Safety and efficacy of tocotrienol supplementation for bone health in postmenopausal women: protocol for a dose-response double-blinded placebo-controlled randomised trial

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ABSTRACT

Introduction: Osteoporosis is a major health concern in postmenopausal women, and oxidative stress contributes to the development of bone loss. Cellular studies and ovariectomised rat model mimicking bone loss in postmenopausal women show the bone-protective effect of tocotrienols (TTs) with antioxidant capability. We aim to access the safety and efficacy of TT consumption for bone health in postmenopausal women.

Methods and analysis: In this 12-week randomised double-blinded placebo-controlled trial for the effects of dietary TT supplementation in postmenopausal women, postmenopausal women aged 45 years and older with at least 1 year after menopause and bone mineral density T-score at the spine and/or hip 2.5 or more below the reference values will be randomly assigned to 3 daily supplements: (1) placebo group receiving 860 mg olive oil, (2) low TT group receiving 430 mg of 70% pure TTs (containing 300 mg TT) and (3) high TT group receiving 860 mg of 70% pure TTs (600 mg TT). The primary outcome measure will be urinary N-terminal telopeptide. The secondary outcome measures will be serum bone-specific alkaline phosphatase, receptor activator of nuclear factor-κB ligand, osteoprotegerin, urinary 8-hydroxy-2'-deoxyguanosine and quality of life. At 0, 6 and 12 weeks, the following will be assessed: (1) primary and secondary outcome measures; (2) serum TT and tocopherol concentrations; (3) physical activity and food frequency questionnaires. Liver function will be monitored every 6 weeks for safety. ‘Intent-to-treat’ principle will be employed for data analysis. A model of repeated measurements with random effect error terms will be applied. Analysis of covariance, χ² analysis and regression will be used for comparisons.

Ethics and dissemination: This study was approved by the Bioethics Committee of the Texas Tech University Health Sciences Center. The findings of this trial will be submitted to a peer-reviewed journal in the areas of bone or nutrition and international conferences.

Trial registration number: NCT02058420; results.
Clinical application of bone densitometry, such as dual-energy X-ray absorptiometry, is generally used to measure bone mineral density (BMD) at the spine, hip and femoral neck to predict the fracture risk, to monitor the natural progression of diseases that affect BMD, or to monitor the therapeutic response to osteoporosis-specific treatments 1–2 years after inception of therapies. In addition to BMD, serum bone turnover markers (bone formation markers and bone resorption markers) can offer another indication of the effectiveness of certain therapies including dietary supplements for restoring BMD. Bone turnover markers have shown to be promising in predicting fractures in the elderly up to 2 years before the event. A higher level of bone resorption rate is significantly associated with faster BMD loss. Decreased vertebral BMD and increased bone turnover have approximately equal power to predict the risk of bone loss and osteoporosis-related fracture rate in postmenopausal women. Therefore, efforts to decrease the occurrence of dilapidation fractures should concentrate on postmenopausal women diagnosed with low BMD. Since bone turnover markers have been increasingly recognised as important for bone metabolism, using bone turnover markers should be the first step to monitor the early response of intervention in this population.

Studies have shown bone loss can be attributed to dysregulation of osteoblast and osteoclast activity mediated by increased oxidative stress through the ageing process. The benefits of antioxidant supplements such as tocopherols (TPs) or tocotrienols (TTs; two major isoforms of vitamin E) to bone health have been shown in cellular and animal studies, including those using the ovariectomised rat model mimicking bone loss in postmenopausal women. In cellular and animal studies, TTs are much more beneficial than TPs in bone protection. Cohort observational studies in humans, however, have shown positive or negative or lack of associations between TP intake/serum TP concentration and BMD, casting doubts on the benefits of TPs. No cohort observational studies or randomised clinical trials have been conducted to address the effect of TTs on bone health in humans, especially in postmenopausal women who are at high risk for bone fracture. Therefore, the objective of this study is to evaluate the safety and efficacy of dietary TT supplements on bone health-related markers and mechanisms in postmenopausal women. We hypothesise that 12 weeks of TT supplements would benefit bone remodelling, as measured by bone markers in postmenopausal women with normal bone status and osteopenia, compared with those receiving placebo only, and the changes in bone markers associated with bone remodelling would be correlated with changes in oxidative stress. This hypothesis is based on the premise that TTs may reduce oxidative stress that adversely affects bone health by increasing differentiation and function of osteoclasts and inhibiting osteoblastic differentiation.

