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1 **Long-Term Testosterone Therapy Improves Liver Parameters and Steatosis in**
2 **Hypogonadal Men: A Prospective Controlled Registry Study**

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18 **Abbreviated title:** Long-term testosterone improves hepatic steatosis

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32

33 **Abstract**

34 Non-alcoholic fatty liver disease (NAFLD) is associated with cardiovascular disease (CVD)
35 and both are prevalent in men with testosterone deficiency. Long-term effects of testosterone
36 therapy (TTh) on NAFLD are not well studied. This observational, prospective, cumulative
37 registry study assesses long-term effects of testosterone undecanoate (TU) on hepatic
38 physiology and function in 505 hypogonadal men (T levels ≤ 350 ng/dL). 321 men received
39 TU 1000mg/12 weeks for up to 12 years following an initial 6-week interval (T-group), while
40 184 who opted against TTh served as controls (C-group). T-group patients exhibited
41 decreased fatty liver index (FLI, calculated according to Mayo Clinic guidelines) (83.6 ± 12.08
42 to 66.91 ± 19.38), γ -GT (39.31 ± 11.62 to 28.95 ± 7.57 U/L), bilirubin (1.64 ± 4.13 to 1.21 ± 1.89
43 mg/dL) and triglycerides (252.35 ± 90.99 to 213 ± 65.91 mg/dL) over 12 years. Waist
44 circumference and body mass index were also reduced in the T-group (107.17 ± 9.64 to
45 100.34 ± 9.03 cm and 31.51 ± 4.32 to 29.03 ± 3.77 kg/m²). There were 25 deaths (7.8%) in the T-
46 group of which 11 (44%) were cardiovascular related. In contrast, 28 patients (15.2%) died in
47 C-group, and all deaths (100%) were attributed to CVD. These data suggest that long-term
48 TTh improves hepatic steatosis and liver function in hypogonadal men. Improvements in
49 liver function may have contributed to reduced CVD-related mortality.

50 **Introduction**

51 In recent decades, the prevalence of hepatic steatosis has increased substantially, reaching
52 10–24% in the overall global population, and is directly associated with the increased
53 incidence of type 2 diabetes (T2D) and obesity that afflicts western societies [1]. Hepatic
54 steatosis is characterised by fat deposition in the liver and is considered the precursor to non-
55 alcoholic fatty liver disease (NAFLD). As NAFLD progresses, liver inflammation and
56 damage can occur ultimately leading to cirrhosis, liver failure and often hepatocarcinoma.
57 Hepatic steatosis linearly correlates with incidence of T2D and metabolic syndrome (MetS),
58 obesity and key features of these diseases including insulin resistance, hyperinsulinemia and
59 dyslipidemia [2-4]. NAFLD prevalence ranges from 50%–75% in patients with T2D [5-8]
60 and from 80%–90% in obese patients [9-11]. Hepatic steatosis increases CVD risk and
61 cardiovascular events in patients with T2D (odds ratio 1.84) [12-15]. Indeed, with their
62 shared aetiology hepatic steatosis and atherosclerosis are often considered tissue specific
63 manifestations of the same cardiometabolic pathology [16].

64 In addition to the characteristic symptoms of functional hypogonadism such as impaired
65 libido, erectile dysfunction, fatigue, increased risk of depression and reduced quality of life
66 (QoL) [17], low testosterone is associated with T2D, MetS and is considered an independent
67 cardiovascular risk factor [18-20]. T2D is prevalent in men with low testosterone levels and
68 increases the risk of cardiovascular disease (CVD) in this population [21,22]. Testosterone
69 therapy (TTh), the primary treatment for alleviating symptoms of functional hypogonadism,
70 has been reported to reduce insulin resistance, dyslipidemia and central adiposity, improve
71 glycaemic control and consequently decrease cardiovascular risk in hypogonadal men with
72 T2D and/or MetS [23-28].

73 A limited number of studies have explored the relationship between hepatic steatosis and
74 testosterone often with conflicting results. Some population-based studies report a correlation
75 between low circulating serum testosterone levels and hepatic steatosis [29-32], whereas
76 others found no association [33]. Hepatic steatosis, defined by sonographic criteria, has been
77 demonstrated to be directly correlated with a total testosterone level lower than 14.2 nmol/L
78 in men in a retrospective cohort study even after adjusting for confounding factors including
79 age, BMI, smoking, diabetes, and visceral adipose tissue [29]. Furthermore, it was reported
80 that low total testosterone levels in healthy Korean men were inversely associated with
81 NAFLD and this association persisted after controlling for the effect of visceral adiposity,
82 insulin resistance and low grade inflammation, indicating an independent correlation [30]. In
83 obese men with sleep apnoea, 18 weeks of TTh led to improvements in insulin sensitivity and
84 reduced liver fat content assessed by CT imaging despite the men not being initially selected
85 for low testosterone [34]. Supporting these clinical findings, several animal studies have also
86 demonstrated beneficial effects of testosterone on hepatic steatosis. Hepatic lipid deposition
87 is elevated in animal models of testosterone deficiency fed a high fat diet [35-37] an effect
88 that is abrogated following TTh [36,38]. The improvement in hepatic physiology as a result
89 of TTh was accompanied by altered expression of important regulatory genes involved in
90 hepatic lipid assembly and secretion [36]. Moreover, TTh ameliorated hepatic steatosis and
91 steatohepatitis by suppressing endoplasmic reticulum stress, inhibiting macrovesicular lipid
92 droplet formation and promoting very-low-density lipoprotein export in castrated male rats
93 [38].

94 Hypogonadism in men with hepatic steatosis is likely to be multifactorial. Patients with low
95 testosterone levels often have comorbidities such as T2D, MetS and CVD which may
96 contribute to hepatic steatosis and can ultimately perpetuate hypogonadism through the
97 hypogonadal-obesity-adipocytokine axis [39]. While experimental and clinical data suggest

98 physiological testosterone, whether endogenous or administered, may prevent or ameliorate
99 hepatic steatosis in males, few studies investigate the direct effects of long-term TTh on
100 hepatic steatosis. The present long-term registry study investigates hepatic steatosis and
101 parameters of liver function in hypogonadal men following 12-years of TTh to establish the
102 role of testosterone in hepatic pathophysiology.

