**Falls predict fractures independently of FRAX probability: A meta-analysis of the Osteoporotic Fractures in Men (MrOS) Study**

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**Abstract**

Although prior falls are a well-established predictor of future fracture, there is currently limited evidence regarding the specific value of falls history in fracture risk assessment, relative to that of other clinical risk factors and BMD measurement. We therefore investigated, across the 3 Osteoporotic Fractures in Men (MrOS) Study cohorts, whether past falls predicted future fracture independently of FRAX®, and whether these associations varied with age and follow-up time. Elderly men were recruited from MrOS Sweden, Hong Kong and USA. Baseline data included falls history (over the preceding 12 months), clinical risk factors, BMD at femoral neck and calculated FRAX probabilities. An extension of Poisson regression was used to investigate the associations between falls, FRAX probability and incident fracture, adjusting for age, time since baseline and cohort in base models; further models were used to investigate interactions with age and follow-up time. Random-effects meta-analysis was used to synthesis the individual country associations. Information on falls and FRAX probability was available for 4365 men in USA (mean age 73.5 years; mean follow-up 10.8 years), 1823 men in Sweden (mean age 75.4 years; mean follow-up 8.7 years), and 1669 men in Hong Kong (mean age 72.4 years; mean follow-up 9.8 years). Rates of past falls were similar at 20%, 16%, and 15% respectively. Across all cohorts, past falls predicted incident fracture at any site [HR: 1.69 (95%CI: 1.49, 1.90)], major osteoporotic fracture (MOF) [HR: 1.56 (95%CI: 1.33, 1.83)] and hip fracture [HR: 1.61 (95%CI: 1.27, 2.05)]. Relationships between past falls and incident fracture remained robust after adjustment for FRAX probability: adjusted HR (95%CI) any fracture: 1.63 (1.45, 1.83); MOF: 1.51 (1.32, 1.73); and hip: 1.54 (1.21, 1.95). In conclusion, past falls predicted incident fracture independently of FRAX probability, confirming the potential value of falls history in fracture risk assessment.

**Key words:** Osteoporosis; epidemiology; FRAX; falls; fracture; interaction

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**Introduction**

Whilst low bone mineral density is a major risk factor for fragility fracture, the majority of such low trauma fracture events occur as a result of a fall from standing height or less([1](#_ENREF_1)). Conversely the number of falls is much greater than the number of consequent fractures with only 5-10% falls in older adults leading to skeletal injury([1](#_ENREF_1)). Interventions aimed at reducing falls have usually been unsuccessful at reducing fractures([2](#_ENREF_2),[3](#_ENREF_3)), probably partly as a consequence of the low falls to injury ratio. Notwithstanding, prior falls have been found to be a risk factor for future fracture in a number of cohorts([4](#_ENREF_4)). With the advent of the FRAX® fracture risk assessment tool, evaluation of an individual’s probability of sustaining a hip or major osteoporotic fracture over a ten-year time period is now readily undertaken using a small number of easily ascertainable clinical risk factors, and BMD if available([5](#_ENREF_5)). FRAX is the most widely used fracture risk assessment tool, incorporated into the majority of assessment guidelines worldwide([6](#_ENREF_6)) but, unlike other tools such as QFracture or the GARVAN calculator([7-9](#_ENREF_7)), does not include falls as a specific input risk factor([4](#_ENREF_4),[5](#_ENREF_5)) due to the inconsistent data across the 12 derivation and 11 validation cohorts([10](#_ENREF_10)). In order for prior falls to be useful in the current context of risk assessment, the associated fracture risk must ideally be independent of FRAX probability and/or BMD. Having demonstrated that the risk of future falls associated with past falls is partly captured by FRAX([11](#_ENREF_11)), we undertook to investigate, across the three Osteoporotic Fractures in Men (MrOS) Study cohorts, whether a history of past falls (in the previous 12 months) independently predicted future fractures, and whether the predictive value varied with follow-up time or age.

**Methods**

*Participants*

Details of the Osteoporotic Fractures in Men (MrOS) International Study have been published previously([12](#_ENREF_12),[13](#_ENREF_13)), but briefly, MrOS is a multicentre study of community-dwelling men aged 65 years or older from three countries, recruited and evaluated using similar criteria. To be eligible for the study, subjects had to be able to walk without aid. In the MrOS Hong Kong Study, 2,000 Chinese men, aged 65–92 years, were enrolled between August 2001 and February 2003([14](#_ENREF_14)). All were Hong Kong residents of Asian ethnicity. Stratified sampling was adopted to ensure that 33% of subjects were included in each of the following age groups: 65–69, 70–74 and ≥75 years. Recruitment notices were placed in housing estates and community centres for the elderly. In the MrOS Sweden Study, 3014 men, aged 69–81 years, were enrolled between October 2001 and December 2004([11](#_ENREF_11),[15](#_ENREF_15)). The cohort comprised men from the cities of Malmo, Gothenburg and Uppsala, identified and recruited using national population registers. More than 99% were of Caucasian ethnicity. The participation rate in the MrOs Sweden Study was 45%. In the MrOS United States Study, 5995 men, aged 65–100 years, were enrolled at 6 sites between March 2000 and April 2002([16](#_ENREF_16),[17](#_ENREF_17)). Each US clinical site designed and customised strategies to enhance recruitment of its population. Common strategies included mailings from the Department of Motor Vehicles, voter registration and participant databases, common senior newspaper features and advertisement and targeted presentations. Self-defined racial/ethnic ancestry was ascertained through questionnaires at baseline.

