were overseen and approved by ethical Committee, faculty of Medicine, Assiut University, Egypt.

**Experimental design:**

Adult mice (Group I) and aged mice (Group II) were equally subdivided into six subgroups (7 per subgroup) as the following:

1) Adult control (Ia) and aged control (IIa) groups: Control animals were fed standard chow diet for 6 weeks.

2) Adult HFD treated (Ib) and aged HFD treated (IIb) groups:

High fat diet treated adult and aged mice were fed HFD composed of standard chow diet supplemented with 1.5\% cholesterol\(^{[10]}\) and 10\% Lard oil\(^{[11]}\) for 6 weeks.

3) Adult artichoke treated (Ic) and aged artichoke treated (Iic) groups:

Groups Ic and Iic were fed HFD for 6 weeks as HFD treated animals. They concomitantly received artichoke extract at a dose of 1.5 gm/kg/day\(^{[12]}\) via gastric tube for 6 weeks.

Under general anesthesia using chloroform, all animals were sacrificed. Hearts were taken, immediately fixed in 10\% Formalin and processed for histological, ultrastructural and morphometric study.

**Light microscopy:**

Specimens from the base of the heart were dissected then processed for general histological examination of left ventricles using Hematoxylin & Eosin (H&E) and Masson trichrome stains\(^{[13]}\).

**Electron microscopy:**

For semithin and ultrathin sections, small specimens from left ventricles were fixed in 2-3\% Glutaraldehyde, kept in the fridge overnight then processed and post fixed in osmium tetra oxide. Semithin sections were cut, stained with Toluidine blue stain, examined, photographed then used to select specific areas for preparing ultrathin-sections of the electron microscope.

Ultrathin sections were cut and stained with uranyl acetate and lead citrate, examined and photographed with a JEOL (JEM-100cx) transmission electron microscope in Assiut University Electron Microscope unit\(^{[14]}\).

**Morphometric study:**

Myocyte number and diameter were estimated as follow:

1) Cardiac myocyte number:

In H & E stained cross sections, myocyte number was estimated by counting nuclei number. Ten sections were used for each group. For each section, six images were scanned and nuclei number was estimated at 100x magnification by using point counting method with a computerized image analyser system software (Leica Q 500 MCO; Leica, Germany\(^{[15]}\)).
2) Cardiac myocyte diameter:

In H & E stained sections, five random regions from each slide at 100x magnification and 10 myocytes were measured from each section. A total of 100 cell was measured per animal. Diameter was measured in (µm) across the nuclei, for those only containing nuclei using a calibrated ocular micrometer scale. Branching sites were excluded from measurement. For calibration, stage micrometer was used[13,16].

Statistical data Analysis (SPSS) of results was done using (ANOVA test) and Mann Whitney U test for pair comparison of means among groups. Results were expressed as mean and standard deviation (mean ± SD). P value less than 0.05 was considered significant.

RESULTS

Light microscope examination:

1) H & E and semithin stained longitudinal sections:

H & E stained sections of left ventricle of adult control group (Ia) revealed individual cardiac myocytes were cylindrical in shape that branch and anastomose forming three-dimensional network. Their sarcoplasm was highly acidophilic, crossly striated and bearing a single central oval and vesicular nucleus. Narrow spaces of endomysium were seen among myocytes. Few fibroblasts having flat dense nuclei were observed at periphery of myocytes residing endomyium (Figs. 1 and 2). In semithin sections, cardiac myocytes were closely packed and branched forming a continuous syncytium separated by narrow endomysial spaces richly supplied in blood capillaries. Sarcoplasm showed obvious cross striations and had single oval vesicular nucleus. Intercalated discs appeared as the most abundant fibroblast pool when compared with all other groups (Fig. 3).

H & E stained sections of left ventricle of adult HFD treated group (Ib) revealed that muscle fibers were remarkably improved compared with other aged groups. No foci of muscle lysis or corrugation or inflammatory cell infiltration were noticed. Some myocytes showed focal degeneration while some others had wavy corrugated appearance. Sarcoplasm was pale, disintegrated and lost its cross striations and nuclei were dense and pyknotic (Figs. 13, 14 and 15). Fibroblast population were noticeably increased and apparently appeared as the most abundant fibroblast pool when compared with all other groups (Fig. 13).

H & E stained and semithin stained sections of left ventricle of aged control group (IIa) showed remarkable changes compared with group Ia. Disturbed architecture was clearly noticed. Muscle fibers became loosely packed with widening of intercellular spaces. Sarcoplasm of most muscle fibers was pale acidophilic, degenerated and lacked its cross striations. Most nuclei were dense and shrunken (Figs. 10, 11 and 12). Many fibroblasts were also noticed unlike group Ia (Fig. 10).

In H&E and semithin stained sections of left ventricle of aged HFD group (IIb), cardiac tissue damage and disruption is more pronounced than groups IIa and Ib. Many muscle fibers were lost with subsequent marked widening of intercellular spaces. Some inflammatory cell infiltrate and congestion of blood vessels were noticed. Some myocytes showed focal degeneration while some others had wavy corrugated appearance. Sarcoplasm was pale, disintegrated and lost its cross striations and nuclei were dense and pyknotic (Figs. 13, 14 and 15). Fibroblast population were noticeably increased and apparently appeared as the most abundant fibroblast pool when compared with all other groups (Fig. 13).

Masson trichrome stained sections of left ventricle of aged artichoke treated group (Iic), cardiac muscle fibers were remarkably improved compared with other aged groups. No foci of muscle lysis or corrugation or inflammatory cell infiltration was noticed. Myocytes appeared branched with narrow intercellular spaces. Sarcoplasm was deeply acidophilic and crossly striated. Some nuclei seemed normal while others were shrunken. Fibroblast population apparently much decreased compared with other aged groups (Figs. 16, 17 and 18). In addition, intercalated discs were seen among myocytes (Fig. 18).