103

104 **Methods**

105 For this observational, prospective, cumulative registry study 505 elderly men (mean age:
106 61.4 ± 9.7 years) with symptoms of functional hypogonadism, where symptoms were defined
107 as at least moderate symptoms on the Aging Males' Symptoms scale, and a total testosterone
108 (TT) ≤ 350 ng/dL (≤ 12.1 nmol/L) were identified for inclusion. Exclusion criteria included
109 active prostate or male breast cancer, indicated desire for paternity (especially in younger
110 patients), allergy to ingredients in the testosterone formulation, acute heart disease within last
111 6 months or uncontrolled blood pressure problems. The study protocol conforms to the
112 ethical guidelines of the 1975 Declaration of Helsinki as reflected in a priori approval by the
113 German Ärztekammer (German medical association), and written informed consent was
114 obtained from each patient included in the study and for receiving their treatment protocol.
115 321 men received 1000 mg testosterone undecanoate (TU) parenterally every 12 weeks,
116 following an initial 6-week interval, for up to 12 years (T-group). TTh was temporarily
117 discontinued in 147 of these men after 5.5 years for 17 months (mean) due to reimbursement
118 issues. TTh was re-established in these patients upon resolution of the issues. The 184 men
119 who opted against TTh comprised the control group (C-group).

120 Total testosterone and SHGB were measured by enzyme-linked immunosorbent assay (Axym
121 System, Abbott Germany. Threshold: 10-73 nmol/l). The effects of long-term TU on hepatic
122 steatosis was assessed biannually by measuring fatty liver index (FLI), calculated based on
123 waist circumference, body mass index (BMI), triglyceride, and gamma-glutamyl-transferase
124 (γ -GT) using the formula published by Bedogni G et al. [40]. A FLI < 30 rules out (negative
125 likelihood ratio: 0. 2) and a FLI ≥ 60 indicates with a positive likelihood ratio of 4.3 that the
126 patient has hepatic steatosis. Parameters of liver function, serum gamma-glutamyl transferase
127 (γ -GT), aspartate transaminase (AST) and alanine transaminase (ALT) were assessed

128 enzymatically on the Abbott Alinity c analytical system. Bilirubin was measured by
129 Photometric Colour Test (Alinity C-Module Abbott).

130

131 *Statistical Analysis*

132 The statistical analysis software Statistical Package for Social Sciences (SPSS) v.11 for
133 Windows (SPSS Inc., Chicago, USA) was used for data processing and analysis. Data are
134 expressed at each timepoint as mean group values with standard deviations. Comparison of
135 clinical parameters between groups across the time points was done using a mixed-effects,
136 repeated-measures model with period, group and their interaction as fixed effects. Analysis of
137 variance (ANOVA) was used to compare continuous variables with significance accepted
138 where $p < 0.05$.

139

140 **Results**

141 Patient baseline characteristics included in this registry are shown in Table 1. The mean
142 baseline age for T-group consisting of 321 patients was 59 ± 9.5 years, with a mean follow-
143 up of 8.3 ± 3.5 years. In the C-group (184 patients), the mean age was 66.1 ± 7.6 years, and
144 mean follow-up of 5.5 ± 1.6 years. The FLI ($p < 0.0001$), bilirubin ($p < 0.05$), triglycerides
145 ($p < 0.0005$), BMI ($p < 0.0001$) and WC ($p < 0.0001$) were significantly higher at baseline in the
146 T-group compared to the C-group. For all subsequent comparisons over the study duration,
147 data were only available for the T-group beyond 8-years.

148 Men in the T-group had a significant elevation in TT levels following TTh at the 1-year
149 follow-up (7.74 nmol/L to 16.11 nmol/L, $p < 0.0001$), which was sustained throughout the 8-
150 year study period (15.98 nmol/L), compared to C-group patients (9.22 nmol/L at baseline to
151 9.24 nmol/L at 8-years) (Figure 1). TT levels continued to increase beyond 8 years in the T-
152 group with a mean concentration of 22.83 ± 2.34 nmol/L at 12 years (Figure 1). SHBG was
153 only measured in the T-group and declined from 36.68 ± 22.45 to 30.61 ± 13.85 at 8-years and
154 then further to 29.91 ± 9.17 at 12 -years. Haematocrit increased in the T-group from
155 42.55 ± 3.94 to 45.96 ± 3.44 at 8-years and declined in the C-group from 45.01 ± 3.78 to
156 42.75 ± 1.52 . Haemoglobin increased in the T-group from 14.14 ± 1.16 to 14.49 ± 0.98 and
157 decreased in the C-group from 13.75 ± 1.16 to 13.61 ± 0.93 .

158 The FLI in the T-group decreased from 83.6 ± 12.08 at baseline to 66.91 ± 19.38 at 8-years
159 ($p < 0.001$), compared to the C-group which increased from 68.67 ± 19.35 to 81.35 ± 16.91
160 ($p < 0.001$); with years 3-8 having a significantly lower FLI in the T-group compared to the C-
161 group ($p < 0.0001$, Figure 2). The decline in the FLI continued in the T-group up to 12-years
162 (59.13 ± 9.52).

