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Dietary protein and bone health across the life-course: an updated systematic review and meta-analysis over 40 years

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Abbreviations:

aBMD, areal Bone Mineral Density ALP, Alkaline Phophatase AP, Animal Protein BAP, Bone Alkaline Phosphatase BCE, Bovine Collagen Equivalents BMC, Bone Mineral Content BMD, Bone Mineral Density BUA, Broadband Ultrasound Attenuation BV, Bone Volume Cr, Creatinine CTX, C-telopeptide of Collagen DPA, Dual Photon Absorptiometry DPYD, Deoxypyridinoline DXA, Dual X-ray Absorptiometry FN, Femoral Neck FNBA, Femoral Neck Bone Area

FNBMC, Femoral Neck Bone Mineral Content FNBMD, Femoral Neck Bone Mineral Density FNvBMD, Femoral Neck volumetric Bone Mineral Density FSBMD, Femoral Shaft Bone Mineral Density HBMD, Hand Bone Mineral Density HCHP, High Calcium High Protein HCLP, High Calcium Low Protein HF, Hip Fracture HPO, Hydroxyproline HR, Hazard Ratio IntertrochBMD, Intertrochanter Bone Mineral Density LCHP, Low Calcium High Protein LCLP, Low Calcium Low Protein LS, Lumbar Spine LSBMC, Lumbar Spine Bone Mineral Content LSBMD, Lumbar Spine Bone Mineral Density LSvBMD, Lumbar Spine volumetric Bone Mineral Density MBP, Milk Basic Protein NTX, N-terminal Peptide of Collagen OC, Osteocalcin OR, Odds Ratio P1NP, Procollagen type 1 N-terminal Propeptide PERI, Perimenopausal PFBMD, Proximal Femur Bone Mineral Density POM, Postmenopausal PRE, Premenopausal PYD, Pyridinoline **RBMC**, Radial Bone Mineral Content RBMD, Radial Bone Mineral Density RR, Relative Risk

SOS, Speed of Sound SP, Soy Protein SSI, Strength-strain Index TBBA, Total Body Bone Area TBBMC, Total body Bone Mineral Content TBBMD, Total body Bone Mineral Density TEI, Total Energy Intake THBMC, Total Hip Bone Mineral Density TP, Total Protein TRAP, Tartrate Resistant Acid Phosphatase TrochBMD, Trochanter Bone Mineral Density TSBMD, Total Spine Bone Mineral Density vBMC, volumetric Bone Mineral Content vBMD, volumetric Bone Mineral Density VP, Vegetable Protein

[1] Supplementary Methods: Data extraction

Correlation coefficients (adjusted and unadjusted), n (number of participants), beta coefficients (standardized and unstandardized) for the relationship between protein intake (g/Kg/d or g/d) and bone outcomes were extracted, as well as bone outcomes by protein intake category. Data for calcium intakes were also extracted. For correlational studies looking at the association between change in bone outcomes over time and baseline protein intake, or assessing associations between protein intake and bone outcomes at different time points, all relevant data were extracted. This included mean and standard deviation (SD) for change in bone indices over time, or else r coefficients or beta coefficients for slope of bone loss in different protein intake groups. It also included bone outcomes by protein intake category.

For studies presenting data on risk of fracture or osteoporosis/osteopenia diagnosis, odds ratios, hazard ratios or relative risk estimates (with 95% confidence intervals) for the highest and lowest categories of protein intakes were extracted, with n and p if available. Mean and SD for protein intakes in cases and controls were extracted if no other data were presented (e.g. no risk estimates, no categories of intake data).

Finally, for the intervention studies, as subjects were randomized at baseline, only the mean, SD and n for follow up measurements were extracted for each relevant outcome in each study arm. Standard errors of the mean (SEM) were converted to standard deviations using the formula (SEM=SD/ \sqrt{n}). Papers not providing complete data to calculate standardized size effects (i.e. not able to calculate the standard deviation, or the standard error of the mean) were excluded from the meta-analysis if this data could not be obtained from the relevant authors. Two authors of relevant articles with missing data were contacted. Neither of the authors replied so their papers were not included in the meta-analysis, but the general findings were included in the systematic review. Two other authors, whose papers had

missing data, or data only available in figures, was not contacted as previous contact when doing the original analysis in 2007 was not successful.

[2] Supplementary Results: Studies reporting correlation or regression coefficients for the relationship between dietary protein and bone indices

Study Characteristics

Of the 74 studies presenting data for correlation or regression coefficients, 18 studies were from South or East Asia (1-17), 21 from Europe(18-38), 2 from the Middle East (39, 40), 6 from Australia, Tasmania or New Zealand(41-46), 25 from USA or Canada (47-71) and 2 from South America(72, 73). Of these 74 studies, 12 were in children or adolescents (3, 17-21, 24, 25, 37, 38, 45, 48), with 1 study combining data from adults and children (44). Also, 13 studies were in premenopausal women (1, 4, 14, 15, 29, 30, 32, 43, 47, 52, 57, 62, 74), 21 in postmenopausal women (2, 6-8, 11, 12, 16, 22, 33, 39-42, 46, 55, 59, 61, 63, 65, 70, 72), 7 in both pre and postmenopausal women(9, 10, 27, 34, 49-51), 2 in both peri and postmenopausal women(5, 53), 5 in men(31, 36, 66, 67, 73), 1 in pre, peri or postmenopausal women(26) and 13 in both men and women(13, 23, 28, 35, 56, 58, 60, 64, 68, 69, 71, 75). Sixty-one studies assessed total protein intake only but 2 studies assessed both soy and total protein(5, 8), one study assessed soy protein only(6), 10 studies assessed animal and/or vegetable protein intake in addition to total protein(7, 15, 16, 19, 20, 34, 47, 54, 58, 71) and one study assessed soy, animal, vegetable and total protein(13).

Systematic Review: Studies reporting correlation or regression coefficients, or protein category data

Seventy-four studies presented correlation coefficient (r) or regression coefficient data (standardized (Beta) or unstandardized (B)) data (Table 1).

Cross-sectional data- BMD

In adults a large number of studies found an association between protein intake and Bone Mineral Density (BMD) at the hip (23, 27, 32, 36, 39, 41-43, 49, 55, 58, 60, 66, 72), radius (4, 49, 57, 58), spine(9, 23, 27, 32, 43, 52, 60, 62, 63, 66, 72), total body (27, 55, 56, 58, 62, 66, 68) or hand (HBMD) (55). Conversely, a large number of studies found that protein intake was not associated with BMD at the hip (1, 2, 9, 11, 22, 28-30, 35, 36, 47, 49, 50, 52, 58, 60, 63, 65, 72, 73), spine (1, 2, 8, 11, 22, 28-30, 35, 39, 47, 49, 50, 54, 58, 60, 65), radius (32, 49, 53, 61) or total body (50, 58, 64). See Supplemental Material for a review of adult studies assessing animal and vegetable protein intake specifically. In children, two studies found that total protein and animal protein intakes were not associated with femoral neck (FN) BMD or lumbar spine (LS) BMD(45). However, two other studies conflicted as to whether or not protein intake was associated with total body BMD (TBBMD) (45, 48).

Cross-sectional data- BMC and Bone Size

In children, seven studies showed that higher protein intake was associated with increased radial periosteal circumference, cortical area, volumetric BMC (vBMC) and polar SSI(18, 38), as well as TBBMC(3, 24, 48), total body bone area (TBBA)(3, 24), total radial BMC (RBMC)(21), radial metaphyseal BMC(21), femoral neck BMC (FNBMC)(21), femoral diaphysis BMC(21) and lumbar spine BMC (LSBMC)(21). Likewise, another study indicated that there was a positive association between dietary protein and forearm cortical BA but not forearm vBMC(25). However, one study found no association between child protein intake and radial diaphysis BMC or total hip BMC (THBMC)(21) and another found that total protein and animal protein intakes were not associated with total body BMC (TBBMC) or lumbar spine BMC (LSBMC)(19). In adults, a large number of studies found a

positive association between total protein intake and RBMC(10, 32, 33, 44, 57, 61), TBBMC(55, 62, 72), Spine BMC (62, 67), LSBMC(32) TBBMC(68) and HipBMC(32), but three studies found no association between protein intake and RBMC(33, 51, 67), humerus BMC(51) or Ulna BMC(51).

Cross-sectional data- quantitative ultrasound (QUS) and bone makers

There were no studies of BUA in children. In adults, four studies found that protein intake was positively associated with calcaneal BUA(15, 34, 41, 52). In one of these studies total protein intake was negatively associated with calcaneal BUA but the association disappeared when adjusting for animal: vegetable protein ratio(34). In terms of bone markers, in children the one study assessing bone markers found a positive association between protein intake and bone alkaline phosphatase (BAP), but found no association with osteocalcin (OC) or c-telopetide of collagen (CTx) (20). In adults, in four studies increased total protein intake was negatively associated with hydroxyproline (HPO) (premenopausal women only) (49), CTX(42), pyridinoline (PYD)(8, 26), deoxypyridinoline (DPYD)(26) and type 1 n-terminal procollagen (P1NP)(42). In contrast, one study found a positive association between total protein intake and CTX(31) and other studies found no association between total protein intake and HPO (postmenopausal women only)(49), OC(8, 12, 27, 49, 50, 59), n-terminal telopeptide (NTX)(12, 50, 59), DPYD(8, 12) or BAP(8, 12).

Systematic Review: Animal, vegetable or soy protein and bone health *Studies in adults*

In four studies, animal protein intake was not associated with FNBMD (7, 47), LSBMD(7, 47), TBBMD(7) LSvBMD(54), THBMD(47) or calcaneal stiffness index (13). One study found a positive association between spine BMD and non-dairy animal protein intake in postmenopausal women aged 50 years or older, but conversely in premenopausal women

found a negative association (71). One study found increasing animal protein was associated with increased THBMD, FNBMD, TSBMD, TBBMD(58). One study found a negative association between BUA of the calcaneus with animal protein, the effect being modified by calcium intake (34). Finally, another study found that increased animal: vegetable protein ratio was a negative predictor of FNBMD (16).

Two studies found that vegetable protein was not associated with FNBMD and LSBMD(7) or calcaneal stiffness index(13). However a negative association was found in three studies between plant protein and spine BMD (16, 71) or TBBMD (7, 47) and hip BMD (47, 58, 71). Interestingly, one study found that higher vegetable protein intake was associated with reduced LSvBMD in persons of White (but not Chinese, Black or Hispanic) ethnicity (54). Increasing vegetable protein was associated in another study with reduced THBMD, FNBMD and TSBMD in women but not men (58). Conversely, one study found a positive association between vegetable protein intake and bone heath for calcaneal BUA (34). In terms of soy protein, four studies showed no association with calcaneal stiffness index (13), LSBMD(5, 6) , FNBMD (5, 6), TBBMC(5, 6, 44), THBMC(5), THBMD(6), or TBBMD(6), Troch/intertroch BMD (6) or leg BMC(44). However, two studies found that increased soy protein intake was associated with lower DPYD(8) or higher LSBMD (8), but no association with PYD, ALP or OC(8).

Studies in children

In children, one study found no association between animal (meat) protein intake and TBBMC or LSBMC, but did find a positive association between BAP and animal protein intake (19). However, there was no association between animal or vegetable protein intake and OC or CTX, or between vegetable protein and BAP(20). Conversely, another paper from the same group found a positive association between animal protein intake and OC, CTX and BAP(19). There were no studies of soy protein intake in children or adolescents.

