**Review Article**

**Nandrolone Decanoate effects on cardiac morphology and function: a structured review and critical appraisal of randomized controlled trials.**

**Abstract:**

**Aims:** The cardiovascular risks associated with anabolic-androgenic steroid (AAS) use, particularly nandrolne decanoate (ND), remain a topic of debate due to conflicting evidence from different types of studies. While observational research frequently reports adverse cardiac effects, findings from randomized controlled trials (RCTs) suggest otherwise. This review aims to critically assess the available RCTs evaluating the impact of ND on cardiac morphology and function, with a focus on their methodological quality and clinical relevance.

**Study Design:** Structured review of randomized controlled trials.

**Method:** A systematic literature search was conducted to identify RCTs that examined the effects of ND on cardiac function. The analysis included study design, sample size, inclusion/exclusion criteria, cardiac evaluation methods, and reported outcomes. The methodological quality of each study was assessed using the Jadad scale to ensure a rigorous evaluation of the evidence.

**Results:** The two reviewed RCTs, which were of moderate to high methodological quality, did not report clinically significant cardiac dysfunction in healthy individuals after short-term exposure to ND. However, these findings contrast with several observational studies, where users—often exposed to uncontrolled conditions and higher doses—frequently present with structural and functional cardiac changes. A key limitation of the RCTs was their small sample sizes, short follow-up durations, and reliance on conventional imaging techniques, which may not be sensitive enough to detect subtle cardiac alterations.

**Conclusion:** While current RCTs do not provide strong evidence of cardiovascular harm from ND, their methodological limitations underscore the need for further research. Future trials with larger sample sizes, higher statistical power, and more advanced cardiac imaging techniques will be crucial in clarifying the true impact of ND on the heart. Until more definitive data are available, routine cardiac monitoring should be considered for individuals using ND, particularly those on prolonged or high-dose regimens.

**Keywords:** Nandrolone decanoate, anabolic steroids, cardiac function, ventricular hypertrophy, cardiovascular risk.

**Introduction:**

The use of anabolic-androgenic steroids (AAS) has become a widespread global phenomenon, raising significant concerns regarding their potential adverse effects [1]. Among these compounds, nandrolone decanoate (ND) is one of the most used, and frequently linked to cardiovascular risks [2]. Observational studies suggest possible cardiac alterations, such as ventricular hypertrophy and diastolic dysfunction, reinforcing concerns that its use may compromise long-term cardiovascular health [3]. However, much of this evidence comes from methodologically weak studies, often lacking proper control of essential variables, making it difficult to draw definitive conclusions [4].

Randomized controlled trials (RCTs), recognized as the gold standard for establishing causality, could provide more reliable insights into the true cardiovascular effects of ND [4]. However, the limited RCTs available suffer from small sample sizes, short follow-up periods, and reliance on conventional cardiac assessment methods, which may not be sensitive enough to detect early or subclinical changes [5,6]. This knowledge gap fuels ongoing uncertainty about ND’s safety, particularly among users exposed to prolonged or high-dose regimens.

Given this scenario, it is imperative to critically evaluate the existing evidence and identify the limitations of studies conducted thus far. This structured review aims to analyze published RCTs on the cardiovascular effects of ND, assessing their methodological quality and the reliability of their findings. A clearer understanding of ND’s actual risks is essential to guide future research, improve clinical safety, and provide more definitive answers regarding its impact on human cardiac morphology and function.

**Method:**

A structured scientific search [7] was conducted in the PubMed database ([www.pubmed.gov](http://www.pubmed.gov/)) using a combination of MeSH terms and synonymous terms, along with Boolean operators (OR, AND, NOT) to identify specific studies addressing the use of Nandrolone Decanoate and mapping its cardiovascular effects in humans, excluding studies conducted on animal models. Both authors independently screened all studies and reached full agreement on the inclusion of the specific studies identified. The complete search strategy and the results obtained are presented in the table below (**Table 1**).

