This is a repository copy of *The timed up and go test predicts fracture risk in older women independently of clinical risk factors and bone mineral density.*

White Rose Research Online URL for this paper:
http://eprints.whiterose.ac.uk/169589/

Version: Published Version

**Article:**
Larsson, B.A.M., Johansson, L., Johansson, H. et al. (9 more authors) (2021) The timed up and go test predicts fracture risk in older women independently of clinical risk factors and bone mineral density. Osteoporosis International, 32 (1). pp. 75-84. ISSN 0937-941X

https://doi.org/10.1007/s00198-020-05681-w

**Reuse**
This article is distributed under the terms of the Creative Commons Attribution-NonCommercial (CC BY-NC) licence. This licence allows you to remix, tweak, and build upon this work non-commercially, and any new works must also acknowledge the authors and be non-commercial. You don’t have to license any derivative works on the same terms. More information and the full terms of the licence here:
https://creativecommons.org/licenses/

**Takedown**
If you consider content in White Rose Research Online to be in breach of UK law, please notify us by emailing eprints@whiterose.ac.uk including the URL of the record and the reason for the withdrawal request.
The timed up and go test predicts fracture risk in older women independently of clinical risk factors and bone mineral density

B. A. M. Larsson¹ · L. Johansson¹,² · H. Johansson¹,³ · K. F. Axelsson¹,⁴ · N. Harvey⁵,⁶ · L. Vandenput³,⁷ · P. Magnusson⁸ · E. McCloskey⁹,¹⁰ · E. Liu³ · J. A. Kanis³,⁹ · D. Sund⁴ · M. Lorentzon¹,³,¹¹

Received: 17 June 2020 / Accepted: 9 October 2020 / Published online: 21 October 2020
© The Author(s) 2020

Abstract

Summary The timed up and go (TUG) test measures physical performance and predicts falls in the elderly. In older women, TUG time predicts the risk of major osteoporotic fracture and hip fracture independently of clinical risk factors and bone mineral density, and has a substantial impact on fracture probabilities.

Introduction The timed up and go (TUG) test measures physical performance and predicts falls in the elderly. A slow TUG has been associated with an increased fracture risk, but it is unclear whether the association is independent of clinical risk factors and bone mineral density (BMD). The aim of this study was to investigate if TUG time was associated with fracture risk independently of clinical risk factors and BMD and to determine its impact on fracture probabilities in older women.

Methods A standardized questionnaire was used to assess information regarding clinical risk factors in the large population-based SUPERB study of 3028 older women (75–80 years). At baseline, the TUG test was performed and BMD measured with DXA. The association between TUG time and the risk of hip fracture and major osteoporotic fracture (MOF) was examined using an extension of Poisson regression.

Results Fracture incidence increased steeply with increasing TUG time up to 12 s and subsequently started to level off. A slow TUG time was therefore defined as TUG > 12 s, a cutoff level then used in Cox models to study the association between slow TUG and fracture risk. A slow TUG time was associated with an increased risk of fracture (MOF 2.39 [1.80–3.18] and hip fracture 2.96 [1.62–5.40]). These associations were slightly attenuated but remained significant after adjustment for clinical risk factors and femoral neck BMD. Depending on BMD, the 4-year fracture probability of MOF increased by a factor of 1.5–1.9 in a 75-year-old woman with slow TUG (> 12 s).

Conclusion The TUG time predicts the risk of MOF and hip fracture independently of clinical risk factors and BMD and has a substantial impact on fracture probabilities, indicating that inclusion of the TUG test in patient evaluation should be considered in order to improve fracture prediction in older women.

Keywords Clinical risk factors and bone mineral density · Fracture risk · The timed up and go test

---

¹ Geriatric Medicine, Department of Internal Medicine and Clinical Nutrition, Institute of Medicine, University of Gothenburg, Gothenburg, Sweden
² Region Västra Götaland, Department of Orthopaedics, Sahlgrenska University Hospital, Mölndal, Sweden
³ Mary MacKillop Institute for Health Research, Australian Catholic University, Melbourne, VIC, Australia
⁴ Region Västra Götaland, Department of Orthopaedic Surgery, Skaraborg Hospital, Skövde, Sweden
⁵ MRC Lifecourse Epidemiology Unit, University of Southampton, Southampton, UK
⁶ NIHR Southampton Biomedical Research Centre, University of Southampton and University Hospital Southampton NHS Foundation Trust, Tremena Road, Southampton, UK
⁷ Centre for Bone and Arthritis Research, Institute of Medicine, University of Gothenburg, Gothenburg, Sweden
⁸ Department of Clinical Chemistry, and Department of Biomedical and Clinical Sciences, Linköping University, Linköping, Sweden
⁹ Centre for Metabolic Bone Diseases, University of Sheffield Medical School, Sheffield, UK
¹⁰ MRC and Arthritis Research UK Centre for Integrated Research in Musculoskeletal Ageing, Mellanby Centre for Bone Research, University of Sheffield, Sheffield, UK
¹¹ Region Västra Götaland, Department of Geriatric Medicine, Sahlgrenska University Hospital, 43180 Mölndal, Sweden
Introduction