In this paper, we present the design and detailed protocol of a double-blinded placebo-controlled and randomised trial, as well as a discussion of the overall challenges of conducting this trial. The results from this trial will be reported at the completion of the study in accordance with the Consolidation of Standards for Reporting Trials guidelines.

METHODS/DESIGN

Study design

This is a 12-week double-blinded placebo-controlled and randomised intervention trial with allocation 1:1:1 for three treatment arms. Women 45 years and older with at least 1 year after menopause but no osteoporosis will be recruited primarily from local independent senior facilities, municipal senior community centres, and obstetrics and gynaecology clinics. A final sample size of 22 participants per group with an expected attrition rate of 15% over 12 weeks of intervention will produce an initial sample size of 78 participants. This sample size will yield a power of 0.80 at α=0.05 for detecting differences in primary outcome (eg, urinary NTX). After screening, qualified participants will be matched for age and randomly assigned to one of the three treatment groups: placebo, low TT and high TT. During the 12-week intervention, all participants will be provided with 500 mg elemental calcium and 400 IU vitamin D daily. The participants in the placebo group will receive two 430 mg olive oil softgels per day (one in the morning and another in the evening) for 12 weeks. The low TT participants will receive one 430 mg olive oil softgel in the morning and one 430 mg, 70% pure TT softgel (representing 300 mg TT) in the evening for 12 weeks. The high TT participants will receive two 430 mg, 70% pure TT softgels per day (representing 600 mg TT; one in the morning and another in the evening) for 12 weeks. The primary outcome measure is concentrations of urinary N-terminal telopeptide (NTX, a bone resorption marker) and the secondary outcome measures are the levels of serum bone-specific alkaline phosphatase (BAP, a bone formation marker), serum receptor activator of nuclear factor-κB ligand (sRANKL), serum osteoprotegerin (OPG), and urinary 8-hydroxy-2′-deoxyguanosine (8-OHdG, an oxidative stress DNA damage marker). Both primary and secondary outcome measures will be taken from the participants at baseline, 6 and 12 weeks. Liver function will also be monitored by assessing the activity of aspartate aminotransferase (AST) and alanine aminotransferase (ALT) at baseline, 6 and 12 weeks. The concentrations of serum TT will be determined at baseline, 6 and 12 weeks for bioavailability determination. Quality of life, physical activity and food frequency questionnaires will be assessed at baseline and 12 weeks. Investigators evaluating the endpoints will be blinded to interventions. The time points of all participant-related actions to be taken during the study period are presented in table 1.
Study setting, study population and recruitment

The participants, women 45 years or older with at least 1-year postmenopausal history and being ambulatory, for this present study will be recruited from Lubbock, Texas and surrounding areas. Ethnicity or race is not a factor in the inclusion of participants. Direct person-to-person solicitation in the obstetrics/gynaecology clinic and health fairs, flyers, non-solicited email system, campus announcements, local radio, newspapers, senior newsletters, and TV scripts will be used to recruit potential participants. In addition, we plan to recruit the minority participants who have limited access to such intervention due to social and cultural factors. We will advertise and host information sessions, particularly at local minority community centres and churches, partnering with leaders identified within these organisations. Advertisement in minority newspaper and radio will also be implemented. Based on our experience, it is not difficult to recruit qualified participants within a reasonable time frame from our existing participant pool with the methods described above. If a longer period is needed to recruit the participants, the intervention may take place in a staggered fashion—a block randomisation strategy will place participants into subgroups for treatments on a first-come, first-served basis.

