Review
Sudden Cardiac Death in Anabolic-Androgenic Steroid Users: A Literature Review

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Abstract: Background and objectives: Anabolic-androgenic steroids (AASs) are a group of synthetic molecules derived from testosterone and its related precursors. AASs are widely used illicitly by adolescents and athletes, especially by bodybuilders, both for aesthetic uses and as performance enhancers to increase muscle growth and lean body mass. When used illicitly they can damage health and cause disorders affecting several functions. Sudden cardiac death (SCD) is the most common medical cause of death in athletes. SCD in athletes has also been associated with the use of performance-enhancing drugs. This review aimed to focus on deaths related to AAS abuse to investigate the cardiac pathophysiological mechanism that underlies this type of death, which still needs to be fully investigated. Materials and Methods: This review was conducted using PubMed Central and Google Scholar databases, until 21 July 2020, using the following key terms: “(Sudden cardiac death) OR (Sudden death)) AND ((androgenic anabolic steroid) OR (androgenic anabolic steroids) OR (anabolic-androgenic steroids) OR (anabolic-androgenic steroids) OR (anabolic-androgenic steroid))”. Thirteen articles met the inclusion and exclusion criteria, for a total of 33 reported cases. Results: Of the 33 cases, 31 (93.9%) were males while only 2 (61%) were females. Mean age was 29.79 and, among sportmen, the most represented sports activity was bodybuilding. In all cases there was a history of AAS abuse or a physical phenotype suggesting AAS use; the total usage period was unspecified in most cases. In 24 cases the results of the toxicological analysis were reported. The most detected AASs were nandrolone, testosterone, and stanozolol. The most frequently reported macroscopic alterations were cardiomegaly and left ventricular hypertrophy, while the histological alterations were foci of fibrosis and necrosis of the myocardial tissue. Conclusions: Four principal mechanisms responsible for SCD have been proposed in AAS abusers: the atherogenic model, the thrombosis model, the model of vasospasm induced by the release of nitric oxide, and the direct myocardial injury model. Hypertrophy, fibrosis, and necrosis represent a substrate for arrhythmias, especially when combined with exercise. Indeed, AAS use has been shown to change physiological cardiac remodeling of athletes to pathophysiological cardiac hypertrophy with an increased risk of life-threatening arrhythmias.

Keywords: AASs; anabolic androgenic steroids; SCD; sudden cardiac death; cardiac damage; adverse effects; cardiac toxicity
1. Introduction

AASs are a group of synthetic molecules derived from testosterone and its related precursors. AASs were developed to minimize the androgenic effects of testosterone and maximize the anabolic effects promoting the growth of skeletal muscles [1–3]. AASs can be administered orally, by intramuscular or subcutaneous injection, by pellet subcutaneous implantation, or by application on the skin.

Only a few AASs are used or proposed for therapeutic use, mainly in replacement treatment of hypogonadism [4,5]. Direct testosterone replacement therapy (TRT) is the only FDA-approved therapy for the treatment of male hypogonadism [6]. Oxandrolone, instead, is used to fight the protein catabolism associated with long-term use of corticosteroids and in the treatment of bone pain due to osteoporosis [7]. Clinical studies have demonstrated the efficacy of oxandrolone in the treatment of acute catabolic disorders, such as severe burns or severe trauma, and chronic catabolic disorders such as AIDS-associated cachexia or neuromuscular diseases such as Duchenne muscular dystrophy [8].

As well as for therapeutic use, AASs are widely used illicitly by adolescents and athletes, especially by bodybuilders, both for aesthetic uses and as performance enhancers to increase muscle growth and lean body mass, in consideration of their significant anabolic effect [9,10].

Although the use of AASs is now widespread around the world, with around ten million users, there are some geographical differences. The Middle East has the highest prevalence rate, with 21.7% of world users, followed by South America (4.8%), Europe (3.8%), North America (3.0%), Oceania (2.6%), Africa (2.4%), and Asia (0.2%). Among developed countries, the highest prevalence is found in Scandinavia, the United States, and countries of the British Commonwealth. The highest overall prevalence rate of AAS use was found in recreational sportspeople (18.4%), followed by athletes (13.4%), prisoners (12.4%), and drug users (8.0%). Non-athletes have the lowest prevalence rate that is estimated to be about 1% [11,12]. The global lifetime prevalence rate of AAS use is estimated to be 3.3%, greater in men than women (6.4% vs. 1.6%). As concerns the age of AAS users, teenagers have a higher overall prevalence rate (2.5%) than people older than 19 years (1.9%). The prevalence rate among high-school students was 2.3% [11,13,14].

While the use of AASs for medical indications is relatively safe, when used illicitly they can damage health and cause disorders affecting several functions (cardiovascular, reproductive, musculoskeletal, endocrine, renal, immunologic, and neuropsychiatric) [3,15–23]. These side effects include cardiac injuries such as fibrosis, cardiac hypertrophy, and dilated cardiomyopathy with an increased risk for myocardial infarction, arrhythmias, and sudden cardiac death [3,24–27].

AASs are typically used in phases referred to as “cycles”. To reduce the dangerous consequences of continuous AAS use at supraphysiological doses, abusers often introduce changes to their intake [28,29]. “Stacking” consumption can also involve nutritional supplements, complements, or other substances [30–32]. The substances most frequently taken at the same time as AASs are alcohol, amphetamines, aspirin, cannabinoids, caffeine, clomiphene citrate, cocaine, codeine, creatine, ephedrine, erythropoietin, furosemide, gamma-hydroxybutyrate (GHB), growth hormone, heroin, insulin, insulin-like growth 1 (IGF-1), melatonin, protein powder, tamoxifen, thyroxine, and tobacco. [33].

Sudden cardiac death (SCD) is generally defined as a sudden unexpected death or arrest from a presumed cardiac cause, which occurs within one hour of symptom onset if witnessed, otherwise within 24 h, in a person without any prior condition that would appear fatal [34–36].

This review aims to investigate the relationship between the use of anabolic-androgenic steroids (AAS) and sudden cardiac death in athletes and identify the possible etiological mechanism.

2. Materials and Methods

2.1. Database Search Terms and Timeline

This review was conducted performing a systematic literature search on online resources (PubMed Central database and Google Scholar) until 21 July 2020, using the following key terms: “((Sudden
cardiac death) OR (Sudden death)) AND ((androgenic anabolic steroid) OR (androgenic anabolic steroids) OR (anabolic-androgenic steroids) OR (anabolic-androgenic steroid))”.

2.2. Inclusion and Exclusion Criteria

The following inclusion criterion were applied: full-text scientific article published in English. The following exclusion criteria were adopted: (1) conference abstracts or reviews and letters to the editor without case reports; (2) animal studies; (3) articles in which the correlation between cardiac death and steroids is not discussed; (4) articles regarding surviving subjects.

2.3. Study Selection

We retrieved 1909 articles (339 from Pubmed Central and 1570 from Google Scholar databases). After excluding all duplicate articles, the reviewers retrieved abstracts and full text of each article independently applying the inclusion and exclusion criteria. Figure 1 summarizes the data obtained after our literature search.

3. Results

The review of the literature using the flow diagram shown in Figure 1 allowed us to identify 13 articles (Table 1) published between 1993 and 2020, for a total of 33 reported cases. The main characteristics of each selected article are summarized in Appendix A.

Of the 33 cases, 31 (93.9%) were males while only 2 (6.1%) were females. The mean age was 29.79 years with an SD of 8.5 years (range 13–54). Twenty-one cases (63.6%) were sportsmen and the most represented sports activity was bodybuilding (13 cases, 39%).