Fractures in older women constitute a serious health risk, and often cause great suffering both in short and long term for the affected individuals [1, 2]. Additionally, patients with hip fracture often experience loss of physical function, decreased social engagement, increased dependence, and worsened quality of life [3–5]. A large proportion of patients who have had a hip fracture are forced to alter their living conditions, which could involve relocating from their home into a residential aged care facility, having extensive impact on the affected person’s autonomy [6–8]. Furthermore, all fractures and especially hip fractures increase the risk of mortality and morbidity in older men and women [9]. The etiology of fracture involves both bone fragility and fall risk. For each standard deviation (SD) decrease in bone mineral density (BMD) as assessed using dual-energy x-ray absorptiometry (DXA) in the femoral neck, the risk of hip fracture is increased nearly 3-fold [10]. Osteoporosis, sarcopenia, and reduced physical performance become more prevalent with increasing age and thereby contribute to the increasing risk of fall injuries and fractures with aging [11, 12]. Over 70% of all fractures affect women older than 65 years [13]. Risk factors for falls such as immobility and previous falls also contribute to the risk of fractures in the elderly [14, 15]. Identifying individuals who will fracture based on BMD alone has low sensitivity [16]. Therefore, fracture risk calculators which incorporate clinical risk factors in addition to BMD have been developed in recent years. Among those, the fracture risk assessment tool FRAX is the most widely used [17]. The timed up and go test (TUG) measures physical performance and has been used to identify frail older individuals. A slow TUG time (> 10 s) was associated with increased risk of fracture in a large study of Australian women, an association independent of BMD and some clinical risk factors [18]. However, the most appropriate cutoff time to define slow TUG time in relation to fracture risk has not been established. It is also not known if assessment of TUG time contributes independently to fracture risk when all clinical risk factors incorporated in FRAX are considered and to what extent TUG time affects fracture probability. The aim of the present work was to study the relationship between TUG time and risk of fracture, to investigate if TUG independently contributes to fracture risk when also considering all clinical risk factors currently included in FRAX and femoral neck BMD, and to determine to which extent TUG time contributes to the overall fracture probability in older women.

Materials and methods

Subjects

SUPERB—Sahlgrenska University Hospital Prospective Evaluation of the Risk of Bone fractures—is a prospective population-based study, carried out in the greater Gothenburg area. The study comprises 3028 women, 75–80 years old [19]. Women were chosen randomly from the Swedish national population register. All subjects signed an informed consent, prior to participation. The study protocol has been approved by the regional Ethics Review Board in Gothenburg. The criteria for being invited to the Study Clinic (Department of Geriatrics, Sahlgrenska University Hospital, Mölndal, Sweden) were as follows: (1) acceptance of the invitation sent by letter and providing a positive response to the follow-up telephone call, (2) be able to attend the clinic visit and being ambulatory, (3) understand Swedish, (4) being a woman between the age of 75 and 80 years.

Anthropometrics and TUG test

Body height was measured using a wall-mounted calibrated stadiometer. Body weight was measured to the nearest 0.1 kg using the same scale in all women. The timed up and go (TUG) test investigates balance and mobility [20, 21]. The participants were timed (in seconds) starting when rising from sitting in a chair, 45 cm high, equipped with armrests, walking 3 m in normal pace, turning around, walking back, and sitting down again. The time for this procedure was recorded. They could use their regular footwear and were allowed to utilize any mobility aids that they normally required. The TUG test was performed by 3004 participants, and 24 could not accomplish the test.

Questionnaires

Data regarding medical and fracture history, physical activity, occurrence of falls in the last 12 months, alcohol consumption, parental history of hip fracture, oral glucocorticoid use, alcohol consumption, and calcium intake were collected using questionnaires. Self-reported fractures after the age of 50 years at any location, except the skull and face, were included in the FRAX score calculations. Current smoking was defined using a validated questionnaire [22]. A small proportion of participants (≈ 1.6%) could not recall if a parent had sustained a hip fracture. A null response was assumed in those cases. A high alcohol consumption was defined as 21 standard drinks per week or more [23]. Mental and physical health (MC12 and PCS12) related quality of life was assessed using SF12 Health Survey [24–26]. The validated questionnaire Physical Activity Scale for the Elderly (PASE) was used to estimate physical activity in the last 7 days before the inclusion in the study [27]. PASE is a self-reported questionnaire which targets individuals over the age of 65 years. Participation (yes/no) or the number of hours spent in an activity is multiplied by given weights, thus giving a total score. The daily calcium intake was assessed and estimated in a validated questionnaire, by adding the food-derived calcium intake to the
amount of calcium provided by supplements [28]. No information regarding vitamin D intake or sunlight exposure was available.

**Dual-energy x-ray absorptiometry measures**

Bone density measurements were performed using the same DXA device for most participants (n = 2995) (Discovery A S/N 86491; Hologic, Waltham, MA, USA). Owing to a temporary machine failure, a few women (n = 33) were measured using another Discovery A Hologic DXA device. A cross-calibration was performed between the two instruments and has been reported elsewhere [19]. The areal BMD (aBMD) (g/cm²) of the femoral neck (FN) and lumbar spine (LS) were used in the analyses. Lateral scan imaging at baseline was performed using DXA with the participant in the supine position, in order to diagnose vertebral fractures, using the software program Physician’s Viewer (Hologic) as previously described [29]. After assessment of the anteroposterior lumbar, vertebral which were fractured and/or contained osteosynthesis materials in the LS (L₁ to L₄) were excluded. The LS aBMD was calculated as the mean of L₁ to L₄ if at least two vertebrae were assessable. The coefficients of variation (CV) were for aBMD FN and aBMD LS, 0.7% and 1.3%, respectively. The 4- and 10-year probabilities of MOF (major osteoporotic fractures; included fractures of the spine, hip, forearm, and proximal humerus) for women 75 and 80 years old were calculated according to femoral neck BMD T-score [30]. The National Health and Nutrition Examination Survey (NHANES) III reference database for total hip and femoral neck in young (20–29-year-old) Caucasian women and the Hologic sample for lumbar spine measurements comprising 30-year-old Caucasian American women were used to calculate the corresponding T-scores [31, 32].