163 γ -GT significantly decreased in the T-group from 39.31 ± 11.62 at baseline to 28.95 ± 7.57 U/L
164 at 8-years ($p<0.0001$). By 12 years γ -GT in the T-group had decreased further to 24.65 ± 3.67 .
165 Whereas it increased from 37.79 ± 29.55 at baseline to 39.5 ± 26.71 U/L at 8-years in the C-
166 group ($p<0.005$; Figure 3A) demonstrating a between group difference from year 1 to year 8
167 ($p\leq 0.005$). A non-significant decline in bilirubin levels was observed in T-group patients
168 from 1.64 ± 4.13 to 1.21 ± 1.89 mg/dL ($p=0.1716$), C-group patients remained unchanged
169 (1.04 ± 7.08 to 1.12 ± 1.96 mg/dL; $p>0.05$, Figure 3B). Although T-group had significantly
170 elevated bilirubin levels at baseline ($p<0.05$) by the 8 year point they were reduced to similar
171 levels to those in C-group. At 12 years in the T-group, bilirubin was 0.9 ± 0.15 which was still
172 not statistically significant vs. baseline. Additionally, AST levels remained unchanged versus
173 baseline at 8 years for both groups and at 12 years for T-group (Figure 4A). Fluctuations
174 occurred throughout with statistically significant increased at 2 ($p<0.0005$), 3 ($p<0.0001$), 4
175 ($p<0.0005$), and 7 years ($p<0.01$), then a significant decrease at 9 ($p<0.0001$) and 10 years
176 ($p<0.001$) but at no other time point for the T-group, while AST significantly increased at 2
177 ($p<0.0005$) and 4 years ($p<0.0005$) for the C-group but not at any other point in time. Between
178 group differences in AST were only demonstrable at 1 ($p<0.0001$), 2 ($p<0.005$) and 4 years
179 ($p<0.005$) with the T-group having lower levels, but these differences were not sustained. ALT
180 levels declined slightly for both patient-groups but not significantly so at the end of the study
181 period (Figure 4B). In the T-group, ALT was significantly increased versus baseline at 2
182 ($p<0.005$), 3 ($p<0.0001$), 4 ($p<0.0005$), and 5 years ($p<0.05$) and decreased at 8 ($p<0.05$), 9
183 ($p<0.0001$) and 10 years ($p<0.0005$), while C-group had an increase statistically significant
184 versus baseline only at 3 years ($p<0.05$). Between group differences showed a T-group ALT
185 level significantly lower than the C-group only at year 1 ($p<0.005$).

186 Triglycerides decreased from 252.35 ± 90.99 at baseline to 213 ± 65.91 mg/dL at 8-years in the
187 T-group and increased from 196 ± 91.31 at baseline to 244.55 ± 61.39 mg/dL at 8-years in the

188 C-group after 8 years; although a significant difference between the groups was reached after
189 the first year and continued up to 8 years ($p<0.0001$, Figure 5). Triglyceride levels continued
190 to decline in the T-group up to 12-years (175.01 ± 37.31 , $p<0.0001$ vs. baseline).

191 Body mass index (BMI) decreased from 31.51 ± 4.32 at baseline to 29.03 ± 3.77 kg/m^2 at 8-
192 years in the T-group and increased from 29.2 ± 3.22 at baseline to 30.68 ± 3.99 kg/m^2 at 8-years
193 in C-group ($p<0.01$, Figure 6A). There was a significant difference of BMI between the
194 groups after 3 years and this trend continued up to 8 years. Furthermore, BMI continued to
195 decline in the T-group up to 12-years (27.57 ± 2.11 , $p<0.0001$ vs. baseline). Waist
196 circumference (WC) was reduced in the T-group from 107.17 ± 9.64 at baseline to
197 100.34 ± 9.03 cm at 8-years and increased from 99.8 ± 9.13 at baseline to 104.65 ± 8.25 cm at 8-
198 years in the C-group ($p<0.05$, Figure 6B). Similarly to BMI, WC continued to decrease
199 significantly in the T-group compared to the C-group after 3 years of TTh. WC also
200 continued to decline in the T-group up to 12-years (93.56 ± 3.05 , $p<0.0001$ vs. baseline).

201 Twenty-seven deaths (14.7%) were recorded in the C-group, all of which were attributed to
202 CVD including myocardial infarction (13, 48%), stroke (7, 27%), heart failure (3, 11%),
203 aortic aneurysm (2, 7%) and lung embolism (2, 7%) (Table 2). In contrast, significantly less
204 recorded deaths of 25 (7.8%, $p=0.0351$) were noted in the T-group, of which a lower
205 proportion of 11 deaths (44%, $p=0.001$) were due to CVD made up of myocardial infarction
206 (5, 20%), stroke (2, 8%), heart failure (2, 8%), aortic aneurysm (1, 4%) and lung embolism
207 (1, 4%) (Table 2).

208

209 **Discussion**

210 Low testosterone levels in men and the resulting hypogonadism correlates with components
211 of MetS and T2D and associated cardiovascular risk [3,4,41-43]. Furthermore, liver steatosis
212 is indicative of metabolic dysfunction and may be considered the hepatic manifestation of
213 MetS common in T2D and increasing CVD risk. Hepatic lipid deposition is independently
214 associated with low testosterone levels in men [31]. TTh is widely reported to improve CVD
215 risk and components of MetS and T2D [44-48], yet the effects of testosterone on hepatic
216 steatosis are not fully understood. In this observational, prospective, registry study, we
217 demonstrate for the first time the beneficial effects of long-term TTh in hypogonadal men on
218 clinical measures of hepatic steatosis and liver function with an associated reduction in
219 cardiovascular death.

220 The majority of studies investigating hypogonadism in the context of liver function
221 have observed a link between low testosterone and NAFLD. In agreement with the present
222 study, Barbonetti et al. [31] reported an independent association between NAFLD and low
223 testosterone levels in male patients with chronic spinal cord injury. Although, the patients
224 included in the study had an increased prevalence of metabolic and lifestyle-related risk
225 factors for NAFLD (such as reduced physical activity, higher BMI and increased insulin
226 resistance and triglyceride levels), within this population it was demonstrated that patients
227 with fatty liver, assessed by sonographic criteria, had reduced total (261.6 ± 159.5 ng/dL) and
228 free testosterone levels (77.4 ± 51.7 pg/ml). The prevalence of NAFLD in men with low
229 testosterone was 85%. Although the relationship was more pronounced in men exhibiting
230 higher BMI and increased insulin resistance, triglycerides and γ -GT levels the association
231 still remained after adjusting for these confounders. Only one, albeit large-scale, population-
232 based cross-sectional study reports an association between low serum testosterone
233 concentrations and hepatic steatosis in men [29]. Two smaller studies report contrasting

234 results in men with hepatic steatosis with one identifying low serum concentrations of
235 testosterone in their patients and the other reporting no significant differences in testosterone
236 levels compared to healthy men [32,33].

