Original Article

High Sensitive CRP is an independent risk factor for all fractures and vertebral fractures in elderly men: The MrOS Sweden Study†

Anna L Eriksson, MD, PhD1, Sofia Movérare-Skrtic, MMSc, PhD1, Östen Ljunggren, MD, PhD2, Magnus Karlsson, MD, PhD3, Dan Mellström, MD, PhD1, Claes Ohlsson, MD, PhD1*

1 Centre for Bone and Arthritis Research, Institute of Medicine, Sahlgrenska University Hospital, Gothenburg, Sweden
2 Department of Medical Sciences, Uppsala University Hospital, Uppsala, Sweden
3 Clinical and Molecular Osteoporosis Research Unit, Department of Clinical Sciences, Lund University and Department of Orthopaedics, Malmö University Hospital, Sweden

* Address for correspondence and reprint requests:
Prof Claes Ohlsson,
Centre for Bone and Arthritis Research
Vita Stråket 11, Sahlgrenska University Hospital
SE-413 45 Gothenburg
Sweden
Phone: +46-31-342 2873
e-mail: claes.ohlsson@medic.gu.se

Disclosures: All other authors state that they have no conflicts of interest.

Keywords: AGING, Osteoporosis < DISEASES AND DISORDERS OF/RELATED TO BONE, General population studies < EPIDEMIOLOGY, Fracture risk assessment < PRACTICE/POLICY-RELATED ISSUES

†This article has been accepted for publication and undergone full peer review but has not been through the copyediting, typesetting, pagination and proofreading process, which may lead to differences between this version and the Version of Record. Please cite this article as doi: [10.1002/jbmr.2037]
Abstract

Epidemiological studies have shown low grade inflammation measured by high sensitivity C-reactive protein (hs-CRP) to be associated with fracture risk in women. However, it is still unclear whether hs-CRP is associated with fracture risk also in men. We therefore measured serum levels of hs-CRP in 2910 men, mean age 75 years, included in the prospective population-based MrOS Sweden cohort. Study participants were divided into tertile groups based on hs-CRP level. Fractures occurring after the baseline visit were validated (average follow-up 5.4 years). The incidence for having at least one fracture after baseline was 23.9/1000 person-years. In cox proportional hazard regression analyses adjusted for age, hs-CRP was related to fracture risk. The HR of fracture for the highest tertile of hs-CRP, compared with the lowest and the medium tertiles combined, was 1.48 (95% CI 1.20-1.82). Multivariate adjustment for other risk factors for fractures had no major effect on the associations between hs-CRP and fracture. Results were essentially unchanged after exclusion of subjects with hs-CRP levels greater than 7.5 mg/L, as well as after exclusion of subjects with a first fracture within three years of follow-up, supporting that the associations between hs-CRP and fracture risk were not merely a reflection of a poor health status at the time of serum sampling. Femoral neck BMD was not associated with hs-CRP, and the predictive role of hs-CRP for fracture risk was essentially unchanged when femoral neck BMD was added to the model (HR 1.37, 95% CI 1.09-1.72). Exploratory sub-analyses of fracture type demonstrated that hs-CRP was clearly associated with clinical vertebral fractures (HR 1.61, 95% CI 1.12-2.29).

We demonstrate, using a large prospective population-based study, that elderly men with high hs-CRP have increased risk of fractures, and that these fractures are mainly vertebral. The association between hs-CRP and fractures was independent of BMD.
**Introduction**

Osteoporosis-related fractures constitute a major health concern not only in women but also in men. In Sweden the lifetime risk of a hip- spine- or forearm fracture, which are common osteoporosis related fractures, at the age of 50 is 46 % for women and 22 % for men (1). These fractures are associated with increased morbidity and mortality, and pose a substantial burden to the individual, the health care system and society in general (2,3). It is thus important to elucidate the pathogenesis of osteoporosis in men, in whom it has been less extensively studied than in women, to aid in prevention and treatment. Risk factors for fractures in men that we know of today include poor neuromuscular function, age, physical inactivity, low levels of estrogens, low BMD, and bone geometry (4).

