Histopathological Effects of Anabolic Androgenic Steroids (Nandrolone Decanoate) on Heart, Liver and Kidney of Male Local Rabbits

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Abstract

Nandrolone decanoate consider as one of the group of anabolic steroids used by athletic with high dose to enhance athletic ability and build muscles. It was found that the use of high dose of these compounds affects the heart, liver and kidneys. Therefore, this study aims to research the histopathological outcome of the drug Nandrolone on the heart, liver and kidneys in adult rabbits. Six adult rabbits were used in this research, and these animals distributed into 2 groups as 3 rabbits in each. The group one considered a control, the second group was given a dose of (10 mg/kg) of nandrolone decanoate IM injection for (15 consecutive days). After (24) hours of giving the last dose, the rabbits were slaughtered and samples were taken from the heart, liver and kidney, and they were prepared for histological study by light microscopy. This study showed clear alteration in the structure of the heart muscle, represented in the infiltration of inflammatory cells, vacuolar degeneration, Zenker’s necrosis and vascular congestion. In the liver also observed inflammatory cells infiltration, vacuolar degeneration, hyperplasia of lining of bile duct, fibrosis and sinus congestion. In the kidneys, vacuolar degeneration, atrophy and congestion of the glomeruli, expansion of the urinary space and cysts formation were noted. We conclude from the current study that the drug nandrolone has a harmful effect on the histological composition of the heart muscle, liver, kidney in adult rabbits, so the current study recommends limiting the excessive use of this drug.

Keywords: Androgenic anabolic steroid, Liver, Heart, Kidney, Rabbits.

Introduction

Anabolic androgenic steroids (AASS) are a kind of synthetic compounds which is resemble testosterone produced naturally by the body [1]. Young people frequently use AASS to increase muscular tone or growth for cosmetic reasons and sports performance [2, 3]. Research have examined the potential negative effects of using or abusing AASs in children [4, 5]. Although anabolic steroids can be utilized for a variety of medical conditions, the use is typically linked by a number of negative side effects. Curative, dose seem to have less side effect, however supraphysiological doses are linked to significant and severe side effect, hence these side effects are typically dose – related [6]. AAS use for a long period of time may cause aberrant endogenous hormone release, leading to either reversible or irreparable harm.

Increased oil secretion from sebaceous glands which results in dermatological conditions such as the acne vulgaris, cases of androgenic alopecia, and the hypertrichosis, is the most frequent adverse effect of AAS [7]. AAS can also cause several sex issues, including male infertility, importance a zoospermia, and testicular atrophy [8, 9]. Abuse of AAS in women can lead to changes in voice, amenorrhea, uterine atrophy, and cortical hypertrophy [9]. AAS abusers are substantially more likely to suffer tendon ruptures, which can be fatal to an athlete’s career [10]. AAS dependence syndrome, headache, irritability, and depression are among the psychobehavioral disorders that long term use of AAS is likely to experiment [11]. In some situations, resulting in violence and suicide [12]. An excessive oral AAS burden can result in damage of the liver or the kidney, as coagulation malfunction, the liver fibrosis, the renal hypertrophy and failure.

A too much oral AAS load is processed via the hepatorenal damage [12,13]. Abuse of AAS also raises cardiovascular risk and poses a major threat to user’s safety (7, 14). Nandrolone decanoate is class II

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androgenic steroids that rapidly spread in the world and clinically used and illicitly composed of 19–nortes–tosterone–derivates [15].

The enzyme 5α-reductase mostly breaks down nandrolone decanoate into 5α-dihydronandrolone, 19–noretiocholanolone. Urine tests can find these metabolites [16]. Despite therapeutic benefits, nandrolone usage that is uncontrolled and prolonged leads to negative effects such as liver toxicity, thyroid function changes, and cardiovascular toxicities [17].

The objective of current study is to find out the impact of nandrolone decanoate management on heart, the liver and the kidney in male local rabbits.

Material and Methods

Ethical Approval

The present research was carried out in the animal house and the scientific research laboratory after approving by the study ethics committee and scientific committee / Dental Basic science / Dentistry College / University of Mosul. No. UOM. Dent.23/30 in 31/5/2023.

