Accepted Manuscript

Relation of Testosterone Normalization to Mortality and Myocardial Infarction in Men with Prior Myocardial Infarction

Olurinde A. Oni MBBS, MPH , Seyed Hamed Hosseini Dehkordi MD , Mohammad-Ali Jazayeri MD , Rishi Sharma MD , Mukut Sharma PhD , Reza Masoomi MD , Ram Sharma PhD , Kamal Gupta MD , Rajat S. Barua MD, PhD

 PII:
 S0002-9149(19)30816-1

 DOI:
 https://doi.org/10.1016/j.amjcard.2019.07.019

 Reference:
 AJC 24071



To appear in: The American Journal of Cardiology

Received date:20 March 2019Revised date:3 July 2019

Please cite this article as: Olurinde A. Oni MBBS, MPH, Seyed Hamed Hosseini Dehkordi MD, Mohammad-Ali Jazayeri MD, Rishi Sharma MD, Mukut Sharma PhD, Reza Masoomi MD, Ram Sharma PhD, Kamal Gupta MD, Rajat S. Barua MD, PhD, Relation of Testosterone Normalization to Mortality and Myocardial Infarction in Men with Prior Myocardial Infarction, *The American Journal of Cardiology* (2019), doi: https://doi.org/10.1016/j.amjcard.2019.07.019

This is a PDF file of an unedited manuscript that has been accepted for publication. As a service to our customers we are providing this early version of the manuscript. The manuscript will undergo copyediting, typesetting, and review of the resulting proof before it is published in its final form. Please note that during the production process errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.

Relation of Testosterone Normalization to Mortality and Myocardial Infarction in Men with Prior Myocardial Infarction

Olurinde A. Oni, MBBS, MPH, † Seyed Hamed Hosseini Dehkordi, MD, ‡ Mohammad-Ali Jazayeri, MD, ‡ Rishi Sharma, MD, † Mukut Sharma, PhD, † Reza Masoomi, MD, ‡ Ram Sharma, PhD, † Kamal Gupta, MD, ‡ Rajat S. Barua MD, PhD *†‡

Running title: Testosterone Normalization in Patients with Prior Myocardial Infarction.

† Division of Cardiovascular Research, Kansas City VA Medical Center, Kansas City, MO

‡ Department of Cardiovascular Medicine, University of Kansas Medical Center,

Kansas City, KS

* Division of Cardiovascular Medicine, Kansas City VA Medical Center, Kansas City, MO Correspondence to Rajat S. Barua, MD, PhD. Division of Cardiovascular Medicine, Kansas City Veterans Affairs Medical Center, 4801 E. Linwood Blvd, Kansas City, MO 64128.

Email: rajat.barua@va.gov

Tel: 816-922-2441

Fax: 816-922-4745

ABSTRACT

The effect of normalization of serum testosterone levels with testosterone replacement therapy (TRT) in patients with a prior history of myocardial infarction (MI) is unknown. The objective of this study was to determine the incidence of recurrent MI and all-cause mortality in subjects with a history of MI and low total testosterone (TT) with and without TRT. We retrospectively examined 1470 males with documented low TT levels and prior MI, categorized into Gp1: TRT with normalization of TT levels (N=755) Gp2: TRT without normalization of TT levels (N=542), and Gp3: no TRT (N=173). The association of TRT with all-cause mortality and recurrent MI was compared using propensity score-weighted Cox proportional hazard models. All-cause mortality was lower in Gp1 vs. Gp2 (HR 0.76, CI 0.64–0.90, P=0.002), and Gp1 vs Gp3 (HR 0.76, CI 0.60–0.98, P=0.031). There was no significant difference in the risk of death between Gp2 vs. Gp3 (HR: 0.97, CI 0.76–1.24, P=0.81). Adjusted regression analyses showed no significant differences in the risk of recurrent MI between groups (Gp1 vs. Gp3, HR 0.79, CI 0.12-5.27, P=0.8; Gp1 vs. Gp2 HR 1.10, CI 0.25-4.77, P=0.90; Gp2 vs. Gp3 HR 0.58, CI 0.08-4.06, P=0.58). In conclusion, in a large observational cohort of male veterans with prior MI, normalization of TT levels with TRT was associated with decreased all-cause mortality compared to those with non-normalized TT levels and the untreated group. Furthermore, in this high-risk population, TRT was not associated with an increased risk of recurrent MI.

Keywords: Testosterone, Testosterone Replacement Therapy, Myocardial Infarction

INTRODUCTION

Approximately 2.4 million men suffer from hypogonadism in the United States (US) (1). In the last decade, there has been a significant increase in the number of prescriptions for testosterone replacement therapy (TRT) (2), but the role of TRT and its effects on cardiovascular disease (CVD) remain controversial. While previous studies evaluating TRT in men did not report adverse cardiovascular outcomes (3-7), more recent studies have contradicted those findings (8-10). Utilizing a large database of US military veterans, we previously found that normalization of TT levels after TRT was associated with a significant reduction in all-cause mortality, myocardial infarction (MI), and stroke (11). However, in that study, patients with a history of MI, a risk factor for recurrent adverse CV events in up to 40% of acute coronary syndrome (ACS) patients (12), were excluded. The impact of TRT on this high-risk population has not been reported previously. In the current study, we examined the effects of TRT in patients with a history of MI and documented low total testosterone (TT) levels.