It is well known that chronic inflammatory diseases such as rheumatoid arthritis, inflammatory bowel disease and chronic obstructive pulmonary disease (COPD) are associated with bone loss and an enhanced fracture risk (5-9), indicating a relationship between the immune system and bone. It has also been shown that certain proinflammatory cytokines are involved in the pathogenesis of osteoporosis (10,11). Moreover, the detrimental effects of estrogen deficiency on bone are to a certain extent mediated via these cytokines (12). Chronic low grade inflammation is involved in the pathophysiology of a large number of conditions including dementia (13), cardiovascular disease (14) and diabetes mellitus (15). Lately the question has arisen as to whether also low-grade inflammation, as estimated by high sensitivity C-reactive protein (hs-CRP), is a risk factor for fractures. There is indeed epidemiological data supporting an association between hs-CRP and fracture risk (16-19). However, these studies have been conducted either in women only, or in mixed cohorts where the number of men with fractures has been too low to clarify the relationship between hs-CRP level and fracture risk in men specifically.
We hypothesized that there is an association between hs-CRP and fracture risk also in men and that such an association can be found if a large study, well powered enough for the purpose, is used. We therefore investigated associations between hs-CRP and fracture risk in MrOS Sweden, which is a large prospective population-based cohort of elderly Swedish men.
Materials and Methods

Study subjects

The Osteoporotic Fractures in Men (MrOS) study is a multicenter, prospective study including older men in Sweden (3014), Hong Kong (~2000), and the United States (~6000). In the present study, associations between serum hs-CRP and fractures that occurred after the baseline visit were investigated in the Swedish cohort (Table 1), which consists of three subcohorts from three different Swedish cities (n = 1005 in Malmö, n = 1010 in Göteborg, and n = 999 in Uppsala). Study subjects (men aged 69 to 81 years) were randomly identified using national population registers, contacted, and asked to participate. To be eligible for the study, the subjects had to be able to walk without assistance, provide self-reported data, and sign an informed consent; there were no other exclusion criteria (20). The study was approved by the ethics committees at the Universities of Göteborg, Lund, and Uppsala. Informed consent was obtained from all study participants.

Assessment of covariates

We used a standardized questionnaire to gather information about self-reported previous fractures after 50 years of age, amount of physical activity, nutritional intake, smoking, use of alcohol, prevalent major diseases (eg, diabetes, stroke, chronic obstructive pulmonary disease (COPD), cancer and rheumatoid arthritis), and medication use (Table 1). Physical activity was the subject's average total daily walking distance, including both walking as a means of exercise and leisure and as a means of outdoor transportation in activities of daily life. Calcium intake was calculated using information from the questionnaires. Use of alcohol was expressed as 3 or more glasses of alcohol-containing drinks per day, calculated from the reported frequency and amount of alcohol use. Grip strength was analyzed using Baseline equipment (Baseline, Chattanooga, TN, USA), and the
average of two consecutive measurements was used in the analyses. Standard equipment was used to measure height and weight.

Assessment of incident fractures

Participants were followed for 5.4 yr on average after the baseline examination. The follow-up time was recorded from the date of the baseline visit to the date of the first fracture or the date of death. When a subject sustained a first fracture at different sites during the follow-up, the various fractures and the follow-up time for each respective first fracture type were included in the analyses. 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 Malmö, Göteborg, and Uppsala were searched for new fractures occurring after the baseline visit, using the unique personal registration number, which all Swedish citizens have. All validated fractures were included in the main analyses, followed by exploratory subanalyses of fracture type. In the latter, we studied the associations between hs-CRP and validated fractures, divided into three main groups: (1) X-ray–verified clinical vertebral fractures, (2) nonvertebral osteoporosis fractures at the major osteoporosis-related locations (defined as hip, distal radius, proximal humerus, and pelvis), and (3) hip fractures (Table 1). Fracture rates were expressed as the number of subjects with first fractures per 1000 person-years (Table 1).
Assessment of BMD (DXA)

Areal BMD (aBMD, g/cm²) of the femoral neck was assessed using the Lunar Prodigy DXA (n = 2004 from the Uppsala and Malmö cohorts; GE Lunar Corp., Madison, WI, USA) or Hologic QDR 4500/A-Delphi (n = 1010 from the Göteborg cohort; Hologic, Waltham, MA, USA). The CVs for the aBMD measurements ranged from 0.5% to 3%. To be able to use DXA measurements performed with equipment from two different manufacturers, a standardized BMD (sBMD) was calculated, as previously described (20).

Serum analyses

C-reactive protein (CRP) was measured by an ultrasensitive particle enhanced immunoturbidimetric assay (Orion Diagnostica, Espoo, Finland). The analyses were performed on a Konelab 20 autoanalyzer (Thermo Fisher Scientific), with a sensitivity of 0.1 mg/L. Inter-assay CV for the Konelab analyses was below 5%.