**Biochemical analyses**

Blood samples were drawn from all study participants, and serum was separated, aliquoted, and stored at –80 °C until analyses. Serum 25-hydroxyvitamin D, calcium, and parathyroid hormone (PTH) were analyzed at the Department of Clinical Chemistry (Swedac accredited no. 1342), Linköping University Hospital, Sweden, and all samples were assayed with reagents from the same batch. Serum 25-hydroxyvitamin D was measured on the DiaSorin LIAISON® XL analyzer with the 25-hydroxyvitamin D total chemiluminescence immunoassay (DiaSorin, Stillwater, MN, USA), which demonstrates 100% cross-reactivity for 25-hydroxyvitamin D₂ and 25-hydroxyvitamin D₃. The assay performances were analytical range 10–375 nmol/L, and total CVs of 8.8%, 6.4%, and 6.8% at levels 25 nmol/L, 68 nmol/L, and 150 nmol/L, respectively. Serum intact PTH was determined using the Elecsys electrochemiluminescence immunoassay on a Roche Cobas e601 platform (Roche Diagnostics Scandinavia AB, Gothenburg, Sweden), which has the following assay performance: analytical range 0.13–530 pmol/L, and total CVs of 4.0%, and 2.9% at levels 1.9 pmol/L and 8.6 pmol/L, respectively.

**Incident fracture assessment**

Incident fractures were verified using radiographs. X-ray reports and/or images were retrieved from a regional digital X-ray archive that included all the 49 municipalities in the Västra Götaland region surrounding Gothenburg. All the radiology reports were initially reviewed by research nurses between the baseline exam until May 24, 2018. All reported fractures were recorded and all radiographs without available radiology reports or reports with uncertain fracture diagnosis were examined by an experienced orthopedic surgeon. Major osteoporotic fractures included clinical spine, hip, forearm, and proximal humerus fractures. Nonvertebral fractures included all fractures, except for fractures of the spine, skull, face, hand, and foot.

**Statistical analyses**

For continuous variables, independent samples t tests were used to examine differences between groups. χ² and Fisher’s exact tests were used for dichotomous variables. The association between TUG as a continuous variable and the risk of fracture was examined using an extension of the Poisson regression model in the whole cohort [33, 34]. The hazard function for major osteoporotic fracture and death was calculated using a modification of the Poisson regression model. For fracture, the variables in the hazard function were current time since baseline, current age, BMI, previous fracture, family history of hip fracture, smoking, corticosteroids, rheumatoid arthritis, alcohol use, and BMD. One additional model was constructed using the variables mentioned before adding slow TUG (0/1). For death, the variables in the hazard function were current time since baseline, current age, BMI, current smoking, per oral corticosteroid use, and BMD. Also, here, one additional model was constructed adding slow TUG (0/1). From the hazard functions for fracture and death, the 10-year probability of major osteoporotic fracture was calculated [35]. Follow-up is approximately 4 years for the SUPERB cohort, so when calculating 10-year probability, the hazard functions were extrapolated in time. It is important to note that the probability models used were based on purpose-built models similar to, but not identical to FRAX. The observation period of each participant was divided in intervals of 1 month. The first fracture per person was counted for each relevant outcome. Covariates included current age and time since start of follow-up. In order to study the association between TUG
and fracture risk in more detail, a spline Poisson regression model was fitted using cohort-specific knots at the 10th, 50th, and 90th percentiles of BMI, as recommended by Harrell [36]. The splines were second-order functions between the breakpoints and linear functions at the tails resulting in a smooth curve. The difference in log likelihood values using the spline and linear models were tested, in order to determine which model provides the most optimal curve-fit. The value considered significant was a $p$ value less than 0.05. Incidence per 1000 person-years was calculated as number of events divided by total follow-up time (until fracture, death, or censored) per 1000 years. Associations between TUG (TUG $\leq 12$ s or $>12$ s) and incident fractures were also studied using Cox proportional hazard models adjusted for age, height and weight as well as additional covariates, including all FRAX clinical risk factors (previous fracture, family history of hip fracture, current smoking, oral glucocorticoid use, rheumatoid arthritis, excessive alcohol intake), osteoporosis medication and history of falls, and censored for death or end of study (May 24, 2018). Hazard ratios (HR) and 95% confidence intervals derived from Cox models are presented. Statistical analyses were performed using SPSS Statistics Version 24 (IBM Corporation, Armonk, NY, USA). The hazard function developed by Fine and Gray was used to assess death as a competing risk for osteoporotic fractures [37]. For the calculations according to Fine Gray, STATA, Statistics/Data Analysis, version 16.0, serial number 401609206078, licensed to the University of Gothenburg, was used.

**Results**

**Baseline characteristics and TUG time**

During a median follow-up of 3.6 (Interquartile range, IQR 1.48) years, 335 women sustained a MOF, 314 a nonvertebral fracture, and 66 women a hip fracture. In the whole cohort ($n=3004$), median TUG time was 8.00 (IQR 2.6) seconds. The relationship between TUG and incident MOF and hip fracture showed a steep increase in fracture risk with TUG up to about 12 s (Fig. 1a and b) and then started to level off. The spline function provided a significantly improved curve-fit in comparison to a linear function for MOF ($p=0.01$), but the improvement was only borderline significant for hip fracture ($p=0.05$).

We therefore choose TUG above 12 s as the cutoff and divided the women with available TUG into two groups, TUG time $\leq 12$ s ($n=2711$) and TUG time $>12$ s ($n=293$). Characteristics of women with TUG $\leq 12$ s and women with TUG $>12$ s at baseline are presented in Table 1. Women with TUG $>12$ s were older, shorter, and heavier than women with TUG $\leq 12$ s. Physical activity and physical and mental health were inferior, while calcium and PTH were higher among those with slow TUG. Prevalence of falls, self-reported prior fracture, rheumatoid arthritis, hyperthyroidism, self-reported osteoporosis, hypertension, stroke, myocardial infarction, angina, heart failure, type 2 diabetes, chronic bronchitis/asthma/emphysema, and women using osteoporosis medication were all more common among those with TUG $>12$ s ($p<0.05$), indicating an increased comorbidity in this group of women.