2) Masson trichrome stained longitudinal sections:

Masson trichrome stained sections of left ventricle of group Ia showed scanty collagen fibers were seen around blood vessels and among muscle fibers (Fig. 19).

In group Ib, Masson trichrome stained sections of left ventricle showed the amount of collagen fibers was remarkably increased among myocytes and around blood vessels compared with group Ia that encroaching adjacent muscle fibers (Fig. 20).

The amount of collagen fibers in Masson trichrome stained sections of group Ic was obviously decreased
compared with group Ib and it seemed almost normal among muscle fibers (Fig. 21).

Masson trichrome stained sections of left ventricle of group IIa revealed increased amount of collagen fibers that encroaching myofibers compared to group Ia (Fig. 22).

In group IIb, Masson trichrome stained sections of left ventricle showed increased amount of collagen fibers replacing many muscle fibers with obvious widening of intercellular spaces (Fig. 23).

Masson trichrome stained sections of left ventricle of group IIc showed decreased amount of collagen fibers among myocytes compared with other aged groups (Fig. 24).

Electron microscope examination:

In adult control group (Ia), most sarcoplasm of muscle fibers was occupied with longitudinal parallel arrays of cylindrical myofibrils. Their nuclei were oval central and euchromatic. Abundant number of large mitochondria were seen juxtaocular and run among and parallel to myofibrils. Mitochondria were oval or rounded and possessed densely packed cristae. Each sarcomere was bounded by two Z lines and was composed of central dark (A) band and peripheral two halves of light (I) band. H band appeared as narrow light zone in the center of A band and was bisected by dark M line. Steplike intercalated discs were seen among myocytes. They were composed of transverse regions containing desmosomes and fascia adherens and longitudinal regions representing gap junctions. Glycogen granules appeared as small dense granules among myofibrils. Sarcolemma was intact, scalloped and showed invaginations of T tubules opposite Z lines. Terminal cisternae of sarcoplasmic reticulum and T tubules constituted diads which were noticed in sarcoplasm at the level of Z line (Figs. 25 and 26).

In adult HFD treated group (Ib), intramyocellular accumulation of lipid vacuoles was observed so close to mitochondria. Mitochondria had partially damaged cristae. Moreover, thinning and splitting of myofibrils and focal myofibrillar lysis were noticed. Sarcomere showed obvious degeneration; A and I bands were not detectable and Z lines were disrupted and hazy. Nuclei appeared shrunken, heterochromatic, had irregular outlines and dilated perinuclear cisternae. Sarcolemma lost its scalloping appearance and appeared mostly smooth lacking its grooves of T tubules (Figs. 27 and 28).

Ultrathin sections of cardiac muscle of adult artichoke treated group (Ic) showed noticeable improvement in architecture compared with group Ib. Myofibrils seemed normal with no foci of degeneration or splitting except for few thin sparse myofibrils. Successive sarcomeres and Z lines appeared normal. Mitochondria had intact densely packed cristae, no lipid vacuoles were noticed and sarcolemma exhibited scalloped appearance similar to adult control group (Figs. 29 and 30).

Ultrathin sections of aged control group (Group IIa) revealed obvious changes compared with group Ia. Abundant large mitochondria were seen apparently occupy most sarcoplasm compared with scanty myofibrils. Many mitochondria appeared large with intact closely packed cristae while other had damaged and loosely packed cristae. Myofibrils were thin and showed obvious splitting of its myofilaments. Sarcomeres had narrow lighter A band and hazy Z lines. Sarcolemma was scalloped at Z line almost like normal. Nucleus was oval, deeply indented and had irregular outlines. It had prominent nucleolus and few heterochromatin clumps (Figs. 31 and 32).

In aged HFD treated group (IIb), ultrathin sections showed the most extensive damage and disruption of architecture of cardiac muscle compared with all other groups in the study. There was complete lysis of many myofibrils with widening of intermyofibrillar space. Other myofibrils were thin or showed splitting of its myofilaments. Noticeably, sarcomeres were disintegrated; A, I and H bands could not be distinguished and Z lines were disrupted in most sections. Simultaneously, complete damage and degeneration of many mitochondria were noticed and others were partially damaged. Sarcolemma was disrupted and lost its grooves in most sections. Nuclei exhibited irregular and indented outlines (Figs. 33 and 34).

Ultrathin sections of aged artichoke treated group (IIc) were improved compared with groups IIa and IIb. Most sarcoplasm was filled with myofibrils unlike other aged groups. Many myofibrils were cylindrical and formed longitudinal parallel arrays with narrow interfibrillar spaces but few of them were still thin and split. Their sarcomeres were improved; A, I and H bands were clearly seen and Z lines were intact. Mitochondria were partially improved; some of them exhibited intact cristae and others were still damaged. Nuclei exhibited regular outlines and had some heterochromatin clumps (Figs. 35 and 36).

Morphometric results:

1) Cell count morphometric analysis

The results of statistical analysis showed significant decrease (p < 0.05) in myocyte number of adult HFD treated group (Ib) compared with adult control (Ia) and artichoke treated (Ic) groups. On the other hand, group Ic showed significant increase in myocyte number compared with group Ib but still significantly less in myocyte count than group Ia. In aged HFD treated group (IIb), there was significant decrease in the number of myocytes compared with both aged control (IIa) and artichoke (Ic) groups. Meanwhile, Group IIc had significant increase in myocyte number compared with other aged groups. All aged groups showed significant decrease in myocyte number compared with the corresponding adult groups (Table 1 and
2) Cell diameter morphometric analysis

In adult HFD treated group (Ib), statistical analysis showed significant increase in myocyte diameter compared with adult control (Ia) and artichoke treated (Ic) groups. Aged HFD treated group (IIB) showed significant increase in the myocyte diameter compared with aged control (IIa) and artichoke (IIc) groups. Moreover, all aged groups showed significant decrease in myocyte diameter when compared with the corresponding adult groups (Table 2 and figure 2').