*Exposure variables*

At baseline, height (centimeters) and weight (kilograms) were measured, and BMI was calculated as kilograms per square meter. The international MrOS questionnaire([16](#_ENREF_16)) was administered at baseline to collect information about current smoking, number and type of medications, fracture history, family history of hip fracture, past medical history (rheumatoid arthritis) and high consumption of alcohol (3 or more glasses of alcohol-containing drinks per day), calculated from the reported frequency and amount of alcohol use. Previous fracture at baseline was documented as all fractures after the age of 50 years, regardless of trauma. For glucocorticoid exposure, this was documented in MrOs as use at least 3 times per week in the month preceding the baseline assessment. Apart from rheumatoid arthritis, there was no information on secondary causes of osteoporosis and the input variable for FRAX probability calculation was set to no for all men. Self-reported falls during the 12 months preceding the baseline were recorded by questionnaire (past falls). Areal bone mineral density (BMD) was measured at the femoral neck using Hologic QDR 4500 A or W (Hologic, Bedford, MA) or Lunar Prodigy (GE Lunar Corp., Madison, WI) depending on the centre, with cross calibration of instruments. A T-score was calculated using NHANES young women as a reference value([18](#_ENREF_18)). 10-year probability of fracture [FRAX major osteoporotic fracture (hip, humerus, vertebral or forearm sites)] was calculated using clinical risk factors described above with and without femoral neck BMD entered into country-specific FRAX models. As the gradients of risk for incident falls were similar with either model, results for the models including femoral neck BMD are presented.

*Fracture and death outcomes*

*Hong Kong(*[*19*](#_ENREF_19)*):* Incident fractures were captured via subject follow-up through phone call or visit to the research centre. All fracture sites (hip, wrist, skull/face, ribs, shoulder, arm, wrist, vertebra, tibia, fibula, foot, metatarsal toes, hand, fingers, and pelvis) were recorded. Pathological fractures were excluded. All incident fractures reported by participants were then confirmed by X-ray or medical record. Deaths were verified by death certificates.

*Sweden(*[*20*](#_ENREF_20)*)*: Central registers covering all Swedish citizens were used to identify the subjects and the time of death for all subjects who died during the study, and these analyses were performed after the time of fracture validation. At the time of fracture evaluation, the computerized X-ray archives in Malmo, Goteborg, and Uppsala were searched for new fractures occurring after the baseline visit using the unique personal registration number allocated to every Swedish citizen. All additional fractures reported by the study subject after the baseline visit were confirmed by physician review of radiology reports. Fractures reported by the study subject but not possible to confirm by radiographic report were not included.

*US(*[*16*](#_ENREF_16)*):* If a participant reported a fracture, study staff conducted a follow-up telephone interview to determine the date and time the fracture occurred, a description of how the fracture occurred, the type of trauma that resulted in the fracture, the participant’s location and activities at the time of the fracture, symptoms just before or coincident with the fracture, and source of medical care for the fracture. All reported fractures were verified by a physician adjudicator through medical records obtained from the participant’s physician. The Clinical Outcomes Committee adjudicated any uncertainties regarding the presence of a fracture. Deaths were verified through state death certificates.

*Statistical methods*

In order to compare the performance of FRAX probability with that of a history of past falls, a dichotomous variable was created such that the percentage of men who had a high fracture risk was similar to the percentage who had had previously fallen (15% for HK, 16% for Sweden and 20% for US). Thus, 15%, 16% and 20% men respectively had a FRAX probability of major osteoporotic fracture, calculated with BMD, above 9.5%, 15.8% and 10.3% and the dichotomised FRAX score was therefore classified as high or low risk. Fracture outcomes considered included: any, osteoporotic (defined according to Kanis et al. 2001([21](#_ENREF_21)) as clinical vertebral, ribs, pelvis, humerus, clavicle, scapula, sternum, hip, other femoral fractures, tibia, fibula, distal forearm/ wrist), major osteoporotic (MOF: hip, clinical vertebral, humerus and wrist/ forearm), osteoporotic fracture without hip fracture (clinical vertebral, humerus and wrist), clinical vertebral, and hip. An extension of Poisson regression models([22](#_ENREF_22)) was used to study the association between FRAX, other risk variables and the future risk of fracture. All associations were adjusted for age and time since baseline. In contrast to logistic regression, the Poisson regression uses the length of each individual’s follow-up period and the hazard function is assumed to be exp(β0 + β1 · current time from baseline + β2 · current age + β3 · variable of interest). The observation period of each participant was divided into intervals of one month. One fracture per person, and time to the first fracture, were counted, and time at risk was censored at the time of first fracture, migration or death. Thus, we investigated the predictive value of prior falls, FRAX (including each individual constituent risk factor), and BMD as individual risk factors, and then in multivariable models to investigate the value of falls independent of FRAX or BMD, and FRAX independent of falls. In further analyses, we explored interactions with age and time since baseline, in which age and time were used as continuous variables and examples given at specific ages and times. Additionally we stratified the analyses by femoral neck BMD T score above or below -2.5. The association between predictive factors and risk of fracture are described as a hazard ratio (HR) or gradient of risk (GR= HR per 1 standard deviation change in predictor in the direction of increased risk). Additionally, we explored the associations between falls and fracture by number of falls reported at baseline (1 vs multiple). Two-sided p-value were used for all analyses and p<0.05 considered to be significant. Analyses were undertaken separately within each cohort and then the β-coefficients from each cohort were weighted according to the variance, and merged to determine the weighted mean of the coefficient and its standard deviation (random-effects meta-analysis). The risk ratios are then given by e(weighted mean coefficient). Whilst there are numerous caveats with the use of Receiver Operator Curve models in this context([23](#_ENREF_23)), we additionally present Area Under the Curve (AUC) values for the predictive models in the Online Supplementary Resources.