Experimental Design

Six male rabbits weighing (1.5–2 kg) and (10–12 months) old made up the sample. Veterinarian will examine them both before and after the operation to assess their condition and overall health. They were given the standard rodent and kept in separate cages with tap water. Animals distributed into two categories at random, with three rabbits in each group.

Group 1: Control group, animals treated by D.W intramuscular injection and served as control.

Group 2: (ND) treated group. The rabbits give nandrolone decanoate(10 mg / kg), IM injection once per day for (15 days).

Tissue harvesting procedures

Animals were sacrificed at the end of the experiment, their organs the heart, the liver and the kidney removed; samples measuring (0.5cm^3) were taken, fixed in 10% neutral formol for 24 hours, washed dehydrates increasing alcohol concentrations, cleared in xylene, and then embedded in paraffin.

After that, specimen sectioned at a thickness of 5 M. The tissues section stained by HandE stain, using the light microscope for histopathological analysis.

Results

After 15 days of the experiment, the histological alterations of the heart, the liver and kidney examined in all groups of rabbits employed in the current investigation.

Control Group

The Heart

The findings of microscopic analysis of cardiac tissue sections from control group rabbits, as well as the typical histological structure of myocardial fibers. The myocardial cytoplasm, a central oral nucleus, and intracellular space with capillaries as show in Fig. (1).

![Microphotograph showing a section of heart tissue of rabbits in control – note elongated cardiac muscle fibers (F) and a central oval nucleus (N) : HandE 400X.](image)

The Kidney

Microscopic analysis of kidney tissue sections from rabbits in the control group revealed that their kidneys had a normal histological structure. The included blood arteries, tubules, interstitial, tissue, and renal corpuscles...
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Large pyramidal cells with an acidic cytoplasm and a big, rounded nucleus line the proximal renal tubules. Microvilli called brush boundary emerge into the tubule lumen from the cells surfaces. The distal tubules in contrast, have a broader lumen, shorter cells, and less acidic cytoplasm as shown in Fig. (2).

Fig. 2. Microphotograph showing a section in kidney tissue of the control rabbits, note renal glomerulus (G), Bowman’s space (BS), Proximal convoluted tubules (P), distal convoluted tubule (D). HandE 100X.

The Liver

Microscopic analysis of liver tissue slices from rabbits in the control group revealed that the liver had a normal histological structure. Hepatic cords or plates radiate from the central vein and are made up of relatively large cells with polyhedral shaped (ribs) arrangements. Contain spherical nuclei and cytoplasm that is acidic. Blood sinusoids that run into the central vein and are bordered with rows of epithelial and kupffer cells separate the hepatic cords as shown in Fig. (3).

Fig. 3. Microphotograph showing a section of liver tissue of control rabbits, shows central vein, with Sinusoids, the hepatic cords, kupffer cells, Hand E 100X.

Anabolic androgenic Steroids (Nandrolone Decanoate)

The Heart

Microscopic analysis of cardiac tissue sections from rabbits treated with anabolic steroids from revealed the development of inflammatory cell infiltration, vacuolar degeneration, and zinker necrosis, as shown in the Fig. (4). As shown in the Fig. (5), further signs of damage include the infiltration of inflammatory cells. Vacuolar degeneration, clogged blood vessels, and Zinker necrosis in the other location. Vacuolar degeneration and blood
vessels congestion were also seen in another location, as shown in the Fig. (6). Infiltration of inflammatory cells vacuolar degeneration, blood vessel congestion, and Zinker necrosis were also seen in some area as shown in figures (7, 8).

Fig. 4. Microphotograph showing a section of the heart tissue of the Nandrolone decanoate group rabbits, note: mild infiltration of inflammatory cells (a), vacuolar degeneration (b), and zinker necrosis (c), HandE, 400X.

Fig. 5. Microphotograph showing a section of the heart tissue of the Nandrolone decanoate group rabbits, note: mild infiltration of inflammatory cells (a), vacuolar degeneration (b), blood vessels congestion (c) and zinker necrosis (arrow), HandE, 400X.