Table 1: The mean ± SD of cell count in the studied groups

<table>
<thead>
<tr>
<th>Age</th>
<th>Group name</th>
<th>Mean ± SD</th>
<th>Range</th>
<th>P-value</th>
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<td>Adult</td>
<td>Group Ia</td>
<td>25±2.06</td>
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<tr>
<td></td>
<td>Group Ib</td>
<td>12.86±2.04</td>
<td>9-16</td>
<td>0.000***</td>
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<tr>
<td></td>
<td>Group Ic</td>
<td>18.58±2.09</td>
<td>14-23</td>
<td></td>
</tr>
<tr>
<td>Aged</td>
<td>Group IIa</td>
<td>11.32±1.8</td>
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<tr>
<td></td>
<td>Group IIb</td>
<td>7.66±1.64</td>
<td>5-11</td>
<td>0.000***</td>
</tr>
<tr>
<td></td>
<td>Group IIc</td>
<td>17±1.57</td>
<td>14-20</td>
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Kruskal-Wallis Test * Statistical significant difference (P < 0.05)
Group Ia vs. Group IIa: 0.000***, Group Ib vs. Group IIb: 0.000***, Group Ic vs. Group IIc: 0.000***, (Mann-Whitney Test), Group Ia vs. Group Ib 0.000***, Group Ia vs. Group Ic 0.000***, Group Ib vs. Group Ic 0.000***, Group IIa vs. Group IIb 0.000***, Group IIa vs. Group IIc 0.000***, Group Ib vs. Group IIc 0.000***, Group Ia vs. Group Ib 0.000***, Group Ia vs. Group Ic 0.000***, Group Ib vs. Group Ic 0.000***, (ANOVA Test)

Table 2: The mean ± SD of cell diameter(µm) in the studied groups

<table>
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<tr>
<th>Age</th>
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<th>Mean ± SD</th>
<th>Range</th>
<th>P-value</th>
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<td>13.2±1.67</td>
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<tr>
<td></td>
<td>Group Ib</td>
<td>16.06±2.36</td>
<td>12-23</td>
<td>0.000***</td>
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<tr>
<td></td>
<td>Group Ic</td>
<td>12.86±1.71</td>
<td>10-15</td>
<td></td>
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<tr>
<td>Aged</td>
<td>Group IIa</td>
<td>9.84±1.012</td>
<td>8-12</td>
<td></td>
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<tr>
<td></td>
<td>Group IIb</td>
<td>14.16±1.59</td>
<td>12-17</td>
<td>0.000***</td>
</tr>
<tr>
<td></td>
<td>Group IIc</td>
<td>11.88±1.26</td>
<td>10-15</td>
<td></td>
</tr>
</tbody>
</table>

Kruskal-Wallis Test * Statistical significant difference (P < 0.05)
Group Ia vs. Group IIa: 0.000***, Group Ib vs. Group IIb: 0.000***, Group Ic vs. Group IIc: 0.000***, (Mann-Whitney Test), Group Ia vs. Group Ib 0.000***, Group Ia vs. Group Ic 0.000***, Group Ib vs. Group Ic 0.000***, Group IIa vs. Group IIb 0.000***, Group IIa vs. Group IIc 0.000***, Group Ib vs. Group IIc 0.000***, (ANOVA Test)
Fig. (1'): The mean ± SD of myocyte count in all studied groups

Fig. 2': The mean ± SD of myocyte diameter (µm) in all studied groups
Fig. 1: A photomicrograph of longitudinal section of left ventricle of group Ia showing cardiac muscle fibers are branching cylindrical with central vesicular oval nuclei (N) and highly acidophilic sarcoplasm (arrow). Minor spaces of endomysium are seen among muscle fibers (*). Small population of fibroblasts with flat dense nuclei are observed (arrow heads) (H & E X400).

Fig. 2: A magnified part of longitudinal section of left ventricle of group Ia showing anastomosing muscle fibers with central elongated vesicular nuclei (N) and transversely striated and highly acidophilic sarcoplasm (arrows). Endomysium is seen among the muscle fibers containing blood capillaries (*). Notice; flat dense nuclei of few fibroblasts (arrowhead) (H & E X1000).

Fig. 3: A photomicrograph of semithin section of left ventricle of group Ia showing branching cardiac muscle fibers that anastomose forming continuous syncytium. Cross striations of sarcoplasm bearing central oval vesicular nuclei (N) is noticed. Intercalated discs (arrows) are visible among adjacent myocytes and many blood capillaries (*) are seen in narrow spaces of endomysium (toluidine blue x1000).

Fig. 4: A photomicrograph of longitudinal section of left ventricle of group Ib showing small foci of muscle fiber loss (*) and local inflammatory cell infiltrate among myofibers (arrows). Loss of sarcoplasmic cross striations is noticed. There are many fibroblasts are seen among myofibers compared with group Ia (arrowheads) (H & E x400).

Fig. 5: A magnified part of longitudinal section of left ventricle of group Ib showing lost sarcoplasmic cross striations, congested blood vessels (bv) and inflammatory cell infiltration among muscle fibers (arrows). Nuclei of some myocytes appear rounded or shrunken (N) (H & E x1000).