**Results**

*Characteristics of participants*

The study cohort consisted of 7857 men who had information on falls, BMD and FRAX risk factors: 4365 men in USA (mean age 73.5 years; mean follow-up 10.8 years); 1823 men in Sweden (mean age 75.4 years; mean follow-up 8.7 years); and 1669 men in Hong Kong (mean age 72.4 years; mean follow-up 9.8 years). Table 1 summarises the baseline characteristics of the individuals by country cohort. Rates of past falls were similar at 20%, 16%, and 15% respectively. Rates of previous fracture were higher in Sweden (33%) than in USA (22%) and Hong Kong (13%). Consistent with the known country-specific epidemiology of fracture, the highest mean FRAX probability was observed in Sweden (11.4% probability of major osteoporotic fracture, calculated with BMD), followed by USA (7.9%) and Hong Kong (6.7%).

*Past falls, FRAX probability and risk of incident fracture*

Table 2 summarises the relationships between past falls or high FRAX probability at baseline, and incident fractures. Online Supplementary Table 1 additionally presents the predictive value of the individual FRAX risk factors, and of falls adjusted for each of these variables. Across all cohorts, past falls predicted any incident fracture [HR: 1.69 (95%CI: 1.49, 1.90)], major osteoporotic fracture (MOF) [HR: 1.56 (95%CI: (1.33, 1.83)] and hip fracture [HR: 1.61 (95%CI: 1.27, 2.05)]. Similar relationships were found for osteoporotic fracture and major osteoporotic fracture without hip fracture, summarised in Table 2. The predictive value of past falls was present within each individual cohort apart from when hip fracture was the outcome, where for Sweden and Hong Kong, the 95% CI included unity. The magnitudes of the gradients of risk were similar in Sweden and US cohorts, and marginally higher in the Hong Kong cohort, albeit with substantially overlapping confidence intervals, and there was thus no statistically significant interaction between falls and centre. For illustrative purposes, Online Supplementary Table 2 demonstrates the fracture incidence amongst the four groups defined by high versus low FRAX probability and falls yes/no; the hazard ratio for major osteoporotic fracture is also given for the remaining three groups relative to the low FRAX probability and no falls groups.

The hazard ratio associated with multiple falls tended to be marginally greater than that associated with a single fall, for example HR 1 fall for MOF = 1.56 (95%CI: 1.33, 1.83) and HR for ≥2 falls = 2.00 (95%CI: 1.35, 2.98). High FRAX probability of major osteoporotic fracture, calculated with BMD, was predictive of all fracture outcomes, with the magnitude of the HR greater than for the equivalent falls-fracture relationships (summarised in Table 2). Thus, across all cohorts, high FRAX predicted any incident fracture [HR: 2.00 (95%CI: 1.73, 2.31)], major osteoporotic fracture (MOF) [HR: 2.35 (95%CI: 1.87, 2.94)] and hip fracture [HR: 2.93 (95%CI: 1.75, 4.88)].

*Independent predictive value of falls, FRAX probability and BMD*

The relationships between past falls and incident fracture remained robust after adjustment for high FRAX probability (MOF): adjusted HR (95%CI) any fracture: 1.63 (1.45, 1.83); MOF: 1.51 (1.29, 1.77); and hip: 1.54 (1.21, 1.95), and for BMD (Table 3). Indeed the hazard ratios and 95% CI were very little altered by adjustment for high FRAX probability or BMD. Similarly, the gradient of risk for fracture outcomes with high FRAX probability were little altered by adjustment for the presence of reported past falls at baseline (Table 3): adjusted HR (95%CI) any fracture: 1.96 (1.69, 2.27); MOF: 2.30 (1.84, 2.88); and hip: 2.86 (1.73, 4.75. The associations with the outcomes of clinical vertebral fracture and osteoporotic fracture without hip fracture (OWH) are documented in Online Supplementary Table 3, demonstrating that both prior falls and FRAX probability were predictive of both outcomes. Online Supplementary Table 4 demonstrates the predictive value of these exposures when femoral neck BMD at baseline is dichotomised above/ below T=-2.5. For illustrative purposes the under the curve (AUC) values for the different prediction models are presented in Online Supplementary Table 5.