**A slow TUG time was associated with a higher fracture incidence**

The incident fractures were divided into three groups; nonvertebral fracture, MOF, and hip fracture. The incidence of nonvertebral fracture, MOF, and hip fracture was substantially higher in women with TUG $>12$ s than in women with faster TUG. Cox proportional hazard models adjusted for age, height, and weight demonstrated that TUG $>12$ s was associated with increased risk of nonvertebral fracture (hazard ratio (HR) and 95% confidence interval (CI) 2.09 [1.53–2.84]), MOF (HR, 95% CI 2.39 [1.80–3.18]), and hip fracture (HR 95% CI 2.96 [1.62–5.40]). These associations were somewhat attenuated but remained significant also after adjustments for clinical risk factors included in FRAX, use of osteoporosis medication, prior falls, and femoral neck BMD (Table 2).

**The impact of TUG $>12$ s on fracture probabilities**

Study subject follow-up time was extrapolated up to 10 years to allow for calculations of 10-year fracture probability. The 10-year probabilities of MOF for women 75 and 80 years old were calculated, setting BMI to 26 kg/m$^2$, previous fracture set to yes, and all other clinical risk factors set to no, according to femoral neck BMD T-score, with or without consideration to TUG ($\leq 12$ s or $>12$ s) in the analysis. For a 75-year-old woman with BMD T-score $-2$, a slow TUG $>12$ s increased the 10-year probability substantially, from 34.3 to 47.5%. The equivalent 10-year probability for an 80-year-old woman with a T-score of $-2$ was 37.3% and 48.7%, for TUG $\leq 12$ s or $>12$ s, respectively (Fig. 2a, b). The 4-year probabilities of MOF for women 75 and 80 years old, with previous fracture, BMD of 26 kg/m$^2$, and no additional clinical risk factors, were also calculated according to femoral neck BMD T-score, with or without consideration to TUG ($\leq 12$ s or $>12$ s) in the analysis. For a 75-year-old woman with BMD T-score $-2$, TUG $>12$ s was associated with a markedly higher 4-year probability (14% vs. 24%). The corresponding 4-year probability for an 80-year-old woman with a T-score of $-2$ was 16% and 26%, for TUG $\leq 12$ s or $>12$ s, respectively (Fig. 2c, d). The ratios between the calculated 4-year probability without considering TUG and with TUG $>12$ s for women 75 and 80 years old, with previous fracture, BMI of 26 kg/m$^2$ but no other clinical risk factors, according to femoral neck BMD T-score, are presented in Fig. 3. The relative importance
The impact of competing risk of death according to Fine and Gray

The association between TUG > 12 and risk for major osteoporotic fracture (subhazard ratio (SHR) and 95% CI 2.31 (1.73–3.09)), hip fracture (SHR 2.85 (1.56–5.22)), and nonvertebral fracture (SHR 2.01 (1.47–2.76)) did not change substantially when a competing risk survival regression model, adjusted for age, height, and weight, was applied.

Discussion

In the present study, we demonstrate that TUG is an independent predictor of nonvertebral fracture, MOF, and hip fracture, and that these associations are independent of clinical risk factors included in FRAX and BMD of the femoral neck. Fracture risk increased progressively with TUG time and started to level off when TUG time exceeded 12 s. Having a slow TUG time (> 12 s) had a substantial impact, on the probability of MOF and hip fracture, indicating that evaluation of TUG could be useful in determining fracture risk in older women.

Although the spline functions describing the relationship between TUG and MOF or hip fracture provided better curve fits than linear models, the proposed and used cutoff of 12 s is clearly not perfect, since it is evident that the risk of fracture progresses further with TUG time slower than 12 s. It should be acknowledged that the spline model herein presented for hip fracture is based on few fractures (n = 66) with resulting large confidence intervals. However, it can be argued that deriving the cutoff using spline models for fracture risk is superior to, as previously done, using age-specific means without any consideration of fracture risk [18, 20].

FRAX is the most widely used fracture risk assessment tool for estimating individualized 10-year probability of hip and major osteoporotic fracture [30, 38–41]. The calculated 10-year fracture probability is based on clinical risk factors with or without BMD, and recommended in many clinical guidelines to calculate the probability of a MOF or hip fracture [41, 42]. In the presently investigated cohort, a slow TUG time (> 12 s) increased the 10-year probability substantially in women of all ages and with low to normal BMD, to risk levels above the 20%, a commonly used treatment threshold [42]. Thus, performing the TUG test and considering TUG time would have a substantial impact on treatment decisions in women in this age group.

TUG performance is known to capture several different aspects of aging, such as poor balance, falls, and disability of daily living [43–46]. TUG time has been shown to be able to predict frailty with high sensitivity and specificity but is to a lesser extent able to discriminate fallers from non-fallers [47, 48]. In the present study, it was apparent that women with a slow TUG time (> 12 s) had a considerably poorer general health than women with normal TUG, as indicated by a higher BMI, worse physical and mental quality of life, as well as more prevalent falls and fractures, type 2 diabetes, stroke, osteoporosis, Parkinson’s disease, myocardial infarction, and heart failure. Thus, a slow TUG time served as a proxy for worse general health and can therefore be used to efficiently identify physical frailty which negatively impacts the risk of falls and fractures.