METHODS

We conducted a retrospective cohort study of male veterans who received medical care through the Veterans Health Administration (VHA) from December 1999 to May 2014. The VHA provides care to veterans at over 1,400 establishments across the United States, and each veteran is assigned a unique identifier in the Corporate Data Warehouse (CDW) database. Data were retrieved from the VHA CDW through the Veterans Administration Informatics and Computing Infrastructure (VINCI) (13) The quality of data from these sources is well documented, and the data have been widely used by investigators for retrospective, longitudinal

studies (14). Data in the CDW are stored in a relational fashion and are definable by attributes such as ICD-9 codes, CPT codes, and other clinical measures, from both inpatient and outpatient encounters. VINCI provides a robust computing environment for researchers to analyze CDW data without moving sensitive health data off secure VA servers. The Institutional Review Board of the Veterans Affairs Medical Center in Kansas City, Missouri approved this study.

The current study examined the risk of recurrent MI among those who had a history of MI before their first low testosterone lab date. There was wide variability in the reporting units and reference ranges for testosterone test results over the extended follow-up period. The lack of standardization of TT levels using stoichiometric measurements has also been documented (15,16). To eliminate the potential for disparities due to the use of multiple assays, we classified a test result as low or normal based on the respective normal laboratory reference range (NLRR) reported with the test result. TT levels less than the lower limit of the normal laboratory reference range (NLRR) for a particular assay were categorized as low TT. This approach obviated the need for a single discrete cut-off value, since facilities used different TT assays, each with their own reference ranges and reporting units (17,18), and enabled us to account for changes in the assays used over time within the same facility. We determined the use of TRT from both inpatient and outpatient prescription records. Patients were considered treated if they received any form of TRT (injection, gel, or patch); otherwise, they were considered as untreated. Treated patients were categorized as normalized or non-normalized depending on whether their TT levels improved to within the NLRR on treatment or remained persistently low.

The primary outcome measures were (1) recurrent MI (ICD-9 410.x0 and 410.x1) and (2) all-cause mortality. Mortality data were obtained from a combination of CDW files and Veteran's Health Administration Vital Status Files, the latter of which contains demographics,

including dates of death obtained from multiple VA and non-VA data sources such as the Beneficiary Identification Records Locator Subsystem death file, the VA Medicare Vital Status File, and the Social Security Administration Death Master File (19,20). Confounding variables that were adjusted for during the analysis were: patient demographics, comorbidities, including hypertension, congestive heart failure, diabetes mellitus, chronic obstructive pulmonary disease, obstructive sleep apnea, low density lipoprotein levels, and peripheral vascular disease, and the use of medications, including aspirin, beta-blockers, and statins. Comorbidities were defined using their respective ICD-9 codes.

We included patients whose first tested TT level was lower than the NLRR who also had a history of MI prior to the first documented low TT. We excluded (1) female patients, (2) those who received TRT before the first available low TT level, and (3) male patients without a history of MI. We used stabilized inverse probability of treatment weights (SIPTW) propensity score matching to ensure a robust analysis. SIPTW propensity score matching significantly corrects for the instability in estimated treatment weights that potentially results from the use of regular IPTW for individuals with a low probability of treatment (21). This method allowed us to maximize the number of patients included in the study after matching, compared to traditional propensity score matching. We computed each subject's propensity score for receiving TRT and adjusted for the covariates in a logistic regression analysis. We computed the incidence of MI and all-cause mortality in each subgroup. Chi-squared test and the Student's t-test were used to compare the normally distributed baseline characteristics of patients. Non-parametric tests were used for non-normally distributed variables. Univariate and multivariate Cox proportional hazard regression analyses were conducted to assess the differences between groups. We reported continuous variables as sample means and their standard deviations (SD), while categorical variables were reported as percentages. SAS Enterprise Guide 7.1 supported on SAS 9.4 (SAS Institute, Inc., Cary, NC, USA) was used for statistical analyses, with TRT as a time-varying exposure variable. The study hypotheses were studied with two-tailed testing. A P value <0.05 was considered statistically significant.

RESULTS

We identified 1560 patients with low TT and a history of MI events prior to their first low TT lab (Figure 1). Of these, we excluded 90 patients who did not have complete data with respect to matching variables. The remaining patients were categorized as follows: Gp1, TRT with normalization of TT levels (N=755); Gp2, TRT without normalization of TT levels (N=542); and Gp3, those who did not receive TRT at any time (N=173).