*Interactions between past falls, age, follow-up time and risk of incident fracture*

In both Sweden and US there was a tendency for the hazard ratio for fracture associated with past falls to reduce with increasing follow-up time (p interaction = 0.12 and 0.15 respectively). In contrast no decline with time was observed in the Hong Kong (p >0.30). The interaction between past falls and follow-up time became close to statistical significance (p = 0.059) when all 3 cohorts were combined (Figure 1). There was no evidence of an interaction with age. No interactions for either follow-up time or age with high FRAX probability were observed.

**Discussion**

In this large combined population cohort of older men, we have demonstrated that previous falls and high FRAX probability independently predict the risk of future fracture. These findings clearly confirm the value of falls in fracture risk assessment, and demonstrate that consideration of past falls yields information over and above that captured by the FRAX algorithm.

The predictive value of past falls for future fracture is well established([24](#_ENREF_24)), but the present study, to our knowledge, provides the first evidence from a large population-based cohort that this risk is independent of that captured by FRAX with or without BMD. It complements our previous findings, from the MrOS Sweden cohort, of similar predictive value of past falls and FRAX probability for future falls([11](#_ENREF_11)), extending this to the key musculoskeletal consequence, namely fracture. Similar to the present study, although risk factors for falls and fracture overlap substantially, and many of which are captured in the FRAX tool, the magnitude of the predictive value of past falls or high FRAX probability was not materially altered by mutual adjustment, indicating that falls history is likely to inform risk not captured by FRAX probability. Interestingly, prior falls predicted incident clinical vertebral fracture as well as the other fracture types. Whilst vertebral fractures in women have largely been thought to result from action such as lifting, rather than from falls([25](#_ENREF_25)), data from the US MrOS cohort suggested that falls were common antecedents of clinical presentation with a vertebral fracture amongst older men([26](#_ENREF_26)).

These findings support the notion that consideration of falls history is likely to add usefully to risk assessment based on the FRAX tool, and as such will be of relevance to a large number of guidelines globally([6](#_ENREF_6)). Whilst falls have been incorporated into risk calculators derived from single cohorts in which these outcomes have been recorded([7-9](#_ENREF_7),[27](#_ENREF_27),[28](#_ENREF_28)), the lack of standardised documentation of falls events across the 23 cohorts used in the development and validation of the FRAX tool has meant that the use of prior falls as a clinical risk factor was not possible([4](#_ENREF_4)). A further consideration is that FRAX input variables were selected on the basis of at least partial independence of BMD, and of constituting a risk amenable to pharmacological therapeutic intervention. Whilst our present findings strongly support the first of these criteria, there is still limited evidence that interventions to reduce falls will also reduce fractures([2-4](#_ENREF_2),[29-34](#_ENREF_29)), or that falls risk is amenable to intervention with pharmacological agents such as bisphosphonates([4](#_ENREF_4),[35](#_ENREF_35)). In one study, baseline risk of falling was not associated with differences in anti-fracture efficacy of clodronate([36](#_ENREF_36)), suggesting efficacy in fallers and non-fallers alike. In contrast, in a trial of risedronate in elderly women selected partly on the basis of high falls risk, the intervention did not lead to statistically significant reductions in fractures([37](#_ENREF_37)).

Recognising the limitations of falls data in the current FRAX cohorts, a report of an International Society for Clinical Densitometry/ International Osteoporosis Foundation Task Force recommended that FRAX probability may be modified to account for a history of prior falls, with the output inflated by 30% (multiplied by 1.3) for each past fall (for up to 5 falls)([4](#_ENREF_4)). This recommendation is based on the univariate hazard ratio (95%CI: 1.1, 1.5) for incident hip fracture associated with a past fall, derived from the Study of Osteoporotic Fractures([38](#_ENREF_38)). Notably in this cohort the fall-fracture relationship became statistically non-significant after adjustment for poor health and markers of poor mobility; furthermore, this study did not investigate the predictive power of falls independent of other clinical risk factors or BMD. Whilst the exact approach to incorporation of prior falls into risk assessment remains to be elucidated, out findings inform clinical care, demonstrating that prior falls indicate increased fracture risk over and above that generated by use of other clinical risk factors and BMD in FRAX. Notwithstanding the inconclusive evidence relating falls interventions to fracture reduction, falls risk should clearly be addressed as part of the risk assessment, in addition to measures specifically aimed at improving bone mineral density.

Our findings of potential time interactions for past falls and incident fractures are intriguing, and echo our previous observation that the predictive value of past falls for incident falls in the MrOS Sweden cohort also waned with increasing follow-up time, and was greater at younger ages([11](#_ENREF_11)). Falls-related risk factors were found to be predictive of fracture risk over a two-year period in a recent US study, but as this investigation did not compare the short-term relationships with those over a longer time period, it is difficult to draw firm conclusions regarding any temporal variation in effect size with regard to falls and follow-up([39](#_ENREF_39)). Whilst it seems intuitively reasonable that falls might mark out particularly unusual individuals relative to the general population at younger ages, where falls overall are less common, it is perhaps counter-intuitive that past falls become less predictive of incident falls with time. It is possible that fallers tend to fracture earlier and become less mobile, and perhaps less exposed to falls risk and thus to fracture risk with time. For the moment, however, particularly since the association just failed to reach statistical significance, and is inconsistent across the 3 cohorts, this observation remains of interest, but requires replication in other populations.