Fig. 6. Microphotograph showing a section of the heart tissue of the Nandrolone decanoate group rabbits, note: vacuolar degeneration(a), blood vessels Congestion (b). HandE stain, 400X.
Fig. 7. Microphotograph showing a section of the heart tissue of the Nandrolone decanoate group rabbits, note: mild infiltration of inflammatory cells (a), vacuolar degeneration (b), blood vessels congestion (c) and zinker necrosis (arrow). HandE stain, 400X.

Fig. 8. Microphotograph showing a section of the heart tissue of the Nandrolone decanoate group rabbits, note: mild infiltration of inflammatory cells (a), vacuolar degeneration (b). HandEstain, 400X.

The Kidney

As can be seen in the figures (9, 10, 11) Microscopic analysis of kidney tissue sections from anabolic steroid exposed rabbits revealed numerous histological alterations, such as vacuolar degeneration, glomerular atrophy, enlarged urinary spaces, and kidney cysts. Mild atrophy, glomeruli congestion, and cyst formation were also seen in another location, as shown in Fig. (12).

Fig. 9. Microphotograph showing a section of the kidney tissue of the Nandrolone decanoate group rabbits, note: mild vacuolar degeneration(a), atrophy of glomeruli (b) urinary space dilatation (arrow) and cyst kidney (c), HandE stain, 400X.
Fig. 10: Microphotograph showing a section of the kidney tissue of the Nandrolone decanoate group rabbits, note: vacuolar degeneration (a), atrophy of glomeruli (b), urinary space dilatation (arrow) and cyst kidney (c). HandE stain, 400X.

Fig. 11. Microphotograph showing a section of the kidney tissue of the Nandrolone decanoate group rabbits, note: atrophy of glomeruli (a), urinary space dilatation (arrow) and cyst kidney (b). HandE stain, 400X.

Fig. 12. Microphotograph showing a section of the kidney tissue of the Nandrolone decanoate group rabbits, note: mild atrophy and congestion of glomeruli and cyst formation (b). HandE stain, 400X.

The Liver
Microscopic analysis of liver tissue sections from rabbits given anabolic steroids revealed hyperplasia, severe vacuolar degeneration, necrosis, and infiltration of inflammatory cells, as shown in Fig. (13).
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Fig. 13. Microphotograph showing a section of the liver tissue of the Nandrolone decanoate group rabbits, note: infiltration of inflammatory cells (a), severe vacuolar degeneration (b), necrosis (c), and hyperplasia of bile ductule (arrow). HandE stain, 100X.

As seen in the figures (14). There was also fibrosis present in that location, along with severe vacuolar degeneration, hyperplasia of the bile ducts, and infiltration of inflammatory cells. There was also modest infiltration of inflammatory cells in the portal area, diffuse vacuolar degeneration, and hyperplasia of the bile duct in some area of the liver tissue, as shown in figures (15). Also as shown in the figures (16). There was sinusoidal congestion.

Fig. 14. Microphotograph showing a section of liver tissue of Nandrolone decanoate group rabbits, note: infiltration of inflammatory cells (a), severe vacuolar degeneration (b), fibrosis (c) and hyperplasia of bile ductule cells. HandE stain, 400X.

Fig. 15. Microphotograph showing a section of liver tissue of Nandrolone decanoate group rabbits, note: infiltration of inflammatory cells in portal area (a), diffuse vacuolar degeneration (b), and hyperplasia of bile ductule (arrow). HandE stain, 400X.
Discussion

Although anabolic steroids can be utilized for a variety of medical conditions, using is typically combined with a number of negative side effects. While therapeutic levels seem to have few side effects, but supraphysiological doses are linked to significant and severe side effects, these adverse effects are typically dose- 

relates [6].

According to National Institute on medicine abuse [18]. Nandrolone Decanote (ND) is one of most popular testosterone anabolic derivatives since it has a moderate androgenic potential and the best anabolic qualities [19]. The heart muscle fibers in this investigation had sever ischemia degradation together with obvious inflammatory infiltration. This may be the reason athletes who use AAS experience deadly arrhythmias. ASS nandrolone in malerate impacted left ventricular remodeling in (2000), according to Woodimiss and others [20], Without causing heart muscle injury. This might be related to the lower dosage they utilized.