Screening

Prescreening will be conducted through a phone interview or medical record review and will cover menstrual history, medication for osteoporosis, age, availability for the study period and most recent BMD results if available. Participants who passed the prescreening will attend an informed consent session and sign consents and Health Insurance Portability and Accountability Act (HIPAA) forms before completing a detailed questionnaire with demographic, health and dietary information. Visits for BMD and fasting blood screenings will follow afterwards.

Inclusion criteria

1. Postmenopausal women with no menses for at least 1 year or a serum follicle-stimulating hormone >30 IU/mL.
2. Age 45 and older.
3. Bone mass with BMD T-score >−2.5 (non-osteoporotic) at measured site (spine and/or hip).3
4. Participants satisfying the above screening criteria will receive BMD measurement of the anteroposterior (AP) lumbar spine (L1–L4) and proximal femur (femoral neck, Ward’s triangle and trochanter) by dual-energy X-ray absorptiometry scan (Norland Excell X-Ray Bone Densitometer, Serial No. 1490). Subject candidates must be diagnosed with no osteoporosis based on BMD T-score at the spine and/or hip >−2.5 below the young normal sex-matched areal BMD of the reference database.3 BMD (gm/cm²) will be determined by dividing bone mineral content (BMC; in grams of calcium hydroxyapatite) by the area (cm²) of interest. Dividing BMC by area partially corrects for differences in bone size.
5. Normal function of thyroid, liver and kidney.
6. Serum 25-hydroxy vitamin D ≥20 ng/mL.
7. No bisphosphonates at all if they had a treatment lasting at least 12 months.

Exclusion criteria

1. History of, or evidence for, metabolic bone disease including recent fractures (other than low BMD).
2. Having received medication (calcitonin, raloxifene or systemic glucocorticoids) within 3 months of the study initiation.
3. Having hormone/hormone-like replacement therapy within 6 months of the study initiation.
4. History of cancer within the past 5 years.
5. History or evidence of endocrine disease or malabsorption syndrome that would be a contraindication to the investigation of TT absorption.
6. Glycated haemoglobin of ≥7% in the past 3 months.
7. History of statin or other drugs for cholesterol control within 3 months of the study initiation.
8. Alcohol intake greater than ‘moderate’ (one drink per day) or use of non-steroidal anti-inflammatory drugs on a regular basis.
9. Cognitive impairment, depression or other medical/eating disorders, likely to move during the trial, lack of transportation to the study site, or being unavailable at sample collection times.
10. Smoking >10 cigarettes/day.
11. Unwillingness to accept randomisation.
12. Taking anticoagulants that may interact with TTs.

Sample size
In this study, we plan to enrol 78 patients with 26 in each of the three arms. Since this is a pilot study to evaluate the effects of TT treatments on bone biomarkers, we use the primary outcome measures (urine NTX) for the sample size calculation. Using Power Analysis and Sample Size software (PASS V.11), a sample size of 22 in each of the placebo, low TT and high TT groups will achieve a power of 80% to detect a clinically significant difference between the group means of urinary NTX level at the end of 12 weeks. This calculation is based on an analysis of covariance (ANCOVA), adjusted for three covariates: baseline NTX, age and body mass index (BMI).

We will perform analysis of variance (ANOVA) analysis of all three groups. If the ANOVA result is significant, we will perform post hoc pair-wise comparison. Therefore, it is not necessary to adjust for multiple testing. PASS software is used to calculate the required sample size based on the assumptions: (1) NTX level change (primary outcome measure) has a mean of 0 nM bone collagen equivalents (BCE)/mM creatinine (Crt) ng/mL in the placebo group with a SD of 12.69 nM BCE/mM Crt.44 (2) Compared with the placebo group, the low TT and high TT groups will have a relative decrease of 15% and 20% from the baseline of 53.5 nM BCE/mM Crt, respectively. (3) We also assume that the adjusted covariates will explain about 20% of the variance in the NTX, that is, a Pearson correlation coefficient of 0.45 (R^²=0.20) between NTX and the adjusted covariates. Taking into consideration an expected attrition rate of 15% and 20% from the baseline of 53.5 nM BCE/mM Crt, respectively, (3) We also assume that the adjusted covariates will explain about 20% of the variability in NTX, that is, a Pearson correlation coefficient of 0.45 (R^²=0.20) between NTX and the adjusted covariates. Taking into consideration an expected attrition rate of 15%, n=26 will be recruited for each group with a total of 78 participants to start the study.