237 Interventional studies investigating TTh in hypogonadal patients have primarily
238 focussed on sexual function restoration and improvements in metabolic parameters such as
239 obesity, insulin resistance, glycaemic control and lean muscle mass rather than on parameters
240 of hepatic steatosis and liver function. Improvements in metabolic parameters including
241 insulin resistance and reduced liver fat are seen following 18 weeks of TTh in obese men
242 with sleep apnea, although no effect on overall body weight was demonstrated [34].
243 Conversely, results from two randomized controlled trials in elderly men with low
244 testosterone levels reported no effect of 6 months of TTh on hepatic fat content as assessed
245 by magnetic resonance imaging (MRI) or magnetic resonance spectroscopy (MRS) [49].
246 Similarly, a randomized placebo-controlled trial of 44 hypogonadal men with T2D reported
247 unchanged hepatic fat content assessed by MRI following 6 months of TTh [50]. These
248 previous studies were relatively short-term in treatment length and had smaller cohorts
249 compared to the current study, where we report the rapid and continuous reduction in the FLI
250 following TTh. This is the first study to suggest that long-term TTh reduces lipid
251 accumulation in the liver and maintains improvements in liver function up to 12 years.

252 The effect of TTh on hepatic steatosis could also be mediated via improvements in the
253 various components of MetS and T2D, as outlined previously [39,48]. The negative
254 correlation of hepatic steatosis with MetS and T2D components such as obesity, insulin
255 resistance and dyslipidemia, observed in previous studies further supports this hypothesis
256 [6,51,52] Furthermore, the improvement in BMI and WC alongside the reduction in
257 triglycerides in the T-group suggests metabolic parameters were improved compared to the
258 C-group. TTh has been shown to reduce BMI and WC in hypogonadal men regardless of

259 testosterone formulation and administration route [53]. This collective continuous
260 improvement of metabolic parameters following TTh highlights the beneficial effects of
261 long-term TTh on body composition and triglycerides that may improve the degree of liver
262 steatosis in hypogonadal men and contribute to cardiometabolic health.

263 Compelling evidence of a direct impact of testosterone upon liver physiology has
264 been obtained from animal models. A greater degree of hepatic steatosis and inflammation
265 was observed in hepatic androgen receptor (AR) knock-out mice [54], 5 α -reductase type 1
266 knock-out mice [55] and testicular feminized (Tfm) mice which have non-functional AR and
267 low circulating levels of testosterone [36]. Similarly, severe androgen deficiency induced by
268 surgical orchiectomy in rodents is associated with the development of hepatic steatosis in
269 male mice and rats fed a high-fat diet, and TTh can significantly reduce the degree of hepatic
270 lipid deposition in these testosterone deficient animals [36,37]. Mechanistically, TTh has
271 been implicated in reducing expression of *Acaca*, *Fasn* and *Srebf1* genes involved in hepatic
272 lipid assembly and secretion [35,36,56] which may therefore contribute to ectopic lipid
273 deposition. It has also been hypothesized that androgens could modulate liver fatty acid β -
274 oxidation and *de novo* lipid synthesis via regulation of SREBF-1c, a key regulator of fatty
275 acid synthesis, and its primary target gene *SCD1* that catalyses the rate-limiting step in the
276 synthesis monounsaturated fatty acids [54]. In further support of this hypothesis,
277 dihydrotestosterone treatment in orchidectomized rats was associated with decreased lipid
278 accumulation, potentially by decreasing fatty acid and cholesterol synthesis and increasing β -
279 oxidation [57]. Nevertheless, mechanistic findings have only arisen from animal studies, and
280 their validity to the relationship between low testosterone levels and hepatic steatosis in
281 humans remains to be established from studies assessing biopsy samples in TTh clinical
282 trials.

283 Improved liver function in the T-group as evidenced by reduced γ -GT and bilirubin
284 levels, suggests improvement of metabolic parameters. γ -GT is a glutathione catalase protein,
285 the major thiol antioxidant. Elevated serum γ -GT is a marker of oxidative stress and is
286 associated with the presence of hepatic steatosis [58]. Elevated serum γ -GT correlates with
287 the presence of CVD and has been identified in atheromatous plaques where it plays a role in
288 LDL oxidation, platelet aggregation, apoptosis and influences plaque rupture [59,60]. Large
289 epidemiological studies suggest an association between elevated γ -GT activity with CVD and
290 CVD-related mortality [60-62]. Furthermore, a study investigating the effects of TTh on 225
291 hypogonadal men with MetS found that AST and ALT levels decline over 5 years [63]. The
292 current study observed moderately increased levels of ALT, exceeding those of AST at
293 baseline in both groups, the typically observed biochemical pattern in hepatic steatosis.
294 However, we report no significant change following TTh, and levels of AST and ALT were
295 within the normal range (8-48 and 7-55 U/L respectively) throughout the study for both
296 groups. Studies have reported that an elevated AST/ALT ratio is significantly associated with
297 increased risk of developing CVD in men [64,65], however another study did not observe this
298 association [66] and indicate that γ -GT is a superior liver function biomarker associated with
299 CVD [60]. Parameters of liver function and steatosis are influenced by alcohol consumption,
300 and while there was no indication of changes in alcohol intake of patients between the
301 groups, the present study did not specifically assess alcohol consumption and should
302 therefore be considered a potential confounding factor and limitation of the observations.

303 Finally, the reduction in CVD related deaths in the T-group agrees with previous
304 reports of improved cardiovascular health and reduced mortality in hypogonadal men that
305 receive TTh [67,68]. It is not certain whether the reduction in CVD related deaths was due to
306 the reduction in hepatic steatosis and improved function. Indeed, patients with NAFLD
307 possess a high risk of developing CVD with shared pathogenic aetiology, and improvement

308 in hepatic steatosis concomitantly decreases cardiovascular risk [16]. Furthermore, long term
309 TTh has been shown to have favourable effects on multiple organ systems [69]. The
310 improvement in body composition and metabolic parameters in the present study did
311 correspond with improved hepatic steatosis, feasibly reducing number of deaths attributed to
312 CVD in the T-group. Although no significant differences in age were reported between the
313 groups, death rates were not corrected for age and therefore we acknowledge that mortality
314 data should be interpreted with caution.

