Hepatotoxicity Associated with Methylstenbolone and Stanozolol Abuse

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Abstract

Background & Objectives: Drug hepatotoxicity is a major cause of liver disease. Many drugs are well known to induce liver damage. Some toxic products, like anabolic androgenic steroids, that are pharmaceutical preparations since they contain pharmaceutically active substance, are available as nutritional supplements. Many patients are used to consume these like dietary stuff.

Methods: We introduce a case series of two patients who developed hepatic damage after the consumption of anabolic-androgenic steroids, accompanied by a detailed bibliographic research on this topic.

Results: We present two young men who developed significant liver damage, both with hyperbilirubinemia pattern after consumption of anabolic-androgenic steroids. This was associated with considerable morbidity, although both recovered without liver transplantation. The two anabolic-androgenic steroids were being marketed as dietary supplements.

Conclusions: Although not well controlled substances in Brazil, anabolic-androgenic steroids are cause of severe hepatotoxicity. Whereas the National Sanitary Surveillance Agency acts in the regulation of such substances, some of these products are still marketed as dietary supplements, requiring a more rigorous surveillance by health professionals.

Keywords: Hepatotoxicity; Anabolic steroids; Methylstenbolone; Stanozolol
**Abbreviation:** AAS: Anabolic-Androgenic Steroids; AP: Alkaline Phosphatase; GGT: Gamma-Glutamyltransferase; BT: Total Bilirubin; DB: Direct Bilirubin; IB: Indirect Bilirubin.

**Introduction**

Testosterone is a long time used medicine substance, with some therapeutic benefits, when used in special medical situations. However, testosterone and its synthetic derivatives have been used and sold, recklessly, like dietary supplements. These products are usually known as anabolic-androgenic steroids (AAS) [1-3]. The most commonly daily used derivatives include nandrolone, oxandrolone, stanozolol, stenbolone and oxymethylol [2]. The AAS have been used for a long time by athletes and body builders in order to boost performance [3,4]. Since 1975, these products are forbidden by the International Olympic Medical Committee and all of them are classified as controlled substances by the Food and Drug Administration [5].

In spite of this, many anabolic steroids are available in natural products stores and can also be purchased easily on the internet. The AAS are often consumed as dietary supplements. To highlight the risks of AAS exposition especially to emphasize hepatic side effects, we report two clinical cases of young men developing potentially fatal hepatotoxicity after consumption of M-STANE-Methylstenbolone (2,17a-Dimethyl-17b-hydroxy-5a-androst-1-en-3-one) which contains ultradrol, a prohormone that, once consumed, is metabolized as a steroid, and WINSTROL (stanozolol), considered a pure anabolic-androgenic steroid [6].

**Patient 1**

A 21-year-old male construction worker was referred to Antônio Pedro University Hospital in January 2014 with a history of diffuse abdominal pain, jaundice, choloria, acholic feces and severe pruritus, as well as an 18-kilogram weight loss in the last three months. There was no history of altered mental status. There was a report of illicit drug use from age 14 to 16, and former smoker 10 packs • year (quit five years ago); without associated diseases.

He admitted to using M-STANE one month prior to the start of his clinical condition, 2 capsules a day (dose recommended by the manufacturer) for a total of four weeks from October to November 2013. M-STANE was obtained from a “natural products” store, according to the patient’s report. The patient denied any prior liver disease, excessive alcohol intake, medication use or travel; as well as family history of liver disease.

At admission the patient was hemodynamically stable, but had a marked conjunctival icterus. His liver was palpable 2 cm below the right costal border. However, there was no evidence of fluid overload or hepatic encephalopathy. Initial laboratory results are shown in Table 1. Viral serologic tests for hepatitis A, B and C, Epstein-Barr virus, cytomegalovirus as well as antinuclear antibody titers, antimitochondrial antibody, and antismooth muscle antibody were all negative. Urea and creatinine were within normal range. Chest radiograph and electrocardiogram were normal. Ultrasonography showed no evidence of biliary obstruction or chronic liver disease. Hepatic biopsy was performed, which histopathology study revealed cholestatic liver disease with marked cholestasis and portal-portal fibrosis, in addition to the presence of portal venous ectasia (Figures 1A &1B). We chose a conservative treatment with ursodeoxycholic acid 900 mg/day.

The patient had jaundice (4+/4+) when he was discharged, without signs of encephalopathy, with a prothrombin time of 100% and in a good physical status. Figures 2A & 2B show the course of hyperbilirubinemia as well as an isolated peak of alkaline phosphatase reaching a level of 701 U/L in the absence of a change in the level of gamma-glutamyl transferase. Jaundice resolved over a period of 8 weeks. At this time ursodeoxycholic acid was finished because patient had no pruritus. At the last follow-up evaluation, 18 months after admission, the total bilirubin level was 0.34 mg/dL with alkaline phosphatase of 122U/L.