Randomisation and allocation concealment
This study has a prospective, randomised, double-blinded, placebo-controlled design. Randomisation is based on the computer-generated table of random digits. All participants who had passed the screening criteria will participate in randomisation where participants will be assigned to one of the three treatment arms with a fixed randomised scheme. Enrolled de-identified patients will be randomly assigned 1:1:1 to receive placebo, low TT and high TT by the study biostatistician. The allocation sequence is sealed in numbered and opaque envelopes to ensure that the sequence is concealed. In order to minimise the consequent bias, a stratified block randomisation method with varying block sizes of 3 and 6 will be employed to assign participants to one of the three groups: placebo, low TT and high TT groups. The strata will be constructed based on BMI ($\geq$30 or $<$30 kg/m²) and age ($\geq$50 or $<$50 years). We will provide de-identified codes of participants to the biostatistician who will randomise group assignment and generate a group assignment number for each participant.

TT and placebo with the same character are prepared by a pharmacist. Patients and all study personnel except the investigative pharmacist will be blind to treatment assignment. The details of the series are unknown to any of the study participants, coordinators and outcome assessors and are contained in a set of opaque, sealed and numbered envelopes.

Intervention
Purchasing and blinding of study agents
Placebo and TT of the same lot, respectively, will be provided by American River Nutrition (Hadley, Massachusetts, USA; Investigational New Drug (IND) number 120761 by the Food and Drug Administration (FDA)). Each placebo capsule of 430 mg olive oil will contain no TT ingredient at detectable levels. Each TT capsule (DeltaGold Tocotrienol 70%) will contain 430 mg TT (90% $\delta$-TT and 10% $\gamma$-TT) with a 70% purity, representing 300 mg TT. The use of placebo to mask the control group is desirable in this study to keep respondents blinded to the TT treatment assignment. Placebo softgels will be made of the same size and colour as the TT softgels for identical appearance and taste. The placebo group will provide a comparison of blood and urinary outcome measures and serve as a basis to assess TT's effect.

Provision of calcium plus vitamin D supplement
The beneficial effects of dietary supplements and exercise on BMD depend on adequate Ca/vitamin D intake.45 According to the results of the National Health and Nutrition Examination Survey (NHANES; 1999–2002),46 the estimated average dietary calcium intake was 687±15 mg/day in postmenopausal women, which was much below the recommended daily intake (1200 mg/day) for this population. Thus, we will provide calcium (500 mg elemental Ca, as Oyster Shell)/vitamin D (400 IU as cholecalciferol) supplement (GlaxoSmithKline) to ensure that recommended daily intake levels of calcium (1000–1200 mg) and vitamin D (600 IU) will be reached, taking into account their dietary intake and sun exposure. Supplementation of calcium and vitamin D has been employed in previous clinical trials to investigate the effects of short-term nutritional intervention on bone turnover markers and bone metabolism in postmenopausal women.47 48 We refrained from providing a higher dose of Ca/vitamin D to avoid blinding the TT effect by the strong antiresorptive activity of Ca/vitamin D.
Treatment arms
The study agent (TT, DeltaGold Annatto Tocotrienol) has generally recognised as safe (GRAS) status. Based on (1) TT fed at 60 mg/kg body weight in rats showed osteoprotective impacts,25 29 30 and (2) the use of body surface area for dose translation from rat (250 g body weight) to human (70 kg body weight),48 the estimated effective dose of TT in humans for osteoprotection is ~680 mg daily. Therefore, we will test two dosages of TT (300 and 600 mg) for 12 weeks in this study. The current recommended dietary allowance (RDA) value of vitamin E, 15 mg/day, for this study population is largely focused on α-TP. No specific recommendation exists for TT. However, no adverse effects were observed with daily intake of 3.2 g or less TT.49

Qualified participants will be randomly assigned into one of the three study groups (placebo, low TT and high TT). Participants in the placebo group will receive 860 mg of olive oil daily (430 mg olive oil softgel×2 per day, morning and evening after meal) for 12 weeks. Participants in low TT group will receive a 430 mg olive oil softgel in the morning and a 430 mg TT softgel in the evening daily for 12 weeks. Participants in high TT group will receive a 430 mg TT softgel in the morning and another in the evening daily for 12 weeks.