315 This observational study is not without inherent limitations. As the current study is
316 not a randomised controlled trial with a placebo arm, it does not allow direct comparison of
317 testosterone versus non-treatment in matched patients, limiting the scope of interpretation.
318 The primary focus of the study was to assess the long-term effectiveness and safety of TU
319 injections in comparison to a voluntarily untreated hypogonadal control group rather than
320 liver steatosis and function. At baseline, several parameters relating to liver function (FLI,
321 bilirubin, triglycerides, BMI and WC) were significantly higher in the T-group compared to
322 the C-group which may indicate a selection bias where patients undergoing treatment have
323 the most pre-existing disease. However, it is notable that men in the T-group were not
324 healthier than the controls as this would represent a bias in terms of cardiovascular events and
325 mortality. This study does present clinically meaningful data of a large patient cohort with
326 long-term follow-up-period of up to 12-years and demonstrates effects that justify further
327 investigation and prospective randomised placebo controlled studies to further delineate the
328 relationship between testosterone and hepatic steatosis. As indicated in the methods, several
329 patients discontinued TTh for a temporary period during the course of follow-up due to
330 reimbursement issues. This interruption in treatment may explain some of the trend reversals
331 demonstrated in treated patients for some parameters, particularly between 6-8 years. This is
332 also demonstrable in the total testosterone levels achieved in the treatment group, although

333 the group mean remained in the physiological range. This additionally may have reduced the
334 inter-group differences in liver function and associated parameters. Indeed, we have
335 previously shown that TTh withdrawal results in a loss of beneficial effects on several
336 cardiometabolic risk factors in hypogonadal men with treatment reinstatement restoring the
337 positive effects [70, 71]. Finally, the current study used measures of total testosterone to
338 indicate androgen status. While SHBG was measured in patients receiving TTh it was not
339 analysed in control patients due to financial reasons and this not being an randomised clinical
340 trial, therefore comparative measures of free and bioavailable testosterone were not assessed.
341 This allows SHBG as a potential confounder in both testosterone action and as a metabolic
342 factor that may influence study outcomes.

343 In conclusion, this study indicates that long term TTh does not worsen liver function,
344 but on the contrary is associated with a possible improvement and highlights the potential
345 beneficial effects of long-term TTh on liver steatosis in hypogonadal men. Consequently, as
346 hepatic steatosis is a cardiometabolic risk factor, this study also suggests this improvement
347 may reduce CVD-related mortality in these patients. Large randomized and placebo-
348 controlled trials are necessary to elucidate the impact of TTh on hepatic function and steatosis
349 in association with cardiovascular risk in hypogonadal men.

350

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358

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559 **Tables**

	Testosterone Group	Control Group
N	321	184
Mean age (years)	59 ± 9.5	66.1 ± 7.6
Follow-up (years)	8.3 ± 3.5	5.5 ± 1.6
Testosterone (nmol/L)	7.7 ± 2.1	9.2 ± 2.4
Waist circumference (cm)	107 ± 10	100 ± 9
Weight (kg)	99 ± 13	91 ± 11
BMI (kg/m²)	31.5 ± 4.3	29.2 ± 3.2
FLI	83.6 ± 12.1	68.7 ± 19.4

560

561 **Table 1.** Baseline characteristics of Testosterone treatment group (T-group) and control group (C-
562 group).

Adverse Events		
	Testosterone Group	Control Group
N	321	184
Deaths (%)	25 (7.8%)	28 (15.2%)
Deaths due to CVD (%)	11 (44%) *	28 (100%)
<i>Myocardial Infarction</i>	5 (20%)	13 (48%)
<i>Stroke</i>	2 (8%)	7 (27%)
<i>heart failure</i>	2 (8%)	3 (11%)
<i>aortic aneurysm</i>	1 (4%)	2 (7%)
<i>lung embolism</i>	1 (4%)	2 (7%)

563

564 **Table 2.** Adverse events observed in Testosterone treatment group (T-group) and control group (C-

565 group). *P<0.001.

566 **Figure Legends**

567 **Figure 1.** Total testosterone (nmol/l) in 321 hypogonadal men on long-term treatment with
568 testosterone undecanoate and 184 untreated hypogonadal controls. * $p < 0.0001$ between groups.

569 **Figure 2.** The Fatty Liver Index (FLI) in 321 hypogonadal men on long-term treatment with
570 testosterone undecanoate and 184 untreated hypogonadal controls. * $p < 0.0001$ between groups.

571 **Figure 3.** γ -GT (a) and bilirubin levels (b) 321 hypogonadal men on long-term treatment with
572 testosterone undecanoate and 184 untreated hypogonadal controls. Significance indicated between
573 groups. * $p < 0.0001$.

574 **Figure 4.** AST (a) and ALT (b) levels in 321 hypogonadal men on long-term treatment with
575 testosterone undecanoate and 184 untreated hypogonadal controls. * $p < 0.0001$ between groups.

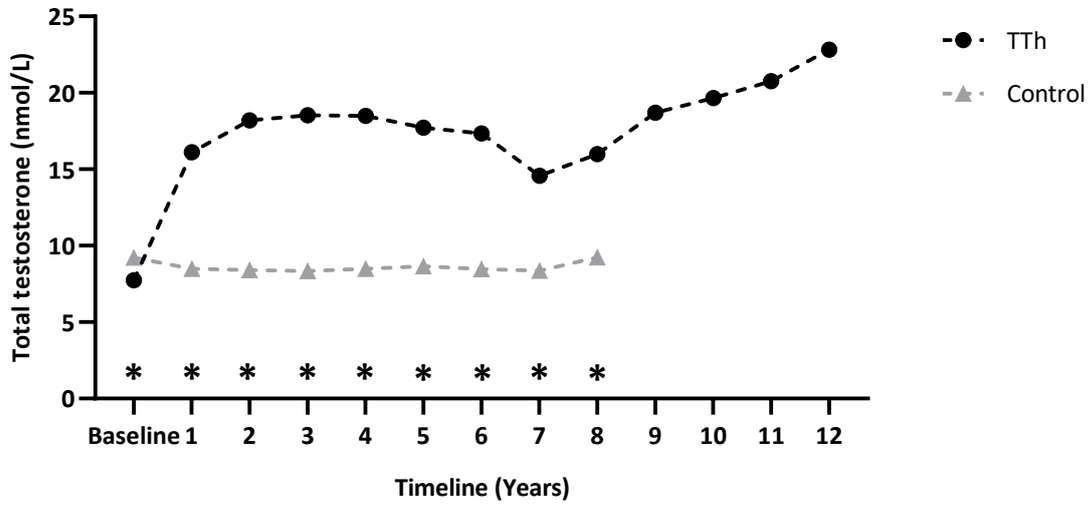
576 **Figure 5.** Triglyceride levels in 321 hypogonadal men on long-term treatment with testosterone
577 undecanoate and 184 untreated hypogonadal controls. * $p < 0.0001$ between groups.