Fig. 6: A photomicrograph of semithin section of cardiac muscle fibers of left ventricle of group Ib showing foci of cardiac myofiber loss (white arrows) and other areas have pale sarcoplasm (*) with obvious widening of intercellular spaces and encroachment of CT replacing parts of the cells (black arrows). Some nuclei appear shrunken (N). Some inflammatory cells are also visible (arrowheads) residing endomysium (toluidine blue x1000).
**Fig. 7:** A photomicrograph of longitudinal section of left ventricle of group Ic showing cardiac myocytes are improved compared with group Ib. No foci of muscle fiber loss or inflammatory cell infiltration can be detected. Nuclei of many myocytes appear more or less normal (N) but few still appear dense (arrow). Fibroblast population appear similar to group Ia (arrowhead) (H & E X400).

**Fig. 8:** A magnified part of longitudinal section of left ventricle of group Ic showing sarcomplasmic cross striations appear more or less similar to control group (arrows). Most nuclei (N) are oval and vesicular while few of them are still affected (arrowheads) (H & E X1000).

**Fig. 9:** A photomicrograph of semithin section of cardiac muscle fibers of left ventricle of group Ic showing muscle fibers appear more or less normal. They are closely packed with minor endomyial spaces in between. Most nuclei seem normal (N) and cross striations of sarcoplasm are obvious. Intercalated discs among fibers are visible (arrows). Blood capillaries mostly look normal (*) (toluidine blue x1000).

**Fig. 10:** A photomicrograph of longitudinal section of left ventricle of group Ila showing cardiac myocytes are loosely packed with widening of intercellular spaces compared with group Ia. Sarcoplasm of most fibers appear pale acidophilic and degenerated with obvious loss of its cross striations (arrows). Nuclei are dense and shrunken (N). Unlike group Ia, many flat dense nuclei of fibroblasts are observed (arrow heads) (H & E x400).

**Fig. 11:** A magnified part of longitudinal section of left ventricle of group Ila showing most muscle fibers have shrunken dense nuclei (N). Sarcoplasm of most cells is pale, degenerated and lacks its cross striations (arrows) (H & E x1000).

**Fig. 12:** A photomicrograph of semithin sections of left ventricle of group Ila showing disturbed architecture of cardiac tissue. Degeneration and lack of cross striations of sarcoplasm (arrows) and pyknosis of nuclei (N) of most cells are observed (toluidine blue x1000).
Fig. 13: A photomicrograph of longitudinal section of left ventricle of group IIb showing cardiac myocyte damage is more exaggerated than groups IIa and Ib. Many myocytes are lost with marked widening of intercellular spaces (black arrows) that harbor some inflammatory cells (#). Some fibers have wavy corrugated appearance (white arrows) 13ad&b. Focal areas of muscle fiber loss (*) are also observed 13c. Nuclei are dense and shrunken (N). Many fibroblasts with flat dense nuclei (arrow heads) are observed apparently more abundant than those noticed in sections of all other groups (H & E x400).

Fig. 14: A magnified part of left ventricle of group IIb showing marked disruption of architecture and more myofiber loss and lysis compared with groups IIa&Ib, leaving very wide intercellular spaces (arrows). Sarcoplasm of remaining muscle fibers appear pale, disintegrated and lost its cross striation (*) and the nuclei are shrunken (N). Notice; intercellular space containing some inflammatory cell infiltrate (arrow heads) (H & E x1000).

Fig. 15: A photomicrograph of semithin sections of left ventricle of group IIb showing muscle fibers have disintegrated sarcoplasm with obvious loss of cytoplasmic striations (*). Their nuclei are pyknotic (N). Marked widening of intercellular spaces, vasodilation and congestion of blood vessels and inflammatory cell infiltrate are noticed (arrows) (toluidine blue x1000).

Fig. 16: A photomicrograph of longitudinal section of left ventricle of group IIc showing cardiac muscle fibers are obviously improved compared with other aged groups. They appear branched and cylindrical more closely paced with narrower intercellular spaces (*) and having more acidophilic cytoplasm (arrows). Fewer population of fibroblasts (arrow heads) are noticed compared with other aged groups. No corrugation or focal degeneration or focal inflammation is observed. Some fibers have vesicular and oval nuclei (N) (H & E x400).

Fig. 17: A magnified part of left ventricle of group IIc showing cardiac myocytes are improved compared with other aged groups. Sarcoplasm appears more acidophilic and have cross striations (arrows). Also, some nuclei appear normal (N) while others are still affected (arrowheads). Small intercellular spaces are observed among muscle fibers (*) (H & E x1000).

Fig. 18: A photomicrograph of longitudinal section of left ventricle of group IIc showing cardiac myocytes are much improved than other aged groups. Sarcoplasm shows cross striations and some nuclei are improved (N). No blood congestion and much smaller intercellular spaces is observed among myocytes (arrow). Notice intercalated disc among myocytes (arrow heads) (toluidine blue x1000).
Fig. 19: A photomicrograph of longitudinal section of left ventricle of group Ia showing scanty collagen fibers are seen around blood vessels (arrow) and among myocytes (arrow heads) (Masson trichrome X400).

Fig. 20: A photomicrograph of longitudinal section of left ventricle of group Ib showing obvious increase in the amount of collagen fibers noticed around congested blood vessels (arrows) and among myocytes (arrow heads) compared with group Ia(Masson Trichrome stain x400). Inset: show remarkable amount of collagen fibers around blood vessels and replacing adjacent myocytes (Masson trichrome X1000).

Fig. 21: A photomicrograph of longitudinal section of left ventricle of group Ic showing decrease in the amount of collagen fibers around blood vessels compared with group Ib (arrow) and almost appear normal among muscle fibers (arrow heads) (Masson trichrome X400).

Fig. 22: A photomicrograph of longitudinal section of left ventricle of group Ila showing greater amount of collagen fibers is observed among myofibers (arrows) compared with group Ia (Masson trichrome X400).