Systematic Review: Studies reporting r values for slope of change in bone mass In children, one study in boys found that baseline protein intake (age 7 years) was not associated with FN vBMD, Total hip vBMD, distal tibia vBMD, FNBA, FN width, total hip aBMD or FN aBMD at age 15 years when physical activity levels were lower (37), but protein intake was positively associated when physical activity levels were higher (37). Another study, in pre-pubertal girls with low calcium intakes, found increased animal protein intake was associated with lower gain in Radial BMC and TBBMC from age 10 to 15 years (17).

In adults, one study found no relationship between baseline intake of total, animal or vegetable protein and 3 year change in hip, spine or TBBMD(47) with another finding no association between protein intake and bone loss, also over 3 years (59). Another study found no association between protein intake and loss of LSBMD, FNBMD, THBMC or TBBMC over 30 months (5).

Similarly, one study found no association between protein intake during adolescence (15 years old) and subsequent LSBMD or FNBMD in young adulthood (20-25y old)(28). Finally, another found no association between BUA, BV or SOS in 18-19 year old women and previous protein intake at 9-11 years old (74). No relationship between protein intake and total hip BMD(75). Sahni 2014 found a negative association in men between protein intake and % change in TrochBMD but not % change in LS or FN BMD in men, but there were no associations at any site for women (60).

However, some studies have found an association between protein intake and change in bone mass. One study found increased protein intake at age 20-25 years was retrospectively

associated with increased gain in TBBMC from peri-adolescence to the present day, in females with adequate calcium intake as well as all males regardless of calcium intake (64). One study found that for every 20% increase in % of total energy from dietary protein, over a 6 year period, there was an increase of 0.003 to 0.004g/cm² for TBBMD, HipBMD and Spine BMD(70). Another study found reduced FNBMD and LS BMD loss (but no change in radial shaft BMD) over a 4 year period in those in with the highest (vs. lowest) quartile of protein intake(69). Finally, another study found an association between higher protein intake and reduced radial bone loss (BMD) in both pre and postmenopausal women(51). Another study in elderly women found a positive association between baseline protein intake and BMC 5 years later with the highest daily protein intake tertile (>87g/d) being associated with higher appendicular and whole body BMC than the lowest quartile (<66g/d)(46). Finally, in one study of men and women aged \geq 50 years old, there was a positive association between baseline total protein intake and BMD 5 years later at the hip, as well as a positive association between protein intake and Spine BMD for both men and women, and with Hip BMD for the women only (71).

Conversely, one study found increased bone loss in men with higher protein intake (60). A negative association between vegetable protein intake and % change in LSBMD, as well as between AP:VP ratio and FNBMD(16). Another study found increased bone loss in women consuming high animal: vegetable protein ratio diets than those consuming low animal: vegetable protein ratio diets (76). One study found that higher quartiles of soy protein intake were associated with protection against loss of TBBMC(5). Another study found no association between soy intake and loss of LSBMD, FNBMD, THBMC or TBBMC over 30 months (5).

Meta-analysis: Funnel Plots and Sensitivity analyses for FNBMD and LSBMD models Funnel plots were conducted for the following models: FNBMD (unadjusted and adjusted); LSBMD (unadjusted and adjusted), BMD in postmenopausal women and BMD in premenopausal women. The FNBMD plot showed all studies within the 95% confidence interval boundaries but there were a lack of smaller studies showing a negative effect estimate. The LSBMD plot had 3 out of 18 studies outside the 95% confidence interval boundary (strong positive effect sizes) showing, as with FNBMD, a lack of smaller studies in the negative effect estimate area. The funnel plot for postmenopausal women BMD showed few small studies with a negative effect, and the premenopausal women BMD plot showed 3 studies out of 10 outside the 95% confidence interval area (strong positive effect sizes) and a lack of medium sized studies in the negative effect estimate area. Taken together, this suggests potential publication bias in terms of smaller to medium studies, particularly those with negative associations between protein and bone health, not being published. In terms of sensitivity analyses, removal of each study in turn for the FNBMD analysis (confounder adjusted data only) had little effect on the above results. Sensitivity analyses show effect sizes as follows when each study excluded in turn: Chan 2009 r(random)=0.07 (-0.04 to 0.18) R^2 =0.005 (0.5%) P=0.21, I²=47% P_{(heterogeneity})=0.15; Cooper 1995 Postmenopausal data $r_{(random)}=0.05$ (-0.06 to 0.17) R²=0.003 (0.3%), P=0.37, I²=63% $P_{\text{(heterogeneity)}}=0.07$; Cooper 1995 Premenopausal data $r_{\text{(fixed)}}=0.01(-0.05 \text{ to } 0.07) \text{ R}^2 = <0.001$ (<0.1%) P=0.33, I²=0% P_(heterogeneity)=0.67; Ho2003 r_(random)=0.05 (-0.08 to 0.19) R²=0.003 (0.3%) P=0.43, I²=62% P_(heterogeneity)=0.07. However, removal of the Cooper et al. (1995)(49) premenopausal data reduced heterogeneity to 0%, suggesting this study contributed strongly to the heterogeneity. Exclusion of this study also reduced the effect size from 0.07 to 0.01, suggesting it was increasing the effect size substantially. However, the R^2 was still very small (<0.1% to 0.5%) and not statistically significant when any of the studies were removed, or all included (0.2%). Therefore removal of studies had little overall effect on the pooled effect size.

In terms of sensitivity analyses for LSBMD (confounder adjusted data only), the following pooled effect sizes were found when each study in turn were excluded: Ho 2003(6) $r_{(fixed)}$ =-0.021 (-0.14 to 0.10) R²=<0.001 (0.1%) P=0.73, I2=0% P_(heterogeneity)=0.39; Cooper 1995(49) Postmenopausal r(fixed)= 0.02 (-0.06 to 0.11) R²<0.001 (0.1%) P=0.60, I²=0% P_(heterogeneity)=0.68; Cooper 1995(49) Premenopausal r_(fixed) = -0.01 (-0.08 to 0.07) R²=<0.001 (0.1%) P=0.88, I²=0% P_(heterogeneity)=0.43. The pooled effect size was unchanged by the elimination of any studies in the model, suggesting no studies were having a strong impact on

the overall effect size.

Meta-analysis: Associations with protein and calcium dose, as well as calcium: protein ratio

Linear, quadratic and cubic regression models (not controlling for confounders) showed that neither the calcium: protein ratio (calcium mg/protein g), nor the protein (g/kg/d) or calcium (mg/kg/d) dose, were associated with correlation (r) coefficients for the association between protein intake and bone health for either LSBMD or FNBMD (P values were P >0.05 or did not survive multiple testing adjustment (revised cutoff of P>0.001)). There was a positive association between protein intake and both FNBMD and LSBMD in the linear model (both P=0.02, b=0.33-0.39) and a negative association in the cubic models (P=0.01 (b= -0.90) for FNBMD and P=0.009 (b= -0.84) for LSBMC) but these models did not survive adjustment for multiple testing (P>0.001). See Supplemental Table 6 for full details of these analyses.

[3] Supplementary Results: Studies reporting fracture or osteoporosis risk

Study Characteristics

Of the 29 studies (including six studies already reported in correlation coefficient or bone slope sections), 4 were from Asia (2, 14, 77, 78), 11 were from the USA or Canada (70, 71, 76, 79-86), 2 were from the Middle East (40, 87) 1 was from New Zealand(42) and 9 were from Europe (36, 88-95). In addition, 2 studies were by authors from the USA but reported data from multiple countries (96, 97).

In terms of study design, 13 studies were cohort studies (70, 71, 76, 78-84, 88, 92, 94), 2 were ecological studies (96, 97), 3 were of cross-sectional design (2, 36, 42) and 11 were case-control studies (14, 40, 77, 85-87, 89-91, 93, 95). Of the latter, only 2 were prospective (nested) case-control studies (89, 93). In terms of fracture type, of the 13 cohort studies, 7 assessed hip fracture only(70, 76, 80-83, 94), 1 assessed hip, spine and forearm/wrist fracture (84), one study assessed hip and forearm/wrist fractures (79), two studies assessed all fragility fractures combined (71, 88) and two studies assessed all fractures (fragility and nonfragility)(78, 92). The 2 ecological studies assessed just hip fracture incidence (96, 97). For the 11 case-control studies, 1 study was in children, assessing dietary protein intake in children with fracture compared with children without fracture (93). The other 10 studies were all in adults, 1 assessing odds of hip fracture(86), 1 assessing odds of all fragility fractures by protein intake(91), 4 assessing differences in protein intake in hip, spine or wrist fracture cases and controls (89) or between osteoporotic/osteopenic cases vs. controls (14, 40, 87), 2 assessing odds of hip fracture only by protein intake(85, 95) and 2 assessing odds of osteoporosis diagnosis by protein intake category(77, 90).

In terms of population studied, 6 of the 13 cohort studies were in postmenopausal women(70, 76, 78, 81, 84, 88), 1 in pre and postmenopausal women(79), 5 in men and women(71, 80, 83, 92, 94) and 1 in men only(82). In the 11 case-control studies, 1 was in children(93), 3

studied men and women(86, 89, 91), 1 studied premenopausal women(14) and 6 studied postmenopausal women(40, 77, 85, 87, 90, 95).

Six cohort studies and the 3 cross-sectional studies assessed total dietary protein only (2, 36, 42, 70, 71, 80, 82, 84, 92), 1 studied animal protein only(94), 1 studied soy protein only(78) and 5 studied total dietary protein, animal protein and vegetable protein(76, 79, 81, 83, 88). The 2 cross cultural studies assessed the relationship between protein intake and fracture risk in 16 (96) to 33(97) countries worldwide, with Abelow et al.(1992)(96) studying animal protein only and Frassetto et al. (2000)(97) studying animal, vegetable and total protein intake. In the case-control studies, 7 studies assessed total protein only(14, 40, 85, 87, 89, 93, 95), whilst 4 assessed total, animal and vegetable protein(77, 86, 90, 91).

Exclusion of studies from fracture risk meta-analysis

Studies were excluded from the quantitative meta-analysis of fracture risk if they were crosscultural studies(96, 97), had only data on risk of osteoporosis or osteopenia (2, 36, 40, 42, 77, 87, 90), had effect statistics that were incompatible with other studies(70, 92) were the only fracture study in children (93) or reporting soy protein(78), only had results that were stratified by calcium (84), had missing data (76) or were case-control studies not reporting odds ratios (14, 89, 95). Case-control studies were analysed separately from cohort studies due to methodological differences. Cohort studies presenting data on RR and HR were analysed separately.

Systematic Review: Studies reporting fracture or osteoporosis risk

Twenty-nine studies reported data on fracture and/or osteoporosis diagnosis (**Table 2**), of which 28 studies were in adults and one in children (93). All studies provided multivariate adjusted estimates adjusted for multiple confounders (see Table 2) except for one cohort

study (76) which controlled for only 2 confounders, 2 ecological studies (96, 97) which were only adjusted for age, 2 cross-sectional studies which were only adjusted for BMI or energy intake(36, 42), 5 of the case-control studies(14, 40, 87, 89, 93) which presented only unadjusted data, and another case-control study which presented categories of intake (non-statistically analysed) so confounder adjustment was not required (90).