578 **Figure 6.** BMI (a) and waist circumference (b) of 321 hypogonadal men on long-term treatment with
579 testosterone undecanoate and 184 untreated hypogonadal controls. * $p < 0.0001$ between groups.

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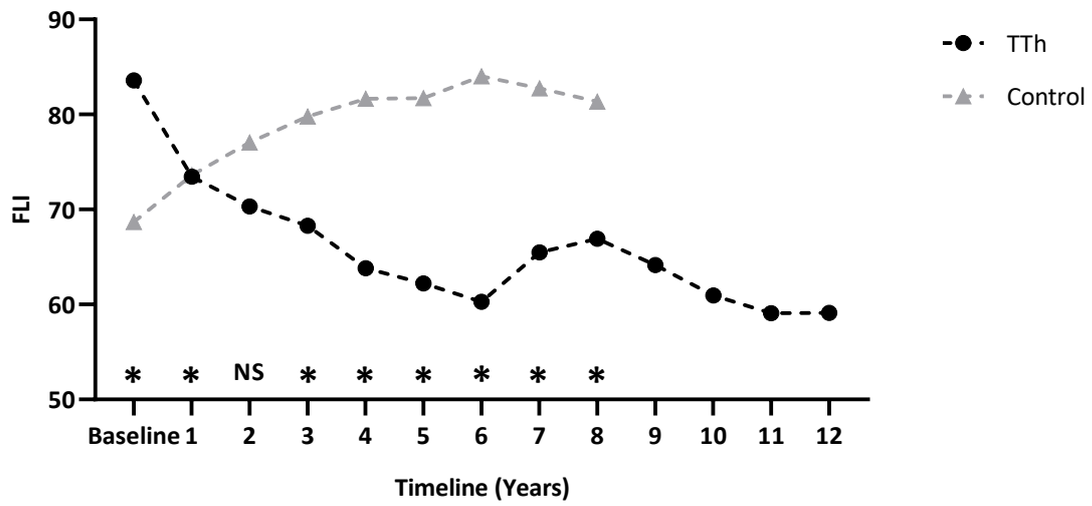
Total Testosterone in 321 Hypogonadal Men on Long-Term Treatment with Testosterone Undecanoate and 184 Untreated Hypogonadal Controls



582

583 **Figure 1.** Total testosterone (nmol/l) in 321 hypogonadal men on long-term treatment with
584 testosterone undecanoate and 184 untreated hypogonadal controls. *p<0.0001 between groups.

The Fatty Liver Index in 321 Hypogonadal Men on Long-Term Treatment with Testosterone Undecanoate and 184 Untreated Hypogonadal Controls

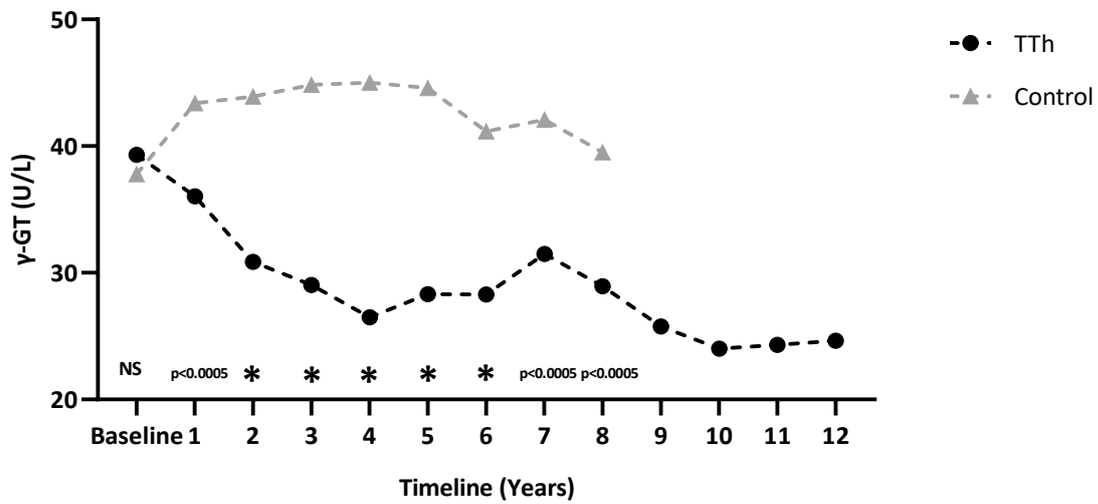


585

586 **Figure 2.** The FLI in 321 hypogonadal men on long-term treatment with testosterone undecanoate and