It can be problematic to identify risk factors in older populations with considerable comorbidities and a high mortality rate, due to the competing risk of death [49]. In the herein presented analysis, we also performed analysis of the relationship between TUG time and incident fracture, using the Fine and Gray method, to adjust for competing mortality. The robust associations between TUG and fracture risk remained when considering competing mortality, supporting the notion that TUG performance is useful to test also in this age group.
Table 1  Characteristics of older women with timed up and go (TUG) ≤ 12 s and > 12 s

| Characteristics                                      | TUG ≤ 12 s (n = 2711) | TUG > 12 s (n = 293) | p value<sup>a</sup> |
|------------------------------------------------------|------------------------|----------------------|---------------------|
| Age (years)                                          | 77.7 ± 1.6             | 78.2 ± 1.6           | < 0.001             |
| Height (cm)                                          | 162.0 ± 5.7            | 160.2 ± 6.5          | < 0.001             |
| Weight (kg)                                          | 68.1 ± 11.5            | 74.4 ± 15.3          | < 0.001             |
| Body mass index (kg/m²)                              | 25.9 ± 4.2             | 29.0 ± 5.5           | < 0.001             |
| FN aBMD (g/cm²)                                      | 0.66 ± 0.10<sup>b</sup> | 0.65 ± 0.12         | 0.041               |
| Hip aBMD (g/cm²)                                     | 0.80 ± 0.11<sup>b</sup> | 0.78 ± 0.13<sup>g</sup> | < 0.001            |
| Lumbar spine aBMD (g/cm²)                           | 0.94 ± 0.16<sup>g</sup> | 0.98 ± 0.18<sup>b</sup> | < 0.001            |
| FRAX MOF w/o BMD (%)                                 | 30.8 ± 12              | 30.3 ± 12            | 0.50                |
| FRAX hip fracture w/o BMD (%)                        | 18.0 ± 12              | 16.9 ± 12            | 0.12                |
| FRAX MOF with BMD (%)                                | 22.7 ± 12              | 25.1 ± 13            | < 0.01              |
| PASE                                                 | 108.4 ± 50.0<sup>b</sup> | 67.9 ± 41.6<sup>g</sup> | < 0.001            |
| PCS                                                  | 46.6 ± 10.1            | 32.9 ± 9.4           | < 0.001             |
| MCS                                                  | 54.0 ± 8.9             | 49.7 ± 11.9          | < 0.001             |
| Calcium intake (from dairy products), mg/day         | 397 ± 362              | 313 ± 381            | < 0.01              |
| Fall accident within the last year, %                | 26.7<sup>j</sup>       | 53.4<sup>j</sup>     | < 0.001             |
| Self-reported prior fracture, %<sup>φ</sup>          | 35.8<sup>k</sup>       | 45.7<sup>k</sup>     | < 0.001             |
| Parental history of hip fracture, %                  | 17.5<sup>m</sup>       | 17.3<sup>n</sup>     | 0.922               |
| Current smoking, %                                    | 5.0                    | 6.5<sup>o</sup>      | 0.263               |
| Excessive alcohol consumption, %<sup>x</sup>         | 0.6<sup>p</sup>        | 0.3<sup>q</sup>      | 1.000<sup>e</sup>   |
| Blood biochemistry (serum)                            |                        |                      |                     |
| Creatinine (μmol/L)                                  | 76.7 ± 15.7            | 85.6 ± 32.5<sup>g</sup> | < 0.001            |
| Calcium (mmol/L)                                     | 2.47 ± 0.1             | 2.49 ± 0.1<sup>g</sup> | 0.016               |
| 25-Hydroxyvitamin D<sub>3</sub> (nmol/L)              | 62.5 ± 21.0            | 60.6 ± 23.4<sup>g</sup> | 0.143               |
| PTH (pmol/L)                                         | 4.90 ± 2.0             | 6.30 ± 3.6           | < 0.001             |
| Medications                                          |                        |                      |                     |
| Glucocorticoid use, %                                 | 3.4<sup>r</sup>        | 2.8<sup>s</sup>      | 0.564               |
| Osteoporosis medication, %<sup>ω</sup>               | 9.7<sup>t</sup>        | 17.2<sup>u</sup>     | < 0.001             |
| Medical history                                      |                        |                      |                     |
| Rheumatoid arthritis, %                              | 3.4<sup>r</sup>        | 7.5<sup>v</sup>      | < 0.001             |
| Hyperthyroidism, %                                   | 4.8<sup>w</sup>        | 7.9<sup>x</sup>      | 0.026<sup>e</sup>   |
| Osteoporosis, %<sup>↓</sup>                          | 18.7<sup>y</sup>       | 34.6<sup>z</sup>     | < 0.001             |
| Hypertension, %                                      | 51.5<sup>n</sup>       | 59.2<sup>β</sup>     | 0.012               |
| Stroke, %                                            | 5.8<sup>v</sup>        | 17.5<sup>u</sup>     | < 0.001             |
| Myocardial infarction, %                             | 4.1<sup>β</sup>        | 11.3<sup>c</sup>     | < 0.001             |
| Angina, %                                            | 4.6<sup>β</sup>        | 12.7<sup>n</sup>     | < 0.001             |
| Heart failure, %                                     | 7.8<sup>β</sup>        | 18.2<sup>a</sup>     | < 0.001             |
| Parkinson’s disease, %                               | 0.52<sup>ε</sup>       | 3.4<sup>κ</sup>      | < 0.001<sup>ε</sup> |
| Type 2 diabetes, %                                   | 8.9<sup>μ</sup>        | 17.1<sup>u</sup>     | < 0.001             |
| Chronic bronchitis, asthma, emphysema, %             | 9.2<sup>α</sup>        | 16.8<sup>n</sup>     | < 0.001             |
| Cancer, %                                            | 20.6<sup>ο</sup>       | 21.2<sup>λ</sup>     | 0.811               |
| Glaucoma, %                                          | 8.0<sup>ν</sup>        | 8.9<sup>η</sup>      | 0.579               |

Values are presented as mean ± standard deviation for continuous variables and as percentage and number for categorical variables. Significance was defined by a p value < 0.05 and significant values are italicized