Statistical analyses

The characteristics of the subjects were compared by tertiles of CRP using linear regression for continuous variables and chi-square tests for categorical variables. Non-normally distributed variables were log transformed.

Associations among variables were examined by Pearson's correlation. Cox proportional hazards models were used to study the associations between hs-CRP and fracture outcomes. The proportional hazard assumption for the main analysis as well as for exploratory sub-analyses was assessed by the graphical log-log method using stphplot in Stata 12. Tertiles of hs-CRP were used in the primary analysis, and in subsequent analyses tertiles 1 and 2 were pooled into one group. Age-adjusted hazard ratios (HRs), with 95% CIs within parentheses, versus the combined group of tertiles 1 and 2, were calculated. Further adjustments for height, weight, calcium intake, physical activity, grip strength, cigarette smoking, use of alcohol, COPD, stroke, diabetes, cancer, rheumatoid arthritis and prevalent fractures and medication use (corticosteroids, statins, thiazide...
diuretics, NSAIDs, osteoporosis medications, testosterone, antidepressants, hypnotics and sedatives, antiandrogens) were made to investigate the independent effects of hs-CRP on fracture outcome. Finally, adjustments were made also for femoral neck BMD. All validated fractures were included in the main analyses, followed by exploratory sub-analyses of fracture type.
Results

Characteristics of the study subjects
Baseline characteristics of the study subjects, including the number and incidence/1000 person-years of validated fractures having occurred after the baseline visit of the older men in the MrOS Sweden cohort, are shown in Table 1. In total, 377 subjects had at least one validated incident fracture, and the average follow-up time of the 2910 subjects was 5.4 years. Study participants were divided into tertile groups based on hs-CRP level. Serum levels of hs-CRP were [<1.81], [≥ 1.81, < 2.76] and [≥ 2.76] mg/L in the lowest, medium and highest tertiles, respectively. CRP was directly associated with baseline weight, BMI, proportion smokers and prevalence of COPD, and inversely associated with grip strength (Table 1).

Hs-CRP as a predictor of fractures
Cox proportional-hazards models demonstrated that high serum hs-CRP was related to an increased risk of first fracture (age adjusted HR 1.25, 95% CI 1.10-1.42 per tertile increase in hs-CRP). The fracture risk for subjects in the highest CRP tertile was clearly increased compared with subjects in the lowest tertile (HR 1.54, 95% CI 1.20-1.97) while the fracture risk was similar in the medium CRP tertile compared with the lowest CRP tertile (HR 1.10 95% CI 0.85-1.43). Therefore the lowest and the medium tertiles were pooled into one group in the subsequent analyses. The HR of fracture for the highest tertile of hs-CRP, compared with the lowest and the medium tertiles combined, was 1.48 (95 % CI, 1.20-1.82, Figure 1a).

Exploratory sub-analyses of fracture type demonstrated that hs-CRP was clearly associated with clinical vertebral fractures (HR 1.61, 95% CI 1.12-2.29, Tertile 3 vs Tertiles 1+2). There was no significant association between hs-CRP and hip fractures and non-vertebral osteoporosis related fractures respectively (Figure 1a).
**Hs-CRP as an independent predictor of fractures**

Multivariate adjustment for other risk factors for fractures (age, height, weight, calcium intake, physical activity, grip strength, cigarette smoking, use of alcohol, COPD, stroke, diabetes, cancer, rheumatoid arthritis, prevalent fractures and medication use) had no major effect on the associations between hs-CRP and fracture (Figure 1b).

Femoral neck BMD was not significantly associated with hs-CRP (p for trend 0.09, Table 1), and the predictive role of hs-CRP for fracture risk was essentially unchanged after adjustment also for femoral neck BMD (HR 1.37, 95% CI 1.09-1.72, Tertile 3 vs Tertile 1+2) (Figure 1c).

Exclusion of subjects with hs-CRP levels greater than 7.5 mg/L (n = 258), had essentially no effect on the results obtained for all fractures (HR 1.39, 95% CI 1.09-1.77, Tertile 3 vs Tertiles 1+2, multivariate adjusted) and for vertebral fractures (HR 1.66, 95% CI 1.10-2.52, Tertile 3 vs Tertiles 1+2, multivariate adjusted).