We studied three well-characterized cohorts drawn from general populations with standardized assessments and prospective recording of fractures. However, there are some limitations that should be considered in the interpretation of our findings([16](#_ENREF_16)). Firstly, the population studied was male, and of a modest age range (64-99 years), so limiting generalizability of our findings. Secondly, the definition of glucocorticoid use differed from those usually specified for incorporation into FRAX. Thirdly there was no information on causes of secondary osteoporosis, and this variable was therefore set to missing. The effect of these considerations on our findings is uncertain, but may have led to an overall underestimation of risk by FRAX. Finally, we did not have information on the severity of a past fall, or whether a past fall was associated with injury, so limiting our ability to identify events potentially most likely to be associated with a fracture outcome.

In conclusion, we have demonstrated that prior falls are a risk factor for incident fracture, independently of FRAX probability calculated with or without BMD. Whilst our findings clearly demonstrate the value of falls history in fracture risk assessment, further prospective studies in cohorts with wider age ranges, other ethnicities, and most importantly women, are now warranted to replicate and extend these findings, ideally to establish the potential for inclusion of falls as a modifier of FRAX probability.

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**Author Roles**

All authors contributed to manuscript drafting, review and finalisation. NCH wrote the first draft of the manuscript and oversaw its preparation; HJ and AO undertook statistical analysis; EO and JL designed and implemented MrOS US, and provided data; MK, BR, OL, CO, DM designed and implemented MrOS Sweden, and provided data; TK designed and implemented MrOS Hong Kong, and provided data; CC contributed expertise on fracture epidemiology; EM and JAK oversee FRAX and provided FRAX methodology; EVM is guarantor.

**Disclosures**

All authors have no disclosures in relation to this manuscript.

**References**

1. Masud T, Morris RO 2001 Epidemiology of falls. Age Ageing **30 Suppl 4:**3-7.

2. Cameron ID, Gillespie LD, Robertson MC, Murray GR, Hill KD, Cumming RG, Kerse N 2012 Interventions for preventing falls in older people in care facilities and hospitals. Cochrane Database Syst Rev **12:**CD005465.

3. Gillespie LD, Robertson MC, Gillespie WJ, Sherrington C, Gates S, Clemson LM, Lamb SE 2012 Interventions for preventing falls in older people living in the community. Cochrane Database Syst Rev **9:**CD007146.

4. Masud T, Binkley N, Boonen S, Hannan MT 2011 Official Positions for FRAX(R) clinical regarding falls and frailty: can falls and frailty be used in FRAX(R)? From Joint Official Positions Development Conference of the International Society for Clinical Densitometry and International Osteoporosis Foundation on FRAX(R). J Clin Densitom **14**(3)**:**194-204.

5. Kanis JA, Hans D, Cooper C, Baim S, Bilezikian JP, Binkley N, Cauley JA, Compston JE, Dawson-Hughes B, El-Hajj FG, Johansson H, Leslie WD, Lewiecki EM, Luckey M, Oden A, Papapoulos SE, Poiana C, Rizzoli R, Wahl DA, McCloskey EV 2011 Interpretation and use of FRAX in clinical practice. Osteoporos.Int. **22**(9)**:**2395-2411.

6. Kanis JA, Harvey NC, Cooper C, Johansson H, Oden A, McCloskey EV 2016 A systematic review of intervention thresholds based on FRAX : A report prepared for the National Osteoporosis Guideline Group and the International Osteoporosis Foundation. Arch Osteoporos **11**(1)**:**25.

7. Hippisley-Cox J, Coupland C 2009 Predicting risk of osteoporotic fracture in men and women in England and Wales: prospective derivation and validation of QFractureScores. BMJ **339:**b4229.

8. Nguyen ND, Frost SA, Center JR, Eisman JA, Nguyen TV 2007 Development of a nomogram for individualizing hip fracture risk in men and women. Osteoporos Int **18**(8)**:**1109-17.

9. Nguyen ND, Frost SA, Center JR, Eisman JA, Nguyen TV 2008 Development of prognostic nomograms for individualizing 5-year and 10-year fracture risks. Osteoporos Int **19**(10)**:**1431-44.

10. McCloskey EV, Kanis JA, Oden A, Johansson H, Diez Perez A, Eisman JA, Nguyen TV, Center JR, Hans D, Krieg MA, Prior JC 2012 A meta-analysis of the association between falls and hip fracture risk. . Osteoporos Int **23**(Suppl. 2)**:**S80-S81.

11. Harvey NC, Johansson H, Oden A, Karlsson MK, Rosengren BE, Ljunggren O, Cooper C, McCloskey E, Kanis JA, Ohlsson C, Mellstrom D 2016 FRAX predicts incident falls in elderly men: findings from MrOs Sweden. Osteoporos Int **27**(1)**:**267-74.