Cohort studies- total protein intake

Three studies found no association between total protein and risk of hip (80, 82) or spine fractures. In terms of fracture type, three studies found no association between total protein intake and either risk of combined fragility and non-fragility fractures (92), just all fragility fractures (71, 88) or hip fracture specifically (79, 81, 94). One study found an increased risk with higher total protein intake (79) and another found the reduced risk with higher protein intake(70). Finally, another study found increased odds of fracture when calcium intake was high and protein intake was low, but this association was attenuated when both calcium and protein were high (84). One study found the relationship between protein intake and fracture risk depended on calcium intake(83).

Case-control, cross-sectional and ecological studies- total protein intake

The one case-control study in children was of prospective (nested within-cohort) design and found no difference in protein intake between fracture cases and controls (93). All case-control studies in adults were of retrospective design, with the exception of 1 study that was prospective (89). Four adult case-control studies found no significant difference between protein intakes in cases with osteoporosis vs. non-osteoporotic controls(90) or

osteoporotic/osteopenic cases vs. healthy controls (14, 40, 87). In contrast, one study found increased odds of osteoporosis diagnosis with higher total protein intake (77). Three studies found no reduction in odds of being a fracture case in persons with higher protein intake compared with low protein intake (85, 91, 95). However, one study found lower total protein intake (non-adjusted for confounders) in cases with fragility fractures than in controls(89), and two studies found an increased odds of fracture(36) or reduced odds of fracture (50-69 year old subgroup) in persons with higher protein intake (86). The two cross-cultural (ecological) studies found a positive association between hip fracture and total protein intake (97) and a positive association between hip fracture and animal protein intake (96), both studies controlling for age only. Two of the 3 cross-sectional studies found that protein intake was a predictor of odds for LS osteopenia(2) or hip BMD below 0.83g/cm2 (men)(36), however the 3rd study found that protein intake was not associated with diagnosis of osteoporosis or osteopenia(42).

Systematic Review: Animal, Vegetable and Soy Protein and Fracture risk

In studies that presented data on animal and vegetable protein, one study found no association between animal or vegetable protein and fragility fracture(88). Similarly, 2 studies found no association between animal or vegetable protein and hip fractures (79, 94), or found an association between animal protein intake and hip fracture in persons with lower calcium intakes only (83). Other studies found no association between hip fracture risk and animal or vegetable protein intake (86), or no difference in odds of being a fragility fracture case in persons with varying animal or vegetable protein intake (91).

However, two other studies did find a significant association between animal protein intake and risk of forearm(79) and hip fractures(81), but one study found no association between vegetable protein intake and fracture risk(81). In addition, 1 study found an association between increased animal protein intake and increased hip fracture risk, as well as increased risk of hip fracture with increased animal: vegetable protein ratio, and a reduced risk of hip fracture with increased vegetable protein intake (76). One study found increased odds of osteoporosis with higher animal protein intake, but lower odds with increased vegetable protein intake(77). Finally, the 2 cross cultural studies(96, 97) found a positive association between animal protein intake and hip fracture incidence as well as a negative association between hip fracture incidence and increasing vegetable protein intake(97). The one cohort study assessing the relationship between soy protein intake and fracture risk (all fractures) found a reduced risk of fracture in the highest intake quintile of soy protein compared with the lowest(78).

Quality Analysis

Twelve cohort studies were assessed for quality using the Newcastle-Ottawa Cohort study assessment tool(98) (scored out of 9). Three studies scored 4-5 (44-55%)(79, 88, 94), 8 studies scored 6-7 (66-78%)(70, 71, 76, 78, 80, 81, 83, 92) and 1 study scored 8 (89%)(82). In addition, 6 case-control studies were assessed using the Newcastle-Ottawa case-control study assessment tool (scored out of 10), with 3 studies scoring 3-5(77, 91, 95) and 2 studies scoring 6-7 (85, 86). The nested case-control study by Samieri et al. (2013) (89) (score=8 out of 9) was assessed using the cohort study tool as it was deemed more appropriate for the study design. As discussed previously, the cross-sectional studies were not analysed for quality due to the very large numbers of studies. It was not possible to formally assess the quality of the 2 ecological studies (96, 97) as there is no specific tool for this.

Fracture risk meta-analysis: Sensitivity and subgroup analysis

Subgroup analysis showed that when removing Dargent-Molina et al 2008, (which was the only study to include non-hip fracture results), pooled estimates were as follows: animal protein intake, (RR (random)=0.83 (0.54 to 1.30, p=0.42, n=3 studies, I^2 =48%

 $P_{(heterogenity)}=0.14)$, vegetable protein intake (RR_(fixed)=1.20 (0.82 to 1.73, p=0.35, n=2 studies, I²=4% P_{(heterogenity})=0.34)), and total protein intake (RR _(random)=0.75 (0.47 to 1.21, p=0.24, n=3 studies, I²=22% P_{(heterogenity})=0.28)).

In terms of sensitivity analysis, the effect sizes when each study were removed in turn were: (see estimates above for removal of Dargent-Molina): Animal Protein: Feskanich: RR (random)=0.91 (0.61 to 1.37, p=0.67, n=3 studies, $I^2=63\%$ P_(heterogenity)=0.07); Meyer: RR (random)=0.93 (0.63 to 1.37, p=0.71, n=3 studies, $I^2=63\%$ P_(heterogenity)=0.07); Munger: RR (fixed)=1.09 (0.97 to 1.21, p=0.73, n=3 studies, $I^2=0\%$ P_(heterogenity)=0.73). For Vegetable Protein: Feskanich: RR (random)=1.13 (0.63 to 2.05, p=0.68, n=2 studies, $I^2=48\%$ P_(heterogenity)=0.68); Munger: RR (fixed)=0.96 (0.86 to 1.08, p=0.51, n=2 studies, $I^2=0\%$ P_(heterogenity)=0.48). For Total Protein: Feskanich: RR (random)=0.76 (0.42 to 1.39, p=0.38, n=3 studies, $I^2=54\%$ P_(heterogenity)=0.12); Munger: RR (fixed)=1.05 (0.93 to 1.17, p=0.43, n=3 studies, $I^2=0\%$ P_(heterogenity)=0.43); Mussolino: RR (random)=0.99 (0.77 to 1.27, p=0.90, n=3 studies, $I^2=33\%$ P_(heterogenity)=0.22). Therefore, for all protein types the removal of Munger rendered the heterogeneity down to zero, suggesting this study was the cause of the heterogeneity observed. Overall statistical significance of the models, for all types of protein, were not affected by the removal of any study.

For the cohort studies reporting hazard ratios, removal of each study in turn led to pooled estimates as follows: Langsetmo (Men): HR $_{(random)}=0.87$ (0.57 to 1.34, p=0.54, n=4 studies, I²=47% P_(heterogenity)=0.13); Langsetmo (Women): HR $_{(random)}=0.82$ (0.47 to 1.44, p=0.50, n=4 studies, I²=50% P_(heterogenity)=0.11); Misra: HR $_{(random)}=0.89$ (0.58 to 1.37, p=0.60, n=4 studies,

 $I^2=41\%$ P_(heterogenity)=0.16); Sahni(High Calcium): HR _(random)=0.84 (0.58 to 1.22, p=0.36, n=4 studies, $I^2=49\%$ P_(heterogenity)=0.12), Sahni(Low Calcium): HR _(fixed)=0.79 (0.64 to 0.97, p=0.02, n=4 studies, $I^2=0\%$ P_(heterogenity)=0.66). The removal of Sahni (Low Calcium) data led to a statistically significant reduction in fracture risk when protein intake was higher. Heterogeneity was also reduced to 0%, suggesting this result was leading to a masking of an association between protein and fracture risk shown in the low calcium arm of the Sahni study and the other studies in the meta-analysis.

For case control studies reporting odds ratios the following effect sizes were obtained when studies were removed as follows: Martinez-Ramirez: OR $_{(random)}=0.65$ (0.26 to 1.65, p=0.36, n=3 studies, I²=73% P_(heterogenity)=0.03); Nieves: OR $_{(random)}=0.57$ (0.23 to 1.44, p=0.23, n=3 studies, I²=47% P_(heterogenity)=0.15); Wengreen (50-69years old: OR $_{(fixed)}=1.10$ (0.53 to 2.26, p=0.81, n=3 studies, I²=0% P_(heterogenity)=0.98); Wengreen (70-89years old: OR $_{(random)}=0.61$ (0.25 to 1.51, p=0.29, n=3 studies, I²=70% P_(heterogenity)=0.04). All results were still not statistically significant when studies were removed in turn. Of note, the removal of Wengreen (50 to 59 years old group) did make heterogeneity go down to zero suggesting this study was the cause of the heterogeneity observed. There were not enough studies to perform funnel plots for each of the fracture meta-analyses.

[4] Supplementary Results: Intervention Studies

Study Characteristics

The 30 intervention studies included 2 studies in peri-menopausal women(99, 100), 6 in premenopausal women(101-106), 12 in postmenopausal women(107-118), 6 in both men and women, of which 4 were elderly groups (119-122) and 2 younger or middle-aged(123, 124), as well as 2 studies in men alone(125, 126) and 2 in children(127, 128). Seventeen of the 30 studies were from USA or Canada (99, 101, 102, 107-109, 112-117, 119, 120, 123-

125), with 2 from Switzerland(121, 122), 2 from Australia(111, 118), 1 from Brazil(110), 1 from Spain(128), 1 from New Guinea(127), 5 from Japan(100, 103, 104, 106, 126) and 1 from China(105). Eight of the 30 studies presented data on soy protein intervention vs. non-soy protein control (99, 107, 111, 112, 114, 116, 117, 125), 6 presented data on milk basic protein (MBP) vs. control (100, 103-106, 126), and 7 studies presented data on other protein types (110, 118, 120-122, 127, 128). Finally, 9 studies assessed dietary interventions, comparing higher vs. lower protein intakes (101, 102, 108, 109, 113, 115, 119, 123, 124).

Jadad Scores

See Supplementary Table x for full details of study quality and risk of bias in the 30 intervention studies. In terms of Jadad scores (0-5, 5=highest quality), 16 studies were scored as 0-2(101, 102, 104-106, 108, 111, 113, 115, 116, 119, 121, 123, 126-128), 7 studies as 3-4(99, 100, 107, 114, 120, 124, 125) and 7 studies as 5(103, 109, 110, 112, 117, 118, 122). Some studies (101, 102, 108, 109, 113, 115, 119, 124) may have scored lower than expected due to being dietary intervention studies, whereby it is difficult to undertake participant blinding as the different diets are difficult to conceal. Indeed, the Jadad scale is really intended for quality assessment of studies were participants and investigators can feasibly be blinded to treatment allocation. Study quality was variable, with many studies having significant methodological flaws.

Intervention Studies

Non-dietary Studies- Bone markers

In soy protein (vs. non-soy protein control) studies, six studies found no effect of soy protein on BAP(99, 107, 114, 125), NTX(114, 117), DPYD or PYD(107, 111, 125). However there

was an effect of soy protein vs. milk protein on reducing both BAP and CTx in one study(112) and in another study there was a positive effect of soy protein on raising BAP and OC, but no effect on NTX(116). Of the three MBP studies reporting bone marker data, one study found no effect of MBP supplementation vs. inactive placebo on NTX and OC concentration(100), another study found a lower NTX and higher OC in the MBP group than the inactive placebo control(126) and the final study found lower NTX and DPYD in the MBP group than inactive placebo(106).