Androgens operate by receptor – dependent, means receptors are current in cardiovascular system, in human vascular endothelium, smooth muscle tissue, macrophage, cardiac myocytes, could be linked to the mechanisms of AAS promotes cardiac muscle injury. Moreover, testosterone directly affects transcription translation, and enzyme function in the myocardium as it is the preferable binding of the androgen receptor (21). In the physiological dose range, AAS has a protecting impact on the cardiovascular system. Nevertheless, supraphysiological AAS doses result in cardio-vascular system toxicity, which sharply raises the risk of cardiovascular disease. The exact cause of AAS toxicity is still unclear. A cording to studies, there are two main mechanisms. One of them is AAS gene regulation, in which AAS or their metabolites bind to AAS and cause these receptors to change conformation [20,21].

The study recorded sever ischemic damage of heart muscle fiber with inflammatory signs, this may be cause of fatal case of arrhythmias in the athletes use AAS [22]. Several research have revealed and suggested that the toxicity of AAS is connected to oxidative damaging and tissue damage [22], by modification of the mitochondrial respiratory chain complexes activity (23). Causing reactive oxygen species ROS, and causing harmful effective on cell. Many reactive oxygen species (ROS) produced can cause oxidative damage, which can injury the cells and eventually lead to cell death and a variety of degenerative diseases [24]. Cytochrom p–450 mono–oxygenses metabolize high dosages of nandorolne decanotae. Reactive oxygen species are produced as a result, which increases the activity of antioxidant enzyme until they are exhausted [25].

Moreover, anabolic steroids boost the lipase enzyme’s action, which increase the availability of long chain fatty acids for mitochondrial oxidation and ATP productions resulting in the developing of lipid peroxidation and the formation of ROS [26]. mechanism for AAS produce cardiac muscle injury by receptor dependent and receptors – independent mechanisms. the Androgen receptor in cardiovascular system, include the human vascular endothelium and smooth muscle fibers, also the cell of macrophages and cardiac myocyte cells. Testosterone is ligand preferred of human androgen receptor in the myocardium and directly modulates transcription, translation, and enzyme function. AAS cause damaging associated with the apoptosis [27, 28, 29]. AAS has a protective impact on cardiac – vascular system in physiological doses. Suprophysiological AAS dose cause toxicity in cardiac vascular system, which substantial increase cardiovascular injury. The mechanisms of toxicity of AAS not fully detected.
Others found two main mechanisms, the first one was the regulation of AAS genes, it means AAS or its metabolite link to ARS, that cause a case of conformational alteration of the receptor. AR dimers then will transport from cytoplasm into nucleus where dimers combined by androgen reacting element in DNA, then regulate the gene transcription in cooperative by active (or in activation) pf coregulations, which result in toxic effects [30]. Other non–gene regulation mechanisms (in the skeletal muscle cell and the prostate cancer cells) is the AR dimers combined by the cytoplasmic protein by a different of signaling pathway which is mainly produce the toxicity no gene transcription pathway[30], toxicity of AAS is connected to a range of aspect like doses, cycle and individual difference.

In (2005), Du Toit et al. [31]; recorded that the supraphysiological dose of AAS, which is taken throughout the exercise training or according to inactive condition elevated myocardial susceptibility to the ischemia or reperfusion damage. The increasing sensitivity linked to steroid – induce increasing in pre–ischemic myocardial CAMP concentration and increasing in both of pre–ischemic and reperusions of the tumor necrosis factor TNF-α concentration. Kahal and Allem. [31], recorded in this study enlargement, sever degenerate ion and the heart muscle damage. The findings support Hassan et al. [32]. Study, which found that " As sustaining produces significant ischemia necrosis and degeneration the heart muscle fibers in male albino rats ". Moreover, Elgend et al. [33], informed on hypertrophy and degenerate of both of cardiac muscle and skeletal muscle, explain by citing its effect on extensively distribute androgen receptor in many type of muscles.