12. Karlsson MK, Ribom E, Nilsson JA, Ljunggren O, Ohlsson C, Mellstrom D, Lorentzon M, Mallmin H, Stefanick M, Lapidus J, Leung PC, Kwok A, Barrett-Connor E, Orwoll E, Rosengren BE 2012 Inferior physical performance tests in 10,998 men in the MrOS study is associated with recurrent falls. Age Ageing **41**(6)**:**740-6.

13. Rosengren BE, Ribom EL, Nilsson JA, Mallmin H, Ljunggren O, Ohlsson C, Mellstrom D, Lorentzon M, Stefanick M, Lapidus J, Leung PC, Kwok A, Barrett-Connor E, Orwoll E, Karlsson MK 2012 Inferior physical performance test results of 10,998 men in the MrOS Study is associated with high fracture risk. Age Ageing **41**(3)**:**339-44.

14. Lau EM, Leung PC, Kwok T, Woo J, Lynn H, Orwoll E, Cummings S, Cauley J 2006 The determinants of bone mineral density in Chinese men--results from Mr. Os (Hong Kong), the first cohort study on osteoporosis in Asian men. Osteoporos.Int. **17**(2)**:**297-303.

15. Mellstrom D, Johnell O, Ljunggren O, Eriksson AL, Lorentzon M, Mallmin H, Holmberg A, Redlund-Johnell I, Orwoll E, Ohlsson C 2006 Free testosterone is an independent predictor of BMD and prevalent fractures in elderly men: MrOS Sweden. J Bone Miner Res **21**(4)**:**529-35.

16. Orwoll E, Blank JB, Barrett-Connor E, Cauley J, Cummings S, Ensrud K, Lewis C, Cawthon PM, Marcus R, Marshall LM, McGowan J, Phipps K, Sherman S, Stefanick ML, Stone K 2005 Design and baseline characteristics of the osteoporotic fractures in men (MrOS) study--a large observational study of the determinants of fracture in older men. Contemp Clin Trials **26**(5)**:**569-85.

17. Blank JB, Cawthon PM, Carrion-Petersen ML, Harper L, Johnson JP, Mitson E, Delay RR 2005 Overview of recruitment for the osteoporotic fractures in men study (MrOS). Contemp Clin Trials **26**(5)**:**557-68.

18. 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)**:**468-89.

19. Kwok T, Khoo CC, Leung J, Kwok A, Qin L, Woo J, Leung PC 2012 Predictive values of calcaneal quantitative ultrasound and dual energy X ray absorptiometry for non-vertebral fracture in older men: results from the MrOS study (Hong Kong). Osteoporos Int **23**(3)**:**1001-6.

20. Ohlsson C, Mellstrom D, Carlzon D, Orwoll E, Ljunggren O, Karlsson MK, Vandenput L 2011 Older men with low serum IGF-1 have an increased risk of incident fractures: the MrOS Sweden study. J Bone Miner Res **26**(4)**:**865-72.

21. Sims I 2017 Many more eligible for bisphosphonates after NICE lowers threshold to 1%, vol. 2017. PULSE.

22. Breslow NE, Day NE 1987 Statistical Methods in Cancer Research. IARC Scientific Publications No 32 **Volume II:**p 131-135.

23. Compston J, Cooper A, Cooper C, Gittoes N, Gregson C, Harvey N, Hope S, Kanis JA, McCloskey EV, Poole KES, Reid DM, Selby P, Thompson F, Thurston A, Vine N 2017 UK clinical guideline for the prevention and treatment of osteoporosis. Arch Osteoporos **12**(1)**:**43.

24. Blain H, Masud T, Dargent-Molina P, Martin FC, Rosendahl E, van der Velde N, Bousquet J, Benetos A, Cooper C, Kanis JA, Reginster JY, Rizzoli R, Cortet B, Barbagallo M, Dreinhofer KE, Vellas B, Maggi S, Strandberg T 2016 A Comprehensive Fracture Prevention Strategy in Older Adults: The European Union Geriatric Medicine Society (EUGMS) Statement. J Nutr Health Aging **20**(6)**:**647-52.

25. Harvey N, Dennison E, Cooper C 2013 The Epidemiology of Osteoporotic Fractures Primer on the Metabolic Bone Diseases and Disorders of Mineral Metabolism. John Wiley & Sons, Inc., pp 348-356.

26. Freitas SS, Barrett-Connor E, Ensrud KE, Fink HA, Bauer DC, Cawthon PM, Lambert LC, Orwoll ES 2008 Rate and circumstances of clinical vertebral fractures in older men. Osteoporos Int **19**(5)**:**615-23.

27. Sambrook PN, Flahive J, Hooven FH, Boonen S, Chapurlat R, Lindsay R, Nguyen TV, Diez-Perez A, Pfeilschifter J, Greenspan SL, Hosmer D, Netelenbos JC, Adachi JD, Watts NB, Cooper C, Roux C, Rossini M, Siris ES, Silverman S, Saag KG, Compston JE, LaCroix A, Gehlbach S 2011 Predicting fractures in an international cohort using risk factor algorithms without BMD. J Bone Miner Res **26**(11)**:**2770-7.