In all cases, there was a history of AAS abuse or a physical phenotype suggesting AAS use, the total period of AAS use was unspecified in 24 cases. In the other 9 cases, the time in which the subjects took AAS varied from 3 months to several years. In 24 cases the results of the toxicological analysis were reported. The tests were negative in 4 subjects. In 8 individuals, the toxicological examination revealed the presence of one or more AASs (Table 2) in blood or urine. The most detected AASs were nandrolone (10 cases), testosterone (9 cases), and stanozolol (7 cases). In two cases the toxicological examination found the presence of other substances used as performance-enhancing drugs (clenbuterol, ephedrine, norephedrine, liothyronine). In five other cases, other drugs of abuse were found such as cocaine, opioids, benzodiazepines, and cannabinoids.
Table 1. Articles included.

| Author                      | Year | Number of Cases | Study Type          |
|-----------------------------|------|-----------------|---------------------|
| Campbell, S.E. et al. [37]  | 1993 | 1               | Case report         |
| Dickerman, R.D. et al. [38] | 1995 | 1               | Case report         |
| Hausmann, R. et al. [39]    | 1998 | 1               | Case report         |
| Fineschi, V. et al. [40]    | 2001 | 2               | Case series         |
| Fineschi, V. et al. [41]    | 2007 | 1               | Case series         |
| Di Paolo, M. et al. [42]    | 2007 | 4               | Letter to the editor|
| Fainton, L. et al. [43]     | 2009 | 6               | Retrospective study |
| Thiblin, I. et al. [44]     | 2009 | 1               | Case report         |
| Montisci, M. et al. [45]    | 2012 | 3               | Case series         |
| Lusetti, M. et al. [46]     | 2015 | 6               | Retrospective study |
| Lichtenfeld, J. et al. [47] | 2016 | 1               | Case report         |
| Lusetti, M. et al. [48]     | 2018 | 5               | Retrospective study |
| Hernandez-Guerra, A.I. et al. [1] | 2019 | 1       | Case report              |

1 case excluded because already present in the previous article; 2 only 6 out of 12 cases died of sudden cardiac death (SCD); 3 3 out of 4 cases died of SCD.

Table 2. Anabolic-androgenic steroids (AASs) found on toxicological analysis.

| Toxicological Findings | Number of Cases | % of Total Cases |
|------------------------|-----------------|------------------|
| Nandrolone             | 10              | 30%              |
| Testosterone           | 9               | 27%              |
| Stanozolol             | 7               | 21%              |
| Boldenon               | 2               | 6%               |
| Norandrostosterone     | 1               | 3%               |
| Mesterolone            | 1               | 3%               |
| Methandienone          | 1               | 3%               |
| Epitestosterone        | 1               | 3%               |
| Nortestosterone        | 1               | 3%               |

In 15 cases it was possible to calculate the weight of the heart as a percentage of body weight (Appendix A). Anamnestic data were present in 24 of the 33 cases examined (72.7%). Of these, in no case was there a personal history of the disease or a family history of heart disease before age 50.

In the 33 examined cases, the most frequent macroscopic alteration was cardiomegaly (11 cases, 33%), based on the weight of the heart as a percentage of body weight, followed by left ventricular hypertrophy (10 cases, 30%). Dilated cardiomyopathy was found in 3 cases (9%). The most frequently reported histological alteration were foci of fibrosis and necrosis of the myocardial tissue, found respectively in 21 (79%) and 17 cases (52%). Other histological alterations reported were atherosclerosis (7 cases, 21%), inflammatory infiltrate (4 cases, 12%), coronary stenosis (3 cases, 9%), and left ventricular apoplexy (2 cases, 6%). The macroscopic and histological findings are summarized in Figure 2.
In all cases, autopsies ruled out causes of extracardiac death, and SCD was correlated with AAS use. Sudden cardiac death (SCD) is generally defined as a sudden unexpected death or arrest from a presumed cardiac cause, which occurs within one hour of symptom onset if witnessed, otherwise within 24 h, in a person without any prior condition that would appear fatal [34–36]. SCD in athletes is an event that profoundly impacts society because athletes are generally seen as a healthy category of people. Although SCD is the most common medical cause of death in athletes, its true incidence is unknown. The risk of SCD in athletes is 2 to 3 times greater than that in the general population. This difference may be due to typical athletes’ demographic factors, such as sex, age, and ethnicity. Potential mechanisms for SCD consist of inflammation, mechanical factors such as ventricular hypertrophy or fibrosis, neurological and metabolic comorbidities, and hereditary factors, arrhythmic mechanisms of abnormal ventricular repolarization, conduction, or autonomic innervation [49]. The etiology of SCD in younger athletes (<35 years of age) is mainly related to inherited cardiac conditions, instead, in older athletes, it is related to atherosclerotic coronary artery disease (CAD) [50–53]. Left ventricular hypertrophy (LVH) has been recognized as an independent risk factor for sudden cardiac death. The high mortality and sudden cardiac death associated with LVH is related to ventricular arrhythmia. Indeed, hypertrophied myocardium has a typical pro-arrhythmic electrophysiological phenotype and predisposes to the presence of myocardial ischemia. The main abnormality is prolongation of the action potential duration and refractoriness, which represents the substrate for arrhythmias [54]. Increased risk of ventricular arrhythmias and SCD, associated with hypertrophy, are related to complex processes involving myocardial cells, interstitium, coronary flow reserve, and neurohumoral activation [55]. SCD in athletes has also been associated with the use of performance-enhancing drugs, both anabolic-androgenic steroids and nonsteroidal agents [56]. AAS users often exhibit signs of neurohumoral activation [55]. SCD in athletes has also been associated with the use of performance-enhancing drugs, both anabolic-androgenic steroids and nonsteroidal agents [56]. AAS users often exhibit signs of neurohumoral activation [55].

Figure 2. Summary of macroscopic and histologic findings. Autopsies ruled out the causes of extracardiac death in all cases. However, in most cases (20.61%) it was not possible to define the exact cardiac cause that led to death, although SCD was correlated with the use of AASs in all cases.

4. Discussion

Data emerging from our study confirm the higher prevalence of AAS assumption among young males (93.9% males compared to 6.1% females, mean age 29.79 years), especially if they are bodybuilders (39%). In none of the cases in which anamnestic data were present was there a personal history of the disease or a family history of heart disease before age 50. In the 33 cases examined, the most frequently reported macroscopic changes were cardiomegaly (33%) and left ventricular hypertrophy (30%). The most frequently reported histological changes were foci of fibrosis (79%) and necrosis (52%) of myocardial tissue. In all cases, autopsies ruled out causes of extracardiac death, and SCD was correlated with AAS use. Sudden cardiac death (SCD) is generally defined as a sudden unexpected death or arrest from a presumed cardiac cause, which occurs within one hour of symptom onset if witnessed, otherwise within 24 h, in a person without any prior condition that would appear fatal [34–36]. SCD in athletes is an event that profoundly impacts society because athletes are generally seen as a healthy category of people. Although SCD is the most common medical cause of death in athletes, its true incidence is unknown. The risk of SCD in athletes is 2 to 3 times greater than that in the general population. This difference may be due to typical athletes’ demographic factors, such as sex, age, and ethnicity. Potential mechanisms for SCD consist of inflammation, mechanical factors such as ventricular hypertrophy or fibrosis, neurological and metabolic comorbidities, and hereditary factors, arrhythmic mechanisms of abnormal ventricular repolarization, conduction, or autonomic innervation [49]. The etiology of SCD in younger athletes (<35 years of age) is mainly related to inherited cardiac conditions, instead, in older athletes, it is related to atherosclerotic coronary artery disease (CAD) [50–53]. Left ventricular hypertrophy (LVH) has been recognized as an independent risk factor for sudden cardiac death. The high mortality and sudden cardiac death associated with LVH is related to ventricular arrhythmia. Indeed, hypertrophied myocardium has a typical pro-arrhythmic electrophysiological phenotype and predisposes to the presence of myocardial ischemia. The main abnormality is prolongation of the action potential duration and refractoriness, which represents the substrate for arrhythmias [54]. Increased risk of ventricular arrhythmias and SCD, associated with hypertrophy, are related to complex processes involving myocardial cells, interstitium, coronary flow reserve, and neurohumoral activation [55]. SCD in athletes has also been associated with the use of performance-enhancing drugs, both anabolic-androgenic steroids and nonsteroidal agents [56]. AAS users often exhibit signs of neurohumoral activation [55]. SCD in athletes has also been associated with the use of performance-enhancing drugs, both anabolic-androgenic steroids and nonsteroidal agents [56]. AAS users often exhibit signs of neurohumoral activation [55].
of performance-enhancing drugs, both anabolic-androgenic steroids and nonsteroidal agents [56]. AAS users often combine the assumption of anabolic substances with other substances such as cocaine, methamphetamine, and smart drugs. These data are in agreement with the results of our review. Mixing two or more substances increases the risk of negative drug interactions, worsening any adverse effects, including SCD [17].