Fig. 23: A photomicrograph of longitudinal section of left ventricle of group IIb showing increased amount of collagen fibers compared with group IIa that replacing many muscle fibers (arrowheads). Notice; marked widening of intercellular spaces (arrows) (Masson trichrome X400).

Fig. 24: A photomicrograph of longitudinal section of left ventricle of group Iic showing decreased amount of collagen fibers among muscle fibers compared with other aged groups (arrows) (Masson trichrome X400).
Fig. 25: An electron micrograph of cardiac muscle fibers of group Ia showing parallel arrays of cylindrical myofibrils (arrows) occupying most of the sarcoplasm. Each myofibril consists of repeated sarcomeres (S) separated at Z lines (Z). Abundant mitochondria (M) with intact densely packed cristae are seen among myofibrils and near nucleus. Intercalated discs (ID) are clearly seen among adjacent cells. Nucleus (N) is oval and euchromatic. Invaginations of sarcolemma are also observed (arrow heads) (X3600).

Fig. 26: An electron micrograph of cardiac myocytes of group Ia showing each sarcomere (S) is limited by two Z lines (Z) and consists of central dark A band (A) and two peripheral halves of light I bands (I). A narrow light H zone (H) is seen in the center of A band and is bisected by M line (arrow). Abundant large mitochondria (M) are observed among and parallel to myofibrils. Mitochondria appear oval and rounded in shape with densely packed intact cristae. Sarcolemma form invaginations (origin of T tubules) opposite Z lines (arrowheads). Diads (Di) are seen at the level of Z lines and glycogen granules (Gl) are seen among myofibrils. Notice; Intercalated disc is observed at the site of Z line connecting adjacent myocytes and has 3 parts; fascia adherents (F), desmosome (D) and gap junction (G) in (26b) (X7200).

Fig. 27: An electron micrograph of cardiac muscle fibers of group Ib showing intramyocellular accumulation of fat vacuoles (V). Many mitochondria have partially damaged cristae (M). Most myofibrils appear thin and split (arrows). Some foci of myofibrillar degeneration with disruption of Z lines (*) are noticed. Nucleus (N) is shrunken with irregular outline and contains many heterochromatin clumps. Sarcolemma appear smooth and lacks most of its invaginations at Z line (arrowheads) (X3600).

Fig. 28: An electron micrograph of cardiac muscle fibers of group Ib showing intermyofibillar accumulation of lipid vacuoles (V). Mitochondria have partially destroyed cristae (M). Some myofibrils appear thin and split (black arrows). Foci of myofibrillarlysis with disruption of Z line (*) are also noticed. A and I bands are not clearly seen (white arrows) and Z lines are either disrupted or appear irregular and hazy (Z). Nucleus is shrunken, have many heterochromatin clumps and exhibits irregular outline with dilated perinuclear cisternae (N) (X7200).
Fig. 29: An electron micrograph of cardiac muscle fibers of group Ic showing most myofibrils are improved (black arrows) compared with group Ib. No intramyocellular fat vaculation or myofibrillar degeneration is observed but few myofibrils still appear thin (white arrow). Successive sarcomeres and Z lines (Z) appear normal. Mitochondria have intact cristae (M). Sarcolemmal invaginations at Z lines are preserved (arrowheads) (X3600).

Fig. 30: An electron micrograph of cardiac muscle fibers of group Ic showing normal appearance of most myofibrils (arrows) that have regular arrangement of sarcomeres (S) and intact Z lines (Z). Mitochondria have intact densely packed crista (M). No lipid vacuoles or focal degeneration are noticed (X7200).

Fig. 31: An electron micrograph of cardiac muscle fibers of group IIa showing myofibrils are disorganized, thin, and their myofilaments are split and loosely packed (black arrows) and Z lines appear hazy (Z). Compared with group Ia, most sarcoplasm is apparently occupied with mitochondria with less number of myofibrils. Many mitochondria appear large with intact closely packed cristae (M). Nucleus (N) has irregular and deeply indented membrane (white arrow) with some heterochromatin clumps. Invaginations of sarcolemma are observed (arrowheads) (X3600).

Fig. 32: An electron micrograph of cardiac muscle fibers of group IIa showing some myofilaments are split (arrow heads), sarcomeres have hazy Z lines (Z) and A bands appear as narrow lighter central areas (black arrows). Many mitochondria (M) are large with intact densely packed cristae occupying most sarcoplasm while others have partially damaged cristae (*). Nucleus has prominent nucleolus (Ne), irregular and indented outline (white arrow) and contains some heterochromatin clumps (X7200).

Fig. 33: An electron micrograph of cardiac muscle fibers of group IIb showing complete lysis of many myofibrils leaving empty spaces with increased intermyofibrillar spaces (*). The remaining myofibrils are thin and split (white arrows). Many mitochondria are completely damaged leaving empty spaces (black arrows). Some mitochondria remain in sarcoplasm and mostly appear damaged (M). Sarcolemma is disrupted and loses almost its invaginations (arrow heads). Nuclei (N) have irregular outlines (X3600).

Fig. 34: An electron micrograph of cardiac muscle fibers of group IIb showing fragmentation and complete lysis of myofibrils with increased intermyofibrillar spaces (black arrows). In 34a Sarcomeres are disintegrated; myofilaments are split and degenerated (*), A, H and I bands are not clearly detected (white arrows) and Z lines (Z) are disrupted. Many mitochondria are completely damaged leaving empty spaces (arrowheads) but others are partially destroyed (M). Nucleus (N) is indented and has irregular outline (X7200).
Fig. 35: An electron micrograph of cardiac muscle fibers of group IIc showing regular parallel arrays of myofibrils with smaller interfibrillar spaces compared with group IIb but still some myofibrils are thin (black arrows). Some mitochondria appear normal (M) but others are damaged (white arrows). Nucleus (N) exhibits regular outline and has scattered heterochromatin clumps. In 35a; invaginations of sarcolemma are observed (arrow head) (X3600).