Table 1 displays baseline variables for the unmatched and SIPTW-matched cohorts. Median ages at enrollment were 64.3 years, 64.5 years, and 66.5 years for Gp1, Gp2, and Gp3, respectively. The cohorts were predominantly obese with mean body mass index (BMI) at enrollment being 31.6 kg/m² (SD 6.1), 32.3 kg/m² (SD 6.1), and 32.3 kg/m² (SD 6.1) in Gp1, Gp2, and Gp3, respectively. Mean follow-up time was longer in the normalized TRT group (Gp1, 4.0 years, SD 3.4) than the non-normalized TRT (Gp2, 3.4 years, SD 3.2) and the untreated (Gp3, 3.3 years, SD 3.1) groups. We adjusted for the differences in age, BMI, and other baseline comorbidities by means of SIPTW matching. Enhancing the yield and validity of Cox proportional hazard regression analyses, all groups were well matched (*P*>0.05) with regard to these covariates, as presented in Table 1.

Mortality rates were among groups were as follows: normalized TRT (Gp1), 101 per 1000 person-years; non-normalized TRT (Gp2), 137 per 1000 person-years; and untreated (Gp3),

163 per 1000 person-years. The risk of death from any cause was significantly lower in Gp1 compared to Gp2 (HR 0.76, 95% CI 0.64–0.90, P=0.002), and in Gp1 compared to Gp3 (HR 0.76, 95% CI 0.60–0.98, P=0.031). There was no significant difference in the risk of all-cause mortality between Gp2 and Gp3 (HR 0.97, 95% CI 0.76–1.24, P=0.81). Kaplan–Meier (KM) survival curves were constructed (Figure 2), which demonstrated that normalization of TRT (Gp1) was associated with significantly greater survival from all-cause death (log-rank P<0.05), compared with non-normalized TRT (Gp2) or untreated (Gp3) groups.

The incidence of recurrent MI in each group was as follows: normalized TRT (Gp1), 1391 per 100,000 person-years; non-normalized TRT (Gp2), 1628 per 100,000 person-years; untreated (Gp3), 1402 per 100,000 person-years. Table 2 presents the results of the Cox proportional hazard regression analysis. There were no significant differences in the risks of recurrent MI between groups: Gp1 vs. Gp3 (HR 0.79, 95% CI 0.12–5.27, P=0.81); Gp1 vs. Gp2 (HR 1.10, 95% CI 0.25–4.77, P=0.90); Gp2 vs. Gp3 (HR 0.58, 95% CI 0.08–4.06, P=0.58). TRT did not appear to be significantly associated with an increased risk of recurrent MI. The KM curves (Figure 3) additionally showed no difference in MI-free survival among the groups (logrank P>0.05).

DISCUSSION

In this study of hypogonadal patients with a history of MI there are several key findings. First, in this population, normalization of TT levels after TRT was associated with a significant decrease in all-cause mortality, compared to those who received TRT without normalization of TT levels and those not treated. Second, there was no statistically significant difference in allcause mortality between the non-normalized TRT group (Gp2) and the untreated group (Gp3).

Finally, there was no significant increase in the incidence of recurrent MI in subjects receiving TRT with subsequent complete or incomplete TT normalization.

Clinical trials examining the effects of TRT historically have been small and underpowered to provide conclusive evidence on the risk of adverse CV events (22). Early trials found that TRT reduced symptoms in patients with chronic stable angina (23, 24). In one study, there were no significant difference in carotid intimal medial thickness or coronary artery calcium scores in over 3 years of follow-up, between patients receiving TRT or placebo (25). In the largest observational study to date, our group previously found that normalization of TT levels after TRT was associated with a significant reduction in all-cause mortality, MI, and stroke in patients without a history of MI or stroke (11). A recent meta-analysis also found no significant association of TRT with adverse CV events (26).

The first major trial describing adverse CV outcomes following TRT was the Testosterone in Older Men (TOM) trial (8), a randomized controlled trial designed to determine the effects of TRT on lower extremity strength and physical function in older men with limited mobility over 6 months. Although the study was not powered to assess CV events, it reported a higher incidence of CV-related events in the treatment arm. These events were diverse and of variable clinical importance (e.g. peripheral edema, ectopy on electrocardiography). In an observational study by Vigen et al., examining data from VA cardiac catheterization laboratories, TRT was significantly associated with a higher number of composite adverse events (MI, stroke, and death; HR 1.29, 95% CI 1.05–1.58, *P*=0.02). In this study, 20% of the population had a history of MI (9), and inclusion of men undergoing coronary angiography might itself have introduced selection bias, resulting in a higher CV risk population (11,26). Recently, Budoff et al. reported that TRT gel administration for 1 year was associated with a significant increase in

coronary artery non-calcified plaque volume measured by coronary computed tomographic angiography (27), but men with a history of MI or stroke within 3 months were excluded.

The present study examined the effect of TRT in patients with a higher CV risk than many prior studies, as all subjects had a history of MI. Despite the increased risk, the results were consistent with our previous findings in a male veteran population, excluding those with a history of stroke or MI, in which normalization of TT levels following TRT was also associated with decreased all-cause mortality compared to those with incomplete normalization (11). In the current study, we found TRT with incomplete or complete TT normalization was not associated with an increase or decrease in recurrent MI. This finding differs from our previous results and is likely due to a lower CV risk in that population (no history of MI and stroke, and only 6% had coronary artery disease [CAD]), as compared to the current study (11).