For total protein (vs non-protein control), protein supplementation was associated with increases in type 1 N-terminal procollagen (P1NP)(120), HPO(124), DPYD(122, 124) and PYD(122, 124). There were conflicting results for CTX and OC with three studies finding no effect (110, 120, 122) one study finding a reduction in CTX (120) and one finding an increase in OC (121). Finally, one study found no statistically significant for an effect of protein supplementation on BAP (110). The only study in children found no statistically significant difference in BAP or OC or tartrate-resistant acid phosphatase (TRAP), but there was a higher increase in CTX over the study period in controls (increase by 6% of baseline value) than in the collagen supplemented group (increase by 3% of baseline value).

Dietary Studies- Bone markers

Seven diet studies found no differential effect of high and low protein diets on CTX (109, 115), OC(101, 102, 109, 115, 119), DPYD(108) or NTX(108). Four studies found a lower NTX (101, 102, 119, 123) and two studies found that DPYD was lower(113) or HPO was higher(115) in those taking a high protein diet (compared with a low protein diet). Finally, two studies conflicted in that they found either a higher BAP(102) or no difference in BAP(123) in those with a low protein compared with those with moderate or high protein diets.

Soy and MBP protein and BMD/BMC

In all supplementation studies no differential effect of soy protein vs. non-soy protein was seen for LSBMD(99, 112, 114, 117), LSBMC(99), FNBMD(112, 114, 117), RBMD(114) or TBBMD(112, 114, 117). Also, in a food based study (107) there was no differential effect of soy or non-soy protein on both BMD and BMC at the LS, TB and TH. (107). For MBP, a statistically significant effect of MBP supplementation in increasing LSBMD was found in two studies by 1 - 1.6%(100, 103) and in one study increasing TBBMD by 2%(105). However other results for MBP were conflicting, with studies finding either increased RBMD(104) or no effect on RBMD LSBMD or TBBMD (105).

Total protein and BMD/Bone size

For total protein, in malnourished New Guinea children aged 7-13 years, one study found an effect of 20g/d milk protein supplementation (vs. no supplement) for increased periosteal breadth, but not endosteal or compact bone breadth (127). In adults, no effect was seen for protein supplementation vs. non-protein control on LSBMD(120-122), THBMD(118, 120), FNBMD(118, 120, 121) FSBMD(121, 122) or TBBMD(122). Finally, one study found no differential effect of high vs low protein supplement drink on TH vBMD or FN vBMD(118).

Meta-analysis: MBP and Soy Protein Sensitivity analysis

Elimination of each MBP study in turn gave the following pooled estimates: Aoe 2005 $MD_{(fixed)}= 0.01 (-0.04 \text{ to } 0.07) \text{ R}^2 < 0.001 \text{ P}=0.69, \text{ I}^2=0\% \text{ P}_{(heterogenity)}=0.69; \text{ Uenishi } 2007$ $MD_{(fixed)}=0.02 (-0.003 \text{ to } 0.04) \text{ R}^2=0.0004 \text{ P}=0.10, \text{ I}^2=0\% \text{ P}_{(heterogenity)}=0.63; \text{ Zou } 2009$ $MD_{(fixed)}= 0.02 (-0.002 \text{ to } 0.04) \text{ R}^2=0.04 \text{ P}=0.07, \text{ I}^2=0\% \text{ P}_{(heterogenity)}=0.07.$ Elimination of each soy protein study in turn gave the following pooled effect sizes: Alekel 2000 $MD_{(random)}=0.02 (-0.07 \text{ to } -0.12, P=0.61) I^2=52\% P_{(heterogenity)}=0.15; Kenny2009 MD_{(fixed)}= -0.03 (-0.07 \text{ to } 0.02, P=0.23) I^2=8\% P_{(heterogenity)}=0.30; Vupadhyahula 2009 MD_{(random)}=0.01 (-0.14 \text{ to } 0.15, P=0.93) I^2=75\% P_{(heterogenity)}=0.04. Removal of Kenny 2009 reduced$ heterogeneity from 51% to 8%, suggesting this study was contributing to the heterogeneity toa large degree. There were not enough studies to produce funnel plots for these metaanalyses.

Table S1: Characteristics and outcomes of 74 cross-sectional and/or longitudinal correlational studies

Study	Mean Protein **	Method	Population	n	Outcome	Coefficient*	Р
Alexy et al, 2005, Germany	Prepubescent (M and F)- 2.0+/-0.3	pQCT	Prepubescent and pubescent	229		Standardized Beta: protein g/d, adjusted for age, sex, energy intake	
	g/Kg/d		boys and girls		Periosteal Circumference	0.170.27	0.0014
	Pubescent (M)-				Cortical Area	0.26	0.0001
	1.6+/-0.3 gKg/d				BMC	0.29	0.0011
	Pubescent(F)- 1.4+/- 0.3 g/Kgd				Polar SSI		<0.0001
Alissa et al, 2011,	1.03 g/Kg/d	DXA	Women aged	122	Protein intake	Mean (SEM): g/d	
Saudi Arabia			46-70 years			Control: 77.5 (3.15) n 61	NS
			old			Osteopenic: 76.6 (2.92) n 61	
Alissa et al, 2014,	71.4+/-1.55 g/d	DXA	Postmenopaus	300		Energy adjusted protein intake:	
Saudi Arabia			al women,			r values:	
			aged 46-88		LSBMD	-0.021	0.722
			years		FNBMD* used pooling	0.182	0.002
					TotalHipBMD	0.244	
							<0.0001
Beasley et al.	TP: 5.7 - 27.6%	DXA	Females aged	560		Tertile of protein intake %total energy	
2010, USA	energy AP:45g/d		14-40 years			BMD:(Mean, 95% CI)	
	VP:19g/d				TP:	T1 (lowest) T3 (highest)	0.94
	e				TotalHipBMD	0.93(0.91,0.95) 0.93 (0.91,0.96)	0.37
					LSBMD	1.00 (0.98,1.02) 1.02(1.00,1.04)	0.98
					TBBMD	1.08(1.07, 1.09) 1.08(1.06,1.10)	
					AP		
					TotalHipBMD	0.93(0.91,0.95) 0.94(0.92,0.96)	0.99
					LSBMD	1.00(0.98,1.02) 1.02(1.00,1.04)	0.40
					TBBMD	1.08(1.07,1.09) 1.08(1.07,1.10)	0.80
					VP		
					TotalHipBMD	0.92(0.90,0.94) 0.94(0.92,0.96)	0.03
					LSBMD	1.00(0.98,1.01) 1.01(0.99,1.04)	0.10
					TBBMD	1.07(1.06,1.11) 1.08(1.06,1.09)	0.04
						Beta for increment of protein as an extra	
						1% energy, adjusted for age, BMI,	

						physical activity, smoking, contraception,	
						energy intake, phosphorus, magnesium.	
					N 224:	TP % energy (Year 3)Beta= -0.0002	
					3 year change in:	Beta= 0.0004	
					e jeur enunge mi	Beta= -0.0012	P value
					HipBMD	<i>Dom</i> 0.0012	i value
					SpineBMD	AP % energy (Year 3)	0.88
					TBBMD	Beta= -0.0002	0.71
					Ibbiib	Beta= 0.0005	0.19
						Beta= -0.0011	0.17
					HipBMD		
					SpineBMD	VP % energy (Year 3)	0.87
					TBBMD	Beta= -0.0023	0.69
					1221112	Beta= -0.0019	0.21
						Beta= 0.0009	0.21
					HipBMD		
					SpineBMD		0.40
					TBBMD		0.50
							0.69
Beasley et al.	15% total energy	DXA	Postmenopaus	144,580		Change in mean BMD per 20% increase in	
2014, USA			al women 50-	,		% of calories from protein:	
,			79 years			1	
			5		TBBMD	At 6 y (n=6552), change in BMD of 0.004	-
						(0.001, 0.007) g/cm2	
					Hip BMD	At 6 y (n=6553) change in BMD of 0.003	-
						(0.000, 0.005)	
					Spine BMD	At 6y (n=6457), change in BMD of	-
						0.003(0.000,0.008)	
Bounds et al,	55g/d (1.9g/Kg/d)	DXA	6-8 year old	25 Boys,		Unadjusted r values- Pearson's	
2005, USA			children	27	TBBMC	0.37	≤0.05
				Females	TBBMD	0.33	≤0.05
					TBBMC	Stand.Beta=2.40*	0.008
					TBBMD	Stand. Beta=0.001**	0.04
						*adjusted for Height, Weight, age and sex	
						**adjusted for Sex	

2.67	Bone	N=81 pubertal	81	TP	Standardized beta: Age and BMI adjusted	
	turnover	boys		sOC microg/L	0.09	0.68
	markers	•		sBAP U/L	0.89	0.01
				sCTX microg/L	<-0.01	0.59
				VP		
				sOC microg/L	0.24	
						0.36
						0.72
				50111 moreg, 2		0.29
				Dairy protein		
				sOC microg/L	-0.45	0.05
				sBAP U/L	0.53	0.16
				sCTX microg/L	-0.01	0.51
				Meat protein		
					0 44	0.11
						0.04
						0.35
TP 12 (Girls) 13	DXA	17-year-olds:	109			0.55
	Diffi		109			0.78
						0.46
		40 00 93		Lobine	0.00	0.40
				ΔP·		
· ·					0.01	0.62
and Doys)						0.78
				LSDIVIC		0.78
77 5 ald Hong Kong		Dramananausa	441			
	DAA	-	441			
05.4g/u Deijing		i women			DIVII)	0.250
					0.102	0.359
						ns
				1 otaiSpineBMD		ns
					-0.094	
1.3 g/Kg/d	DXA	Older men and	2217	Energy adjusted protein	B coefficient (adjusted for age, weight,	
		women		intake		
					physical activity, calcium supplement,	
	TP: 1.2 (Girls), 1.3 (Boys) AP: 0.4 (Girls), 0.5 (Boys) DP: 0.4 (Both Girls and Boys) 77.5g/d Hong Kong 65.4g/d Beijing	TP: 1.2 (Girls), 1.3 DXA (Boys) AP: 0.4 (Girls), 0.5 (Boys) DP: 0.4 (Both Girls and Boys) 77.5g/d Hong Kong DXA 65.4g/d Beijing	TP: 1.2 (Girls), 1.3DXA17-year-olds: 63 girls and 46 boys(Boys)63 girls and 46 boysAP: 0.4 (Girls), 0.546 boys(Boys)DP: 0.4 (Both Girls and Boys)77.5g/d Hong Kong 65.4g/d BeijingDXAPremenopausa l women1.3 g/Kg/dDXAOlder men and	TP: 1.2 (Girls), 1.3DXA17-year-olds:109(Boys)63 girls andAP: 0.4 (Girls), 0.546 boys(Boys)DP: 0.4 (Both Girlsand Boys)77.5g/d Hong KongDXAPremenopausa4411.3 g/Kg/dDXAOlder men and2217	SCTX microg/L VP sOC microg/L sBAP U/L sCTX microg/L Dairy protein sOC microg/L sBAP U/L sCTX microg/L sAP U/L sCTX microg/L sBAP U/L sCTX microg/L sBAP U/L sCTX microg/L sBAP U/L sCTX microg/L sBAP U/L sCTX microg/L sAP U/L sCTX microg/L sBAP U/L sCTX microg/L sBAP U/L sCTX microg/L sBAP U/L sCTX microg/L sCTX microg/L s	SCTX microg/L <-0.01