**Table 1** – Database, Search Strategy and Results

|  |  |
| --- | --- |
| **Database** | **PubMed** |
| **Search Strategy** | ("Nandrolone"[Mesh] OR (19-Nortestosterone) OR (17beta-Hydroxy-19-Nor-4-Androsten-3-One) OR (17beta Hydroxy 19 Nor 4 Androsten 3 One) OR Estrenolone OR Nortestosterone OR (17-Hydroxy-Estr-4-Ene-3-One) OR Norandrostenolone OR (17 beta Hydroxyestr 4 en 3 one decanoate) OR (19 Nortestosterone Decanoate) OR (19 nor 4 Androstene 17 beta ol 3 one 17 decanoate) OR Retabolil OR Retabolyl OR Decadurabolin OR Decadurobolin)) AND ((randomized controlled trial[pt] OR controlled clinical trial[pt] OR randomized[tiab] OR placebo[tiab] OR drug therapy[sh] OR randomly[tiab] OR trial[tiab] OR groups[tiab] NOT (animals [mh] NOT humans [mh]))) |
| **Results:** Nandrolone Decanoate + Randomized Controlled Trials + Humans | Total: 913 studies  Selected / included: 2 studies |

**Results:**

As seen at the end of the previous section (**Table 1**), only two randomized studies were identified in the search, Hartgens F, et al. (2003) [5] and Chung T, et al. (2007) [6], and they will be described in detail below.

In the first study, Hartgens F, et al. (2003) published the results of two studies in a single article [5]. The first was an uncontrolled study that evaluated 32 bodybuilders who self-administered AAS, with varying cycles, including differences in doses, combinations, and durations of use. The second study, which will be used for the purposes of this article, was a randomized, placebo-controlled trial that assessed the cardiovascular effects (morphological and functional measures) through two-dimensional Doppler echocardiography in 16 healthy men, before and after the administration of 200 mg/week of Nandrolone Decanoate for 8 weeks [5].

To minimize potential confounders, all subjects underwent a comprehensive medical examination to assess their health status and to exclude any relevant medical conditions, such as hypertension, diabetes mellitus, liver disease or abnormal liver enzyme levels, hereditary hypercholesterolemia, elevated serum total cholesterol (>6.5 mmol/L), infertility, and smoking. Additionally, diet, training, and anti-doping controls (to detect the presence of other substances potentially used inadvertently by participants) were rigorously maintained, with no differences observed between the placebo and nandrolone decanoate groups throughout the study, as measured at baseline, week four, and week eight. As results, the researchers did not observe any clinically significant morphological or functional changes in any of the echocardiographic parameters assessed [5].

In a second study, Chung T, et al. (2007) [6] evaluated the comparative effects of testosterone (n = 10) and nandrolone decanoate (n = 10) on cardiac function in 30 healthy young men (placebo group: n = 10) in a randomized, double-blind, placebo-controlled clinical trial. Participants received 200 mg/week of testosterone, nandrolone, or placebo for 4 weeks, with detailed assessments and hemodynamic monitoring performed through transthoracic Doppler echocardiography.

Moreover, the sample was carefully selected to avoid potential confounding factors related to testosterone or ND effects in cardiac function. Thus, the exclusion criteria adopted by the researchers were: (i) contraindications to testosterone administration (breast or prostate cancer); (ii) previous or current use of androgens; (iii) use of disallowed drugs (including illicitly obtained substances) that may interfere with androgen absorption, distribution, metabolism, excretion, or action (e.g., androgens, anti-androgens, finasteride); (iv) any chronic medical condition requiring regular medication or likely to interfere with safe participation; and (v) significant cardiac disease or abnormal baseline echocardiogram [6].