The higher prevalence of AAS use among athletes, especially non-professionals, can be explained by their determination to achieve a perfect body and to improve performance and self-esteem. Indeed, the positive effects of AAS use are the increase of muscle mass, strength, energy and concentration, and the reduction of fat mass [57,58]. However, the use of anabolic-androgenic steroids has also many negative effects. Many of these are mild and transient (fluid retention, acne, agitation, gynecomastia, aggressiveness), but others are more serious and can damage multiple organs and functions, such as cardiovascular, reproductive, musculoskeletal, endocrine, renal, immunologic, and neuropsychiatric functions [2,57–60]. The cardiovascular system is one of the most affected by the side effects of AAS use. AAS use enhances vascular resistance and blood pressure, pro-inflammatory biomarker profile, and sympathetic tone alters serum lipoproteins and produces direct myocardial toxicity [53,61]. The adverse cardiovascular events reported are: impaired left ventricular function, arterial thrombosis, pulmonary embolism, and left ventricular hypertrophy, associated with myocytolysis and fibrosis [1]. It is reported that AAS abuse can promote cardiac tissue growth, leading to hypertrophic cardiomyopathy, followed by apoptotic cell death. This phenomenon is associated with ventricular remodeling, cardiomyopathy, myocardial infarction, and SCD and can explain how AAS may lead to cardiac death without coronary thrombosis or atherosclerosis [62,63]. AASs cause cardiac hypertrophy by a direct action on cardiac androgen receptors and these effects are directly proportional to the dose, time, and duration of administration [45]. Melchert and Welder proposed at least four hypothetical models explaining how AASs cause cardiovascular side effects. The atherogenic model concerns the alterations on lipoprotein serum levels caused by AASs, increasing the risk of atherosclerosis. The thrombosis model regards enhancing platelet aggregation and polycythemia that increase the risk of thrombus formation. The third model involves vasospasm caused by nitric oxide release induced by anabolic agents. The direct myocardial injury model concerns direct myocardial toxicity causing apoptosis, with increased collagen deposition, fibrosis, and altered microcirculation resulting in chronic ischemic damage. All of these mechanisms associate AAS use with a high risk of SCD [64,65].

A recent study showed that chronic nandrolone treatment with or without severe training causes a significant increase in beta–myosin heavy chain (β-MHC) gene expression, calcium/calmodulin-dependent protein kinaseII (CaMKIIδ), and monoamine oxidase (MAO) activities in the heart tissue of male Wistar rats [66]. Cardiac hypertrophy has a genetic substrate too; ND, in adult rats, reduces cardiac contractile performance through enhancing β-MHC mRNA expressions, causing alterations of pressure-overload cardiac hypertrophy [67]. In the 9 cases of SCD, the most representative macroscopic alterations were cardiomegaly and left ventricular hypertrophy. Cardiomegaly was diagnosed by comparing the weight of the heart with the body weight and BMI of the subject [68–70]. Histologically, the most representative alterations were fibrosis and necrosis. These results are in agreement with what has been reported by many authors, according to whom myocardial necrosis and focal myocardial fibrosis, are highly significant alterations in the hearts of athletes who abuse AAS and may be responsible for atrioventricular conduction abnormalities and provide a substrate for the occurrence of potentially lethal arrhythmias and SCD [40,71–73]. It must be taken into account that, regardless of AAS abuse, increased cavity dimensions, wall thickness, and left ventricular mass are typical consequences of high-intensity exercise training and are included in the physiological cardiac remodeling of the “athlete’s heart”. A modest amount of fibrosis may be present in physiological cardiac remodeling associated with lifelong endurance training. This fibrosis and hypertrophy represent a substrate for arrhythmias [74,75]. When combined with exercise, AAS use has been shown to change physiological cardiac remodeling
of the athlete to pathophysiological cardiac hypertrophy with an increased risk of life-threatening arrhythmias [1,76].

It is difficult to distinguish the etiology of these changes from histological findings alone, and it becomes essential to evaluate the subject’s clinical history and physical characteristics in all cases of sudden cardiac death in which AAS abuse is suspected. The physical phenotype of a male who abuses AASs includes some characteristics such as muscle hypertrophy, prominence striae above the pectoralis or biceps muscle, breast development in men (gynecomastia), testicular atrophy, and severe acne that may suggest AAS abuse [1,77]. In women, signs of AAS use also include hirsutism, deepening of the voice, and masculinization of secondary sexual characteristics [78,79].

Long-term use of AASs causes cardiac alterations affecting the conductive system, as demonstrated by subjects who have undergone signal-averaging electrocardiography (SAECG). SAECG is an inexpensive, safe, and highly reproducible technique that records low-amplitude electrical activity in the myocardium and provides information on the presence of a monomorphic TV substrate. SAECG performed on AAS users shows alterations in myocardial electrophysiology such as significantly longer QTc interval and greater QT dispersion, at rest and after moderate exercise, and attenuated heart rate recovery after exercise compared to subjects who do not use AASs. Abnormal SAECG indicates a re-entry mechanism for arrhythmias and, because sympathetic thrust during acute exercise lowers the ventricular fibrillation threshold, these individuals will be at increased risk of tachyarrhythmia and potential SCD following exercise [73,80,81]. Despite evidence linking the use of AAS abuse to SCD, reports in the literature are most likely underestimated due to the few autopsy data and because of the study of the pathophysiological mechanisms that lead to sudden cardiac death in subjects using AASs is severely limited [44,48,53]. In fact, information on the modalities and doses relating to the abuse of AASs is generally self-reported. Furthermore, most of the data in the literature on the effects of AAS administration derive from animal studies, as the administration of high doses of AASs in humans would be unethical, given the serious health risks [82].

Because of the high prevalence of AAS use among athletes, toxicological investigations are therefore fundamental in those cases of sudden death in subjects suspected of consuming AASs [83,84]. To date, there are still few studies published in the literature that correlates SCD in athletes with AAS use, highlighting the pathophysiological mechanisms that cause it and that are mostly based on experimental models derived from animal experiments [73,76,82,85–91]. It could be important to investigate new research fields to define the exact mechanism of action. For this reason, in recent years some studies have been carried out on miRNAs, a family of non-coding nucleotides that control gene expression and that appear to be related to numerous diseases. Mir-133a and mir-1, for example, appear to increase the risk of arrhythmia in the ischemic heart and may, in the future, play a role as prognostic biomarkers [25,92,93]. Nowadays, there are no studies in the literature that link the expression of miRNAs with SCD in AAS abusers.

5. Conclusions

Because of the high prevalence of AAS use among athletes, toxicological investigations are therefore fundamental in those cases of sudden death in which there is suspicion of AAS consumption. The cardiovascular system is one of the most affected by the side effects of AAS use. AAS use enhances vascular resistance and increases blood pressure, pro-inflammatory biomarker profile, sympathetic tone, alters serum lipoproteins, and produces direct myocardial toxicity. In agreement with the evidence in the literature, the most reported macroscopic heart changes reported in our review were cardiomegaly and hypertrophy, and the main histological changes were necrosis of myocardial tissue and foci of fibrosis. Hypertrophy, fibrosis, and necrosis represent a substrate for arrhythmias, especially when combined with exercise. AAS use has been shown to change physiological cardiac remodeling of athletes to pathophysiological cardiac hypertrophy with an increased risk of life-threatening arrhythmias. The evaluation of the parameters of electrocardiographic repolarization at rest and post-exercise, using SAECG, could provide diagnostic and prognostic information on the risk of cardiac
arrhythmias and SCD in apparently healthy subjects who chronically use supraphysiological doses of AAS [27,81].