Almost all anabolic steroids experience many side effects as the liver toxicity [34]. Because of the secrecy around the use of large doses of androgens, limited data on the relationship of adverse effects to doses or substances used are available. While some symptoms are caused by androgenic or estrogenic actions, others may be caused by the toxic (non-hormonal) consequences of the androgen or its metabolites, particularly when the dose is very high [35]. More hepatocytes were harmed by vascular degeneration, particularly those near the central vennles, since these cells were more exposed to hypoxia than those near the portal area [36]. Vacular degeneration affected nearly all hepatocytes, and areas of localized necrosis were frequently infiltrated with monoclear inflammatory cells. This is consistent with the findings of Boada, who discovered centriflobular degeneration and lipid vacuolization in male rats given large doses of anabolic androgenic steroids. Vieira et al. [37], discovered that subchronic treatment with nandrolone decanoate, primarily at higher doses, is potentially toxic to the liver and causes fibrosis. [38].

A cause of spontaneous hepatic rupture with life-threatening bleeding has been linked to a history of anabolic steroid usage. In (2011), Tousson et al., [39,40] showed that sinusoid and centrolobular necrosis occurs as a result of anabolic androgenic steroids’ ability to cause hepatocyte injury via their effect on mitochondria, specifically the mitochondrial membrane, and inhibit mitochondrial respiration, resulting in mitochondrial swelling and ALT and AST leakage. This study found fibrosis in the portal area, which means that the liver goes to healing the damaged cells caused by fibrosis in the portal area rather than regeneration of other hepatocytes because the damage is powerful and strong, and regeneration may take more than 15 days because anabolic steroids remain in the tissue after administration is stopped [41]. Proposed mechanisms of liver injury include anabolic steroid-induced infiltration of inflammatory cells in the hepatic tissue and activation of kupffer cells, which results in the production of inflammatory cytokines and collagen deposition; increased oxidative stress and reactive oxygen species, which leads to mitochondrial degeneration in liver cells; and stimulation of intracellular androgenic steroid receptors, which leads to uncontrolled growth of hepatocytes [30].

An in vitro study published in 2021 found that mitochondrial respiratory chain processes are disrupted early in the pathophysiology of liver injury caused by supraphysiologic dosages of nandrolone, an anasynthetic AAS extensively used for performance enhancement. It particularly inhibits respiratory chain complexes I and III, causing reactive oxygen species buildup and oxidative damage [42]. Animal investigations revealed a decrease in glutathione and enzymes that act as free radical scavengers (superoxide dismutase and catalase), as well as an increase in thiobarbituric acid - a reactive compound indicating poor redox equilibrium [34, 44].

Oxidative stress disrupted mitochondrial membrane potential dynamics, resulting in decreased cell energy supply, hepatocyte dysfunction, and necrosis [45]. Chronic AAS use was known to produce a number of changes. Renal illnesses have received less attention among these conditions, most likely because they are less common and hepatotoxicity has been thoroughly studied [46, 47].

Nephrotoxic side effects have been documented, such as renal failure or Wilms tumours [48, 49]. AAS illicitly used by athletes for decades to increase muscle mass and decrease body fat [50], are emerging as podocyte toxins in long term abusers [48]. Until now, the effects of AAS on kidney structure have often included the following: glomerular and tubular atrophy, vascular degeneration, congestion, necrosis, interstitial fibrosis, inflammatory interstitial nephritis, and segmental glomerulosclerosis [51, 49].
The current investigation found significant histological damage to the rabbit renal cortex after 2 weeks of AAS treatment. Congestion in both glomerular capillaries and interstitial blood vessels has been practically universal in this group. Similarly, Kahal and Allem, 2018 reported congestion of renal blood vessels and interstitial hemorrhage following AAS a base in adult male mice and added that it persisted and even worsened after stopping treatment [50]. The observed congestion could also be attributed to the AAS- that induced PCT which increase in width and luminal deprivation [51,52], congest ion was accompanied by a rise in PCT diameter and obstruction of their Lumina by swelling cells and ischemia-associated cell debris.