The results demonstrated that neither of the two androgens (Testosterone and ND) caused clinically significant changes in cardiac function. The testosterone group showed alterations that, although statistically significant, were minor, affecting diastolic function and the ECG. However, all these changes remained within normal range limits and were not considered clinically relevant by the authors. In contrast, nandrolone did not produce any significant effects on the evaluated cardiac function or ECG [6].

The studies cited and their respective results are summarized in the following table (**Table 2**).

**Table 2:** Descriptive characteristics of the studies by Hartgens F, et al. (2007), and Chung T, et al. (2007).

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **Author**  **Year** | **Study**  **Type** | **ND**  **group** | **ND dose** | **Duration** | **CV Outcome** |
| Hargens F, et al.  2003  [5] | RCT | 16 healthy young man | 200 mg / week | 8 weeks | No clinically significant adverse effects in morphology and function |
| Chung T, et al.  2007  [6] | RCT | 10 healthy young man | 200 mg / week | 4 weeks | No clinically significant adverse effects in morphology and function |

The assessment of the methodological quality, by Jadad Scale [8], of the two randomized controlled trials identified through the structured search with a systematic approach in PubMed indicated that they exhibited moderate to high quality (3/5 and 5/5 points). The details are presented in **Table 3** below.

**Table 3:** Jadad Scale Evaluation of the Studies

|  |  |  |
| --- | --- | --- |
| **Criterion** | **Hartgens F, et al. (2003)** | **Chung T, et al. (2007)** |
| **Study described as randomized** | **1** | **1** |
| **Randomization method described and appropriate** | **0** | **1** |
| **Study described as double-blind** | **1** | **1** |
| **Blinding method described and appropriate** | **0** | **1** |
| **Description of withdrawals and dropouts** | **1** | **1** |
| **Total Score** | **3** | **5** |

**Discussion:**

The main finding observed in this review was that no clinically significant deleterious changes in cardiac morphology or function were found in healthy young individuals in randomized controlled trials assessed with moderate to high methodological quality, following the short-term administration of nandrolone decanoate [5,6].

This finding differs from what is generally postulated in the scientific literature [2], and these discrepancies, between findings from observational studies and randomized controlled trials (RCTs) can, at least to some extent, be attributed to fundamental methodological differences [5,6,7,9]. RCTs are conducted in a highly controlled environment, where participants are carefully selected, undergo baseline assessments to exclude preexisting conditions, and have their individual, familial, and lifestyle-related risk factors balanced across groups [7,9]. Furthermore, in these studies, the substances under investigation are standardized, sourced from the pharmaceutical industry, administered in moderately supraphysiological doses, and used for a relatively short duration [5,6].

In contrast, observational studies often reflect a far more complex and uncontrolled reality. These studies lack randomization, participants may have diverse health backgrounds without a structured baseline assessment, and risk factors are unevenly distributed between groups [9,10]. Additionally, these studies frequently involve individuals who use multiple substances simultaneously (abuse of licit and illicit drugs), including drugs obtained from the underground market (adulteration, contamination), often in extremely high doses—sometimes up to 30 times the therapeutic levels—and for prolonged, uninterrupted periods [10,11]. Such methodological discrepancies must be considered when interpreting findings, as direct extrapolations between these different types of evidence can lead to misinterpretations [10,11].

Still within this context of methodological quality and its differences, Goldman A. and Basaria S. (2018) [4] emphasize that the most severe adverse effects, particularly cardiovascular ones, are often reported in studies with methodological limitations that lack the rigor necessary to establish clear causal relationships. Case reports, case series, retrospective case-control studies, cross-sectional analyses, and uncontrolled cohort studies frequently present significant constraints, making their conclusions less reliable [4,9]. Furthermore, as highlighted by Fanaroff AC et al. (2020) [9], when comparing observational data with randomized clinical trials, the absence of proper randomization can introduce considerable bias, hindering an accurate assessment of the risks and benefits of an intervention.