Since the pathophysiological mechanisms that lead to SCD in subjects who use AAS have not yet been fully clarified, the link between AAS abuse and SCD is probably underestimated, considering the few data in the literature.

Toxicological investigations, performed on different matrices, such as blood, urine, and hair, can confirm the use of AAS or other drugs that may have played a role in the death. A complete autopsy with histological and immunohistochemical studies, with a particular regard to the organs in which anabolic adverse events occur most frequently, is mandatory to evaluate the relationship between AAS use and SCD.

Given the young age of the subjects who usually use AASs and given the importance of the consequences related to their abuse, the identification of new tools to study AAS use, such as miRNAs, could be an important goal for the scientific community.

Nowadays, clinicians must pay attention to indicative signs of AAS use, considering those physical and epidemiological characteristics that can lead to the suspicion of abuse of these drugs to implement primary prevention measures of the serious adverse effects of AAS use. An interesting challenge would be to further investigate these findings to be able to use these biomarkers both to facilitate the post-mortem diagnosis of sudden deaths related to AAS abuse and as a screening method in living subjects to prevent fatal consequences.

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## Appendix A

| Author (Year) | Age (Years); Sex; Height; Weight | BMI | Personal and Family Medical History | Kind of Sport Activity | ASS Reported Use—Time of Assumption | Use of Other Drugs | Circumstance of Death | Macroscopic Heart Findings | Histological Heart Findings | Toxicological Analysis | Cause of Death |
|---------------|----------------------------------|-----|-----------------------------------|------------------------|------------------------------------|------------------|----------------------|--------------------------|----------------------------|-------------------------|---------------|
| Campbell, S. E. et al. (1993) | 21; M NR | Absence of significant diseases | Bodybuilder | Testosterone; nandrolone—several months | NR | Collapse during a weight-lifting workout at the gym | 530 g—Marked left and right ventricular hypertrophy | Extensive perivascular fibrosis of intramural coronary arteries—interstitial fibrosis | NR | Unspecified | SCD |
| Dickerman, R. D. et al. (1995) | 20; M; 180 cm; 100.7 kg 31.08 | No past or family history of cardiac disease | Bodybuilder | Methylone enanthate—just complete a 3-month cycle | NR | Sudden witnessed death | 515 g (0.51% of body weight)—Signs of concentric left ventricular hypertrophy | Mild atherosclerosis | NR | Unspecified | SCD |
| Hausmann, R. et al. (1998) | 23; M; 192 cm; 94 kg 25.50 | NR | Bodybuilder | Testosterone cyclopentilpropionate; methenolone enanthate; mestanolone—9 months | Other performance-enhancing drugs (lithotrycine hydrochloride, clenbuterol hydrochloride) | Found unconscious at home in his bed | 500 g (0.53% of body weight)—Cardiac hypertrophy, right ventricle dilatation, focal induration of endocardium | Enlargement and nuclear polymorphism of the left ventricular muscle fibers. Disseminated focal necrosis and interstitial fibrosis | Urine: Mesterolone, methandienone, testosterone, nandrolone, clenbuterol | Unspecified | SCD |
| Fineschi, V. et al. (2003) | 32; M; 189 cm; 90 kg 25.20 | No history of disease | Bodybuilder | Testosterone propionate; nandrolone—several months | NR | Sudden loss of consciousness during a weight lifting workout | 450 g (0.50% of body weight)—Normal heart measures (14 × 14 × 4 cm)—Normal valves, endocardium, and coronary arteries—one grayish zone in the left ventricle myocardium | Infarct necrosis corresponding to the grayish zone—some foci of contraction band necrosis and fibrosis | Urine: Metabolites of nandrolone, metabolites of stanozolol | SCD most likely related to adrenergic stress |
| Fineschi, V. et al. (2005) | 29; M; 166 cm; 72 kg 26.13 | His medical history was unremarkable | Bodybuilder | Nandrolone; stanozol—several months | NR | Found unconscious at home in his bed | 390 g (0.54% of body weight)—Normal heart measures (13 × 10 × 5 cm)—Normal valves, endocardium, and coronary arteries | Occasional isolated myocardial cells with contraction band and segmentation | Urine: Metabolites of nandrolone, metabolites of stanozolol | Unspecified | SCD |
| Fineschi, V. et al. (2009) | 30; M; 178 cm; 90 kg 28.41 | NR | Bodybuilder | Nandrolone decanoate—6 months | Unspecified other drugs | Sudden collapse at home | 400g (0.44% of body weight)—Scattered fatty streaks in coronary arteries | Focal myocardial fibrosis | Urine: Norandrostenedione. Blood: nandrolone | Unspecified | SCD |
| Author (Year) | Age (Years); Sex; Height; Weight | BMI | Personal and Family Medical History | Kind of Sport Activity | ASS Reported Use—Time of Assumption | Use of Other Drugs | Circumstance of Death | Macroscopic Heart Findings | Histological Heart Findings | Toxicological Analysis | Cause of Death |
|---------------|----------------------------------|-----|-------------------------------------|------------------------|-------------------------------------|------------------|---------------------|--------------------------|--------------------------|--------------------------|-------------------|
| Di Paolo, M. et al. (2007) | 29; M; 190 cm; 127 kg | 35.2 | No prior history of disease. No family history of cardiac disease under the age of 50 | Bodybuilder | History of use of unspecified AAS—unspecified | NR | Sudden loss of consciousness during the first minutes of a spin bike lesson | 490 g (0.39% of body weight)—Normal heart wall thickness, normal valve, normal coronary arteries | Severe epicardial interstitial fibrosis, small vessel disease | Negative | Unspecified SCD |
| | 27; M; 190 cm; 108 kg | 25.8 | No prior history of disease. No family history of cardiac disease under the age of 50 | Bodybuilder | History of use of unspecified AAS—unspecified | NR | Sudden illness while he was at a night club | 360 g (0.36% of body weight)—Normal heart wall thickness, normal valve, normal coronary arteries | Mild focal epicardial interstitial fibrosis, small vessel disease | Urine: Stanozolol, testosterone | Unspecified SCD |
| | 37; F; 161 cm; 71 kg | 27.4 | No prior history of disease. No family history of cardiac disease under the age of 50 | Bodybuilder and weight lifter | History of use of unspecified AAS—unspecified | NR | Found dead in her car | 310 g (0.44% of body weight)—Normal heart wall thickness, normal valve, normal coronary arteries | Moderate focal epicardial interstitial fibrosis, small vessel disease | Negative | Unspecified SCD |
| Fanton, L. et al. (2009) | 31; M; 175 cm; 79 kg | 25.8 | No prior history of disease. No family history of cardiac disease under the age of 50 | Bodybuilder | History of use of unspecified AAS—unspecified | NR | Found dead in his bedroom: alive 7 h before | 400 g (0.51% of body weight)—Normal heart wall thickness, normal valve, normal coronary arteries | Moderate epicardial interstitial fibrosis, small vessel disease | Urine: Stanozolol | Unspecified SCD |
| | 19; M | NR | No history of cardiac disease | Weight lifter | History of use of unspecified AAS—unspecified | NR | SD during training | 360 g—Left ventricle apoplexy | Multiple focal areas of necrosis, myolysis, scarring fibrosis | NR | Unspecified SCD |
| | 22; M | NR | No history of cardiac disease | PE teacher | History of use of unspecified AAS—unspecified | NR | SD during training | 520 g—Left ventricle apoplexy | Multiple focal areas of necrosis, myolysis, scarring fibrosis | NR | Unspecified SCD |
| | 25; M | NR | No history of cardiac disease | Bodybuilder | History of use of unspecified AAS—unspecified | NR | SD during training | 460 g—Disseminated myocarditis | Multiple focal areas of necrosis, myolysis, scarring fibrosis | NR | Unspecified SCD |
| | 28; M | NR | No history of cardiac disease | Soccer player | History of use of unspecified AAS—unspecified | NR | SD during training | 380 g—Disseminated myocarditis | Multiple focal areas of necrosis, myolysis, scarring fibrosis | NR | Unspecified SCD |
| | 54; M | NR | No history of cardiac disease | Marathon runner | History of use of unspecified AAS—unspecified | NR | SD during training | 410 g—Coronary thrombosis and dilated cardiomyopathy | Multiple focal areas of necrosis, myolysis, scarring fibrosis | NR | Unspecified SCD |