28. Cooper C, Harvey NC 2012 Osteoporosis risk assessment. BMJ **344:**e4191.

29. Miake-Lye IM, Hempel S, Ganz DA, Shekelle PG 2013 Inpatient fall prevention programs as a patient safety strategy: a systematic review. Ann Intern Med **158**(5 Pt 2)**:**390-6.

30. Martin JT, Wolf A, Moore JL, Rolenz E, DiNinno A, Reneker JC 2013 The effectiveness of physical therapist-administered group-based exercise on fall prevention: a systematic review of randomized controlled trials. J Geriatr Phys Ther **36**(4)**:**182-93.

31. Karlsson MK, Magnusson H, von Schewelov T, Rosengren BE 2013 Prevention of falls in the elderly--a review. Osteoporos Int **24**(3)**:**747-62.

32. Pfortmueller CA, Lindner G, Exadaktylos AK 2014 Reducing fall risk in the elderly: risk factors and fall prevention, a systematic review. Minerva Med **105**(4)**:**275-81.

33. Lovarini M, Clemson L, Dean C 2013 Sustainability of community-based fall prevention programs: a systematic review. J Safety Res **47:**9-17.

34. Gillespie LD, Robertson MC, Gillespie WJ, Lamb SE, Gates S, Cumming RG, Rowe BH 2009 Interventions for preventing falls in older people living in the community. Cochrane Database Syst Rev (2)**:**CD007146.

35. Kanis JA, Hans D, Cooper C, Baim S, Bilezikian JP, Binkley N, Cauley JA, Compston JE, Dawson-Hughes B, El-Hajj Fuleihan G, Johansson H, Leslie WD, Lewiecki EM, Luckey M, Oden A, Papapoulos SE, Poiana C, Rizzoli R, Wahl DA, McCloskey EV 2011 Interpretation and use of FRAX in clinical practice. Osteoporos Int **22**(9)**:**2395-411.

36. Kayan K, Johansson H, Oden A, Vasireddy S, Pande K, Orgee J, Kanis JA, McCloskey EV 2009 Can fall risk be incorporated into fracture risk assessment algorithms: a pilot study of responsiveness to clodronate. Osteoporos Int **20**(12)**:**2055-61.

37. McClung MR, Geusens P, Miller PD, Zippel H, Bensen WG, Roux C, Adami S, Fogelman I, Diamond T, Eastell R, Meunier PJ, Reginster JY 2001 Effect of risedronate on the risk of hip fracture in elderly women. Hip Intervention Program Study Group. N Engl J Med **344**(5)**:**333-40.

38. Cummings SR, Nevitt MC, Browner WS, Stone K, Fox KM, Ensrud KE, Cauley J, Black D, Vogt TM 1995 Risk factors for hip fracture in white women. Study of Osteoporotic Fractures Research Group. N Engl J Med **332**(12)**:**767-73.

39. NICE 2008 TA161: Alendronate, etidronate, risedronate, raloxifene, strontium ranelate and teriparatide for the secondary prevention of osteoporotic fragility fractures in postmenopausal women. National Institute for Health and Care Excellence, London.

**Figure legends**

**Figure 1:** Interaction between past falls and follow-up time, and risk of any incident fracture.

**Table 1:** Baseline characteristics of MrOS participants by country cohort.

|  |  |  |  |
| --- | --- | --- | --- |
|  | **Hong Kong** | **Sweden** | **USA** |
| Proportion of whole cohort | 83% | 61% | 73% |
| n | 1669 | 1823 | 4365 |
| Person-years | 16423 | 15878 | 47044 |
| Age [mean (range)], years | 72.4 (65-91) | 75.4 (70-81) | 73.5 (64-99) |
| BMI | 23.5±3.2 | 26.3±3.6 | 27.42±3.9 |
| Previous fracture | 13% | 33% | 22% |
| Family history hip fracture | 5% | 13% | 17% |
| Smoker | 12% | 8% | 3% |
| Steroid | 1% | 2% | 2% |
| Rheumatoid arthritis | 1% | 1% | 5% |
| Excess alcohol | 1% | 3% | 4% |
| BMD FN T-score | -1.4±0.9 | -0.9±1.0 | -0.6±1.1 |
| Fall at BL | 15% | 16% | 20% |
| No falls at BL |  |  |  |
|  0: 0 times  | 1426 (85%) | 1538 (84%) | 3478 (80%) |
|  1: 1 time | 192 (12%) | 162 (9%) | 519 (12%) |
|  2: 2-3 times | 42 (3%) | 85 (5%) | 305 (7%) |
|  3: 4-5 times | 6 (0.4%) | 14 (0.8%) | 41 (0.9%) |
|  4: 6+ times | 3 (0.2%) | 13 (0.7%) | 22 (0.5%) |
| Mean±SD |  |  |  |
| FRAX MOF without BMD | 6.9±2.9 | 13.5±6.2 | 9.2±5.0 |
| FRAX hip without BMD | 3.4±2.6 | 7.5±5.5 | 3.7±4.0 |
| FRAX MOF with BMD | 6.7±3.3 | 11.4±6.8 | 7.9±4.8 |
| FRAX hip with BMD | 3.1±2.7 | 5.6±6.1 | 2.5±3.6 |
| High FRAX (ost with BMD) | 15% | 16% | 20% |
| Threshold for high FRAX (%) | 9.50 | 14.00 | 10.30 |
|  |  |  |  |
| FU (hip fx: mean (SD), years | 9.8 (2.9) | 8.7 (2.8) | 10.8 (3.8) |
| Any fx | 11% | 23% | 19% |
| Osteoporotic fx | 9% | 19% | 14% |
| MOF fx | 7% | 16% | 10% |
| Hip fx | 3% | 7% | 4% |

Fx=fracture; Ost=osteoporotic; MOF=Major Osteoporotic Fracture

**Table 2:** Relationships between past falls, FRAX and risk of new fracture. Data are hazard ratio (95%CI) adjusted for age and time since baseline. HK=Hong Kong; SW=Sweden; US=United States of America.