Specifically, regarding ND, in a comprehensive review conducted by Patanè FG et al. (2020) [3], studies were presented investigating potential mechanisms through which ND abuse could induce cardiovascular damage, ranging from alterations in the cardiac protein expression of myosin isoforms, enzymatic and autonomic changes, to indirect effects such as increased hematocrit, blood pressure, and dyslipidemia. However, the cited references by the authors in the cardiovascular adverse effects section are mostly derived from other review studies, and pathophysiological mechanisms research conducted in animal models, and based on the latter, making direct extrapolation to humans inappropriate, despite being explanatory [3].

In this regard, an important point for reflection is that animal models, frequently employed in the investigation of adverse effects of AAS in various aspects, including, therefore, ND [12,13], play a crucial role in biomedical research, allowing for the exploration of physiological and pathological mechanisms due to their cellular and molecular similarities with humans [14,15]. However, the direct extrapolation of these findings to human clinical practice must be approached with caution, as interspecies differences can significantly impact the observed effects [16,17]. As highlighted in a publication in *Arquivos Brasileiros de Cardiologia*, by Carvalho VO. & Guimarães GV (2010) the physiology of the cardiovascular system in small rodents presents substantial differences compared to humans, which may compromise the applicability of experimental results to clinical practice [14].

Having established these essential considerations regarding the discussion of different scenarios—controlled and uncontrolled, human and animal models—and their potential limitations in inferring direct and specific causality of ND in humans, it is also necessary to address some key aspects of the RCTs presented.

In this regard, Kindermann W. and Urhausen A. (2004) [18] highlighted that the study by Hartgens F. et al. (2003) [5] did not assess diastolic function, a crucial parameter for the early detection of pathological myocardial hypertrophy. More recent studies, although observational, suggest that detrimental alterations in both systolic and diastolic function can be found in AAS users and that these changes may become permanent once established, persisting even after discontinuation of use [2,19].

Furthermore, the randomized clinical trials analyzed relied on conventional echocardiography for cardiac function assessment [5,6]. However, with advancements in imaging technology, more sensitive and refined methods—such as cardiac magnetic resonance imaging and speckle-tracking analysis—have become available and are increasingly employed in recent observational studies. These techniques provide a more detailed and comprehensive evaluation of cardiac morphological and functional changes, allowing for the early detection of potential adverse effects [2,19].

Another relevant methodological and statistical consideration is the small sample size of both RCTs, which collectively assessed only 26 individuals [5,6]. Additionally, only the study by Chung T. et al. (2007) [6] included a sample size calculation and statistical power analysis, with an 80% power to detect an absolute increase or decrease of 6.5%. Therefore, given these potential methodological and statistical limitations, the validity of the findings may be impacted, particularly in detecting early clinical small relevant effects.

Although the study by Hartgens et al. (2003) [5] was conducted as a randomized, double-blind, placebo-controlled trial, certain methodological shortcomings limit its overall robustness. The lack of a detailed description of the randomization process makes it difficult to assess whether an appropriate method was used, leaving room for potential selection bias. Similarly, while the study was classified as double-blind, it does not specify how blinding was maintained for both participants and investigators, raising concerns about potential performance and detection bias. These methodological gaps resulted in a two-point deduction on the Jadad scale, placing the study within a moderate quality range [5].

Moreover, other potentially important factors related to cardiovascular outcomes, such as alterations in lipid profile (particularly increased LDL and reduced HDL), hematological changes (elevated hematocrit), autonomic dysfunction, and dysregulation of the renin-angiotensin-aldosterone system (RAAS), were not assessed in these studies [20, 21].

Considering all these aspects and controversies, it appears that a knowledge gap persists regarding the specific cardiovascular effects of ND. On one hand, we have RCTs with methodological limitations—particularly concerning assessment accuracy—that fail to demonstrate clinically significant adverse effects [5,6]. On the other hand, we have uncontrolled observational studies conducted in a complex real-world setting, which employed more advanced analytical techniques and identified significant deleterious alterations [2,22,23].