						energy adjusted calcium and vitamin D	
					% change Hip BMD	intakes)	0.147
					% change FNBMD	Men:	0.006
					C	B= -0.007	
						B= -0.013	
						Women: data not reported (all ns)	
Chevalley et al.	47.3 g/d, 1.78	DXA	Prepubertal	232		Protein intake g/d:	Р
2008, Switzerland	g/Kg/d		boys			r (not adjusted)	
,	0 0				Radial Metaphysis BMC	0.26	0.0001
					Radial Diaphysis BMC	0.21	0.002
					Total Radius BMC	0.27	0.0001
					FNBMC	0.20	0.002
					Total Hip BMC	0.18	0.002
					Femoral Diaphysis BMC	0.23	0.0003
					LSBMC	0.24	0.0003
					LSDIMC	0.24	0.0002
						Standardized Beta (adjusted for physical	
						activity and calcium intakes)	
					Radial Metaphysis BMC	0.201	0.013
					Radial Diaphysis BMC	0.120	0.146
					Total Radius BMC	0.199	0.013
					FNBMC	0.187	0.028
					Total Hip BMC	0.122	0.136
					Femoral Diaphysis	0.190	0.025
					LSBMC	0.217	0.009
						Data for <median activity="" only<="" physical="" td=""><td></td></median>	
						shown:	
					Mean(SD)	Protein>median vs. <median:< td=""><td></td></median:<>	
					Radial Metaphysis BMC	649(82) vs. 663(103)	-
					Radial Diaphysis BMC	919(104) vs. 937(104)	-
					Total Radius BMC	2679(379) vs. 2807(422)	-
					FNBMC	1980(321) vs. 1988(321)	-
					Total Hip BMC	10342(1958) vs. 10535(1973)	-
					Femoral Diaphysis	17575(3698) vs. 18431(3486)	-
					LSBMC	15652(2080) vs. 15839(2505)	-

Chevalley et al. 2014, Switzerland	Age 7: 1.8; Age 15: 1.1 (g/Kg/d)	High resolution pQCT	Adolescent boys	176	Bone outcomes at 15 years: FN vBMD TotalHipvBMD FNBA FN width FNBMD (DXA) TotalHipBMD (DXA) DistalTibia Total vBMD	Protein intake at Age 7 years: Higher (n=36) vs lower protein (n=52) (lower physical activity) 4645±788 vs. 4411±795 36389±7995 vs. 34381±7493 5.28±0.50 vs. 5.18±0.47 3.49±0.33 vs. 3.43±0.31 879±109 vs. 846±112 976±127 vs. 937±130 276±39 vs. 259±44	0.176 0.233 0.341 0.341 0.178 0.169 0.063
					FN vBMD TotalHipvBMD FNBA FN width FNBMD (DXA) TotalHipBMD (DXA) DistalTibia Total vBMD	Higher (n=49) v s lower protein(n=38) (higher physical activity) 5075±894 vs. 4405±858 40913±8451 vs. 35303±7863 5.46±0.36 vs. 5.26±0.47 3.61±0.24 vs. 3.48±0.31 932±139 vs. 834±122 1011±140 vs. 929±144	0.0006 0.002 0.030 0.030 0.0009 0.009 0.336
Chiu et al, 1997, Taiwan	1.09	DPA (BMD)	Older post F	258	LSBMD FNBMD	273±41 vs. 263±53 r Protein g/d (unadjusted- Pearson's values) 0.107 0.085	0.09 0.18
Coin et al, Italy, 2008	75.8+/-22.1 g/d Weight=74.2+/-13.4 So 1.02 g/Kg/d	DXA	Males, mean age 73.9+/-5.6 years	136	Male data only for protein (no data for females) n=136 Total Hip BMD	0.12(non adj) p<0.001 0.06(adj) p<0.01 r(adj)=0.25	Controlling for BMI, albumin, skeletal muscle, age
					FNBMD* chosen for pooling men as same as other studies	0.03(nonadj) p<0.05 0.01(adj) p>0.05 r(adj)=0.1	
					TrochBMD	0.10(nonadj)p<0.001 0.08(adj) p<0.01 r(adj)=0.28	

	(BMD)	Pre (72) and post (218) F	290		Adjusted for age, weight, physical activity	P for unadj data Ns
	· /	Γ		LSBMD(pre)	0.20 adj=0.07 ns	< 0.01
				TrochBMD (pre)	0.36 $adj=0.35 p<0.01$	< 0.05
						<0.01
						< 0.05
				.		Ns
						Ns
						<0.01
						<0.001
						<0.01
					5	<0.01
						<0.001
						0.001
				i obiiib (post)		
				HPO(pre)*		
70a/d	DYA	184 men and	18/	OC(post)		
1)g/u	DAA		104		Tertile protein make, <i>w</i> of energy	
	3	years old) in		TRRMD	$T_{artile} = 1 + 1 + 12(0 + 13)$	
				IBDND		20
						ns
					10100 - 5 - 1.07(0.14)	
				ENDMD	$T_{out} = 1, 0, 80(0, 14)$	
				FINDIVID		20
		on that				ns
					1000000000000000000000000000000000000	
				LSBMD	Tertile 1 1.17(0.23)	
				202112		ns
						115
					Tertile 2 1.11 (0.23)	
				sOC (nmol/L)	Tertile 1 1.1(0.3)	
						ns
				uNTX (nmol)		
					Tertile 1 231(172)	
						ns
	79g/d	79g/d DXA	women(>=65	women(>=65 years old) in placebo (inactive) arm of calcium supplementati	79g/d DXA 184 men and 184 women(>=65 years old) in TBBMD 79g/d DXA 184 men and 184 women(>=65 years old) in TBBMD 79g/d DXA 184 men and 184 women(>=65 years old) in TBBMD	FNBMD(Pre) 0.26 adj=0.27 p<0.05 DRBMD (pre) 0.35 adj=0.28 p<0.01 MRBMD(pre) 0.27 adj=0.21 p<0.05 FSBMD(pre) 0.22 adj=0.16 ns adj=0.05 ns FNBMD(post) 0.13 adj=-0.05 ns 1000 ns FNBMD(post) 0.20 adj=-0.06 ns 1000 ns FNBMD(post) 0.21 adj=-0.05 ns 1000 ns FNBMD(post) 0.21 adj=-0.06 ns 1000 ns FNBMD(post) 0.21 adj=-0.05 ns 1000 ns FNBMD(post) 0.21 adj=-0.05 ns 1000 ns FSBMD(post) 0.24 adj=0.01 ns 1000 ns FSBMD(post) 0.24 adj=0.01 ns 1000 ns FSBMD(post) 0.25 p<0.01

Devine et al, 2005, Australia	1.2	DXA , QUS	Elderly F mean age 75y+/-3y	1077	TP: Total Hip BMD BUA calcaneus	r values (unadjusted) 0.138 0.136	-
			Caucasian		TP:	Unstandardized Beta (SE):	
					TotalHipBMD	0.31 (0.07)	<0.01
					FNBMD	0.26 (0.07)	<0.05
					TrochBMD	0.32 (0.08)	<0.01
					IntertrochBMD	0.32 (0.06)	<0.05
					BUA calcaneus	0.02 (0.08)	<0.01
					SOS	Not shown	NS
					Stiffness	0.02 (-0.06)	NS
					Sumess	0.02 ((0.00)	110
						BUA (db/Hz): <66 g/d: 99.6 ± 0.4 (n = 357)	
						$66-87$ g/d: 100.8 ± 0.4 (n = 337)	
						>87 g/d: 101.2 ± 0.4 (n = 341)	
						BMD at Hip Sites (mg/cm ²)	
						Tertile 1 <66 g/d (n = 374)	
						Tertile 2 66-87 g/d (n = 350)	
						Tertile 3 >87 g/d (n = 351)	
						Mean SE 95%CI	
					Total Hip BMD	Tertile 1 0.798 0.006 0.79, 0.81	
						Tertile 2 0.815 0.006 0.80, 0.83	
						Tertile 3 0.823 0.006 0.81, 0.84	
					Femoral Neck BMD	Tertile 1 0.679 0.005 0.67, 0.69	
						Tertile 2 0.695 0.005 0.69, 0.71	
					Trochanter BMD	Tertile 30.7010.0050.69,0.71Tertile10.6250.0050.62,0.64	
					Hochanter BMD	Tertile 2 0.640 0.005 0.63, 0.65	
						Tertile 3 0.649 0.005 0.64, 0.66	
					Intertrochanter BMD	Tertile 1 0.937 0.007 0.92, 0.95	
						Tertile 2 0.957 0.007 0.92, 0.95 Tertile 2 0.957 0.007 0.94, 0.97	
						Tertile 3 0.964 0.007 0.95, 0.98	
Ekbote et al,	18.6g/d-normal and	DXA	2-3 year old	71	Normal children:		
2011, India	malnourished		children		TBBMC	0.62*	<0.01
	children combined				TBBA	0.65*	<0.01

					Malnourished Children:		
					TBBMC	0.44*	
					TBBA	0.57*	< 0.05
							< 0.01
					All children:		
					TBBMC	0.55*	
					TBBA	0.58*	< 0.01
						*remained statistically significant when	< 0.05
						adjusted for energy intake	
Fairweather-Tait	81.3g/d	DXA	Postmenopaus	2464	Energy adjusted protein	Beta(adjusted for age, BMI, smoking,	P (adjusted for
et al, 2011, UK	C		al female twin	pairs	intake (g):	physical activity),	multiple
, ,			pairs	1		Including variables for individual diet and	comparisons)
			(Monozygotic			twin difference:	1 /
			or dizygotic				0.651
			twins)		LSBMD (n=1232 pairs)	0.029 (-0.014, 0.072)	0.964
			,		HipBMD(n=1218 pairs)	-0.013 (-0.047, 0.022)	0.365
					FNBMD (n=1019 pairs)	-0.033 (-0.071, 0.005)	
						Individual intakes only in model:	0.502
					LSBMD (n=1232 pairs)	0.012 (-0.023, 0.046)	0.738
					HipBMD(n=1218 pairs)	-0.005 (-0.036, 0.025)	0.102
					FNBMD (n=1019 pairs)	-0.027 (-0.060, 0.005)	
Freudenheim et	1.02	SPA	Pre and post F,	84 (17		r, p	Adjusted for bone
al, 1986, USA			35-65y,	pre F, 67	RBMC (pre)	0.384, 0.128	width
, ,			Caucasian	post F)	HumBMC (pre)	0.157, 0.546	
				1	UBMC (pre)	0.282, 0.272	
					RBMC (post)	-0.017, 0.889	
					HumBMC (post)	0.138, 0.267	
					UBMC (post)	0.044, 0.725	
						Slope of bone loss:	
					TP	Pre F: r, p	
						RBMD 0.742, 0.022	
						HuBMD 0.518, 0.153	
						Ulna 0.428, 0.250	
l					TP	Post F: r, p	
					11	RBMD 0.493, 0.004	
						HuBMD 0.258, 0.147	
						110DIVID 0.230, 0.17/	