Figure 1A: Cholestatic liver disease (H.E. stain 20x), bile- plug dark arrow.

anabolic use, in addition to coexisting lobular and interface hepatitis suggestive of autoimmune hepatitis, possibly induced by the substance. In addition, there was the presence of portal fibrosis with short fibrous septa (F2). He was discharged with cholestyramine, hydroxyzine, ursodeoxycholic acid; the use of latter two being suspended by the patient himself two weeks later, due to diarrhea.

The total bilirubin level reached a peak of 37.5 mg/dL (Figure 4). Later on, hyperbilirubinemia showed a gradual decrease. At the last clinic appointment, one month after initial evaluation, total serum bilirubin levels were 2.55 mg/dL and transaminases were within normal limits.

Discussion

The two patients mentioned presented an important hepatic injury due to the use of AAS. Our first patient was consuming M-STANE (Methylstenbolone), which contains a prohormone called ultradrol that causes a high hormonal load, being like an AAS. Our second patient used Winstrol. It contains a synthetic steroid derived from testosterone called Stanozolol, which increases free testosterone in bloodstream [6]. Because of the serological and epidemiological exclusion of other causes and the compatibility in liver histopathology, the AAS consumed by these two patients was the most likely cause of hepatotoxicity. Initially, other causes of liver disease were excluded. Imaging studies revealed no evidence of biliary obstruction or vascular disease. The second patient had a history of alcohol consumption, but the biopsy was not compatible with an alcohol-induced injury. We should consider that alcohol may have increased susceptibility to hepatic injury induced by AAS.
in our second patient [2]. The hepatic biopsy in both patients was consistent with AAS-induced hepatotoxicity, and in addition transaminases results showed spontaneous improvement after discontinuation of the substance. Our patients didn’t show evidence from another kind of tissue damage [7].

Currently, AASs are classified as controlled substances, and it’s illegal to possess, manufacture and distribute these products, except in case of medical prescription. However, these products are easily available on the internet and in “natural products” stores as dietary supplements. Misuse of AAS is a public health issue. Under the influence of the body builder culture and the body image valorization, we see the indiscriminate consumption of these substances. Studies show that anabolic steroid use among high school seniors increased in the last years in the United States [6,8-10]. Also, men are more likely to use AAS compared to women [11]. Muscle dysmorphia is the new term used by Psychiatric researchers to describe people with an excessive fitness behavior, which can include toxic products consumption, with health risks [4,12,13].

AAS users practice unsafe injections and almost all report subjective side effects after the use of these drugs. Case reports linking AAS with other side effects have been reported including a bad lipid profile and atherogenesis [14], suppressed testicular function, cardiotoxicity, growth retardation, neuropsychiatric effects, nephrotoxicity including rare Wilms’ tumors, acute kidney injury, acute renal failure as a complication of rhabdomyolisis [11,12,14-16].

Also a handful of animal studies provides evidence that the use of these pharmaceutical substance could lead to alterations of renal structure and function and a many cardiovascular complications such as focal fibrosis, inflammatory infiltrations of cardiac tissue [17,18]. Hepatic complications include peliosishepatis, hepatic adenomas, hepatocellular carcinomas, and hepatotoxicity [17,18]. These side effects abides a multi-factorial, partly irreversible effect that includes augmented tissue oxidative status [15,19,20]. In our patients, there was no evidence of cardiac or renal damage, but they developed cholestasis and hepatocellular injury [17,18].

Cholestasis has been described as benign, but fatal cases have been reported. In most of cases, cholestasis induced by AAS is a result of a change in hepatocyte biliary secretion and generally is not associated with hepatocellular damage [21-23]. However, hepatitis may occur. Studies in rats showed that hepatotoxicity and increased levels of liver enzymes were drug-specific changes and that AAS had direct toxic effects on hepatocytes and that oxidative stress play a role, in impairment of the canalicular bile salt export pump [17,18,22-25]. Structural changes induced by AAS include degenerative effects on mitochondria and lysosomes [26]. Animal studies exhibited that also cardiotoxicity and nefrotoxicity are associated with an increase of oxidative stress biomarkers [27]. In the present study, both patients demonstrated cholestasis that improved after AAS discontinuation [17,18].

The differential diagnosis of hepatocellular injury with cholestasis includes viral hepatitis, drugs including alcohol, autoimmune disease and biliary duct obstruction. Family history of hepatic diseases was also analyzed. In our study, we investigated these diseases and there was no other explanation for the cholestasis and hepatic damage besides the use of AAS.

Conclusion

In conclusion, AAS consumption is a current tendency around the world, encouraged by body building culture. Although AAS-induced cholestasis is not very common, it is potentially fatal and associated with significant morbidity. Health professionals need to maintain a high level of vigilance over this scenario. Frequently, patients don’t tell about AAS consumption. It is important to regularly inquire patients about the consumption of AAS or dietary supplements. Also, it is imperative to educate our patients and health care professionals about the hepatotoxicity and other risks associated with the use of these products. Also, public health authorities must find strategies to control AAS market and consumption.

References