From a therapeutic standpoint, in diseased individuals, multiple RCTs conducted in humans—providing potentially more extrapolable findings than those derived from animal models—have already investigated the use of ND at comparable doses (at least 200 mg/week) for periods between 12 weeks and 12 months [24-28]. As reported by the authors, although the assessment of cardiac morphology and function was not the specific objective, these studies did not observe increased cardiovascular adverse events (**Table 4**), even among individuals with higher baseline risk (such as chronic kidney disease patients) [24-28], reinforcing and, over a longer duration of use, the non-clinically significant findings of the studies by Hartgens F et al. (2003) [5] and Chung T et al. (2007) [6].

**Table 4:** Summary of RCT Studies on therapeutic use of Nandrolone Decanoate

| **Author / Year** | **Disease** | **Nandrolone Dose** | **Study Duration** | **Adverse Cardiovascular Events** |
| --- | --- | --- | --- | --- |
| **Cattran et al. (1977)** | Uremic anemia | 200 mg/week | 12 months | No serious events reported |
| **Johansen et al. (2006)** | Hemodialysis patients | 100 mg (women) / 200 mg (men) per week | 12 weeks | No serious events reported |
| **Gascón et al. (1999)** | Anemia in elderly hemodialysis patients | 200 mg/week | 6 months | No serious events reported |
| **Williams et al. (1973)** | Anemia in chronic renal failure | 200 mg/week | 13 weeks | No serious events reported |
| **Navarro et al. (2002)** | Patients on peritoneal dialysis | 200 mg/week | 6 months | No serious events reported |

Thus, in our view, well-designed RCTs with appropriate sample size calculations, sufficient statistical power, and the incorporation of more advanced echocardiographic and imaging techniques would be highly valuable and strongly recommended to either corroborate or refute the findings of the two studies reviewed here, regarding the use of ND in healthy young man.

Finally, until more robust, detailed and adequate quality evidence is available regarding the true influence, safety and efficacy, of high doses of ND, it would be prudent to ensure that every user or patient undergoes comprehensive cardiac function and morphology assessment, facilitating the early detection of cardiovascular any damage.

**Conclusion:**

The available RCTs, conducted with moderate to high methodological quality, did not demonstrate clinically significant adverse effects on cardiac morphology or function in young, healthy individuals following short-term administration of ND [5,6]. However, these findings contrast with those from observational studies, which frequently report deleterious cardiovascular changes in chronic AAS users [2,22,23]. This discrepancy likely stems from fundamental methodological differences, such as the absence of randomization, heterogeneous participant profiles, and the use of substantially higher doses in uncontrolled settings [9,11].

Despite these differences, it is important to recognize the limitations of the analyzed RCTs, particularly their small sample sizes and reliance on conventional echocardiographic methods, which may not be sufficiently sensitive to detect early or subtle cardiac changes. The integration of more advanced imaging techniques, such as cardiac magnetic resonance imaging and speckle-tracking analysis, could enhance the precision of future assessments. Moreover, while therapeutic clinical trials have suggested the safety of ND at similar doses (200 mg/week) over longer periods, extrapolating these findings to recreational users requires caution due to factors such as polypharmacy and individual susceptibilities.

Given these considerations, there is a pressing need for well-designed RCTs with larger sample sizes, appropriate statistical power calculations, and advanced cardiac imaging methodologies to further clarify the cardiovascular effects of ND. Until stronger and more definitive evidence becomes available, it would be prudent to ensure that any individuals using high doses of ND undergo comprehensive cardiac evaluations, facilitating the early detection of potential risks and optimizing long-term cardiovascular monitoring.

COMPETING INTERESTS DISCLAIMER:

Authors have declared that they have no known competing financial interests OR non-financial interests OR personal relationships that could have appeared to influence the work reported in this paper.

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