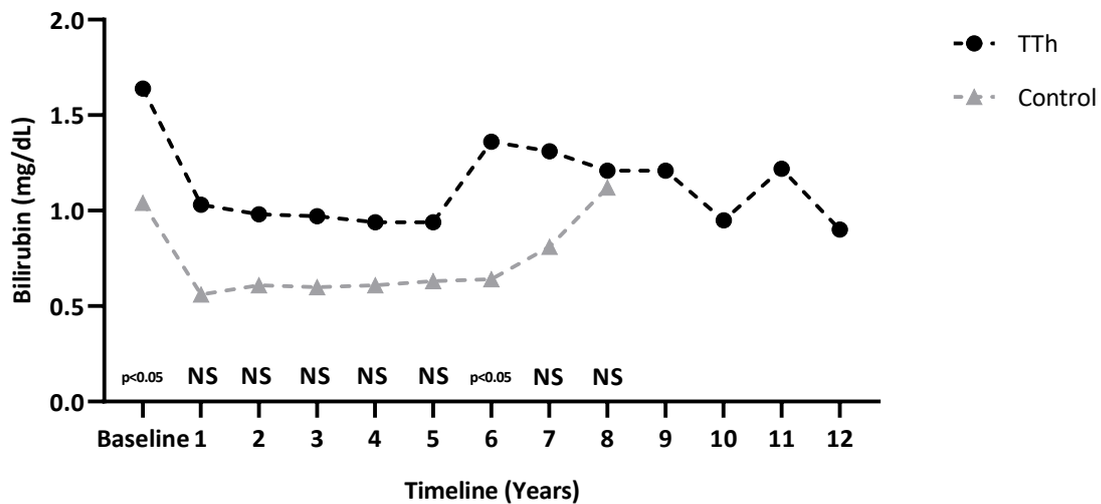
587 184 untreated hypogonadal controls. * $p < 0.0001$ between groups.

A) γ -GT levels in 321 Hypogonadal Men on Long-Term Treatment with Testosterone Undecanoate and 184 Untreated Hypogonadal Controls



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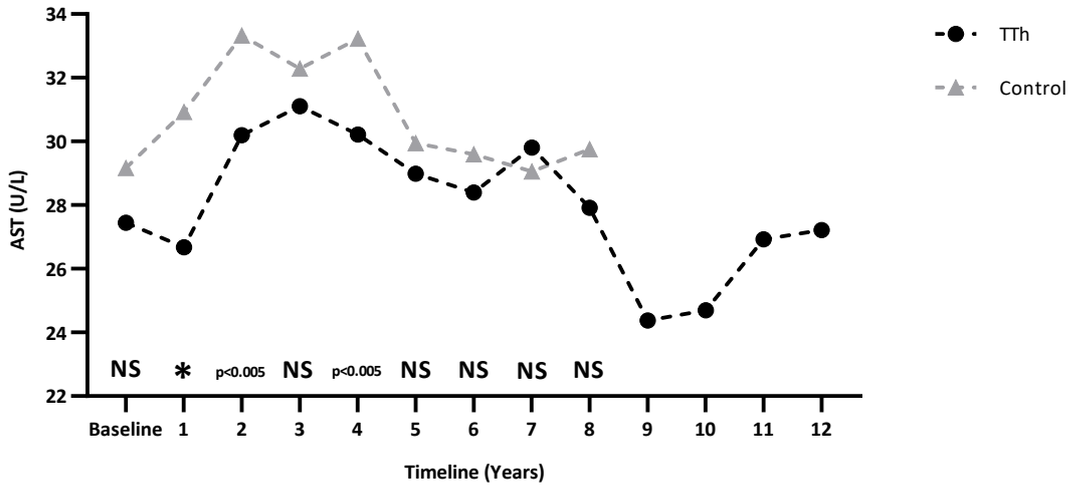
B) Bilirubin levels in 321 Hypogonadal Men on Long-Term Treatment with Testosterone Undecanoate and 184 Untreated Hypogonadal Controls



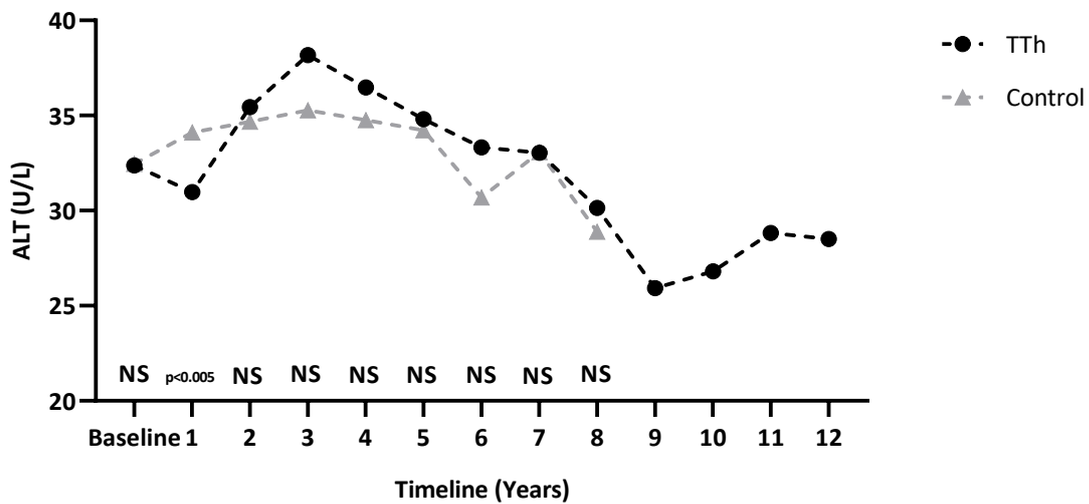
589

590 **Figure 3.** γ -GT (a) and bilirubin levels (b) 321 hypogonadal men on long-term treatment with
 591 testosterone undecanoate and 184 untreated hypogonadal controls. Significance indicated between
 592 groups. *p<0.0001.