						Ulna -0.095, 0.597	
Geinoz et al,	Mean Intake in g/d	DXA	Elderly M and	74	Gender, protein intake	<u>Mean +/-SD</u>	<u>P</u>
1993, Switzerland	by group: 37.8-59.4		F		E > 1 - NZ - /4		
	57.8-39.4		Mean age 82y(F); 80(M)		<u>F,>1g/Kg/d</u> FNBMD	0.679+/-0.09	Ns
			$02y(\Gamma), 00(101)$		FSBMD	1.288+/-0.35	Ns
					SpineBMD	0.935+/-0.24	Ns
					SpineBMD	0.933+7-0.24	188
					<u>F<1g/Kg/d</u>		
					FBBMD	0.574+/-0.13	p<0.05
					FSBMD	1.120+/-0.33	ns
					SpineBMD	0.877+/-0.36	ns
					M,>1g/Kg/d		
					FNBMD	0.761+/-0.12	ns
					FSBMD	1.516+/-0.19	ns
					SpineBMD	1.094+/-0.26	ns
					M < 1 - M - 14		
					<u>M,<1g/Kg/d</u> FNBMD	0.643+/-0.14	≈ <0.05
					FSBMD	1.318+/-0.34	p<0.05
					Spine BMD	0.847+/-0.18	ns p<0.05
Comoro et al	66-11	DXA	Women over	200	Spille BMD		1
Genaro et al, 2015, Brazil	66g/d	DXA	65 years old	200		Protein:g/Kg/d <0.8 (n=73)	<u>P for trend:</u>
2013, D1a2ii			05 years old		TBBMCg/cm2	0.988	0.011
					LSBMD g/cm2	0.903	0.014
					FNBMD g/cm2	0.760	0.017
					TrochBMD g/cm2	0.679	0.071
					Total Femur BMD g/cm2	0.807	0.026
						0.8-1.2 (n=84)	Posthoc tests:
					TBBMCg/cm2	1.025	>1.2 vs. <0.8
					LSBMD g/cm2	0.965	P < 0.05 at all sites
					FNBMD g/cm2	0.795	1 (0.05 at all sites
					TrochBMD g/cm2	0.689	>1.2 vs 0.8-1.2
					Total Femur BMD g/cm2	0.833	>1.2 vs 0.8-1.2 P<0.05 for
					rotar remui Divid g/clil2	0.035	TBBMC,
						>1.2 (n=43)	LSBMD and
					TPPMCalam2	>1.2 (n=43) 1.039	ESBMD and FNBMD
					TBBMCg/cm2		LINDMD
					LSBMD g/cm2	0.983	

					FNBMD g/cm2 TrochBMD g/cm2 Total Femur BMD g/cm2	0.813 0.727 0.868	
Gregg et al, 1999, USA	0.9	QUS	Middle aged (premenopaus al) F- mean age= 45.5y	393	BUA Calc SOS Calc LSBMD FNBMD	Unadjusted coefficients (non adjusted) 3.15 0.96 0.015 0.010	0.0008 0.02 0.02 0.09
					Dietary protein: per 87kcal BUA	Controlling for lean body mass, physical activity, race, menopausal status, BMI: 0.14 SD increase	0.004
Gunn et al, 2014, New Zealand	79g/d	Bone markers, DXA	Postmenopaus al women, 60 years of age	142	FNBMD FN T-Score	Energy adjusted protein: (not adjusted for other confounders) 0.19 0.17	<0.05 <0.05
Hannan et al. 2000, USA	68g/d (16% of total energy) 0.97 g/kg/d	DXA	Older men and women	615	sCTX sP1NP <u>TP</u>	-0.18 -0.23 Change in BMD by protein quartile:	<0.05 <0.05 *= p<0.05 Q1 compared with
2000, USA	chergy) 0.97 g/kg/u		women		FNBMD QI	-4.61 +/- 070*	Q4
					Q4 TrochBMD	-2.32 +/-0.74	Adjusted for total energy intake, age, sex, weight,
					Q1 Q4	-8.00 +/- 0.84 -6.65+/-0.90	weight change, height, alcohol intake and
					Wards BMD Q1 Q4	-7.05+/-1.0 -4.39+/-1.1	smoking (current or former).
					LSBMD Q1	-3.72+/-0.97	
					Q4 RBMD	-1.11+/-1.1	
					Q4	-4.31+-0.76	
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					AP		
					FNBMD		
					QI	-3.95 +/- 0.69*	
					Q4	-2.15+/-0.73	
					TrochBMD		
					Q1	-2.57+/-0.86	
					Q4	-1.95+/-0.92	
					Wards BMD		
					Q1	-4.02+/-1.0	
					Q4	-1.97+/-1.1	
					LSBMD		
					Q1	-3.79+/-0.99	
					Q4	-1.65+/-1.1	
					RBMD		
					Q1	-4.60+/-0.71	
					Q4	-4.52+/-0.76	
Henderson et al,	1.0	DXA	Pre F- mean	115		Unadjusted r values	
1995 , Australia			age=18y		FNBMD	0.22	p<0.05
					IntertrochBMD	0.19	p<0.05
					TrochBMD	0.27	p<0.005
					DTB BMD	0.05	p>0.05
					TFBMD	0.21	p<0.05
					FSBMD	0.09	p>0.05
			D	201	LSBMD	0.05	p>0.05
Hernandez et al,	76g/d	SPA	Pre- and	281	Ultradistal R BMD	Beta=0.0108 SE=0.259 (unstandardized	NS
1993, USA			Perimenopaus			beta)	
			al Women			Adjusted for dietary nutrients, alcohol and	
			(50-60 years old)			caffeine.	
Hirota et al, 1992,	1.13	SPA	Young pre F:	161	Forearm BMD	r=0.0017 (adjusted for sports, BMI, milk	0.03
Japan		(BMD)	19-25y			intake in childhood, dieting, skipping	Adjusted for
						meals)	sports, BMI,

						Dietary intakes g/d by Forearm BMD category (BMD % of mean) <=85% 50.7+/-13.6 86-100% 56.8+/-13.3 101-114% 60.1+/-18.2* >=115% 64.2+/-19.7* *significantly different from the <=85% group (lowest)	childhood milk intake, dieting, skipping meals
Ho et al, 2003, 1.(China Soy protein	1.01 SP	DXA	<12y post F(48-62y), Asian	454 (269 <4 y post F 185 >4 y post F)	ALL WOMEN Spine BMD FNBMD TrochBMD IntertrochBMD TotalHipBMD TBBMD TBBMD	Quartile of soy protein intake:Q1Q4 0.825 ± 0.118 0.844 ± 0.133 0.668 ± 0.103 0.694 ± 0.099 $0.581\pm0.098*$ 0.606 ± 0.095 $0.945\pm0.145*$ 0.981 ± 0.130 $0.781\pm0.118*$ 0.815 ± 0.111 0.958 ± 0.088 0.966 ± 0.084 1601 ± 255 1649 ± 228	
					Spine BMD FNBMD TrochBMD IntertrochBMD TotalHipBMD TBBMD TBBMC	Standardized beta (SE) 0.0034(0.005) 0.0048(0.004) 0.0056(0.004) 0.0069(0.005) 0.0070(0.004) 0.0071(0.004) 5.974 (8.784)	0.497 0.200 0.119 0.162 0.087 0.842 0.497
Ho et al, 2008, China	5.2g/d SP 48.6 g/d TP	DXA	Pre and perimenopaus	438		Controlling for soy protein intake quartile, weight, years since menopause, calcium intake quartile, soy protein- calcium interaction, total protein intake, and energy intake r(adj)=adjusted for age-menopause stage and energy intake	

al women 45-	TP:		
55 years old	LSBMD	r=0.064	n 0
55 years old	LSBMD	r(adj)=0.016	ns
	FNBMD	r=0.088	ns
	FNBMD	r(adj)=0.037	ns
	TotalHipBMC	r=0.084	ns
	Тоганирымс	r(adj)=0.053	ns
	WBBMC	r=0.075	ns
	WBBINE	r(adj)=0.024	ns ns
		1(auj)=0.024	115
	SP:		
	LSBMD	r= -0.043	ns
		r(adj) = -0.05	ns
	FNBMD	r=0.020	ns
		r(adj) = -0.004	ns
	TotalHipBMC	r= -0.001	ns
		r(adj) = -0.027	ns
	WBBMC	r = -0.002	ns
		r(adj) = -0.017	ns
	Quartile of soy protein	Standardized beta (SE)	
	intake:	Q1(<1.07)=Reference	-
	WBBMC	Q2(1.07-)=0.19 (0.3282)	ns
		Q3(2.85-)=0.73 (0.3340)	< 0.05
		Q4(5.72+)=0.73 (0.3225)	<0.05
	Change from baseline:	Quartile of soy protein intake vs TBBMC:	
	(30 months)	Unst. Beta* SE	
	TBBMC		
	Q1 <1.07 g/d Reference	1.0000	
	Q2 1.07-2.84 g/d	0.1932 0.3282	ns
	Q3 2.85-5.72 g/d	0.7306 0.3340	< 0.05
	Q4 >5.72 g/d	0.7303 0.3225	< 0.05
		*Controlling for baseline BMC, lean mass,	
		change in weight, number of pregnancies,	
		walking and menopausal status.	

Ho-Pham et al,	g/d:	DXA	105 Post F	210		AP: +10g Beta	
2009, Vietnam	$\overline{\text{TP}}=35.4(11.6)$		Buddhist		FNBMD	0.008 (0.006)	0.175
,	Vegans, 62.6(18.3)		vegan Nuns		LSBMD	0.013 (0.008)	0.108
	Omnivores		and 105 omnivorous		TBBMD	0.006 (0.006)	0.313
	AP=2.1(3.2)		women 62+/-			VP:+10g Beta	
	Vegans, 34.6(15.8)		10 years old		FNBMD	-0.008 (0.007)	0.261
	Omnivores		•		LSBMD	0.014 (0.009)	0.128
					TBBMD	-0.014(0.006)	0.033
	VP=33.2(11.6)Vega ns, 28.0(8.4) Omnivores						
Ho-Pham et al,	TP:	DXA	105 Buddhist	181		Beta: (adjusted for age, anthropometry, fat	
2012, Vietnam	36g/d Vegans		vegan Nuns		Change in:	intake)	
,	62g/d Omnivores		and 105		LSBMD	VP: -0.075(0.035)	0.036
			omnivorous women Mean(SD) age		FNBMD	AP:VP ratio: -0.244 (0.094)	0.01
		DUA	61(9.2)	105			.0.001
Hoppe et al.	82g/d (Boys)	DXA	10 year old	105	WBBMC	0.327 (unadjusted r values)	< 0.001
2000, Denmark	73g/d(Girls)		children		WBBA	0.311 Has linear regression but only p values, not effect size p1027	<0.01
Horiuchi et al,	Total- 62.5g/d	DXA	Post F, 52-83y	85	Soy	r values (unadjusted)	
2000, Japan	Soy-12.6g/d	Diffi	10001,02000	00	LSBMD	0.251	p<0.05
2000, Jupun	509 12.0g/a				Osteocalcin	-0.097	ns
					ALP	-0.017	ns
					Pyridinoline	-0.132	ns
					Deoxypyd	-0.229	p<0.05
					Total Protein		r
					LSBMD	0.223	p<0.05
					Pyridinoline	-0.229	p<0.05
					Deoxypyd	-0.218	ns
					Osteocalcin	-0.131	ns
					ALP	-0.09	ns
					Z score for LSBMD	Beta (standardized):	
					Total protein	-0.03 (no SE or p value given)* used in SR	ns
					Soy protein	0.225 (no SE given)	0.038