| | 48; M | NR | No history of cardiac disease | Marathon runner | History of use of unspecified AAS—unspecified | NR | SD during training | 430 g—Left ventricle hypertrophy | Multiple focal areas of necrosis, myolysis, scarring fibrosis | NR | Unspecified SCD |
| Author (Year) | Age (Years); Sex; Height; Weight | BMI | Personal and Family Medical History | Kind of Sport Activity | ASS Reported Use—Time of Assumption | Use of Other Drugs | Circumstance of Death | Macroscopic Heart Findings | Histological Heart Findings | Toxicological Analysis | Cause of Death |
|--------------|----------------------------------|-----|------------------------------------|------------------------|-------------------------------------|---------------------|----------------------|--------------------------|--------------------------|--------------------------|--------------------------|
| Thiblin, I. et al. (2009) | 29; F; 172 cm; 76 kg | 25.7 | No history of disease | Fitness athlete | History of use of unspecified AAS—unspecified | Unspecified other drugs | Found naked in a prone position on the floor beside her bed, with a pillow partly under her body | 331 g (0.44% of body weight)—Normal heart measures—Normal coronary arteries, with an isolated flat area of fatty thickening in the proximal part of the left anterior descending (LAD) coronary artery. | Lymphocytic infiltration around several middle-sized and small intramural vessels—minimal myocardial necrosis | Blood ephedrine, non-ephedrine. Urine testosterone, metabolites of stanozolol, boldenone | Sudden cardiac arrhythmia, possibly related to the combination of an otherwise unspecified inflammatory process in the heart and the acute influence of ASS and ephedrine |
| Montisci, M. et al. (2012) | 32; M; 180 cm; 110 kg | 33.95 | Bodybuilder | History of use of unspecified AAS—7 years (recently withdraw) | NR | Found dead at home in his bed | 450 g (0.41% of body weight)—11 × 9.5 cm—cardiomegaly, concentric left ventricular hypertrophy, normal valve, normal coronary arteries | Hypertrophic myocytes, focal disarray, interstitial and replacement fibrosis, foci of lymphoplasmacytic cellular infiltrates (CD3+), with edema and patchy necrosis | Negative | Concentric left ventricular hypertrophy, focal acute myocarditis. |
| Montisci, M. et al. (2012) | 32; M; 178 cm; 94 kg | 29.67 | Bodybuilder | History of use of unspecified AAS—several years | NR | SD after a dentistry visit | 580 g (0.62% of body weight)—12.5 × 11 cm—Cardiomegaly, hypertrophy, biventricular dilatation, normal valve, non-obstructive LAD stenosis | Hypertrophic myocytes, foci of necrosis, replacement fibrosis, LAD 50% stenosis, fibrofatty replacement | Negative | Inflammatory dilated cardiomyopathy with subacute-chronic stages, hemorrhagic pulmonary infarction |
| 25; M; 185 cm; 125 kg | 36.52 | Bodybuilder | An ECG performed 5y before death was normal | Bodybuilder | Circumstantial finding of unspecified use of AAS—unspecified | Unspecified other performance-enhancing drugs | SD while sleeping | 390 g (0.31% of body weight)—10.5 × 9.5 cm—normal heart wall thickness, normal valve, normal coronary arteries | Inflammatory infiltrate, myocyte necrosis | Urine: Testosterone, epitestosterone, nortestosterone | Eosinophilic myocarditis |
| Author (Year) | Age (Years); Sex; Height; Weight | BMI | Personal and Family Medical History | Kind of Sport Activity | ASS Reported Use—Time of Assumption | Use of Other Drugs | Circumstance of Death | Macroscopic Heart Findings | Histological Heart Findings | Toxicological Analysis | Cause of Death |
|--------------|----------------------------------|-----|-----------------------------------|------------------------|------------------------------------|-------------------|----------------------|---------------------------|----------------------------|------------------|-----------------|
| Lusetti, M. et al. (2015) | 39 (mean age); M (All 6 cases) | NR  | NR | NR | History of use of unspecified AAS—unspecified | NR | Sudden unwitnessed death | Normal heart wall thickness, normal valve, normal coronary arteries. In one case: 490 g (0.54% of body weight) | Intermittent fibrosis (6 cases); perivascular fibrosis (4 cases); perineural fibrosis within the left ventricle (2 cases); fibroadipous metaplasia (2 cases); contraction band necrosis (2 cases); Myocyte segmentation (2 cases); Intercalated disc widening (2 cases); myocyte hypertrophy (3 cases); coronary intimal and media thickening (4 cases) | Blood: Ethanol (1 case). Urine and hair: nandrolone (3 cases), Testosterone (3 cases) | Sudden cardiac arrhythmia |
| Lichtenfeld, J. et al. (2016) | 13; M | NR | No prior history of disease. An episode of syncope with exertion 1 week before cardiac arrest. No family history of sudden death, hypertrophic cardiomyopathy, or heart rhythm abnormalities | Sprinter | Physical Phenotype suggesting AAS use | NR | Sudden cardiac arrest while performing timed wind sprints at a competitive sports camp | 465 g—Cardiomegaly, marked LV Hypertrophy | Foci of myofilibrillar disarray, the proliferation of fibroblasts consistent with early fibrosis, and enlarged myofibers with the heterogeneity of nuclear size including “box-car” nucleus | NR | Sudden cardiac arrest followed by brain death |
| Author (Year) | Age (Years); Sex; Height, Weight | BMI | Personal and Family Medical History | Kind of Sport Activity | ASS Reported Use—Time of Assumption | Use of Other Drugs | Circumstance of Death | Macroscopic Heart Findings | Histological Heart Findings | Toxicological Analysis | Cause of Death |
|--------------|----------------------------------|-----|-----------------------------------|-----------------------------|-----------------------------------|-------------------|------------------------|-----------------------------|-----------------------------|---------------------------|------------------|
| Lusetti, M. et al. (2018) | 32; M NR | | No "officially" medically prescribed drug treatment at the time of death. | NR | History of use of unspecified AAS—unspecified | NR | unspecified SD | 380 g—Left ventricular hypertrophy | Myocardial fibrosis | Urine: Nandrolone, Testosterone. Blood: Methadone, Citalopram, Clozapine, Venlafaxine, Lorazepam, Phenoobarbital, THC. | Unspecified SCD |
| | 32; M NR | | No "officially" medically prescribed drug treatment at the time of death. | NR | History of use of unspecified AAS—unspecified | NR | unspecified SD | 360 g | Fatty streaks, intima, and media thickening within the coronary arteries | Urine: Boldenone, Clomiphene, Methandienone, Oxandrolone, Stanazolol. Blood: Lorazepam, THC. | Unspecified SCD |
| | 33; M NR | | No "officially" medically prescribed drug treatment at the time of death. | NR | History of use of unspecified AAS—unspecified | NR | unspecified SD | 425 g—Left ventricular hypertrophy | Myocyte necrosis | Urine: Testosterone. Blood: Methadone, Cocaine | Unspecified SCD |
| | 39; M NR | | No "officially" medically prescribed drug treatment at the time of death. | NR | History of use of unspecified AAS—unspecified | NR | unspecified SD | 480 g—Left and right ventricular hypertrophy | Myocyte necrosis, Myocardial fibrosis | Urine: Nandrolone. Blood: Morphone, THC. | Unspecified SCD |
| | 29; M NR | | No "officially" medically prescribed drug treatment at the time of death. | NR | History of use of unspecified AAS—unspecified | NR | unspecified SD | 340 g | NR | Urine: Nandrolone, Testosterone. Blood: morphone, THC, Ethanol | Unspecified SCD |
| Author (Year)         | Age (Years); Sex; Height; Weight | BMI | Personal and Family Medical History | Kind of Sport Activity | ASS Reported Use—Time of Assumption | Use of Other Drugs | Circumstance of Death | Macroscopic Heart Findings | Histological Heart Findings | Toxicological Analysis | Cause of Death     |
|----------------------|----------------------------------|-----|------------------------------------|------------------------|-------------------------------------|--------------------|-----------------------|-----------------------------|-----------------------------|-------------------------|---------------------|
| Hernandez-Guerra, A. I. et al. (2019) | 24; M; 178 cm; 85 Kg | 26.8 | No past or family history of cardiac disease. One episode of precordial pain some months before. | NR | stanozolol, testosterone, mesterolone, nandrolone—6 months | tamoxifen | Sudden death at home | 420 g (0.49% of body weight)—Cardiomegaly, Normal ventricular thickness, >75% | Stenosis of the left main trunk and the LAD, areas of scarring located at the intersection between the posterior wall and the posterior third of the septum | Blood: Ethanol, Stanozolol, Nandrolone | Acute myocardial infarction |