<sup>a</sup>Categorical variables χ² test;<sup>b</sup>2699;<sup>c</sup>2673;<sup>d</sup>2709;<sup>e</sup>Fisher’s exact test;<sup>f</sup>2693;<sup>g</sup>292;<sup>h</sup>285;<sup>i</sup>1724;<sup>j</sup>156;<sup>k</sup>967;<sup>l</sup>135;<sup>m</sup>133;<sup>n</sup>469;<sup>o</sup>49;<sup>p</sup>16;<sup>q</sup>1;<sup>r</sup>92;<sup>s</sup>8;<sup>t</sup>262;<sup>u</sup>50;<sup>v</sup>22;<sup>w</sup>129;<sup>x</sup>23;<sup>y</sup>506;<sup>z</sup>101;<sup>α</sup>1395;<sup>β</sup>173;<sup>γ</sup>157;<sup>δ</sup>110;<sup>ε</sup>33;<sup>ζ</sup>125;<sup>η</sup>37;<sup>θ</sup>373;<sup>ι</sup>53;<sup>κ</sup>14;<sup>λ</sup>10;<sup>μ</sup>242;<sup>π</sup>248;<sup>ρ</sup>559;<sup>ς</sup>62;<sup>σ</sup>216;<sup>τ</sup>26.<sup>φ</sup>After 50 years of age, fractures of the skull and face are excluded,<sup>χ</sup>21 or more units per week;<sup>ψ</sup>daily oral treatment with at least 5 mg prednisolone (or equivalent) for 3 months or more ever;<sup>ω</sup>current treatment with bisphosphonates, teriparatide, or denosumab;<sup>↓</sup>Self-reported—been diagnosed with osteoporosis, TUG timed up and go, FN femoral neck, PASE Physical Activity Scale for the Elderly, MCS Mental health Component Scale, PCS Physical Health Component Scale
In a randomized controlled trial of older women (age 75 years) investigating the effect of calcium supplementation on fracture risk, TUG was associated with incident fractures after adjusting for calcium treatment and several other risk factors and covariates, including BMD [18]. In a recent, very large study (n = 1,070,320) of male and female Koreans, 66 years old, a slow TUG time (≥10 s) was found to be associated with a modest 21\% increased risk of hip fracture and a 7\% increased risk in vertebral fracture, compared to those with a faster TUG time [50]. In contrast to the analysis in the present study, neither of these previous studies attempted to investigate and identify the appropriate threshold, most strongly associated with an elevated fracture risk. Furthermore, all risk factors presently included in FRAX were not considered and the impact of TUG on the over-all fracture probability was not assessed in these studies [18, 50].

This study has some limitations. The inclusion criteria required that women were older (75–80 years old), were ambulatory, and able to understand Swedish. Thus, the results may not apply to women in other age groups, residing in nursing homes or with other ethnic backgrounds. Although women with walking aids were included, women with more severe disability (not able to walk at all) were excluded. The TUG test was not available in all women in this study, but a very large proportion were able to complete the test (3004 out of 3028) demonstrating its usefulness in this population. The prevalence of a slow TUG is likely higher in women with a higher prevalence of disability but lower in younger women. The importance of TUG as predictor of fracture in such populations could be different from the herein investigated population.

This study also has strengths. It is a large population-based, prospective study, comprising over 3000 older women, a population with very high fracture risk. All identified fractures were confirmed using x-rays or radiology reports, ensuring high quality of the fracture data. It is the first study to

| Table 2 | Associations between TUG time > 12 s and fracture risk in older women |
|-----------------|-----------------|-----------------|-----------------|-----------------|
|                | TUG ≤ 12 s (n = 2711) | TUG > 12 s (n = 293) |
| Nonvertebral fracture\(^a\) | | |
| No. (%)        | 262 (9.7)      | 52 (17.7)      |
| Rate per 1000 person-years | 28.6          | 54.5          |
| Time at risk, median (IQR), years | 3.4 (1.68) | 3.3 (1.54) |
| HR (95\% CI)   |                |                |
| Adjusted for age, height, weight (model 1) | 1 [Reference] | 2.09 [1.53–2.84] |
| + Clinical risk factors (model 2) | 1 [Reference] \(^b\) | 1.61 [1.16–2.23] \(^e\) |
| + FN BMD (model 3) | 1 [Reference] \(^d\) | 1.45 [1.04–2.02] \(^e\) |
| Major osteoporotic fracture | | |
| No. (%)        | 271 (10.0)      | 64 (21.8)      |
| Rate per 1000 person-years | 29.7          | 67.9          |
| Time at risk, median (IQR), years | 3.4 (1.68) | 3.3 (1.62) |
| HR (95\% CI)   |                |                |
| Adjusted for age, height, weight (model 1) | 1 [Reference] | 2.39 [1.80–3.18] |
| + Clinical risk factors (model 2) | 1 [Reference] \(^b\) | 1.96 [1.45–2.65] \(^e\) |
| + FN BMD (model 3) | 1 [Reference] \(^d\) | 1.76 [1.30–2.40] \(^e\) |
| Hip fracture   | | |
| No. (%)        | 51 (1.9)        | 15 (5.1)       |
| Rate per 1000 person-years | 5.5          | 14.6          |
| Time at risk, median (IQR), years | 3.6 (1.28) | 3.5 (1.61) |
| HR (95\% CI)   |                |                |
| Adjusted for age, height, weight (model 1) | 1 [Reference] | 2.96 [1.62–5.40] |
| + Clinical risk factors (model 2) | 1 [Reference] \(^b\) | 2.66 [1.40–5.08] \(^e\) |
| + FN BMD (model 3) | 1 [Reference] \(^d\) | 1.93 [1.00–3.71] \(^e\) |

Associations were studied using Cox proportional hazard models. Hazard ratios (HR) and 95\% confidence intervals (CI) are presented. Model 1: adjusted for age, height, and weight. Model 2: adjusted for age, height, weight, and clinical risk factors included in FRAX (previous fracture, family history of hip fracture, current smoking, oral glucocorticoid use, rheumatoid arthritis, excessive alcohol intake), osteoporosis medication, and history of falls. Major osteoporotic fractures included fractures of the spine, hip, forearm, and proximal humerus. Model 3: adjusted all covariates used in model 2 with the addition of FN BMD

\(^a\) Excluding fractures of the skull, face, hand and foot; \(^b\) 2661; \(^c\) 279; \(^d\) 2649; \(^e\) 278
investigate and present results on the impact of TUG performance on fracture probability, also after considering all currently used FRAX clinical risk factors and BMD. However, additional similar studies and meta-analyses will be required to determine if TUG performance could provide additional value to future FRAX-models.