The lack of reduction in MI after TRT in patients with prior MI suggests normalization of TT alone may not be sufficient to reduce MI events in this high-risk population. Given the multifactorial nature of atherosclerotic CAD and complex pathophysiology of ACS, a comprehensive risk factor reduction strategy is needed to reduce the risk of recurrent MI. Unlike the study of Vigen et al., in which 20% of the population had a history of MI (9), our study examining an entirely post-MI cohort did not show an association between TRT and adverse CV outcomes, Based on the mean TT levels reported by Vigen et al., a number of patients likely did not achieve normalization of TT levels following TRT and may therefore have been at increased risk compared to a cohort with normalized TT levels after TRT. While further study is needed to better define the appropriate role for TRT in patients with a history of CVD, in carefully selected hypogonadal patients there may be a role for TRT. Despite the controversies surrounding TRT (8,9), current guidelines and scientific consensus statements indicate the totality of evidence

favors a lack of adverse CV events due to TT normalization with TRT (28-31). Our study fortifies this position with evidence regarding the effects of TRT in a high-risk population.

There were a number of limitations in the current study. Our study results are specific to the population studied and may not be generalized. Given its observational nature, residual confounding cannot be entirely ruled out. Inclusion criteria and outcomes were determined using ICD-9 and CPT codes. CDW does not capture the reasons for initiating/withholding TRT, so we cannot rule out the possibility that TRT was preferentially offered to healthier subjects. In our study, data regarding clinical response to TRT were also not available. Similarly, we were unable to study quality of care and/or poor compliance as reason(s) for persistent low testosterone levels observed in some individuals.

In conclusion results from our study illustrate that in a high CV risk population with a history of MI, TRT with normalization of TT levels is associated with decreased all-cause mortality. Exposure to TRT, resulting in normalized or non-normalized TT levels, had no significant effect on recurrent MI compared to untreated subjects, suggesting the safety of TRT this population. These findings and our understanding of the cardiovascular effects of TRT will be further enhanced with adequately powered, prospective clinical trials featuring long-term follow-up.

ACKNOWLEDGMENT

None.

CONFLICTS OF INTEREST

None of the authors have a conflict of interest regarding the contents of the paper. Views expressed in this article are those of authors and do not necessarily reflect the position and policy of the Department of Veterans Affairs or the United States Government

REFERENCES

- Araujo AB, O'Donnell AB, Brambilla DJ, Simpson WB, Longcope C, Matsumoto AM, McKinlay JB. Prevalence and incidence of androgen deficiency in middle-aged and older men: estimates from the Massachusetts Male Aging Study. *J Clin Endocrinol Metab* 2004;89:5920–5926.
- 2- Baillargeon J, Urban RJ, Ottenbacher KJ, Pierson KS, Goodwin JS. Trends in androgen prescribing in the United States, 2001 to 2011. JAMA Intern Med 2013;173(15):1465-6.
- 3- Haddad RM, Kennedy CC, Caples SM, Tracz MJ, Boloña ER, Sideras K, Uraga MV, Erwin PJ, Montori VM. Testosterone and cardiovascular risk in men: a systematic review and meta-analysis of randomized placebo-controlled trials. *Mayo Clin Proc* 2007;82(1):29-39.
- Shores MM, Smith NL, Forsberg CW, Anawalt BD, Matsumoto AM. Testosterone treatment and mortality in men with low testosterone levels. *J Clin Endocrinol Metab* 2012;97:2050–2058.
- 5- Maggi M, Wu FC, Jones TH, Jackson G, Behre HM, Hackett G, Martin-Morales A, Balercia G, Dobs AS, Arver ST, Maggio M, Cunningham GR, Isidori AM, Quinton R, Wheaton OA, Siami FS, Rosen RC; RHYME Investigators. Testosterone treatment is not associated with increased risk of adverse cardiovascular events: results from the Registry of Hypogonadism in Men (RHYME). *Int J Clin Pract* 2016;70(10):843-852.
- 6- Cheetham TC, An J, Jacobsen SJ, Niu F, Sidney S, Quesenberry CP, VanDenEeden SK.
 Association of Testosterone Replacement With Cardiovascular Outcomes Among Men
 With Androgen Deficiency. JAMA Intern Med 2017;177(4):491-499.