Acute myocardial infarction
References

1. Hernández-Guerra, A.I.; Tapia, J.; Menéndez-Quintanal, L.M.; Lucena, J.S. Sudden cardiac death in anabolic androgenic steroids abuse: Case report and literature review. *For. Sci. Res.* 2019, 4, 267–273. [CrossRef] [PubMed]

2. Piacentino, D.; D Kotzalidis, G.; Del Casale, A.; Rosaria Aromatario, M.; Pomara, C.; Girardi, P.; Sani, G. Anabolic-androgenic steroid use and psychopathology in athletes. A systematic review. *Curr. Neurupharmacol.* 2015, 13, 101–121. [CrossRef] [PubMed]

3. Al-harbi, F.F.; Gamaleddin, I.; Alsubaie, E.G.; Al-Surimi, K.M. Prevalence and Risk Factors Associated with Anabolic-androgenic Steroid Use: A Cross-sectional Study among Gym Users in Riyadh, Saudi Arabia. *Oman Med. J.* 2020, 35, e110. [CrossRef] [PubMed]

4. Bhasin, S.; Cunningham, G.R.; Hayes, F.J.; Matsumoto, A.M.; Snyder, P.J.; Swerdloff, R.S.; Montori, V.M. Testosterone therapy in men with androgen deficiency syndromes: An endocrine society clinical practice guideline. *J. Clin. Endocrinol. Metab.* 2010, 95, 2536–2559. [CrossRef] [PubMed]

5. Nieschlag, E. Current topics in testosterone replacement of hypogonadal men. *Best Pract. Res. Clin. Endocrinol.* 2015, 29, 77–90. [CrossRef] [PubMed]

6. Carrasquillo, R.; Chu, K.; Ramasamy, R. Novel therapy for male hypogonadism. *Curr. Urol. Rep.* 2018, 19, 63. [CrossRef]

7. Orr, R.; Singh, M.F. The anabolic androgenic steroid oxandrolone in the treatment of wasting and catabolic disorders. *Drugs* 2004, 64, 725–750. [CrossRef] [PubMed]

8. Wu, C.; Kovac, J.R. Novel uses for the anabolic androgenic steroids nandrolone and oxandrolone in the management of male health. *Curr. Urol. Rep.* 2016, 17, 72. [CrossRef] [PubMed]

9. Frati, P.; Busardo, F.; Cipolloni, L.; De Dominicis, E.; Fineschi, V. Anabolic-androgenic steroid (AAS) related deaths: Autoptic, histopathological and toxicological findings. *Curr. Urol. Rep.* 2018, 19, 63. [CrossRef]

10. Pereira, E.; Moyses, S.J.; Ignácio, S.A.; Mendes, D.K.; Da Silva, D.S.; Carneiro, E.; Johann, A.C.B.R. Anabolic steroids among resistance training practitioners. *PLoS ONE* 2019, 14, e0223384. [CrossRef]

11. Reyes-Vallejo, L. Current use and abuse of anabolic steroids. *Actas Urol. Esp.* 2020, 44, 309–313. [CrossRef]

12. Fineschi, V.; Neri, M.; Di Donato, S.; Pomara, C.; Riezzo, I.; Turillazzi, E. An immunohistochemical study in a fatality due to ovarian hyperstimulation syndrome. *Int. J. Legal Med.* 2006, 120, 293–299. [CrossRef]

13. Sagoe, D.; Molde, H.; Andreassen, C.S.; Torsheim, T.; Pallesen, S. The global epidemiology of anabolic-androgenic steroid use: A meta-analysis and meta-regression analysis. *Ann. Epidemiol.* 2014, 24, 383–398. [CrossRef]

14. Kanayama, G.; Pope Jr, H.G. History and epidemiology of anabolic androgens in athletes and non-athletes. *Mol. Cell Endocrinol.* 2018, 464, 4–13. [CrossRef]

15. Kicman, A.T. Pharmacology of anabolic steroids. *Br. J. Pharmacol.* 2008, 154, 502–521. [CrossRef]

16. Bertozzi, G.; Salerno, M.; Pomara, C.; Sessa, F. Neuropsychiatric and behavioral involvement in AAS abusers. A literature review. *Medicina* 2019, 55, 396. [CrossRef]

17. Sessa, F.; Salerno, M.; Cipolloni, L.; Bertozzi, G.; Messina, G.; Di Mizio, G.; Pomara, C. Anabolic-androgenic steroids and brain injury: miRNA evaluation in users compared to cocaine abusers and elderly people. *Aging* 2020, 12, 15314. [CrossRef] [PubMed]

18. Agriesti, F.; Tataranni, T.; Pacelli, C.; Scrina, R.; Laurenzana, I.; Ruggieri, V.; Sani, G. Nandrolone induces a stem cell-like phenotype in human hepatocarcinoma-derived cell line inhibiting mitochondrial respiratory activity. *Sci. Rep.* 2020, 10, 1–17. [CrossRef] [PubMed]

19. Bertozzi, G.; Sessa, F.; Maglietta, F.; Cipolloni, L.; Salerno, M.; Fiore, C.; Pomara, C. Immunodeficiency as a side effect of anabolic androgenic steroid abuse: A case of necrotizing myofasciitis. *For. Sci. Med. Pathol.* 2019, 15, 616–621. [CrossRef] [PubMed]

20. Bertozzi, G.; Sessa, F.; Albano, G.D.; Sani, G.; Maglietta, F.; Roshan, M.H.; Salerno, M. The role of anabolic androgenic steroids in disruption of the physiological function in discrete areas of the central nervous system. *Mol. Neurobiol.* 2018, 55, 5548–5556. [CrossRef]

21. Pomara, C.; Neri, M.; Bello, S.; Fiore, C.; Riezzo, I.; Turillazzi, E. Neurotoxicity by synthetic androgen steroids: Oxidative stress, apoptosis, and neuropathology: A review. *Curr. Neuropharmacol.* 2015, 13, 132–145. [CrossRef]
22. Pomara, C.; Barone, R.; Marino Gammazza, A.; Sangiorgi, C.; Barone, F.; Pitruzzella, A.; Locorotondo, N.; Di Gaudio, F.; Salerno, M.; Maglietta, F.; et al. Effects of nandrolone stimulation on testosterone biosynthesis in Leydig cells. *J. Cell Physiol.* 2016, 231, 1385–1391. [CrossRef]

23. Albano, G.D.; Sessa, F.; Messina, A.; Monda, V.; Bertozzi, G.; Maglietta, F.; Giugliano, P.; Vacchiano, G.; Gabriella, M.; Salerno, M. AAS and organs damage: A focus on Nandrolone effects. *Acta Med. Mediter.* 2017, 6, 939–946.

24. Joukar, S.; Yoosefnia, M.; Naderi-Boldaj, V.; Nasri, H.; Rafie, F. Heart reaction to nandrolone decanoate plus two different intensities of endurance exercise: Electrocardiography and stereological approach. *Addict. Health 2018*, 10, 180.