**Long term effects of hs-CRP on fracture risk**

To explore the possibility that hs-CRP only reflected a poor health status at the time of serum sampling, we performed the above mentioned analyses excluding all fractures occurring within the first three years of follow up. The start time in these exploratory subanalyses was adjusted to begin at three years after study start. Hs-CRP remained a significant predictor for fractures in this model. Interestingly, hs-CRP tended to be an even stronger predictor for all fractures when long term effects were investigated (HR 1.61, 95% CI 1.18-2.20, multivariate adjusted Tertile 3 vs Tertiles 1+2). This was true also for vertebral fractures (HR 1.89, 95% CI 1.19-3.02, multivariate adjusted HR 2.29, 95% CI 1.37-3.81, Tertile 3 vs Tertiles 1+2).  Inclusion of femoral neck BMD in the
models had no influence on the associations between hs-CRP and long term risk of fracture (Table 2).

Discussion

As hypothesized, in this prospective population based study, men with slightly elevated levels of hs-CRP, suggesting a low grade inflammation, had an increased risk of fractures. The associations between hs-CRP and fracture risk in our cohort were largely independent of other known risk factors for fracture including physical performance, smoking, use of alcohol, previous fractures, several chronic conditions and medication use. The associations between hs-CRP and fractures were also independent of BMD.

Previous epidemiological research on associations between hs-CRP and incident fractures has been carried out primarily in women. Pasco and colleagues showed a dose-response relationship between hs-CRP and fractures in Australian women (median age 77 years, 96 fractures) (16). In a cohort of Japanese women (mean age 74 years, 50 limb or vertebral fractures), Nakamura et al found an increased risk for fractures in the medium and highest tertiles compared to the lowest tertile of hs-CRP (18). In a recent study of American women (mean age 46 years, 194 fractures), published by Ishii and colleagues, fracture hazard increased significantly for values of hs-CRP above 3 mg/L (19).

Schett and colleagues studied Italian women and men (mean age 59 years, 69 hip or vertebral fractures). Individuals in the highest tertile of hs-CRP had a clearly increased risk of non-traumatic fracture. However, only 19 of the fractures occurred in men making it difficult to draw firm conclusions on the relationship between hs-CRP and fractures in men from this study (17). A mixed cohort was used also by Cauley and colleagues (US men and women, mean age 74 years, 253 fractures) but in this study results were not statistically significant for hs-CRP. Borderline significance was reached in a multi adjusted model comparing the highest quartile of hs-CRP (Q4) to Q 1,2 and 3 combined. Results for men were not reported separately (21). Thus, there has definitively been a lack of well-powered studies on the associations between hs-CRP and fracture
risk in men, and this is where our large cohort including as many as 377 men with at least one fracture adds new knowledge.

Several risk factors for fracture such as smoking, low physical activity, malignancies and chronic inflammatory diseases are associated with elevated levels of hs-CRP (22-24), and the associations between hs-CRP and fracture risk could thus be mediated via these conditions. Our data are very robust however; adjustments for multiple known risk factors for fracture including smoking, physical activity, previous fractures, concomitant diseases and medication use had no major effects on the results. The same was true when individuals with hs-CRP above 7.5 mg/L as well as when fractures occurring during the first three years of follow up, were excluded from the analyses. This makes it highly unlikely that our results are due to increased fracture risk in individuals with frailty or undiagnosed concomitant diseases associated with elevated hs-CRP levels.

Moreover, the high number of fractures enabled us to study individual fracture types, and it appears that in men, hs-CRP mainly is a predictor of fractures in the vertebrae, as the associations with vertebral fractures prevail after all the above mentioned adjustments.

In our study, there were no associations between hs-CRP and BMD, and the predictive role of CRP for fracture risk was independent of BMD. This is in line with findings from those of the other studies on hs-CRP and fracture where BMD by DXA (16-18,21) or bone ultrasonographic data at the heel (17,21) were measured. However, there are also studies indicating that there actually is an association between hs-CRP and BMD, including a large study by Koh et al in healthy Korean women where a negative association between hs-CRP and BMD was found (25), and a study by Ding et al (26) where there were negative associations between hs-CRP and longitudinal change in BMD, while others report no associations (27-29). All of these studies were conducted in women.
The mechanism behind the associations between hs-CRP and fracture risk in our study is thus not an effect on BMD as measured by DXA. However, it is still possible that aspects of bone quality other than BMD, such as quality of the collagenous matrix, bone microarchitecture or bone size, mediate the effects of low grade inflammation on fracture risk. In the study by Ishii and colleagues, femoral neck composite strength indices were calculated using DXA-derived measurements. CRP was inversely associated with these strength indices, as well as with bone size factors, but not with BMD. Some, but not all, of the association between high hs-CRP and increased fracture risk in their study was explained by the decrement in composite strength indices with high hs-CRP (19). It thus appears that low grade inflammation can affect the balance between bone strength and load.