- Anderson JL, May HT, Lappé DL, Bair T, Le V, Carlquist JF, Muhlestein JB. Impact of Testosterone Replacement Therapy on Myocardial Infarction, Stroke, and Death in Men With Low Testosterone Concentrations in an Integrated Health Care System. *Am J Cardiol* 2016;117(5):794-799.
- Basaria S, Coviello AD, Travison TG, Storer TW, Farwell WR, Jette AM, Eder R, Tennstedt S, Ulloor J, Zhang A, Choong K, Lakshman KM, Mazer NA, Miciek R, Krasnoff J, Elmi A, Knapp PE, Brooks B, Appleman E, Aggarwal S, Bhasin G, Hede-Brierley L, Bhatia A, Collins L, LeBrasseur N, Fiore LD, Bhasin S. Adverse events associated with testosterone administration. *N Engl J Med* 2010;363(2):109-122.
- 9- Vigen, R., O'Donnell, C. I., Barón, A. Grunwald GK, Maddox TM, Bradley SM,
 Barqawi A, Woning G, Wierman ME, Plomondon ME, Rumsfeld JS, Ho PM.
 Association of testosterone therapy with mortality, myocardial infarction, and stroke in men with low testosterone levels. *JAMA* 310, 1829-1836.
- 10- Finkle WD, Greenland S, Ridgeway GK, Adams JL, Frasco MA, Cook MB, Fraumeni JF Jr, Hoover RN. Increased risk of non-fatal myocardial infarction following testosterone therapy prescription in men. *PLoS ONE* 2014;9(1):e85805.
- Sharma R, Oni OA, Gupta K, Chen G, Sharma M, Dawn B, Sharma R, Parashara D, Savin VJ, Ambrose JA, Barua RS. Normalization of testosterone level is associated with reduced incidence of myocardial infarction and mortality in men. *Eur Heart J* 2015;36(40):2706-2715.
- 12- Motivala AA, Tamhane U, Ramanath VS, Saab F, Montgomery DG, Fang J, Kline-Rogers E, May N, Ng G, Froehlich J, Gurm H, Eagle KA. A prior myocardial infarction:

how does it affect management and outcomes in recurrent acute coronary syndromes? *Clin Cardiol* 2008;31(12):590-596.

- 13- VA Informatics and Computing Infrastructure (VINCI). Accessible at:
 http://www.hsrd.research.va.gov/for_researchers/vinci/default.cfm. Last accessed May 26, 2019.
- Byrd JB, Vigen R, Plomondon ME, Rumsfeld JS, Box TL, Fihn SD, Maddox TM. Data quality of an electronic health record tool to support VA cardiac catheterization laboratory quality improvement: the VA Clinical Assessment, Reporting, and Tracking System for Cath Labs (CART) program. *Am Heart J* 2013;165(3):434-440.
- 15- Wang C, Catlin DH, Demers LM, Starcevic B, Swerdloff RS. Measurement of total serum testosterone in adult men: comparison of current laboratory methods versus liquid chromatography-tandem mass spectrometry. *J Clin Endocrinol Metab* 2004;89(2):534-543.
- Vesper HW, Botelho JC. Standardization of testosterone measurements in humans. J
 Steroid *Biochem Mol Biol* 2010;121(3-5):513-519.
- 17- Lazarou S, Reyes-vallejo L, Morgentaler A. Wide variability in laboratory reference values for serum testosterone. J Sex Med 2006;3(6):1085-1089.
- 18- Rosner W, Auchus RJ, Azziz R, Sluss PM, Raff H. Position statement: Utility, limitations, and pitfalls in measuring testosterone: an Endocrine Society position statement. *J Clin Endocrinol Metab* 2007;92(2):405-413.
- 19- The VA Medicare Vital Status File. Accessible at *http://www.virec.research.va.gov* Last accessed May 26, 2019

- 20- The Social Security Administration (SSA) Death Master File. Accessible at: https://www.ssa.gov/dataexchange/request_dmf.html. Last accessed May 26, 2019.
- Austin PC. An Introduction to Propensity Score Methods for Reducing the Effects of Confounding in Observational Studies. *Multivariate Behav Res* 2011;46(3):399-424.
 Jones TH, Kelly DM. Randomized controlled trials - mechanistic studies of testosterone and the cardiovascular system. *Asian J Androl* 2018;20(2):120-130.
- 22- English KM, Steeds RP, Jones TH, Diver MJ, Channer KS. Low-dose transdermal testosterone therapy improves angina threshold in men with chronic stable angina: A randomized, double-blind, placebo-controlled study. *Circulation* 2000;102(16):1906-1911.
- 23- Malkin CJ, Pugh PJ, Morris PD, Kerry KE, Jones RD, Jones TH, Channer KS. Testosterone replacement in hypogonadal men with angina improves ischaemic threshold and quality of life. *Heart* 2004;90(8):871-876.
- Basaria S, Harman SM, Travison TG, Hodis H, Tsitouras P, Budoff M, Pencina KM, Vita J, Dzekov C, Mazer NA, Coviello AD, Knapp PE, Hally K, Pinjic E, Yan M, Storer TW, Bhasin S. Effects of Testosterone Administration for 3 Years on Subclinical Atherosclerosis Progression in Older Men With Low or Low-Normal Testosterone Levels: A Randomized Clinical Trial. *JAMA* 2015;314(6):570-581.
- Alexander GC, Iyer G, Lucas E, Lin D, Singh S. Cardiovascular Risks of Exogenous
 Testosterone Use Among Men: A Systematic Review and Meta-Analysis. *Am J Med* 2017;130(3):293-305.
- 26- Budoff MJ, Ellenberg SS, Lewis CE, Mohler ER 3rd, Wenger NK, Bhasin S, Barrett-Connor E, Swerdloff RS, Stephens-Shields A, Cauley JA, Crandall JP, Cunningham GR,

Ensrud KE, Gill TM, Matsumoto AM, Molitch ME, Nakanishi R, Nezarat N, Matsumoto S, Hou X, Basaria S, Diem SJ, Wang C, Cifelli D, Snyder PJ. Testosterone Treatment and Coronary Artery Plaque Volume in Older Men With Low Testosterone. *JAMA* 2017;317(7):708-716.