A recent study found a relationship between glomerular damage and tubular interstitial injury, with podocyte damage causing substantial plasma protein leakage into the tubules. These proteins were reabsorbed by the PCT, resulting in the release of mediators such as monocyte chemotactic protein type 1, platelet-derived growth factor, and endothelin into the interstitium. These mediators cause an inflammatory response that results in interstitial fibrosis and tubular degeneration [53]. Similarly, Hasso, [45], observed vacular degeneration of the PCT lining epithelium with sloughing and necrosis of the epithelial cells and engorgement of numerous interstitial blood vessels in male rabbits following ND treatment. According to Sood et al. [55], necrotic nuclei and brush border separation indicate functional impairment of urine reabsorption.

This necrosis could be linked to mitochondrial injury and subsequent cellular ATP depletion [55]. Furthermore, mitochondrial damage can result in intracellular oxidative stress and higher ROS levels due to disruption of the mitochondrial electron transport chain [56]. Benu et al., [57]. Suprophysiological dosages of ND increase the production of inflammatory cytokines, interleukin-1 beta (1L-713), and tumor necrosis factor-β (TNF-β) in human peripheral blood lymphocyte cultures in vitro [57].

TNF - has been identified as a cytotoxic that produces apoptosis in numerous cells, including renal tubular cells, either through association with its membrane - bound receptor or by triggering mitochondrial membrane disruption and cytochrome C release [58]. Long-term administration of ND. resulted in a decrease in antioxidant enzyme activity, which resulted in oxidative renal cell injury due to its decreased ability to scavenge harmful hydrogen peroxide and lipid peroxides [59].

Anabolic androgenic steroids may contribute to the development of CKD by generating reactive oxygen species and causing oxidative stress [60]. Testosterone can increase endotheline synthesis directly or indirectly (via RAAS). As a result, testosterone may have a role in both icnedy fibrosis and ischemia-reperfusion injury via both local and systemic endoltheline effects [61]. Other laboratories have shown that renal tissues are more sensitive to oxidative tissue injury due to the high content of long-chain polyunsaturated fatty acids [62]. Furthermore, extremely high metabolic rates result in excessive free radical production [63]. Renal tissue damage is caused by reactive oxygen species produced by kidney cells or toxic substances introduced to the kidney, both of which contribute to a number of nephron-degeneration disorders [64]. Androgen receptors (ARS) are expressed in many human tissues, including the kidney. AAS operates by binding to these receptors to boost endogenous testosterone synthesis. The increased testosterone production has an effect on the glucocorticoid receptors, which decrease glucose synthesis and protein catabolism. As a result, renal tissue expands and distorts [65- 67]. Memudu and Dongo, recently reported that the pathophysiology of nephron toxicity caused by a hazardous chemical is characterized by increased lipid peroxidation and decreased SOD level in the presence of AAS-induced oxidative tissue damage, which resulted in the loss of renal tissue cell membrane integrity [68].

More research is needed to overcome this study's limitations and examine the impact of anabolic steroids on overall health.

Conclusion

In rabbits, nandrolone discount at a dose of 10 mg/kg for 15 days had a notable effect on the heart, liver, and kidney. It can cause significant disruption in tissue architecture.

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The effects of anabolic androgenic steroids are well documented, with studies showing their impact on various organs in the body. In a recent study published in *Archives of Renal Diseases Management*, researchers investigated the histopathological effects of anabolic androgenic steroids on renal morphology in rats. The study concluded that these steroids had a detrimental effect on renal tissue, leading to changes in renal architecture and function. These findings are significant as they highlight the potential risks associated with the use of anabolic androgenic steroids, particularly in the context of sports performance enhancement.

In another study, the authors examined the role of testosterone in the pathogenesis, progression, and comorbidity of chronic kidney disease. Their findings suggested that testosterone levels may be a contributing factor to the development of chronic kidney disease. This research is important as it provides a new perspective on the role of hormonal factors in kidney disease, potentially opening up new avenues for treatment.

Furthermore, the study also highlighted the role of oxidative stress in kidney disease. Several studies have shown that oxidative stress is a significant contributor to the progression of kidney disease. The authors discussed the mechanisms by which oxidative stress could exacerbate kidney disease and suggested potential strategies for mitigating its effects.

Overall, these studies demonstrate the complex interplay between anabolic androgenic steroids and kidney health. They emphasize the need for further research to better understand the mechanisms behind these effects and to develop effective therapeutic strategies to mitigate their impact.

**References**