The associations between bone microarchitecture and hs-CRP were recently investigated by Rolland et al in a cohort of 1149 men aged 19-87. As in several other studies, there were no associations between hs-CRP and aBMD. Interestingly however, in men aged ≥ 72, but not in younger men, there was an association between bone microarchitecture as measured by high resolution pQCT (HR-pQCT), and hs-CRP. In the distal radius, men in the highest quartile of hs-CRP had 6.6% lower trabecular volumetric BMD and 4.5% lower trabecular number compared with the other quartiles combined. They also had a higher trabecular spacing, and more heterogenous trabecular distribution than men with lower hs-CRP (30). Inflammation upregulates osteoclasts and downregulates osteoblasts (31) and it is thus not surprising to see an effect of low grade inflammation in the metabolically more active trabecular bone compartment as in the study by Rolland and colleagues. Fracture prevalence increased with increasing hs-CRP concentration in the study by Rolland at al, but in contrast to our prospective study, this was a cross-sectional study where peripheral fractures were self-reported and only vertebral fractures were X-ray verified.

Thus, the two above mentioned studies give suggestions on how the association between high hs-CRP and an increased fracture risk are mediated. Because data on neither bone microarchitecture
nor bone strength indices are available for the men in our study we do not know if these parameters were affected in our cohort. There are yet other possible explanations not investigated so far including bone composition, falls and neuromuscular function.

Our study has a number of strengths. The population based nature, the large number of study subjects, the complete follow-up and the X-ray validation of fractures all contribute to the validity of the results. There were however also limitations. No other inflammatory markers were measured and therefore the potential causal mechanism underlying the associations between hs-CRP and fractures could not be studied. Hs-CRP was measured only once, and that might underestimate the effect size. Moreover, markers of bone turnover were not measured and thus it is not possible to investigate associations between bone turnover and hs-CRP.

In summary we show that in elderly Caucasian men, higher levels of hs-CRP are associated with an increased risk of fractures. We thus propose that low grade inflammation leads to an increased risk for fractures in men. Our study is the first to convincingly show the association between hs-CRP and fractures in men.

Disclosures

All other authors state that they have no conflicts of interest.
Acknowledgments

This work was supported by the Swedish Research Council, the Swedish Foundation for Strategic Research, the ALF/LUA Research Grant in Gothenburg, the Lundberg Foundation, the Torsten and Ragnar Söderberg’s Foundation, the Åke Wiberg Foundation and the Novo Nordisk Foundation.

Authors’ roles: Study design: AE, CO. Data Collection: MK, ÖL, DM. Data analysis: AE, CO, SMS. Data interpretation: AE, CO. Drafting manuscript: AE, CO. Revising manuscript content: AE, SMS, MK, ÖL, DM, CO. Approving final version of manuscript: AE, SMS, MK, ÖL, DM, CO. AE takes responsibility for the integrity of the data analysis.
References

1. Kanis JA, Johnell O, Oden A, Sembo I, Redlund-Johnell I, Dawson A, De Laet C, Jonsson B Long-term risk of osteoporotic fracture in Malmo. Osteoporos Int 2000; 11:669-74.
2. Center JR, Nguyen TV, Schneider D, Sambrook PN, Eisman JA Mortality after all major types of osteoporotic fracture in men and women: an observational study. Lancet 1999; 353:878-82.
3. Johnell O The socioeconomic burden of fractures: today and in the 21st century. Am J Med 1997; 103:20S-25S; discussion 25S-26S.
4. Khosla S, Amin S, Orwoll E Osteoporosis in men. Endocr Rev 2008; 29:441-64.
5. Spector TD, Hall GM, McCloskey EV, Kanis JA Risk of vertebral fracture in women with rheumatoid arthritis. BMJ 1993; 306:558.
6. Bernstein CN, Blanchard JF, Leslie W, Wajda A, Yu BN The incidence of fracture among patients with inflammatory bowel disease. A population-based cohort study. Ann Intern Med 2000; 133:795-9.
7. Weiss RJ, Wick MC, Ackermann PW, Montgomery SM Increased fracture risk in patients with rheumatic disorders and other inflammatory diseases -- a case-control study with 53,108 patients with fracture. J Rheumatol 37:2247-50.
8. Dennison EM, Compston JE, Flahive J, Siris ES, Gehlbach SH, Adachi JD, Boonen S, Chapurlat R, Diez-Perez A, Anderson FA, Jr., Hooven FH, LaCroix AZ, Lindsay R, Netelenbos JC, Pfeilschifter J, Rossini M, Roux C, Saag KG, Sambrook P, Silverman S, Watts NB, Greenspan SL, Premaor M, Cooper C Effect of co-morbidities on fracture risk: findings from the Global Longitudinal Study of Osteoporosis in Women (GLOW). Bone 2012; 50:1288-93.
9. Graat-Verboom L, van den Borne BE, Smeenk FW, Spruit MA, Wouters EF. Osteoporosis in COPD outpatients based on bone mineral density and vertebral fractures. J Bone Miner Res 2011; 26:561-8.