Dietary intake, physical activity, concomitant medication assessment and quality of life
A food frequency questionnaire and a physical activity log will be collected at the baseline and 12-week visits to account for any drastic changes in the intake of macronutrients and micronutrients and deviations from the usual activities that could affect outcome measures. Prescription and over-the-counter medications and dietary supplement, as well as the reasons for taking these substances, will be recorded for the study period.

General health status will be measured with the Medical Outcomes Study 36-item short form Health Survey (SF-36, V2) at baseline and 12 weeks of study. SF-36 has been reported to have good validity, internal consistency and reliability in the assessment of physical and mental health status of participants and their progression.50 51 The SF-36 consists of eight dimensions of health (physical function, bodily pain, general health, vitality, mental health, social function, and role of physical and emotional health) in the conduct of daily activity.52

Blinding and unblinding
Study participants and investigators including biostatistician, coordinators, and measurement and site personnel will be blinded to intervention allocation throughout this study. The investigators will be supplied with a blind code-breaker envelope for each participant. The blind code will not be broken except in a medical emergency or a potential study-related adverse event determined by the principal investigator. Medical emergencies may include abnormal elevation in liver function (ALT, AST) according to the regulation of FDA for TT intake.

Sample collection
Blood will be drawn between 8:00 and 10:00 from a superficial arm vein will be allowed to clot in a vacutainer at room temperature. Within 2 hours of collection, blood samples will be centrifuged at 1500×g for 10 min and aliquoted. Urine samples will be collected in acid-washed polyethylene containers and aliquoted. Simultaneously collected blood and urine samples will be stored in −80°C freezers prior to biochemical analyses for primary and secondary outcome measures.

Outcomes measures
In the present study, the primary outcome is urinary NTX. The secondary outcomes include serum BAP, sRANKL, OPG, TT and TP concentrations, urinary 8-OHdG, and quality of life. In addition, we will monitor adherence and compliance, adverse events, food intake and physical activity throughout the study period.

Evaluation of adherence and compliance
We will measure adherence to the intervention by counting the softgels consumed and determine compliance by the percentage of all softgels ingested. We will also measure serum concentrations of TT and TP at baseline and 12-week visit as additional evidence for adherence and compliance.

Evaluation of adverse events
Despite the GRAS reports that TT at the doses used in this study has minimal liver toxicity in humans, we will monitor the liver function by measuring AST and ALT at 0, 6 and 12 weeks. Adverse effects associated with dietary supplement treatments, if any, will be self-reported by the participants and will be monitored by observing liver function every 6 weeks in the course of the intervention trial. All adverse events, whether observed by investigators or voluntarily disclosed by participants, will be recorded on the adverse event form throughout the study.

Measurements
Every participant will be evaluated at 0 (prior to starting intervention) 6 and 12 weeks of intervention.

Bone biomarkers
Rationale: Biochemical markers of bone turnover are promising in predicting fractures in the elderly for up to 2 years prior to the event.14 Serum BAP (a bone formation biomarker) and urinary NTX (a bone resorption biomarker) are more thorough clinical indicators of bone status than BMD in predicting skeletal response to a dietary supplement and in monitoring bone resorption changes as early as 3 months following initiation of intervention.53 On the other hand, studies have demonstrated that sRANKL and its decoy receptor OPG constitute a complex physiological mediator system involved in the regulation of bone resorption and may be responsible for the homoeostatic mechanism of normal bone...
remodelling. The serum BAP, RANKL, OPG and urine NTX are commonly used by bone researchers to monitor the changes in bone remodelling due to treatments.