In conclusion, the present study demonstrates that TUG time is strongly associated with hip, MOF, and nonvertebral fractures in older women, also after adjustments for all FRAX clinical risk factors and BMD. These results indicate that TUG performance could be included as a routine clinical assessment in order to improve fracture prediction in older women.
Acknowledgments  We would like to thank the late Associate Professor and MD Anna Nilsson for her dedicated work, for being an inspiration, and for her invaluable contributions to the planning of this study, collection of data, and in the interpretations of results. We also express our sincere gratitude to all the study participants in the SUPERB cohort.

Funding  Open access funding provided by University of Gothenburg.

Compliance with ethical standards  All subjects signed an informed consent, prior to participation. The study protocol has been approved by the regional Ethics Review Board in Gothenburg.

Conflicts of interest  Professor Lorentzon has received lecture fees from Amgen, Lilly, Meda, Renapharma, UCB Pharma, and consulting fees from Amgen, Radius Health, UCB Pharma, Renapharma, and Consilient Health. J. A. Kanis has received grant support from Amgen, Lilly, and Radius Health; N. Harvey has received consultancy, lecture fees, and honoraria from Amgen, AstraZeneca, Consilient Healthcare, Fresenius Kabi, GSK, Hologic, Internis, Lilly, Merck, Novartis, Pfizer, Roche, Sanofi-Aventis, Servier, Synexus, UCB, Unilever, and Warner Chilcott.

Open Access  This article is licensed under a Creative Commons Attribution-NonCommercial 4.0 International License, which permits any non-commercial use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by-nc/4.0/.