A) AST levels in 321 Hypogonadal Men on Long-Term Treatment with Testosterone Undecanoate and 184 Untreated Hypogonadal Controls



B) ALT levels in 321 Hypogonadal Men on Long-Term Treatment with Testosterone Undecanoate and 184 Untreated Hypogonadal Controls

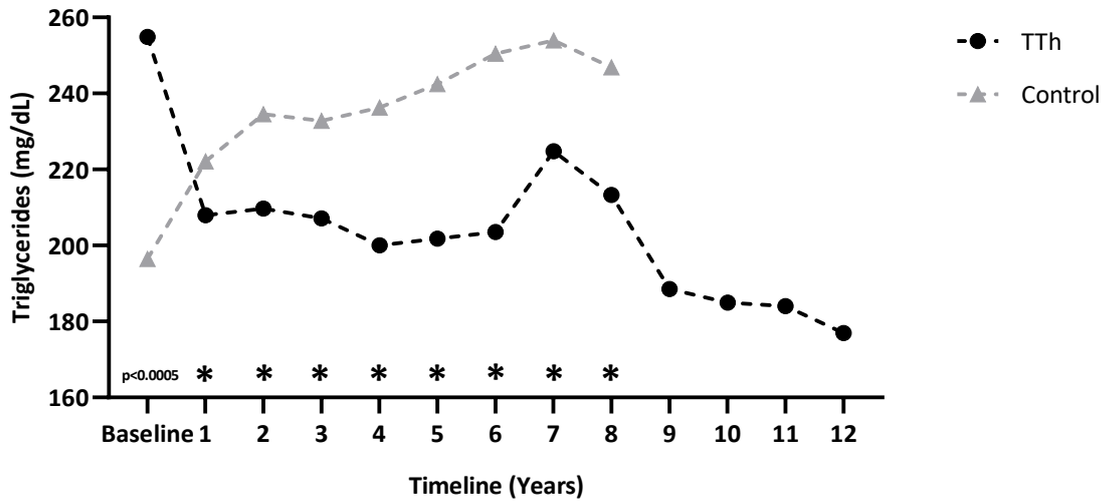


593

594 **Figure 4.** AST (a) and ALT (b) levels in 321 hypogonadal men on long-term treatment with
 595 testosterone undecanoate and 184 untreated hypogonadal controls. *p<0.0001 between groups.

596

Triglyceride levels in 321 Hypogonadal Men on Long-Term Treatment with Testosterone Undecanoate and 184 Untreated Hypogonadal Controls



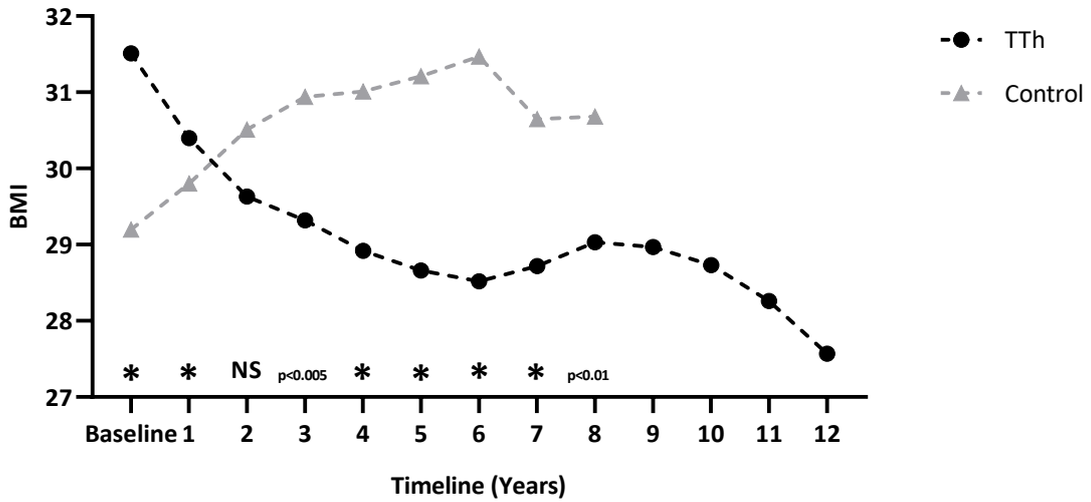
597

598 **Figure 5.** Triglyceride levels in 321 hypogonadal men on long-term treatment with testosterone

599 undecanoate and 184 untreated hypogonadal controls. * $p < 0.0001$ between groups.

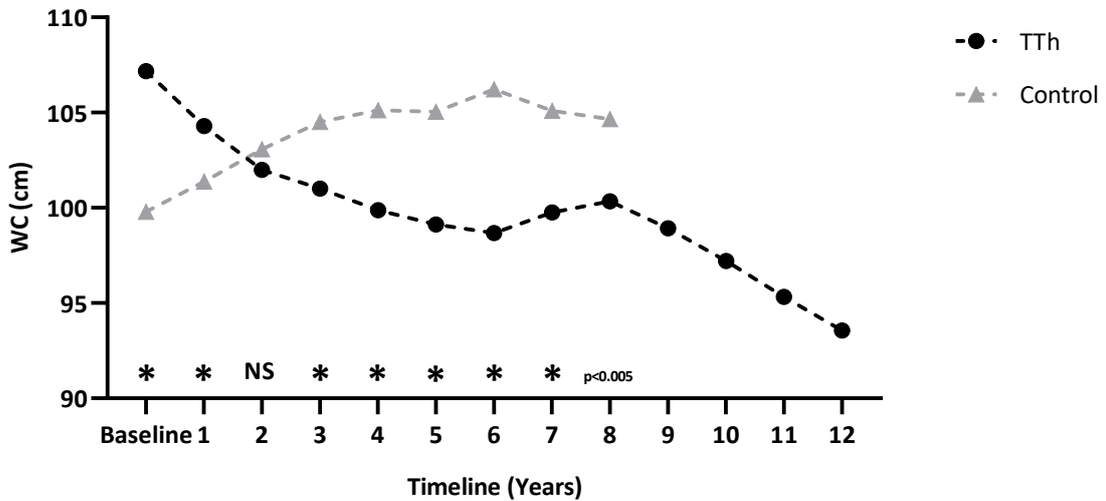
A)

BMI of 321 Hypogonadal Men on Long-Term Treatment with Testosterone Undecanoate and 184 Untreated Hypogonadal Controls



B)

Waist Circumference of 321 Hypogonadal Men on Long-Term Treatment with Testosterone Undecanoate and 184 Untreated Hypogonadal Controls



600

601

602 **Figure 6.** BMI (a) and waist circumference (b) of 321 hypogonadal men on long-term treatment with

603 testosterone undecanoate and 184 untreated hypogonadal controls. *p<0.0001 between groups.

604

605