**Methods:** The serum concentration of bone formation biomarker, BAP, will be measured using Metra BAP immunoassay kits (Quidel Corporation, San Diego, California, USA): the intra-assay and interassay coefficient of variation (CVs) are 5.2% and 5.0%, respectively. The concentration of bone resorption biomarker, NTX, in urine will be quantified using a commercial kit with a monoclonal antibody specific for the urinary N-terminal telopeptide (Alere, Providence, Rhode Island, USA). The intra-assay and interassay CVs are 2.2% and 3.0%, respectively. The concentrations of creatinine in urine will be measured using MicroVue Creatinine Assay Kit (Quidel Corporation). The final NTX will be normalised by urine creatinine. The serum concentration of sRANKL will be measured using sRANKL (total) human ELISA (BioVendor, LLC, Asheville, North Carolina, USA); the intra-assay and interassay CVs are 9.38% and 11.99%, respectively. The serum concentration of OPG will be quantified using a commercial MicroVue OPG kit (Quidel Corporation). The intra-assay and interassay CVs are 2.8% and 5.1%, respectively. In order to avoid the interassay variation, the samples from 0-week, 6-week and 12-week visits of the same patients will be measured for bone biomarkers within the same assay each time.

**Urinary 8-OHdG level**

**Rationale:** Urinary 8-OHdG measurement provides a sensitive and non-invasive way to evaluate the efficacy of dietary antioxidant supplements, such as TT, reactive oxygen species (ROS)-mediated macromolecule oxidation produces C8-OH-adduct radical with hydroxylation of guanine at the C8 position; C8-OH-adduct radical is subsequently converted to 8-OH-Guanine (8-OH-Gua) by a one-electron oxidation. Unlike damaged lipids and proteins, impaired DNA cannot be removed by metabolic turnover of molecules; instead, damaged DNA has to be repaired in situ or destroyed by apoptotic processes to avoid mutations. In humans, 8-OH-Glu glycosylase performs a short-patch base-excision repair to remove 8-OH-Gua, which is later converted to 8-OHdG and excreted into urine without further metabolism. The stability of urinary 8-OHdG renders the molecule a putative biomarker for oxidative stress and DNA damage.

**Methods:** The serum concentrations of 8-OHdG will be measured by OxiSelect Oxidative DNA Damage ELISA Kit (Cell Biolabs, San Diego, California, USA) following the manufacturer’s instruction. The intra-assay and inter-assay CVs were 9.38% and 11.99%, respectively.

**Serum TT and TP concentrations**

**Rationale:** The bioavailability and accumulation of TT is considered when we quantitatively evaluate the biological effects of TT intervention. Therefore, we will measure serum TT and TP concentrations of participants at 0, 6 and 12 weeks.

**Methods:** Internal standard (IS) retinyl acetate will be dissolved into methanol. We will mix 500 µL of serum and IS, then add 2 mL of 1% ascorbic acid in ethanol (wt/vol) and 25 µL of butylated hydroxytoluene (BHT) (10 mg/100 mL), and 2 mL of n-hexane. After vortex for 1 min, the tube will be centrifuged at 1500×g at 4°C for 5 min to facilitate phase separation. The supernatant will be transferred into a new tube. We will add additional 2 mL of n-hexane into serum sample tube, and repeat vortex and centrifugation as described above. The supernatant will be transferred and combined with the previous extraction and will be dried under nitrogen. We will add methanol to the tube and transfer the extraction to an ultracentrifuge filter before the filtrate transfer to an high-performance liquid chromatography (HPLC) phial. Extracted samples will be stored at ~20°C prior to HPLC analysis. Serum concentrations of TT will be measured using a HPLC system (Waters Corporation) equipped with a Waters1525 binary pump, 2707 autosampler with refrigeration unit, 2489 dual wavelength ultraviolet-visible detector, 1525 multi λ fluorescence detector and in-line degasser. The detection wavelength of TTs and TPs is 295/325 nm, and wavelength for IS is 325 nm. We will use a Phenomenex Kinetex PFP column (2.6 µm, 150×4.6 mm) with the mobile phase composed of methanol and water (7:1, v/v) and a flow rate of 0.8 mL/min. We will measure the four isomers of TPs (α, β, δ and γ) and four isomers of TT (α, β, δ and γ) in sera of study participants.