25. Wadthaisong, M.; Witayavanitkul, N.; Bupha-Intr, T.; Wattanapermpool, J.; De Tombe, P.P. Chronic high-dose testosterone treatment: Impact on rat cardiac contractile biology. *Physiol. Rep.* 2019, 7, e14192. [CrossRef]

26. Climstein, M.; O’Shea, P.; Adams, K.J.; DeBeliso, M. The effects of anabolic-androgenic steroids upon resting and peak exercise left ventricular heart wall motion kinetics in male strength and power athletes. *J. Sci. Med. Sport* 2003, 6, 387–397. [CrossRef]

27. Pomara, C.; D’Errico, S.; Riezzo, I.; De Cillis, G.P.; Fineschi, V. Sudden cardiac death in a child a...[CrossRef]

28. Fanton, L.; Belhani, D.; Vaillant, F.; Tabib, A.; Gomez, L.; Descotes, J.; Timour, Q. Heart lesions associated with anabolic steroid abuse: Comparison of post-mortem findings in athletes and norethandrolone-induced lesions in rabbits. *Exp. Toxicol. Pathol.* 2009, 61, 317–323. [PubMed]
44. Thiblin, I.; Mobini-Far, H.; Frisk, M. Sudden unexpected death in a female fitness athlete, with a possible connection to the use of anabolic androgenic steroids (AAS) and ephedrine. *For. Sci. Int.* 2009, 184, 7–11. [CrossRef]

45. Montisci, M.; El Mazloum, R.; Cecchetto, G.; Terranova, C.; Ferrara, S.D.; Thiene, G.; Basso, C. Anabolic androgenic steroids abuse and cardiac death in athletes: Morphological and toxicological findings in four fatal cases. *Forensic Sci. Int.* 2012, 217, 13–18. [CrossRef]

46. Lusetti, M.; Licata, M.; Silingardi, E.; Bonetti, L.R.; Palmieri, C. Pathological changes in anabolic androgenic steroid users. *J. For. Leg. Med.* 2015, 33, 101–104. [CrossRef] [PubMed]

47. Lichtenfeld, J.; Deal, B.J.; Crawford, S. Sudden cardiac arrest following ventricular fibrillation attributed to anabolic steroid use in an adolescent. *Cardiol. Young* 2016, 26, 996–998. [CrossRef]

48. Lusetti, M.; Licata, M.; Silingardi, E.; Bonsignore, A.; Palmiere, C. Appearance/image- and performance-enhancing drug users: A forensic approach. *Am. J. For. Med. Pathol.* 2018, 39, 325–329. [CrossRef]

49. Narayan, S.M.; Wang, P.J.; Daubert, J.P. New concepts in sudden cardiac arrest to address an intractable epidemic: JACC state-of-the-art review. *J. Am. Coll. Cardiol.* 2019, 73, 70–88. [CrossRef]

50. Wasfy, M.M.; Hutter, A.M.; Weiner, R.B. Sudden cardiac death in athletes. *Methodist DeBakey Cardiovasc. J.* 2016, 12, 76. [CrossRef]

51. Harmon, K.G.; Asif, I.M.; Klossner, D.; Drezner, J.A. Incidence of sudden cardiac death in National Collegiate Athletic Association athletes. *Circulation* 2011, 123, 1594–1600. [CrossRef]

52. Ackerman, M.; Atkins, D.L.; Triedman, J.K. Sudden cardiac death in the young. *Circulation* 2016, 133, 1006–1026. [CrossRef]

53. Sheppard, M.N. Aetiology of sudden cardiac death in sport: A histopathologist’s perspective. *Br. J. Sports Med.* 2012, 46, i15–i21. [CrossRef]

54. Wolk, R. Arrhythmogenic mechanisms in left ventricular hypertrophy. *Europace* 2000, 2, 216–223. [CrossRef]

55. Aro, A.L.; Reinier, K.; Phan, D.; Teodorescu, C.; Uy-Evanado, A.; Nichols, G.A.; Chugh, S.S. Left-ventricular geometry and risk of sudden cardiac arrest in patients with preserved or moderately reduced left-ventricular ejection fraction. *Europace* 2017, 19, 1146–1152. [CrossRef] [PubMed]

56. Montagnana, M.; Lippi, G.; Franchini, M.; Banfi, G.; Guidi, G.C. Sudden cardiac death in young athletes. *Intern. Med.* 2008, 47, 1373–1378. [CrossRef]

57. Smit, D.L.; De Hon, O.; Venhuis, B.J.; Den Heijer, M.; De Ronde, W. Baseline characteristics of the HAARLEM study: 100 male amateur athletes using anabolic androgenic steroids. *Scand. J. Med. Sci. Sports* 2020, 30, 531–539. [CrossRef] [PubMed]

58. De Ronde, W.; Smit, D.L. Anabolic androgenic steroid abuse in young males. *Endocronol. Connect.* 2020, 9, 102–111. [CrossRef] [PubMed]

59. Salerno, M.; Cascio, O.; Bertozzi, G.; Sessa, F.; Messina, A.; Monda, V.; Pomara, C. Anabolic androgenic steroids and carcinogenicity focusing on Leydig cell: A literature review. *Onco Targets* 2018, 9, 19415. [CrossRef]

60. Monda, V.; Salerno, M.; Sessa, F.; Bernardini, R.; Valenzano, A.; Marsala, G.; Zammit, C.; Avola, R.; Carotenuto, M.; Messina, G.; et al. Functional changes of orexinergic reaction to psychoactive substances. *Mol. Neurobiol.* 2018, 55, 6362–6368. [CrossRef]

61. Rothman, R.D.; Weiner, R.B.; Pope, H.; Kanayama, G.; Fifer, M.A.; Baggish, A.L. Anabolic androgenic steroid-induced myocardial toxicity: An evolving problem in an ageing population. *BMJ Case Rep.* 2011. [CrossRef] [PubMed]

62. Youssef, M.Y.; Alqallaf, A.; Abdella, N. Anabolic androgenic steroid-induced cardiomyopathy, stroke and peripheral vascular disease. *BMJ Case Rep.* 2011. [CrossRef]

63. Baggish, A.L.; Weiner, R.B.; Kanayama, G.; Hudson, J.I.; Lu, M.T.; Hoffmann, U.; Pope, H.G., Jr. Cardiovascular toxicity of illicit anabolic-androgenic steroid use. *Circulation* 2017, 135, 1991–2002. [CrossRef] [PubMed]

64. Melchert, R.B.; Welder, A.A. Cardiovascular effects of androgenic-anabolic steroids. *Med. Sci. Sports Exerc.* 1995, 27, 1252–1262. [CrossRef]

65. Monda, V.; Salerno, M.; Fiorenzo, M.; Villano, I.; Viggiano, A.; Sessa, F.; Triggiani, A.; Cibelli, G.; Valenzano, A.; Marsala, G.; et al. Role of sex hormones in the control of vegetative and metabolic functions of middle-aged women. *Front. Physiol.* 2017, 8, 773. [CrossRef]
66. Shirpoor, A.; Heshmatian, B.; Tofighi, A.; Eliasabad, S.N.; Kheradmand, F.; Zerehpoosh, M. Nandrolone administration with or without strenuous exercise increases cardiac fatal genes overexpression, calcium/calmodulin-dependent protein kinase II, and monoamine oxidase activities and enhances blood pressure in adult wistar rats. *Gena* 2019, 697, 131–137. [CrossRef]

67. Vanderheyden, M.; Mullens, W.; Delrue, L.; Goethals, M.; De Bruyne, B.; Wijns, W.; Geelen, P.; Verstreken, S.; Wellens, F.; Bartunek, J. Myocardial gene expression in heart failure patients treated with cardiac resynchronization therapy responders versus nonresponders. *J. Am. Coll. Cardiol.* 2008, 51, 129–136. [CrossRef]

68. Kitzman, D.W.; Scholz, D.G.; Hagen, P.T.; Ilstrup, D.M.; Edwards, W.D. Age-related changes in normal hearts during the first 10 decades of life. Part II (maturity): A quantitative anatomic study of 765 specimens from subjects 20 to 99 years old. *Mayo Clin. Proc.* 1988, 63, 137–146. [CrossRef]