- Bhasin S, Brito JP, Cunningham GR, Hayes FJ, Hodis HN, Matsumoto AM, Snyder PJ,
 Swerdloff RS, Wu FC, Yialamas MA. Testosterone Therapy in Men With
 Hypogonadism: An Endocrine Society Clinical Practice Guideline. *J Clin Endocrinol Metab* 2018;103(5):1715-1744.
- 28- Khera M, Adaikan G, Buvat J, Carrier S, El-Meliegy A, Hatzimouratidis K, McCullough A, Morgentaler A, Torres LO, Salonia A. Diagnosis and Treatment of Testosterone Deficiency: Recommendations From the Fourth International Consultation for Sexual Medicine (ICSM 2015). J Sex Med 2016;13(12):1787-1804.
- 29- Channer KS. Endogenous testosterone levels and cardiovascular disease in healthy men. *Heart* 2011 97:867–869
- Mulhall JP, Trost LW, Brannigan RE, Kurtz EG, Redmon JB, Chiles KA, Lightner DJ,
 Miner MM, Murad MH, Nelson CJ, Platz EA, Ramanathan LV, Lewis RW. Evaluation
 and Management of Testosterone Deficiency: AUA Guideline. *J Urol.* 2018;200(2):423-432.
- 31- Kapoor D, Goodwin E, Channer KS, Jones TH. Testosterone replacement therapy improves insulin resistance, glycaemic control, visceral adiposity and hypercholesterolaemia in hypogonadal men with type 2 diabetes. *Eur J Endocrinol* 2006;154(6):899-906.

Figure 1. Methodology and patient selection process.



Figure 2. Kaplan-Meier curve comparing the all-cause-mortality in the non-normalized treated vs. normalized treated group (a), untreated vs. normalized treated groups (b), and untreated vs. non-normalized treated groups (c).



Figure 3. Kaplan-Meier curve comparing recurrent myocardial infarction in the non-normalized treated vs. normalized treated group (a), untreated vs. normalized treated groups (b), and untreated vs. non-normalized treated groups (c).

Figure 3.





Figure: 3b











 Table 1. Baseline variables for the unmatched and SIPTW-matched cohorts of all patients in the study.

		Table	1		λ	
Variable	UNMATCHED COHORT (Stabilized- IPTW			SITY MATCH OHORT lized- IPTW)	IED	
	Norm	alized-treated	Vs. Unt	reated (ref = l	U ntreated)	
	Normalized Treated (N=755)	Untreated (N=173)	<i>P</i> -value	Normalized Treated (N=755)	Untreated (N=176)	<i>P</i> -value
Age ≥50 years	736 (97.5%)	172 (99.4%)	0.1133	739 (97.8%)	169 (96.2%)	0.2142
Age, Median (Years)	64.3	66.5		64.4	64.3	
Body mass index ≥30 kg/m ²	439 (58.2)	98 (56.7)	0.7189	437 (57.8)	104 (59.4)	0.7072
Body mass index kg/m ² , mean (SD)	31.6 (6.1)	31.5 (6.6)		31.5 (6.1)	31.9 (6.5)	
Follow up time (years), mean (SD)	4.0 (3.4)	3.3 (3.1)		4.0 (3.4)	3.7 (3.6)	
Hypertension	630 (83.4%)	156 (90.2%)	0.0266	639 (84.7%)	144 (82.2%)	0.4214
Diabetes mellitus	323 (42.8%)	82 (47.4%)	0.2694	329 (43.6%)	73 (41.6%)	0.6355
Chronic obstructive pulmonary disease	48 (6.4%)	16 (9.3%)	0.1759	52 (6.9%)	12 (6.5%)	0.8735
Obstructive sleep apnea	81 (10.7%)	20 (11.6%)	0.7512	82 (10.9%)	18.3 (10.4%)	0.8675
Congestive heart failure	173 (22.9%)	49 (28.3%)	0.1325	181 (24.0%)	42 (23.9%)	0.9894