10. Nanes MS. Tumor necrosis factor-alpha: molecular and cellular mechanisms in skeletal pathology. Gene 2003; 321:1-15.

11. Baker-Lepain JC, Nakamura MC, Lane NE. Effects of inflammation on bone: an update. Curr Opin Rheumatol.

12. Pacifici R. The immune system and bone. Arch Biochem Biophys 503:41-53.

13. Kuo HK, Yen CJ, Chang CH, Kuo CK, Chen JH, Sorond F. Relation of C-reactive protein to stroke, cognitive disorders, and depression in the general population: systematic review and meta-analysis. Lancet Neurol 2005; 4:371-80.

14. Danesh J, Wheeler JG, Hirschfield GM, Eda S, Eiriksdottir G, Rumley A, Lowe GD, Pepys MB, Gudnason V. C-reactive protein and other circulating markers of inflammation in the prediction of coronary heart disease. N Engl J Med 2004; 350:1387-97.

15. Thorand B, Lowel H, Schneider A, Kolb H, Meisinger C, Frohlich M, Koenig W. C-reactive protein as a predictor for incident diabetes mellitus among middle-aged men: results from the MONICA Augsburg cohort study, 1984-1998. Arch Intern Med 2003; 163:93-9.

16. Pasco JA, Kotowicz MA, Henry MJ, Nicholson GC, Spilsbury HJ, Box JD, Schneider HG. High-sensitivity C-reactive protein and fracture risk in elderly women. JAMA 2006; 296:1353-5.

17. Schett G, Kiechl S, Weger S, Pederiva A, Mayr A, Petrangeli M, Oberhollenzer F, Lorenzini R, Redlich K, Axmann R, Zwerina J, Willeit J. High-sensitivity C-reactive protein and risk of nontraumatic fractures in the Bruneck study. Arch Intern Med 2006; 166:2495-501.
18. Nakamura K, Saito T, Kobayashi R, Oshiki R, Oyama M, Nishiwaki T, Nashimoto M, Tsuchiya Y C-reactive protein predicts incident fracture in community-dwelling elderly Japanese women: the Muramatsu study. Osteoporos Int 22:2145-50.

19. Ishii S, Cauley JA, Greendale GA, Crandall CJ, Danielson ME, Ouchi Y, Karlamangla AS C-reactive protein, bone strength, and 9-year fracture risk: Data from the study of women's health across the nation (SWAN). J Bone Miner Res 2013.

20. Mellstrom D, Johnell O, Ljunggren O, Eriksson A, Lorentzon M, Mallmin H, Holmberg A, Redlund-Johnell I, Orwoll E, Ohlsson C Free Testosterone is an Independent Predictor of BMD and Prevalent Fractures in Elderly Men - MrOs Sweden. J Bone Miner Res 2006; 21:529-35.

21. Cauley JA, Danielson ME, Boudreau RM, Forrest KY, Zmuda JM, Pahor M, Tylavsky FA, Cummings SR, Harris TB, Newman AB Inflammatory markers and incident fracture risk in older men and women: the Health Aging and Body Composition Study. J Bone Miner Res 2007; 22:1088-95.

22. Tonstad S, Cowan JL C-reactive protein as a predictor of disease in smokers and former smokers: a review. Int J Clin Pract 2009; 63:1634-41.

23. Plaisance EP, Grandjean PW Physical activity and high-sensitivity C-reactive protein. Sports Med 2006; 36:443-58.

24. Deichmann M, Benner A, Waldmann V, Bock M, Jackel A, Naher H Interleukin-6 and its surrogate C-reactive protein are useful serum markers for monitoring metastasized malignant melanoma. J Exp Clin Cancer Res 2000; 19:301-7.