References

1. Covinsky KE, Palmer RM, Fortinsky RH, Counsell SR, Stewart AL, Kresevic D, Burant CJ, Landefeld CS (2003) Loss of independence in activities of daily living in older adults hospitalized with medical illnesses: increased vulnerability with age. J Am Geriatr Soc 51(4):451–458. https://doi.org/10.1046/j.1532-5415.2003.51152.x
2. von Friesendorff M, McGuigan FE, Wizert A, Rogmark C, Holmberg AH, Woolf AD, Akesson K (2016) Hip fracture, mortality risk, and cause of death over two decades. Osteoporosis Int 27(10):2945–2953
3. Adib Hajbaghery M, Abbassinia M (2013) Quality of life for the elderly after hip fracture surgery: a case-control study. J Caring Sci 2(1):53–59. https://doi.org/10.5681/jcs.2013.007
4. Gesar B, Hommel A, Hedin H, Baath C (2017) Older patients’ perception of their own capacity to regain pre-fracture function after hip fracture surgery - an explorative qualitative study. Int J Orthop Trauma Nurs 24:50–58. https://doi.org/10.1016/j.ijotn.2016.04.005
5. Alegre-Lopez J, Cordero-Guevara J, Alonso-Valdivielso JL, Fernandez-Melon J (2005) Factors associated with mortality and functional disability after hip fracture: an inception cohort study. Osteoporos Int 16(7):729–736. https://doi.org/10.1007/s00198-004-1740-0
6. Johansen A, Mansor M, Beck S, Mahoney H, Thomas S (2012) Outcome following hip fracture: long term mortality and post-discharge residence. Osteoporos Int 23:5545–5545
7. Salkeld G, Cameron ID, Cumming RG, Easter S, Seymour J, Kurrel SE, Quine S (2000) Quality of life related to fear of falling and hip fracture in older women: a time trade off study. Brit Med J 320(7231):341–345
8. Cumming RG, Klineberg R, Katelaris A (1996) Cohort study of risk of institutionalisation after hip fracture. Aust N Z J Public Health 20(6):579–582
9. Center JR, Nguyen TV, Schneider D, Sambrook PN, Eismann JA (1999) Mortality after all major types of osteoporotic fracture in men and women: an observational study. Lancet 353(9156):878–882
10. Kanis JA, Johnell O, De Laet C, Johansson H, Oden A, Delmas P, Eisman J, Fujiwara S, Garnero P, Kroger H, McCloskey EV, Mellstrom D, Melton LJ, Pols H, Reeve J, Silman A, Tenenhouse A (2004) A meta-analysis of previous fracture and subsequent fracture risk. Bone 35(2):375–382. https://doi.org/10.1016/j.bone.2004.03.024
11. Tieland M, Trouwborst I, Clark BC (2018) Skeletal muscle performance and ageing. J Cachexia Sarcopenia Muscle 9(1):3–19. https://doi.org/10.1007/s12325-022-012238
12. Lorentzon M, Cummings SR (2015) Osteoporosis: the evolution of a diagnosis. J Intern Med 277(6):650–661. https://doi.org/10.1111/joim.12369
13. Sambrook P, Cooper C (2006) Osteoporosis. Lancet 367(9527):661–668. https://doi.org/10.1016/j.ijotn.2016.04.005
14. Nilsson M, Eriksson J, Larsson B, Oden A, Johansson H, Lorentzon M (2016) Fall risk assessment predicts fall-related injury, hip fracture, and head injury in older adults. J Am Geriatr Soc 64(11):2242–2250. https://doi.org/10.1111/jgs.14439
15. Deandrea S, Lucenteforte E, Bravi F, Foschi R, Vecchia C, Negri E (2010) Risk factors for falls in community-dwelling older people: a systematic review and meta-analysis. Epidemiology (Cambridge, Mass) 21:658–668. https://doi.org/10.1097/EDE.0b013e3181e89905
16. Cranney A, Jamal SA, Tsang JF, Josse RG, Leslie WD (2007) Low bone mineral density and fracture burden in postmenopausal women. Can Med Assoc J 177(6):575–580
17. Kanis JA, Harvey NC, Johansson H, Oden A, Leslie WD, McCloskey EV (2017) FRAX update. J Clin Densitom 20(3):360–367. https://doi.org/10.1016/j.jocd.2017.06.022
18. Zhu K, Devine A, Lewis JR, Dhaliwal SS, Prince RL (2011) ‘Timed up and go’ test and bone mineral density measurement for fracture prediction. Arch Intern Med 171(18):1655–1661. https://doi.org/10.1001/archinternmed.2011.434
19. Lorentzon M, Nilsson AG, Johansson H, Kanis JA, Mellstrom D, Sundh D (2019) Extensive undertreatment of osteoporosis in older Swedish women. Osteoporos Int 30(6):1297–1305. https://doi.org/10.1007/s00198-019-04872-4
20. Bohannon RW (2006) Reference values for the timed up and go test: a descriptive meta-analysis. J Geriatr Phys Ther 29(2):64–68. https://doi.org/10.1519/00139143-20060800-00004
21. Podsiadlo D, Richardson S (1991) The timed “Up & Go”: a test of basic functional mobility for frail elderly persons. J Am Geriatr Soc 39(2):142–148. https://doi.org/10.1111/j.1532-5415.1991.tb01616.x
22. Vartiainen E, Seppala T, Lillsunde P, Puska P (2002) Validation of self-reported smoking by serum cotinine measurement in a
community-based study. J Epidemiol Community Health 56(3): 167–170. https://doi.org/10.1136/jech.56.3.167
23. Bergman H, Kallmen H (2002) Alcohol use among Swedes and a psychometric evaluation of the alcohol use disorders identification test. Alcohol Alcohol 37(3):245–251. https://doi.org/10.1093/alcalc/37.3.245
24. Resnick B, Nahm ES (2001) Reliability and validity testing of the revised 12-item Short-Form Health Survey in older adults. J Nurs Meas 9(2):151–161
25. Vilagut G, Forero CG, Pinto-Meza A, Haro JM, de Graaf R, Bruffaerts R, Kovess V, de Girolamo G, Matschinger H, Ferrer M, Alonso J, Investigators E (2013) The mental component of the short-form 12 health survey (SF-12) as a measure of depressive disorders in the general population: results with three alternative scoring methods. Value Health 16(4):564–573
26. Luo XM, George ML, Kakouras I, Edwards CL, Pietrobon R, FRAX and the assessment of fracture probability in men and women from the UK. Osteoporos Int 19(4):385–397. https://doi.org/10.1007/s00198-004-1780-5
27. Washburn RA, McAuley E, Katula J, Mihalko SL, Boileau RA (2011) Is the timed up and go test a useful predictor of risk of falls in older adults? J Am Geriatr Soc 59(5):887–892
28. Angbratt M, Moller M (1999) Questionnaire about calcium intake: can we trust the answers? Osteoporos Int 9(3):220–225
29. Johansson L, Sundh D, Zoulakis M, Rudang R, Darelid A, Brisby EM, Kallmen H, Nilsson AG, Mellstrom D, Lorentzon M (2018) The prevalence of vertebral fractures is associated with reduced hip bone density and inferior peripheral appendicular volumetric bone density and structure in older women. J Bone Miner Res 33(2):250–260
30. kanis JA, ghizer CC (2000) An update on the diagnosis and assessment of osteoporosis with densitometry. Committee of Scientific Advisors, International Osteoporosis Foundation. Osteoporos Int 11(3):192–202. https://doi.org/10.1007/s001980050281
31. Kelly T (1990) Bone mineral reference databases for American men and women. J Bone Miner Res 5:249
32. Looker AC, Wahner HW, Dunn WL, Calvo MS, Harris TB, Heyse SP, Johnston CC Jr, Lindsay R (1998) Updated data on proximal femur bone mineral levels of US adults. Osteoporos Int 8(5):468–489. https://doi.org/10.1007/s001980050093
33. Breslow NE, Day NE (1987) Statistical methods in cancer research. Volume II—the design and analysis of cohort studies. IARC Sci Publ 82:1–406
34. Albertsson-Wikland K, Martensson A, Savendahl L, Niklasson A, Bang P, Dahlgren J, Gustafsson J, Kristrom B, Norgren S, Pehson NG, Oden A (2016) Mortality is not increased in recombinant human growth hormone-treated patients when adjusting for birth characteristics. J Clin Endocrinol Metab 101(5):2149–2159. https://doi.org/10.1210/jc.2015-3951
35. kanis JA, Johnell O, Oden A, Johansson H, McCloskey E (2008) FRAX and the assessment of fracture probability in men and women from the UK. Osteoporos Int 19(4):385–397. https://doi.org/10.1007/s00198-007-0543-5
36. Harrell FJ (2001) General aspects of fitting regression models: regression modeling strategies. Springer Science+Business Media Inc, New York
37. Fine JP, Gray RJ (1999) A proportional hazards model for the subdistribution of a competing risk. J Am Stat Assoc 94(446):496–509
38. kanis JA, Harvey NC, Johansson H, Liu E, Vandeven H, Lorenzton M, Leslie WD, McCloskey E (2003) Five years of FRAX: how has it changed the management of osteoporosis? Aging Clin Exp Res 32:187–196. https://doi.org/10.1007/s00520-003-0232-y
39. kanis JA, Borgstrom F, De Laet C, Johansson H, Johnell O, Jonsson B, Oden A, Zethraeus N, Pfeifer B, Khaltava N (2005) Assessment of fracture risk. Osteoporos Int 16(6):581–589. https://doi.org/10.1007/s00198-004-1780-5
40. kanis JA, Cooper C, Rizzoli R, Reginster JY, Soc SABE, Aspects CE, Soc CSAN (2019) European guidance for the diagnosis and management of osteoporosis in postmenopausal women (vol 30, pg 3, 2019). Osteoporos Int 30:3–44