Fig. 36: An electron micrograph of cardiac muscle fibers of group IIc showing cylindrical myofibrils with regular sarcomeres, tightly packed myofilaments, A and Ib and are clearly demarcated but A band appear narrow like group Ila (black arrow). Unlike other aged groups, myofibrils apparently fill most of sarcoplasm. Some mitochondria have damaged cristae (white arrow) while others appear with intact cristae (M) (X7200).

DISCUSSION

Previous efforts have described aging related cardiac muscle changes[17]. Some researchers also have demonstrated lipotoxic effects of HFD on cardiac muscle function[18] but such studies are still limited. The coexistence of aging and obesity has been associated with serous cardiac morbidity, which is mostly fatal[17]. However, the study of associated cardiac structural alterations in these cases are not clear and need further investigations. Up to our knowledge, so far, the present study is the first detailed histological, morphometric and ultrastructural investigation that reporting high fat induced structural changes in cardiac of adult mice, in addition to, addressing and comparing the effect of adiposity in aged cardiac muscle. Moreover, the current study evaluated the efficacy of concomitant treatment of artichoke in mitigating HFD induced cardiac myopathy in both adult and aged mice. In the current study, mice model was used as commonly done in previous cardiovascular researches. Mice are considered an ideal model in the current work especially in studying aging groups, as they do not develop hypertension with aging or senescence[17]. This allowed us to evaluate aging related cardiac changes independent of vascular effects. Moreover, it has been reported that cardiac aging in mice closely recapitulates human aging process[19].

In the current study, cardiac myocytes of adult HFD treated group showed remarkable structural changes and disturbance of architecture. Intramyocellular accumulation of lipid droplets, partial destruction of mitochondrial cristae, Pyknosis and shrinkage of nuclei, and loss of sarcolemmal invaginations of T tubule system were obviously reported. Loss of sarcoplasmic cross striations, consistent with myofibrillar alterations seen by electron microscope, was also noticed. Increased fibrous tissue content of cardiac tissue and inflammatory cell infiltrate were also observed. There were diverse myofibrillar changes were recorded including thinning and splitting of myofibrils, focal myofibrillar lysis and disruption of the organization and normal appearance of sarcomeres. Statistically, myocytes of adult HFD treated mice showed significant increase in myocyte diameter associated with significant decrease in myocyte number compared with other adult groups.

Our findings are consistent with previous reports of intramyocellular accumulation of lipid droplets[20] and mitochondrial damage and cristolysis[18] noticed in hyperlipidemic rat cardiac muscle. In the current work, the observed cardiac muscle damage can be attributed to serious effects of hyperlipidemia that can be evoked at molecular and cellular levels. In this regard, previous efforts have revealed that high fat intake can lead to abnormal and inappropriate intramyocellular deposition of triglyceride in cardiac muscle, such condition is referred to be as cardiac steatosis[20]. Overaccumulation of triglyceride is associated with increased intramyocellular expression of lipogenic enzymes resulting in increased esterification and impaired oxidation of the raised pool of fatty acids[21]. Subsequently, the intracellular pool of fatty acyl-coenzyme A enlarges, providing substrate for non-oxidative metabolic pathways that lead to oxidative stress, inflammation, apoptosis, fibrosis and cellular dysfunction[18,22,24].

Interestingly, the current statistical analysis of adult HFD group showed significant decrease in myocyte number concomitant with significant increase in myocyte diameter, which indicates cardiac muscle loss and hypertrophy respectively. The present results are similar to recent reporton myocardium of obese mouse[22]. As explained in previous studies, lipotoxicity is characterized by both cardiomyocytes loss as a consequence of increased apoptosis with fibrotic tissue replacing lost cells and hypertrophy of the remaining myocytes[23,24]. In fact, myocyte loss is associated with diverse cardiac dysfunction.
mainly decrease in contractility, arrhythmia, wall dilatation. Cardiac arrhythmia can be attributed to disruption in gap junctions and electrical coupling as a result of lost myocytes.[27] In this regard, we reported obvious impairment of myofibrils and their constituting myofilaments in adult hyperlipemic mice, which represent the contractile machinery of cardiac muscle. Accordingly, we can add that at subcellular level, the observed myofibrillar loss and disruption can also contribute to obesity associated cardiac contractility disorders. Earlier efforts recorded impaired myocardial contractility and function in hyperlipemic rats. The authors owed the observed cardiac dysfunction to activation of apoptotic pathway "lipoapoptosis" secondary to increased ceramide level, a mediator of apoptosis, with subsequent increased expression of nitric oxide synthase "iNOS" and overproduction of nitric oxide "NO" resulting in impaired cardiac contractility and function.[20].

In the present work, a remarkable finding in myocardium of adult hyperlipemic mice is the increase in amount of collagen and fibroblast pool, which indicates cardiac fibrosis. Fibrosis ultimately contributes to more deterioration and loss of cardiac function secondary to obesity. Zhou YT and his colleagues (2000)[29] reported significant increase in the ratio of hydroxyproline amount compared to the total protein content of cardiac tissue in adult obese rats, which is consistent with our record. It was concluded that cardiac steatosis, like hepatic and insular steatosis, is ultimately followed by progressive fibrosis[20].