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
|  |  | **Any fx** | **Ost fx** | **MOF** | **Hip fx** |
| Falls at baseline | HK | 1.93 (1.38, 2.70) | 1.83 (1.25, 2.68) | 2.01 (1.32, 3.05) | 1.71 (0.92, 3.21) |
|  | SW | 1.61 (1.27, 2.03) | 1.50 (1.16, 1.94) | 1.50 (1.13, 1.98) | 1.34 (0.85, 2.09) |
|  | US | 1.67 (1.43, 1.94) | 1.54 (1.29, 1.84) | 1.50 (1.21, 1.86) | 1.74 (1.27, 2.38) |
|  | Total | 1.69 (1.49, 1.90) | 1.56 (1.36, 1.79) | 1.56 (1.33, 1.83) | 1.61 (1.27, 2.05) |
|  |  |  |  |  |  |
| High FRAX (MOF with BMD) | HK | 2.45 (1.78, 3.38) | 3.04 (2.14, 4.32) | 3.20 (2.17, 4.72) | 5.27 (3.07, 9.05) |
| SW | 1.76 (1.40, 2.21) | 1.83 (1.43, 2.34) | 1.98 (1.52, 2.57) | 1.82 (1.21, 2.74) |
| US | 2.01 (1.74, 2.33) | 2.13 (1.80, 2.52) | 2.29 (1.87, 2.79) | 2.84 (2.11, 3.81) |
|  | Total | 2.00 (1.73, 2.31) | 2.21 (1.75, 2.79) | 2.35 (1.87, 2.94) | 2.93 (1.75, 4.88) |

Fx=fracture; Ost=osteoporotic; MOF=Major Osteoporotic Fracture

**Table 3:** Past falls adjusted for FRAX probability, and FRAX probability adjusted for past falls, as predictors of incidence fracture. Data are hazard ratio (95%CI) adjusted for age and time since baseline. HK=Hong Kong; SW=Sweden; US=United States of America.

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
|  |  | **Any fx** | **Ost fx** | **MOF** | **Hip fx** |
| Falls at baseline adjusted for FRAX | HK | 1.87 (1.34, 2.62) | 1.76 (1.20, 2.59) | 1.94 (1.28, 2.96) | 1.47 (0.78, 2.78) |
| SW | 1.56 (1.23, 1.97) | 1.45 (1.12, 1.88) | 1.44 (1.09, 1.90) | 1.29 (0.82, 2.01) |
| US | 1.61 (1.39, 1.88) | 1.49 (1.25, 1.78) | 1.45 (1.17, 1.80) | 1.69 (1.23, 2.31) |
| Total | 1.63 (1.45, 1.83) | 1.51 (1.32, 1.73) | 1.51 (1.29, 1.77) | 1.54 (1.21, 1.95) |
|  |  |  |  |  |  |
| Falls at baseline adjusted for femoral neck BMD | HK | 1.92 (1.38, 2.69) | 1.82 (1.24, 2.67) | 2.00 (1.31, 3.03) | 1.68 (0.89, 3.14) |
| SW | 1.64 (1.29, 2.07) | 1.52 (1.17, 1.96) | 1.50 (1.14, 1.99) | 1.31 (0.84, 2.05) |
| US | 1.69 (1.45, 1.96) | 1.56 (1.31, 1.86) | 1.54 (1.24, 1.90) | 1.82 (1.33, 2.48) |
| Total | 1.71 (1.51, 1.92) | 1.58 (1.38, 1.81) | 1.58 (1.35, 1.85) | 1.64 (1.29, 2.08) |
|  |  |  |  |  |  |
| High FRAX (MOF with BMD) adjusted for falls | HK | 2.41 (1.74, 3.33) | 3.00 (2.11, 4.26) | 3.15 (2.13, 4.65) | 5.13 (2.98, 8.85) |
| SW | 1.72 (1.36, 2.16) | 1.80 (1.41, 2.30) | 1.94 (1.49, 2.52) | 1.79 (1.19, 2.71) |
| US | 1.97 (1.70, 2.28) | 2.10 (1.78, 2.48) | 2.25 (1.85, 2.75) | 2.79 (2.08, 3.75) |
| Total | 1.96 (1.69, 2.27) | 2.17 (1.72, 2.74) | 2.30 (1.84, 2.88) | 2.86 (1.73, 4.75) |

Fx=fracture; Ost=osteoporotic; MOF=Major Osteoporotic Fracture