Hu et al, 2014, USA	TP: 12.0-19.0 % energy intake (F) TP: 11.6-20.4% energy intake (M)	QCT	801 women and 857 men enrolled on the Multi-Ethnic Study of	1658	LS vBMDmg/cm ³ n1658	Quartil Q1 11 Q2 11 Q3 11 Q4 11	5+/-38 6+/-42	P=0.88
			Atherosclerosi s (age 62+/-10		LSBMD ZScore: n801 TP:	Standa	rdized Beta, p	
			years)		White	-0.06	0.75	
					Chinese	-0.07	0.84	
					Black	0.35	0.2	
					Hispanic	0.16	0.55	
					AP:			
					White	-0.13	0.51	
					Chinese	-0.65	0.06	
					Black	0.29	0.37	
					Hispanic	0.40	0.16	
					VP:			
					White	0.44	0.02	
					Chinese	0.22	0.49	
					Black	-0.36	0.27	
					Hispanic	0.06	0.9	
						activity educati menop energy	adjusted for age, BMI, physical y, sedentariness, smoking, ion, hormone therapy use, age at ause, and intakes of total , dietary carbohydrate as a tage of energy, Ca, P, Mg and l.	
					LS trabecular vBMD	Quartil CI)	es of protein intake: Q4 mean (96%	
					Female, AP White Chinese Black	107.5 (134.9 (7.7, 106.8) (68.0, 146.9) (117.3, 152.4) (105.2, 135.8)	

					· · ·		
					Hispanic		
					Male AP White Chinese Black Hispanic	110.3 (100.8, 119.7) 115.9 (99.5, 132.2) 155.7 (135.3, 176.1) 128.8 (117.3, 140.3)	
					Female, VP White Chinese Black Hispanic	107.4 (98.6, 116.2) 139.4 (102.5, 176.2) 125.8 (110.3, 141.4) 112.8 (100.5, 125.2)	
Ilich et al, 2003, USA	1.04	DXA	Older F, >5 post, Caucasian,	136	Male VP White Chinese Black Hispanic	105.0 (97.1, 112.9) 109.6 (94.8, 124.4) 141.6 (122.9, 160.3) 123.6 (112.6, 134.6) Unstandardized. Beta (adjusted for age, lean body mass, total body fat, and height (in TBBMC model), past physical activity,	
			mean age 68.7+/-7.1y		TBBMD TBBMC	present mode of walking, and energy intake 1 x10 ⁻³ (also adjusted for Ca intake) 2.9	0.027 0.03
Iuliano-Burns et al, 2005, Australia	76g/d	DXA	7-20 year old Male twins (Monozygotic	56	WBMD HBMD	1.4x10 ⁻³ (also adj for ca and vit C intake) 4.1x10 ⁻⁴ (also adjusted for ca intake) Beta coefficient: Within pair difference in protein intake, adjusted for anthropometric and lifestyle factors.	0.021 0.021
			n=30) and Dizygotic (n=26)		TBBMC (g) Arm BMC (g) Leg BMC (g) LSBMC (g)	1.3 0.7 0.3 0.0	ns <0.05 ns ns
Jaime et al, 2006, Brazil	1.2	DXA	Men- Over 50y	277 (n=31 Black and	FNBMD (Black) FNBMD (White)	Energy adjusted protein intake (not adjusted for other confounders) 0.359 0.055	0.040 0.505

0.261
0.299
>0.05
>0.05
>0.05
0.017
ns
ns
<0.05
0.0001
0.529
0.536
-
-
-
0.04
0.05
_

Hip BMD	-0.004 -0.015, 0.006
HipBMD change:	0.001 -0.002, 0.005
LSBMD	-0.012* -0.024, 0.000
LSBMD change:	0.001 -0.005, 0.003
Men 50+ y	
Hip BMD	-0.002 -0.011, 0.006
HipBMD change:	0.001 -0.001, 0.004
LSBMD	0.000 -0.011, 0.011
LSBMD change:	0.000 -0.003, 0.004
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Postmenopausal Women	
50+ y	
Hip BMD	0.004 -0.001, 0.009
HipBMD change:	0.000 -0.002, 0.002
LSBMD	0.010* 0.003, 0.016
LSBMD change:	0.001 -0.001, 0.003
Men 25-49 y	
Hip BMD	-0.010 -0.024, 0.003
HipBMD change:	0.001 -0.003, 0.005
LSBMD	-0.013 -0.027, 0.001
LSBMD change:	0.000 -0.005, 0.005
Premenopausal Women	
25-49	
Hip BMD	-0.011* -0.022, -0.001
HipBMD change:	-0.003 -0.006, 0.001
LSBMD	-0.005 -0.017, 0.007
LSBMD change:	-0.002 -0.006, 0.003
Men 50+	
Hip BMD	-0.007 -0.016, 0.001
HipBMD change:	0.001 -0.002, 0.003
LSBMD	-0.009 -0.020, 0.002
LSBMD change:	0.001 -0.002, 0.005
_	
Postmenopausal Women	
50+ y	-0.006* -0.011, -0.001
Hip BMD	0.000 -0.002, 0.002

					HipBMD change:	-0.012* -0.019, -0.005	
					LSBMD LSBMD change:	-0.003* -0.005, 0.000	
					U	*=CI indicates exclusion of null effect	
Lau et al, 1998,	0.65 (vegetarians)	DXA	Post F, 70-89y	76		Unadjusted r values	-
China					LSBMD	0.09	-
					FNBMD	0.13	-
					IntertrochBMD	0.084	-
					WBMD	0.042	
						Beta coefficient- adjusted for energy	
						intake, age, calcium intake, urinary Na:Cr ratio	
					LSBMD	7.9x10-4	>0.05
					FNBMD	-6.8x10-4	>0.05
					IntertrochBMD	-3.6x10-3	>0.05
					WBMD	-1.0x10-3	>0.05
Libuda et al.	1.3 g/Kg/d	pQCT	Children and	228		Standardized (Beta) coefficient: Total	
2008, Germany			adolescents 8-		BMC	Protein	0.02
			14 years old			1.02	0.03
					Cortical Area: PC:	0.97 0.28	0.01 0.02
					SSI:		0.02
					551:	5.23	0.01
Libuda et al.	Median protein:	pQCT	Pre-pubertal	107 N. 57	Diaphyseal bone	Coefficients: Controlling for muscle area	
2011, Germany	Boys: 46.1g/d		children	(N=57 Boys	Forearm :	and androstenediol	
	Girls: 42.7g/d			воуs N=50	vBMC mg/mm	Beta=1.49	0.073
	01118. 42.7g/u			Girls)	VBINC Ing/IIIII	Beta (stand)= 0.11	0.075
				UIIS)	Cortical Area mm ²	Beta (stand)=0.11 Beta=1.37	
					Contical Alta IIIII	Beta= 1.57 Beta(stand)= 0.11	0.056
Loenekke et al.	91.3+/- 45.15 g/d	DXA	Males and	27		r values, controlling for body mass:	0.050
2010, USA	71.5T/- T J.15 g/u	DAA	Females,	<i>∠</i> 1	TBBMD	0.607	0.001
2010, UDA	71.72 +/- 13.95 kg		22+/-3 years		TBBMC	0.557	0.001
MacDonald et al,	79.4g/d	Bone	45-54y	5119	ibbiic	Mean (g/d) by Quartile (Q)	ANCOVA:
2005, UK	12.16.0	Markers,	women, pre,	5117	DPD/Cr n=2929	Q1 69.0	(confounder
2000, UIX		DXA	peri or post		$D_1 D_1 C_1 \Pi = L / L /$	Q2 76.4	adjusted)
		21111	menopausal			Q3 84.3	uajustea)

Meng et al. 2009 Australia	80.6g/d	DXA	862 elderly women 75 ± 3	862	PYD/Cr n=2929 LSBMD n=3226 FNBMD n=3226 TBBMC	Q4 99.3 Data not shown Data not shown Data not shown r=0.15 Unadjusted correlation between baseline	P=0.02 P=0.01 ns ns <0.001
		(SD) yr of age of white origin.		TBBMC AppendicularBMC	protein intake and 5 year BMC Q1 n=287, <66g/d 1352±236 1 388±242		
					TBBMC AppendicularBMC	Q2 n=287, 66-87g/d 1433±262 888±162	
					TBBMC AppendicularBMC	Q3 n=288, >87g/d 918±164 942±177	
						Whole body BMC (mg/cm2, headless) <66 g/d: 1357 ± 17 (n = 287) 66-87 g/d: 1387 ± 13 (n = 287) >87 g/d: 1429 ± 18 (n = 288)	
Metz et al, 1993,	1.24	SPA	Pre F	38		Appendicular BMC (mg/cm2) <66 g/d: 889 ± 11 (n = 287) 66-87 g/d: 917 ± 9 (n = 287) >87 g/d: 942 ± 12 (n = 288) Unstandardized B (SEM) adjusted for	
USA			Caucasian (24-28y)		DRBMC DRBMD MRBMC MRBMD	calcium intake, physical activity, lean body mass -0.450 (0.183) -0.434 (0.194) -0.503 (0.180) -0.251(0.214)	0.019 0.032 0.009 0.248
Michaelsson et al, 1995, Sweden	59g/d	DXA (Dietary Records	F 28-74y, Caucasian	175		Standardized Beta (adjusted for BMI, energy intake, physical activity, menopausal status, menopausal age,	

		data used,				smoking, diabetes, cortisone, HRT, athletic	
		not FFQ)				activity	
						Dietary records	
					TBBMD	0.00086	0.28
					LSBMD	-0.0010	0.51
					FNBMD	0.0028	0.04
						Food frequency records:	
					TBBMD	0.0020	0.005
					LSBMD	0.0013	0.36
					FNBMD	0.0024	0.06
						Unadjusted r values:	
					TBBMD	0.189	0.018
					LSBMD	0.058	0.474
					FNBMD	0.117	0.151
					OC	-0.036	0.669
Nakamura et al,	1.29	Bone	Elderly post F,	43	oc	Unadjusted r values:	0.009
	1.29	markers		43	OC	-0.197	p>0.05
2004, Japan		markers	mean				
			age=68.3y,		Bone ALP	-0.039	p>0.05
			range 43-79		Deoxypyd.	-0.241	p>0.05
					NTX	-0.205	p>0.05
Neville et al, 2002,	98g/d (M) and	DXA	238 M and	443	Young adult (20-25y old)	Standardized Beta(adjusted for dietary,	
UK	66g/d(F)		205 F, at both 15 and 20-25		BMD:	anthropometric and lifestyle parameters):	
			years of age		MALES:	Young adult protein intake:	
			-		LSBMD	-0.62	0.13
					FNBMD	-0.57	0.16
					FEMALES:	Young adult protein intake:	
					LSBMD	-0.11	0.61
					FNBMD	-0.04	0.87
					Young adult (20-25y old) BMD:	Standardized Beta(adjusted for dietary, anthropometric and lifestyle parameters)	