Peripheral vascular disease	69 (9.1%)	15 (8.7%)	0.8464	68 (9.0%)	15 (8.5%)	0.8295				
Depression	217 (28.7%)	44 (25.4%)	0.3827	212 (28.1%)	47 (26.9%)	0.7535				
LDL >100mg/dl	239 (31.7%)	54 (31.2%)	0.9102	238 (31.5%)	52 (29.3%)	0.5737				
Concomitant therapy with:					\mathbf{k}					
Antiplatelet Agents (ASA	677 (89.7%)	152 (87.9%)	0.4873	675 (89.4%)	158 (90.0%)	0.8047				
B-blockers	712 (94.3%)	162 (93.6%)	0.7369	711 (94.2%)	167 (94.8%)	0.7616				
Statins	719(95.2%)	162 (93.6%)	0.3896	717 (95.0%)	168 (95.8%)	0.6700				
	Normaliz	Normalized-treated Vs. Non-normalized-treated (ref = Non- normalized treated)								
	Normalized Treated (N=755)	Non- normalized- treated (N=542)	<i>P</i> -value	Normalized Treated (N=755)	Non- normalized- treated (N=542)	<i>P</i> -value				
Age ≥50 years	736 (97.5%)	528 (97.4%)	0.9402	736 (97.5%)	528 (97.4%)	0.9770				
Age, Median (Years)	64.3	64.5		64.3	64.5					
Body mass index ≥30 kg/m ²	439 (58.2)	335 (61.8)	0.1848	451 (59.7)	324 (59.7)	0.9987				
Body mass index kg/m ² , mean (SD)	31.6 (6.1)	32.3 (6.1)		31.8 (6.2)	32.1 (6.1)					
Follow up time (years), mean (SD)	4.0 (3.4)	3.4 (3.2)		4.0 (3.3)	3.4 (3.2)					
Hypertension	630 (83.4%)	456 (84.1%)	0.7401	632 (83.7%)	454 (83.7%)	0.9867				
Diabetes mellitus	323 (42.8%)	267 (49.3%)	0.0208	343 (45.5%)	246 (45.5%)	0.9918				
Chronic obstructive pulmonary disease	48 (6.4%)	52 (9.6%)	0.0312	59 (7.8%)	42 (7.7%)	0.9960				
						-				

Congestive heart failure	173 (22.9%)	142 (26.2%)	0.1736	183 (24.3%)	131 (24.2%)	0.9788			
Peripheral vascular disease	69 (9.1%)	52 (9.6%)	0.7811	70 (9.2%)	50 (9.2%)	0.9788			
Depression	217 (28.7%)	164 (30.3%)	0.5542	222 (29.4%)	159 (29.3%)	0.9817			
LDL >100mg/dl	239 (31.7%)	172 (31.7%)	0.9760	239 (31.7%)	171 (31.6%)	0.9790			
Concomitant therapy with:					R,				
Antiplatelet Agents (ASA)	677 (89.7%)	491 (90.6%)	0.5844	680 (90.0%)	488 (90.0%)	0.9875			
B-blockers	712 (94.3%)	510 (94.1%)	0.8738	712 (94.3%)	511 (94.3%)	0.9887			
Statins	719(95.2%)	521 (96.1%)	0.4387	722 (95.6%)	519 (95.7%)	0.9875			
	Non-Normalized-treated Vs. Untreated (ref = Untreated)								
	Non- normalized-	Untreated	Р-	Non- normalized-	Untreated	P-			
	treated (N=542)	(N=173)	value	treated (N=542)	(N=174)	value			
Age≥50 years	treated (N=542) 528 (97.4%)	(N=173) 172 (99.4%)	value 0.1091	treated (N=542) 531 (97.9%)	(N=174) 169 (97.2%)	0.5960			
Age ≥50 years Age, Median (Years)	treated (N=542) 528 (97.4%) 64.5	(N=173) 172 (99.4%) 66.5	value 0.1091	treated (N=542) 531 (97.9%) 64.8	(N=174) 169 (97.2%) 64.3	0.5960			
Age ≥50 years Age, Median (Years) Body mass index ≥30 kg/m ²	treated (N=542) 528 (97.4%) 64.5 335 (61.8)	(N=173) 172 (99.4%) 66.5 98 (56.7)	value 0.1091 0.2266	treated (N=542) 531 (97.9%) 64.8 328 (60.4)	(N=174) 169 (97.2%) 64.3 104 (59.9)	0.5960 0.9043			
Age ≥50 years Age, Median (Years) Body mass index ≥30 kg/m ² Body mass index kg/m ² , mean (SD)	treated (N=542) 528 (97.4%) 64.5 335 (61.8) 32.3 (6.1)	(N=173) 172 (99.4%) 66.5 98 (56.7) 31.5 (6.6)	value 0.1091 0.2266	treated (N=542) 531 (97.9%) 64.8 328 (60.4) 32.2 (6.1)	(N=174) 169 (97.2%) 64.3 104 (59.9) 31.9 (6.6)	0.5960 0.9043			
Age ≥50 years Age, Median (Years) Body mass index ≥30 kg/m ² Body mass index kg/m ² , mean (SD) Follow up time (years), mean (SD)	treated (N=542) 528 (97.4%) 64.5 335 (61.8) 32.3 (6.1) 3.4 (3.2)	(N=173) 172 (99.4%) 66.5 98 (56.7) 31.5 (6.6) 3.3 (3.1)	value 0.1091 0.2266	treated (N=542) 531 (97.9%) 64.8 328 (60.4) 32.2 (6.1) 3.4 (3.2)	(N=174) 169 (97.2%) 64.3 104 (59.9) 31.9 (6.6) 3.6 (3.4)	0.5960 0.9043			
Age ≥50 years Age, Median (Years) Body mass index ≥30 kg/m ² Body mass index kg/m ² , mean (SD) Follow up time (years), mean (SD) Hypertension	treated (N=542) 528 (97.4%) 64.5 335 (61.8) 32.3 (6.1) 3.4 (3.2) 456 (84.1%)	(N=173) 172 (99.4%) 66.5 98 (56.7) 31.5 (6.6) 3.3 (3.1) 156 (90.2%)	value 0.1091 0.2266 0.0488	treated (N=542) 531 (97.9%) 64.8 328 (60.4) 32.2 (6.1) 3.4 (3.2) 463 (85.6%)	(N=174) 169 (97.2%) 64.3 104 (59.9) 31.9 (6.6) 3.6 (3.4) 146 (84.1%)	0.5960 0.9043 0.6407			
Age ≥50 years Age, Median (Years) Body mass index ≥30 kg/m ² Body mass index kg/m ² , mean (SD) Follow up time (years), mean (SD) Hypertension Diabetes mellitus	treated (N=542) 528 (97.4%) 64.5 335 (61.8) 32.3 (6.1) 3.4 (3.2) 456 (84.1%) 267 (49.3%)	(N=173) 172 (99.4%) 66.5 98 (56.7) 31.5 (6.6) 3.3 (3.1) 156 (90.2%) 82 (47.4%)	value 0.1091 0.2266 0.0488 0.6695	treated (N=542) 531 (97.9%) 64.8 328 (60.4) 32.2 (6.1) 3.4 (3.2) 463 (85.6%) 264 (48.7%)	(N=174) 169 (97.2%) 64.3 104 (59.9) 31.9 (6.6) 3.6 (3.4) 146 (84.1%) 82 (47.4%)	0.5960 0.9043 0.6407 0.7614			