25. Koh JM, Khang YH, Jung CH, Bae S, Kim DJ, Chung YE, Kim GS Higher circulating hsCRP levels are associated with lower bone mineral density in healthy pre- and postmenopausal women: evidence for a link between systemic inflammation and osteoporosis. Osteoporos Int 2005; 16:1263-71.
26. Ding C, Parameswaran V, Udayan R, Burgess J, Jones G Circulating levels of inflammatory markers predict change in bone mineral density and resorption in older adults: a longitudinal study. J Clin Endocrinol Metab 2008; 93:1952-8.

27. Shea MK, Dallal GE, Dawson-Hughes B, Ordovas JM, O'Donnell CJ, Gundberg CM, Peterson JW, Booth SL Vitamin K, circulating cytokines, and bone mineral density in older men and women. Am J Clin Nutr 2008; 88:356-63.

28. Nabipour I, Larijani B, Vahdat K, Assadi M, Jafari SM, Ahmadi E, Movahed A, Moradhaseli F, Sanjdideh Z, Obeidi N, Amiri Z Relationships among serum receptor of nuclear factor-kappaB ligand, osteoprotegerin, high-sensitivity C-reactive protein, and bone mineral density in postmenopausal women: osteoimmunity versus osteoinflammatory. Menopause 2009; 16:950-5.

29. Bhupathiraju SN, Alekel DL, Stewart JW, Hanson LN, Shedd KM, Reddy MB, Hanson KB, Van Loan MD, Genschel U, Koehler KJ Relationship of circulating total homocysteine and C-reactive protein to trabecular bone in postmenopausal women. J Clin Densitom 2007; 10:395-403.

30. Rolland T, Boutroy S, Vilayphiou N, Blaizot S, Chapurlat R, Szulc P Poor trabecular microarchitecture at the distal radius in older men with increased concentration of high-sensitivity C-reactive protein--the STRAMBO study. Calcif Tissue Int 2012; 90:496-506.

31. Baker-LePain JC, Nakamura MC, Lane NE Effects of inflammation on bone: an update. Curr Opin Rheumatol 2011; 23:389-95.
Figure legend

Figure 1

Forest plot of Cox proportional hazard ratio (HR) and 95% CI of fracture by hs-CRP (highest tertile vs medium and lowest tertiles combined). Figure 1a is adjusted for age. Figure 1b is adjusted for age, height, weight, calcium intake, physical activity, grip strength, cigarette smoking, use of alcohol, chronic obstructive pulmonary disease (COPD), stroke, diabetes, cancer, rheumatoid arthritis, prevalent fractures and medication use (corticosteroids, statins, thiazide diuretics, non-steroidal anti-inflammatory drugs (NSAIDs), osteoporosis medications, testosterone, antidepressants, hypnotics and sedatives, antiandrogens). Figure 1c is further adjusted for femoral neck sBMD.
## Tables

### Table 1 Baseline Characteristics by tertiles of hs-CRP

| Characteristic                      |              | All subjects (n=2910) | 1 (Low) (n=971) | 2 (Medium) (n=970) | 3 (High) (n=969) | P value |
|-------------------------------------|--------------|-----------------------|------------------|---------------------|------------------|---------|
| Age (yr)                            |              | 75.4 ± 3.2            | 75.4 ± 3.2       | 75.4 ± 3.2          | 75.5 ± 3.1       | 0.36*   |
| Height (cm)                         |              | 174.8 ± 6.6           | 175.2 ± 6.5      | 174.7 ± 6.4         | 174.4 ± 6.7      | 0.007*  |
| Weight (kg)                         |              | 80.7 ± 12.1           | 78.1 ± 11.1      | 81.4 ± 11.7         | 82.7 ± 13.1      | <0.001* |
| BMI (kg/m²)                         |              | 26.4 ± 3.6            | 25.4 ± 3.2       | 26.6 ± 3.5          | 27.2 ± 3.8       | <0.001* |
| Femoral neck BMD (g/cm²)            |              | 0.83 ± 0.13           | 0.82 ± 0.13      | 0.84 ± 0.13         | 0.83 ± 0.14      | 0.09*   |
| Physical activity (km)              |              | 3.9 ± 3.1             | 4.2 ± 3.2        | 4.1 ± 3.2           | 3.5 ± 3.0        | <0.001* |
| Grip Strength (kg)                  |              | 39.9 ± 7.5            | 40.4 ± 7.2       | 40.0 ± 7.9          | 39.3 ± 7.4       | 0.003*  |
| Smoking (%)                         |              | 246 (8.5)             | 60 (6.2)         | 70 (7.2)            | 116 (12.0)       | <0.001* |
| Alcohol 3 or more units per day (%) |              | 76 (2.6)              | 19 (2.0)         | 25 (2.6)            | 32 (3.3)         | 0.18*   |
| Calcium intake (mg)   | 898 ± 435 | 893 ± 403 | 895 ± 429 | 906 ± 470 | 0.51* |
|----------------------|-----------|-----------|-----------|-----------|-------|
| Hs-CRP               | 2.17 (1.67-3.26) | 1.51 (1.31-1.67) | 2.17 (1.98-2.40) | 4.31 (3.26-7.92) | NA    |