In the current study, obvious changes were detected in aged cardiac muscle compared with adult myocardium. There was significant decrease in cardiac myocyte number and diameter compared with adult control group. Myofibers became loosely packed with widening of intercellular spaces. Lack of sarcoplasmatic reticulum, apparent increase in fibroblast population and increase collagen fiber content were noticed. At ultrastructural level, many large mitochondria, some of them with partially damaged cristae, were occupying most of sarcoplasm compared with apparently scanty number of myofibrils. Thinning and splitting of myofibrils, and disruption of sarcomeres were reported but sarcosomal T tubules invaginations were still preserved. Indentation, shrinkage and pyknosis of nuclei was also noticed. Similar observations were previously reported in aged myocardium in both humans and rats[28]. In the current work, we observed widening of intermyofiber spaces, obvious increase and encroachment of fibrous tissue that replacing adjacent myocytes associated with significant decrease in myocyte numbers of aged myocardium. Previous histological analysis has revealed that aging is associated with progressive cardiomyocytes loss due to both apoptotic and necrotic cell death[20]. Furthermore, Anna B and Nikolas G (2011) have demonstrated that aging is characterized by continuous and progressive circuit of cardiac fibrosis and myocyte loss. They hypothesized that the thickening of the extracellular matrix occur around aged cardiomyocytes results in mismatch between supply and demand of nutrients, and leads to myocyte death. Intersitial fibroblasts respond to signals released by dying cardiomyocytes by synthesizing new matrix aimed at replacing the damaged cells resulting in more myocyte loss. Interestingly, previous researches on aged myocardium have used the term cardiac sarcopenia to describe the aging related continuous decline in myocyte numbers which associated with reduced cardiac function mainly contractile performance[31]. Our observations of myocyte loss are also in accordance with these reports. Moreover, at ultrastructure level and unlike adult control myocardium, we reported apparently scanty abnormal myofibrils within the aged cardiac myocytes compared with more abundant pool of enlarged mitochondria that occupy most sarcoplasm. In fact, our observation of obvious disturbance in contractile elements of aged myocytes in addition to their reduced numbers can both contribute to aging related reduced cardiac function. Collectively, for the first time, we can suggest that, the term aging related sarcopenia should not only used to refer to and to describe the aging related decrease in myocyte numbers but should also include the accompanied decrease and disruption in myofibrils and contractile machinery within aged myocytes seen at ultrastructure level.

In the present study, aged myocytes showed significant decrease in myocyte diameter compared with adult group, which is opposite to previous reports of myocyte hypertrophy in aged myocardium[30,31]. Actually, our record showed that all aged groups had significant decrease in both cell diameter and number when compared with the corresponding adult groups. In this regard, our finding of apparently abnormal large abundant mitochondria which fill most sarcoplasm compared with scanty myofibrils may provide an evidence of the decrease in myocyte diameter. Aged cardiac myocytes seem under stress due to reduction in their number, and size, and contractile elements so they need more energy and effort trying to maintain their normal function, thereby there was abnormal compensatory increase in size and number of mitochondria in aged cardiac myocytes. Unfortunately, mitochondria is vulnerable to mitochondrial oxidative stress with aging that is associated with increased production of reactive oxygen species "ROS". This could explain the noticeable partial crisisolysis of some mitochondria in the current work. This fact has been confirmed by previous report of increased production of superoxide radicals from mitochondria extracted from aging rat hearts[32]. In addition, another study on mice has documented increased mitochondrial DNA mutations and deletions, and has concluded that the production of mitochondrial ROS contributes to cardiac myocyte damage with aging[32].

An important finding in our study of aged myocardium, is the apparent increase in collagen fibers and fibroblast pool, which indicates myocardial fibrosis. Earlier efforts have documented increased collagen deposition and aging associated myocardial fibrosis[30,31]. Indeed, the
have documented that the coupling of aging and obesity myocaridan damage. In this regard and in accordance which ultimately can lead to death due to extensive profound and progressive deterioration in cardiac function, aging together is highly serious situation that can lead to accentuate and aggravate myocardial damage. In other the coexistence of aging and high fat diet synergistically arrangement and appearance were obviously disrupted. The profound myocyte loss was associated with marked decrease in myocardium changes. We noticed maximum damage and disruption of architecture of myocardium in aged HFD treated group when compared with the adult HFD treated and control aged mice. Statistically, there were significant decrease in myocyte number compared with other aged groups and adult HFD group. The myocyte diameter was significantly increased compared with other aged groups but significantly smaller than that of adult HFD group. The profound myocyte loss was associated with marked widening of intermyofiber space, corrugation, focal degeneration and loss of cross striations of myocytes. In addition, local inflammatory cell infiltrate, shrinkage, irregularity and pyknosis of nuclei, extensive deposition and displacement of connective tissue of adjacent myocytes and marked increase in fibroblast population were observed. At ultrastructural level, there was pronounced myofibrillar lysis and degeneration accompanied by simultaneous complete destruction of many mitochondria so many regions of sarcoplasm appeared empty. The remaining myofibrils were split and thin and sarcomere arrangement and appearance were obviously disrupted. Taken together, these observations provide evidence that the coexistence of aging and high fat diet synergistically accentuate and aggravate myocardial damage. In other words, we can hypothesize that coexistence of obesity and aging together is highly serious situation that can lead to profound and progressive deterioration in cardiac function, which ultimately can lead to death due to extensive myocardial damage. In this regard and in accordance with our observation and conclusion, previous reports have documented that the coupling of aging and obesity factors contributes to increase the production of ROS[34]. Other efforts have added that the aging process enhances the metabolic and vascular effects evoked by HFD through augmenting oxidative stress and increase the production of ROS[35]. Furthermore, a recent study in skeletal muscle showed loss of myonuclei and myofiber area in HFD aged mice and the authors hypothesized that high-fat feeding in conjunction with aging generates signals that induce myofiber atrophy, most probably through ubiquitin proteasome system[36]. In fact, further investigations are still needed and highly recommended to define the underlying mechanisms behind the synergistic destructive effect of both aging and obesity on myocardium that we noticed in the current study.