69. Mandal, R.; Loeffler, A.G.; Salamat, S.; Fritsch, M.K. Organ weight changes associated with body mass index determined from a medical autopsy population. *Am. J. Fore. Med. Pathol.* 2012, 33, 382–389. [CrossRef]

70. Neri, M.; Riezzo, I.; Pomara, C.; Schiavone, S.; Turillazzi, E. Oxidative-nitrosative stress and myocardial dysfunctions in sepsis: Evidence from the literature and postmortem observations. *Mediat. Inflamm.* 2016, 2016, 3423450. [CrossRef] [PubMed]

71. Kennedy, M.C.; Lawrence, C. Anabolic steroid abuse and cardiac death. *Med. J. Aust.* 1993, 158, 346–348. [CrossRef]

72. Sullivan, M.L.; Martinez, C.M.; Gennis, P.; Gallagher, E.J. The cardiac toxicity of anabolic steroids. *Prog. Cardiovasc. Dis.* 1998, 41, 1–15. [CrossRef]

73. Sculthorpe, N.; Grace, F.; Jones, P.; Davies, B. Evidence of altered cardiac electrophysiology following prolonged androgenic anabolic steroid use. *Cardiovasc. Toxicol.* 2010, 10, 239–243. [CrossRef]

74. Shirpoor, A.; Heshmatian, B.; Tofighi, A.; Eliasabad, S.N.; Kheradmand, F.; Zerehpoosh, M. Nandrolone administration with or without strenuous exercise increases cardiac fatal genes overexpression, calcium/calmodulin-dependent protein kinase II, and monoamine oxidase activities and enhances blood pressure in adult wistar rats. *Gena* 2019, 697, 131–137. [CrossRef]

75. Papamitsou, T.; Barlagiannis, D.; Papaliagkas, V.; Kotanidou, E.; Dermentzopoulou-Theodoridou, M. Testosterone-induced hypertrophy, fibrosis and apoptosis of cardiac cells—an ultrastructural and immunohistochemical study. *Med. Sci. Monit.* 2011, 17, 266. [CrossRef]

76. Riezzo, I.; Di Paolo, M.; Neri, M.; Bello, S.; Cantatore, S.; D’Errico, S.; Dinucci, D.; Parente, R.; Pomara, C.; Rabozzi, R.; et al. Anabolic steroid-and exercise-induced cardio-depressant cytokines and myocardial β1 receptor expression in CD1 mice. *Curr. Pharm. Biotechnol.* 2011, 12, 275–284. [CrossRef] [PubMed]

77. Kanayama, G.; Hudson, J.I.; Pope, H.G., Jr. Anabolic-androgenic steroid use and body image in men: A growing concern for clinicians. *Psychother. Psychosom.* 2020, 89, 65–73. [CrossRef]

78. Pope, H.; Brower, K.J. Treatment of anabolic-androgenic steroid related disorders. In *The American Psychiatric Publishing Textbook of Substance Abuse Treatment*; American Psychiatric Publishing: Washington, DC, USA, 2008; pp. 237–246.

79. Sessa, F.; Salerno, M.; Bertozzi, G.; Cipolloni, L.; Messina, G.; Aromatario, M.; Polo, L.; Turillazzi, E.; Pomara, C. miRNAs as novel biomarkers of chronic kidney injury in anabolic-androgenic steroid users: An experimental study. *Front. Pharmacol.* 2020, 11, 1454. [CrossRef]

80. Gatzoulis, K.A.; Arsenos, P.; Trachanas, K.; Dilaveris, P.; Antoniou, C.; Tsiachris, D.; Toussoulis, D. Signal-averaged electrocardiography: Past, present, and future. *J. Arrhythm.* 2018, 34, 222–229. [CrossRef] [PubMed]

81. Maior, A.S.; Menezes, P.; Pedrosa, R.C.; Carvalho, D.P.; Soares, P.P.; Nascimento, J.H.M. Abnormal cardiac repolarization in anabolic androgenic steroid users carrying out submaximal exercise testing. *Clin. Exp. Pharmacol. Physiol.* 2010, 37, 1129–1133. [CrossRef] [PubMed]

82. Ozdemir, O.; Bozkurt, I.; Ozdemir, M.; Yavuz, O. Side effect of metenolone enanthate on rats heart in puberty: Morphometrical study. *Exp. Toxicol. Pathol.* 2013, 65, 745–750. [CrossRef] [PubMed]

83. Strano-Rossi, S.; Fiore, C.; Chiarotti, M.; Centini, F. Analytical techniques in androgen anabolic steroids (AASs) analysis for antidoping and forensic purposes. *Mini Rev. Med. Chem.* 2011, 11, 451–458. [CrossRef]

84. Sessa, F.; Franco, S.; Picciocchi, E.; Geraci, D.; Chisari, M.G.; Marsala, G.; Polito, A.N.; Sorrentino, M.; Tripi, G.; Salerno, M.; et al. Addictions substance free during lifespan. *Acta Med. Mediter.* 2018, 34, 2081–2087.
85. Moacir, M.; Silva-Neto, J.A.; Neto, O.B. Acute interruption of treatment with nandrolone decanoate is not sufficient to reverse cardiac autonomic dysfunction and ventricular repolarization disturbances in rats. Steroids 2018, 132, 12–17.
86. Olivares, E.L.; Silveira, A.L.; Fonseca, F.V.; Silva-Almeida, C.; Cortes, R.S.; Pereira-Junior, P.P.; Nascimento, J.H.M.; Reis, L.C. Administration of an anabolic steroid during the adolescent phase changes the behavior, cardiac autonomic balance and fluid intake in male adult rats. Physiol. Behav. 2014, 126, 15–24. [CrossRef]
87. Marocolo, M.; Maior, A.S.; Katayama, P.L.; Mota, G.R.D.; Neto, O.B.; Lauria, A.D.A.; Santos, E.L. Anabolic steroid treatment induces cardiac autonomic dysfunction in rats: Time-course of heart rate variability. Am. J. Biomed. Eng. 2013, 3, 54–62.
88. Tanno, A.P.; Cunha, T.S.; Fernandes, T.; Guzzoni, V.; da Silva, C.A.; de Oliveira, E.M.; Costa Sampaio Moura, M.J.; Marcondes, F.K. Effects of nandrolone and resistance training on the blood pressure, cardiac electrophysiology, and expression of atrial β-adrenergic receptors. Life Sci. 2013, 92, 1029–1035.
89. Medei, E.; Marocolo, M.; de Carvalho Rodrigues, D.; Arantes, P.C.; Takiya, C.M.; Silva, J.; Rondinelli, E.; dos Santos Goldenberg, R.C.; Campos de Carvalho, A.C.; Nascimento, J.H.M. Chronic treatment with anabolic steroids induces ventricular repolarization disturbances: Cellular, ionic and molecular mechanism. J. Mol. Cell Cardiol. 2010, 49, 165–175. [CrossRef]
90. Phillis, B.D.; Abeywardena, M.Y.; Adams, M.J.; Kennedy, J.A.; Irvine, R.J. Nandrolone potentiates arrhythmogenic effects of cardiac ischemia in the rat. Toxicol. Sci. 2007, 99, 605–611. [CrossRef] [PubMed]
91. Binayi, F.; Joukar, S.; Najafipour, H.; Karimi, A.; Abdollahi, F.; Masumi, Y. The effects of nandrolone decanoate along with prolonged low-intensity exercise on susceptibility to ventricular arrhythmias. Cardiovasc. Toxicol. 2016, 16, 23–33. [CrossRef]
92. Sessa, F.; Salerno, M.; Di Mizio, G.; Bertozzi, G.; Messina, G.; Tomaiuolo, B.; Pisanelli, D.; Maglietta, F.; Ricci, P.; Pomara, C. Anabolic androgenic steroids: Searching new molecular biomarkers. Front. Pharmacol. 2018, 9, 1321. [CrossRef]
93. Sessa, F.; Maglietta, F.; Bertozzi, G.; Salerno, M.; Di Mizio, G.; Messina, G.; Montana, A.; Ricci, P.; Pomara, C. Human brain injury and mirnas: An experimental study. Int. J. Mol. Sci. 2019, 20, 1546. [CrossRef]

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