Obstructive sleep apnea	68 (12.6%)	20 (11.6%)	0.7312	67 (12.3%)	21 (11.9%)	0.8897
Congestive heart failure	142 (26.2%)	49 (28.3%)	0.5824	145 (26.8%)	47 (27.1%)	0.9415
Peripheral vascular disease	52 (9.6%)	15 (8.7%)	0.7167	51 (9.4%)	16 (9.4%)	0.9978
Depression	164 (30.3%)	44 (25.4%)	0.2238	158 (29.1%)	49 (28.2%)	0.8287
LDL >100mg/dl	172 (31.7%)	54 (31.2%)	0.8980	171 (31.6%)	52 (29.9%)	0.6831
Concomitant therapy with:						
Antiplatelet Agents (ASA)	491 (90.6%)	152 (87.9%)	0.2990	487 (89.9%)	156 (89.7%)	0.9476
B-blockers	510 (94.1%)	162 (93.6%)	0.8268	509 (94.0%)	164 (94.2%)	0.9018
Statins	521 (96.1%)	162 (93.6%)	0.1689	518 (95.5%)	166 (95.7%)	0.9070
		22				

22

 Table 2. Unadjusted and adjusted hazard ratios for all-cause mortality and recurrent myocardial

infarction.

Table 2							
Comparing Normalized-treated Vs. Untreated (ref = Untreated)							
Model	Al	ll-cause mort	ality	Myocardial infarction			
	HR	95% CI	Р	HR	95% CI	P	
Univariate	0.568	0.450-	<.0001	0.496	0.096-	0.4031	
NI 755 172		0.717			2.568		
N = 755 vs. 175							
Propensity Matched (Stabilized	0.764	0.598-	0.0312	0.790	0.118-	0.8073	
IPTW)	01701	0.976	0.0012		5.267	0.0072	
N=755 vs. 176							
Comparing Normalized-trea	ted Vs. No	n_normalized	l-treated (ref	Non-norr	nalized treate	d)	
Model	$\frac{1}{1}$					ction	
	HR	95% CI	P	HR	95% CI	P	
Univariate	0.722	0.611-	0.0001	1 027	0 245-	0.9710	
	0.722	0.854	0.0001	1.027	4.307	0.9710	
N= 755 vs. 542			r				
D	0.750	0.00	0.0017	1.000	0.050	0.0000	
Propensity Matched (Stabilized	0.759	0.039-	0.0017	1.099	0.253-	0.9000	
		0.901			4.764		
N= 755 vs. 542							
Comparing Non	-normalize	ed-treated Vs	. Untreated (ref=Untreat	ed)		
Model		l-cause mort	ality	Myo	cardial infar	ction	
	HR	95% CI	Р	HR	95% CI	Р	
Univariate	0.808	0.636-	0.0791	0.474	0.079-	0.4133	
N= 542 vs. 173		1.025			2.836		
Propensity Matched (Stabilized	0.970	0.757-	0.8084	0.575	0.082-	0.5794	
IPTW)		1.243			4.061		
N = 542 ys + 174							
11- 5+2 15. 174							
				L	1		
>							