Interestingly, the most striking findings noticed in the current work, were related to the impact of HFD on aged myocardium. Indeed, up to our knowledge, the current study is the first reporting of the impact of HFD on aged myocardium at myocellular level. We reported that the coexistence of adiposity has exaggerated the aging induced myocardium changes. We noticed maximum damage and disruption of architecture of myocardium in aged HFD treated group when compared with the adult HFD treated and control aged mice. Statistically in both adult and aged artichoke groups, myofibrils appeared occupying most of sarcoplasm. Compared with other aged groups, myofibrils and sarcomeres exhibited normal appearance and organization. Compared with other aged groups, myofibrils appeared occupying most of sarcoplasm. Mitochondria were improved, more obvious in adult mice. Statistically in both adult and aged artichoke groups, there were significant increase in the number of cardiac myocytes associated with significant decrease in cell size compared with the corresponding HFD groups. Indeed, the noticeable improvement of myocardium in the current study reinforces previous reports done on other organs.
Artichoke was found to be effectively and significantly decrease the level of cholesterol and triglyceride and also have hepatoprotective effect in both human and animal[38]. According to the current observations, we can suggest that artichoke has both powerful hypolipidemic and antiaging properties on cardiac muscle, which could be attributed to its antioxidant properties. In accordance with that, earlier investigations have demonstrated that artichoke has powerful antioxidant effects[9]. It has been found to be highly effective in decreasing the production of reactive oxygen species, the oxidation of low-density lipoproteins, lipid peroxidation and protein oxidation, in addition to scavenging free radicals before they can damage cells[40]. In this regard, it is found that artichoke is highly rich in many antioxidant ingredients such as mono- and dicaffeoylquinic acid (cynarin and chlorogenic acid), caffeic acid and flavonoids including the glycosides luteolin-7-β-rutinoside (scyoside), luteolin-7-β-D-glucoside, and luteolin-4-β-D-glucoside with chlorogenic acid is reported as the most active antioxidant ingredient[41]. Actually, recent efforts have investigated the possible underlying mechanisms behind the hypolipidemic nature of artichoke[8,42]. It has been found that artichoke can interact with Hydroxy-Methyl-Glutaryl-Coenzyme A (HMG-CoA) reductase, as well as with liver sterol regulatory element-binding proteins (SREBPs) and acetyl-CoA C-acetyltransferase (ACAT), leading to reduce cholesterol level[43]. Moreover, artichoke might reduce hypercholesterolemia by increasing the fecal excretion of bile acids and also by directly influencing biosynthesis of cholesterol[38,43]. Indeed, the current reporting of obvious and remarkable improvement in the myocardium of both adult and aged artichoke groups may give very promising insights for using artichoke as natural hypolipidemic and antiaging agent that is effective in protecting and/or treating obesity and/or aging associated myocardial diseases.

CONCLUSION

Our findings demonstrated that aged myocardium is more susceptible to and has more extensive lipotoxic damage compared with adult one. Lipotoxic insults include disruption of powerhouse and myofibrillar contractile machinery, thereby, could impair cardiac function. There could be synergistic cross talk between obesity and aging that leads to profound cardiac damage. Further investigations are highly recommended to explain the underlying mechanisms in case of such coupling. Concomitant treatment of artichoke is markedly improved myocardial damage in both adult and aged mice so it could be used as an effective cardioprotective against hyperlipidemia and aging insults.

CONFLICT OF INTEREST

There are no conflicts of interest.

REFERENCES


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المملوكة: إن تواجد فرط الدهون بالدم والشيخوخة معا من أشد الحالات خطورة والتي تؤدي إلى تدهور هائل في وظيفة القلب. يعتبر الخرشوف أحد مضادات الأكسدة المعروفة والتي يشيع استخدامها في النظام الغذائي للبلدان طويلة العمر. وقد أظهرت الأبحاث الحديثة مدى فاعليته كخافض للدهون وكذلك مضاد للشيخوخة.

الفوانيس: يهدف هذا العمل إلى استكشاف الآثار المرضية للحمية عالية الدهون على عضلة القلب لدى الفئران البالغة، وكذلك تحديد الهدف ومقارنة ذلك بمدى تلف عضلة القلب لدى الفئران المسنة وتقييم الآثار المفيدة للخرشوف في تخفيف هذه الآثار الضارة.

الطرق والمواد المستخدمة: استخدم في هذه الدراسة عدد 42 من الفئران البالغة (المجموعة الأولى) والمسنة (المجموعة الثانية) والتي بلغ من العمر 3+26 شهرا على التوالي وقد تم تقسيمها إلى ست مجموعات فرعية متساوية (7 فئران لكل مجموعة): المجموعات المختبرية: IIb و Ib المجموعة المعتمدة على حمية عالية الدهون والتي تتغذى على وجبات مزودة بنسبة 10% من الدهون و10% من زيت شحم الخنزير الذي تم تغذية الفئران البالغة والمسنة بجرعة 1.5 جم / كجم مرة واحدة يوميًا عبر أنبوب معد.

النتائج: أظهرت خلايا عضلة القلب للفئران البالغة المعتمدة على حمية عالية الدهون فقدانها المخططات العضلية وزيادة الكولاجين، وكذلك تحلل بؤري للخلايا العضلية، وتكون الليفية لدى الخلايا الهيكلية وتكون الفجوات الشقوقية. وكانت تأثيرات الخرشوف على هذه التغييرات في البيئة عالية الدهون. وقد أدت تناول الخرشوف إلى تحسين هذه التأثيرات بشكل كبير بفضل تبكية عالية في تحسين هذه التأثيرات.

الاستنتاج: قد أدى تناول الخرشوف للحمية عالية الدهون والشيخوخة إلى حدوث تأثيرات تعتبر الأشذ والأكثر آثرا على خلايا عضلة القلب. وقد وجد أن الخرشوف ذو كفاءة عالية في تحسين هذه التأثيرات.