**Major prevalent diseases**

| Cancer (%)           | 450 (15.5) | 151 (15.6) | 157 (16.2) | 142 (14.7) | 0.65  |
| COPD (%)             | 245 (8.5)  | 63 (6.5)   | 59 (6.1)   | 123 (12.8) | < 0.001 |
| Diabetes (%)         | 276 (9.5)  | 90 (9.3)   | 87 (9.0)   | 99 (10.2)  | 0.63  |
| Stroke (%)           | 189 (6.5)  | 55 (5.7)   | 69 (7.1)   | 65 (6.7)   | 0.41  |
| Rheumatoid arthritis (%) | 43 (1.5) | 12 (1.2)   | 12 (1.2)   | 19 (2.0)   | 0.31  |
| Fractures > 50 years (%) | 501 (17.3) | 160 (16.6) | 172 (17.9) | 169 (17.6) | 0.73  |

**Subjects with validated incident fractures**

| All fractures        | 377 (23.9) | 110 (20.4) | 116 (21.6) | 151 (30.1) | 0.01  |
| Non vertebral osteoporosis fractures | 159 (9.7) | 51 (9.1) | 49 (8.8) | 59 (11.2) | 0.57  |
| Hip fractures        | 89 (5.4)   | 27 (4.8)   | 28 (5.0)   | 34 (6.4)   | 0.61  |
| Clinical vertebral fractures | 125 (7.6) | 39 (6.9) | 34 (6.1) | 52 (9.9) | 0.11  |
Values are given as mean ± SD, median (inter quartile range) or n (percent). For fractures, the numbers of subjects with first fractures are given, with the incidence/1000 person-years shown within parentheses. Some subjects, included in the group of “all fractures,” had more than one type of first fracture, and therefore, these subjects were included in more than one of the different subtypes of fractures. Non-vertebral osteoporosis fractures are defined as fractures in hip, distal radius, proximal humerus, and pelvis.

sBMD, standardized BMD; hs-CRP, high sensitivity CRP; NA, data not applicable; COPD, chronic obstructive pulmonary disease

* p for trend
Table 2 Cox proportional hazard ratio (HR) and 95% CI of fracture by hs-CRP, excluding all fractures during the first three years of follow up

|                      | All fractures | Hip                | Non-vertebral | Vertebral   |
|----------------------|---------------|--------------------|---------------|-------------|
| **Age-adjusted, HR (95 % CI)** | 1.62 (1.22-2.16) | 1.23 (0.70-2.16) | 1.00 (0.63-1.60) | 1.89 (1.19-3.02) |
| **Multivariate adjusteda, HR (95 % CI)** | 1.61 (1.18-2.20) | 1.23 (0.66-2.29) | 1.05 (0.62-1.75) | 2.29 (1.37-3.81) |
| **Multivariate adjustedb, HR (95 % CI)** | 1.58 (1.15-2.16) | 1.14 (0.61-2.13) | 0.98 (0.58-1.64) | 2.22 (1.33-3.69) |

*a* Highest tertile vs medium and lowest tertiles of hs-CRP combined. Adjusted for age, height, weight, calcium intake, physical activity, grip strength, cigarette smoking, use of alcohol, chronic obstructive pulmonary disease (COPD), stroke, diabetes, cancer, rheumatoid arthritis, prevalent fractures and medication use (corticosteroids, statins, thiazide diuretics, non-steroidal anti-inflammatory drugs (NSAIDs), osteoporosis medications, testosterone, antidepressants, hypnotics and sedatives, antiandrogens)

*b* Further adjusted for femoral neck sBMD
Figure 1