**Statistical analysis**

In the study, the primary outcome is NTX at 12 weeks and intervention is TT at three levels: placebo, low dose and high dose. The outcome and other cofactors will be summarised using appropriate summary measures as per the type and distribution of the variable according to each intervention level. The baseline cofactors will be compared among the three groups using one-way ANOVA or Fisher’s exact test as appropriate.

To assess the effect of intervention on primary outcome (urine NTX) at 12 weeks, ANCOVA will be conducted after adjusting the baseline NTX and other significant cofactors, if any. A post hoc comparison among the three intervention groups will be made using Tukey’s test if there is a significant effect of intervention. The assumptions of the ANCOVA model will be assessed and appropriate transformations will be made if needed for inferential data analysis. An ‘intention-to-treat’ analysis will be carried out for efficacious end point. The results will be summarised using mean differences and their 95% CIs.

Similarly, the same analysis approach for NTX will be employed for serum BAP, RANKL, OPG, T Ts and 8-OHdG. For secondary measures, the Pearson/Spearman rank correlation between change in NTX and change in the secondary biomarker (BAP, TTs, RANKL,
OPG, and 8-OHdG) will be computed and reported in tabular form. To evaluate quality of life, the ordinal score on the SF-36 test will be used, and therefore the Kruskal-Wallis test will be employed to detect any significant differences in the distribution of this score among the three groups followed by post hoc multiple comparisons if needed. p Value <0.05 will be considered as significant results. All the statistical analysis will be carried out using SAS V9.3.

Data management and data collection
Researchers will keep information obtained in this study confidential except as required by law. All participant questionnaires, records and data will be coded and properly stored in locked cabinets. Only study investigators will have access to those files linking a person’s study number to his/her name. Those forms will include the informed consent form, HIPAA authorisation form, all recruitment questionnaires and all other related materials.

The principle investigator will be responsible for oversight of data management of the trial. A certified monitor from Clinical Research Institute, independent from research team and sponsor, will assist the principle investigator in monitoring data collection before the data sets are delivered to study biostatistician. Questionnaire data will be entered on various forms, verified using computerised data entry forms and then entered into the Excel database. Laboratory data will also be verified and entered into an Excel database. Data of eligibility, medical records, attrition rate and compliance rate will all be entered into an Excel database. Data queries including missing values will be referred to the principle investigator. The principle investigator will have access to the final trial data set and disclosure of contractual agreements. The principle investigator will also incorporate any correction or addition into the datasets. Finally, a clean data set without any identification will be generated and delivered to the biostatistician for statistical analysis.

Ethics
An informed consent form will be signed by a participant before enrolling in the study. Any modifications to the protocol, which may affect the conduct of the study, potential benefits to the study participants, or their safety will be reported to the ethics committee for all necessary amendments. All study-related information will be stored securely at the study site in local cabinets, in an area with limited access (databases will be secured with a password-protected access system).

Dissemination
The findings of this trial will be submitted to a peer-reviewed journal in the areas of nutrition or bone. Abstracts will be submitted to relevant national and international conferences.

CONCLUSIONS
The effectiveness of TT for bone health in postmenopausal women is still unknown. Our study, carried out at a research centre with experience in conducting independent, investigator-initiated FDA-IND clinical trial, intends to address a gap in the field and will test the safety and efficacy of TT supplementation for bone markers in postmenopausal women.

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Contributors C-LS conceptualised the study. C-LS, HM, SY, SW, CKF and INS designed this trial. C-LS developed the first draft of the manuscript and the rest of coauthors participated in the revision of subsequent drafts. All authors approved the final draft of the manuscript. C-LS made final decision to submit the report for publication.

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