					MALES: LSBMD FNBMD FEMALES: LSBMD FNBMD	Adolescent protein intake: 0.53 -0.08 Adolescent protein intake: 0.12 0.47	0.13 0.83 0.76 0.27
New et al, 1997, UK	81+/-22 g/d	DXA	Women aged 44-50 years (Premenopaus al)	994	LSBMD FNBMD TrochBMD WardsBMD	Energy adjusted protein intake 0.03 0.02 0.04 0.02:	P>0.05 ns P>0.05 ns P>0.05 ns P>0.05 ns
Oh et al, 2013, Korea	TP 52.3g/d (Men) 45.0g/d(Women) AP 15.8g/d(Men) 12.0g/d(Women) SP 3.1g/d(Men) 2.8g/d(Women) VP 35.5g/d(Men) 32.2g/d(Women)	Ultrasound Calcaneal bone density (stiffness index only)	Men and Postmenopaus al Women aged 50-70 years	3330 (2575 in analysis)	Bone Stiffness	Spearmans Rho (Adjusted for age, energy intake, BMI, alcohol, smoking, HRT use, exercise, calcium intake): Men: TP 0.027 AP 0.044 VP -0.026 SP -0.013 VP:AP ratio -0.036 Women: TP 0.030 AP 0.035 VP -0.012 SP -0.014 VP:AP ratio -0.027	0.347 0.136 0.379 0.656 0.220 0.257 0.195 0.657 0.592 0.318

Orozco et al, 1998, Spain	TP: 73.4(17.9) g/d AP: 49.7(15.3)g/d VP: 23.7(8.7)g/d	DXA	Premenopausa l women aged 42years old	76	LS BMD FN BMD TrochBMD IntertrochBMD WardsBMD Normal (n=64) vs. Osteopenic (n=12): LSBMD	Unadjusted r values -0.03 -0.03 -0.04* chosen for troch-intertroch analysis -0.08 -0.05 73.5(18.1) g/d vs. 72.8(17.4)g/d	ns ns ns ns 0.9
					Normal (n=64) vs. Osteopenic (n=10): TotalHipBMD	72.8(18.4)g/d vs 77.0(17.7)g/d	0.5
Orwoll et al, 1987, USA	-	CT (vertebrae) , SPA (radius)	Study 1: Men Study 2: Men 30-90y	62 92	PRBMC- 1 DRBMC- 1 Vertebral BMC-1 DRBMC- 2 PRBMC- 2 Vertebral BMC-2	Unadjusted r values: 0.20 0.03 *chosen for radius pooled analysis 0.27 * chosen for men BMC analysis 0.22* chosen for radius pooled analysis 0.15 0.30*chosen for men BMC analysis	Ns Ns <0.05 Ns Ns <0.01
Pearce et al. 2010, UK	Median: 87.7g/d	Bone Markers	Men aged 49- 52 years	412	sCTX	r (95% CI) 0.04 (0.001, 0.1) Unadjusted linear regression coefficient, daily protein intake (per 100g)	0.04
Promislow et al, 2002 USA	72.5g/d	DXA	M/F 55-92y 572F 388M	960		Standardized Beta coefficients (95% CI) Controlling for age, body mass index, calcium intake, years menopausal (women only), diabetes status, current exercise, and current use of estrogen (women only), steroids, cigarettes, alcohol, thiazides, and thyroid hormones	
					TP: per 15g THBMD(F) FNBMD(F) TotalSpineBMD(F) TBBMD(F)	0.0094 (-0.0025, 0.0214) 0.0063 (-0.0039, 0.0165) 0.0084 (-0.0090, 0.0258) 0.0081 (-0.0017, 0.0179)	0.12 0.22 0.34 0.11
					THBMD(M) FNBMD(M) TotalSpineBMD(M)	-0.0003 (-0.0180, 0.0174) -0.0045 (-0.0202, 0.0112) -0.0095 (-0.0345, 0.0155)	0.97 0.57 0.45

					TBBMD(M)	-0.0078 (-0.0212, 0.0057)	0.26
					AP: per 15g THBMD(F)	0.0162 (0.0049, 0.0275)	0.005
					FNBMD(F)	0.0115 (0.0019, 0.0211)	0.005
					TotalSpineBMD(F)	0.0149 (-0.0016, 0.0314)	0.02
					TBBMD(F)	0.0098 (0.0005, 0.0191)	0.08
					$I DDIVID(\Gamma)$	0.0098 (0.0003, 0.0191)	0.04
					THBMD(M)	0.0059(-0.0112,0.0230)	0.50
					FNBMD(M)	0.0007 (-0.0145,0.0159)	0.93
					TotalSpineBMD(M)	-0.0007(-0.0249,0.0235)	0.96
					TBBMD(M)	-0.0036(-0.0167,0.0095)	0.59
					VP: per 5g		
					THBMD(F)	-0.0133 (-0.0219, -0.0047)	0.002
					FNBMD(F)	-0.0102(-0.0175, -0.0028)	0.002
					TotalSpineBMD(F)	-0.0129 (-0.0255, -0.0003)	0.04
					TBBMD(F)	-0.0047 (-0.0121, 0.0003)	0.20
					$I DDIVID(\Gamma)$	-0.0047 (-0.0121, 0.0020)	0.20
					THBMD(M)	-0.0206(-0.0357,-0.0054)	0.01
					FNBMD(M)	-0.0131 (-0.0267, 0.0006)	0.06
					TotalSpineBMD(M)	-0.0327(-0.0542,-0.0112)	0.003
					TBBMD(M)	0.0124 (-0.0243,-0.0004)	0.04
Quintas et al,	1.4g-1.7	DPA	Pre F	164		Unadjusted r values	
2003, Spain	11.8 11,	2111		101	RBMC	0.236	P<0.05
2000, Spain					RBMD	0.070	ns
					LSBMC	0.434	p<0.05
					HipBMC	0.412	<0.05
					LSBMD	0.317	<0.05
					HipBMD	0.301	<0.05
Rapuri et al,	53.7-71.2	DXA	Post F-	473	тарына	r values:	
2003, USA			65-77y		MRBMD	0.097	0.036
,			-		FNBMD	0.092	0.047
					TrochBMD	0.155	0.001
					TFBMD	0.136	0.003
					LSBMD	0.065	0.163
					TBBMD	0.129	0.005
					NTX	-0.022	0.641
					OC	0.01	0.832

					Baseline:	Protein tertile data:	
					OC (g/L)	4.07±0.012 Q1	0.50
						3.74±0.012 Q2	
						3.81±0.012 Q3	
						3.57±0.012 Q4	
					NTX:Cr ratio	56.2±2.45 Q1	0.50
						51.82±2.45 Q2	
						50.56±2.47 Q3	
						44.35±2.46 Q4	
Rubinacci et al,	Recent menopause	SPA	Post F	120		N=81, recent menopause, unadjusted r	
1992, Italy	(less than 9 years				Total Protein Intake:	values	<0.001
· •	ago, median age				DRBMC	0.305* used for pooling	ns
	51y)- 83+/-21.7 g/d				DRBMC/BW	-0.062	< 0.05
					Ultradistal RBMC	0.281	
	Distant menopause						
	(more than 15 years					N=39 distant menopause, unadjusted r	ns
	ago, median age				DRBMC	values	ns
	68y) - 68+/-17.6 g/d				DRBMC/BW	0.041 * used for pooling	ns
					Ultradistal RBMC	-0.031	
						-0.111	
Sahni et al. 2013,	81g/d (Men)	DXA	1,280 men and	2919		Model 2- adjusted for energy intake, age,	
USA	77g/d (Women)		1,639 women			height, weight, dietary vitamin D intake	
	e ()		,			(IU/d), vitamin D supplement use (yes/no),	
						Ca supplement use (yes/no), dietary Ca	
						intake	
						(,800 mg/d or \$800 mg), current smoking	
						(yes/no), menopausal status (yes/no),	
						current oestrogen use (yes/no) in women	
						alone, caffeine intake (g/d),	
						Physical Activity Scale in the Elderly	
						(PASE), osteoporosis medication use	
						(yes/no) and alcohol intake (none,	
						moderate and heavy intake	
					Cross sectional data:	Standardized coefficients:	
					Cross sectional data.	Standardized coefficients.	
					FNBMD	MEN (N=1268):	0.31
							0.31 0.28

						Beta(SE)=0.00065 (0.001)	
					FNBMD TrochBMD LSBMD	WOMEN (N=1614): Beta (SE)=0.00185 (0.001) Beta(SE)=0.00200(0.001) Beta(SE)=0.00280 (0.001)	0.04 0.02 0.04
						Model 2- adjusted for energy intake, age, height, weight, dietary vitamin D intake (IU/d), vitamin D supplement use (yes/no), Ca supplement use (yes/no), dietary Ca intake (,800 mg/d or \$800 mg), current smoking (yes/no), menopausal status (yes/no), current oestrogen use (yes/no) in women alone, caffeine intake (g/d), Physical Activity Scale in the Elderly (PASE), osteoporosis medication use (yes/no) and alcohol intake (none, moderate and heavy intake). Beta(SE)	
					Bone change data: FNBMD TrochBMD LSBMD	MEN (N=493): Beta (SE)=-0.0052(0.019) Beta(SE)=-0.0498 (0.020) Beta(SE)=-0.0062(0.019)	0.78 0.01 0.75
					FNBMD TrochBMD LSBMD	WOMEN (N=673): Beta (SE)=-0.0131(0.017) Beta(SE)=-0.0288(0.21) Beta(SE)=0.0042 (0.018)	0.44 0.21 0.81
Tanaka et al, 2001, Japan	1.3	Ultrasonic Bone Absorptio metry	Pre F- 18-22y	965		Regression B (Unstandardized) Coefficient, adjusted for age, weight, height, exercise, menstrual	

					OSI calcaneus	status and daily nutrient intakes (energy, Ca, Phosphorus, Sodium) 0.234	0.009
Teegarden et al,	1.21	DXA	Young pre F	215		Unadjusted r values:	
1998, USA			01		TBBMD	0.11	Ns
,					RBMD	0.16	< 0.05
					LSBMD	0.19	< 0.05
					FNBMD	0.08	Ns
					TrochBMD	0.10	Ns
					WBMD	0.08	Ns
					TBBMC	0.12	Ns
					RBMC	0.08	Ns
					Spine BMC	0.23	<0.05
						Unstandardized B (SE) adjusted for postmenarchal age, lean and fat mass,:	
					TBBMD	0.0016+/-0.0006	<0.05
					TBBMC	6.95+/-2.09	<0.05
					SpineBMD	0.0029+/-0.0013	<0.05
					SpineBMC	0.1823+/-0.068	<0.05
Thorpe et al,	74.7g/d	DXA	Postmenopaus	161	LSBMD	-0.01	0.94
2008, USA	74.7g/u	DAA	al women	101	TotalHipBMD	0.08	0.30
2008, USA			mean age		(non adjusted,	0.08	0.50
			68+/-6 years		Spearmans)		
			00+7-0 years		Spearmans)	Unstandardized correlation coefficient:	
					LSBMD	B(SE): controlling for body weight and	
					LSDWD	sulphur intake	0.04
						1.35×10^{-3} (6x10-4)	0.04
Tylavsky and	1.01	SPA	60-98y elderly	375		\Box coefficient	
Anderson, 1988,	1.01	SIA	F	515	DRBMC	2.72	0.03
USA			1,		DRBMD	0.63	0.03
USA					MRBMC	2.96	0.23
					MRBMD	1.36	0.003
Votonnonost st al	20-25 years: 68+/-	DXA	Voung adulta	133	WINDIVID	Unstandardized Beta+/-SE (adjusted for	0.00
Vatanparast et al,	•	DAA	Young adults	133			
2007, Canada	22(F) and 119+/-53		(59 males, 74 females).			sex, current height and weight, physical	
	(M)					activity level, and other dietary nutrients)	
	Dania dala		Measured at			Comment another in (1) () and (1)	
	Periadolescence:		both		TDDMC	Current protein intake (young adult)	
	64.2+/-17 (F) and		periadolesence		TBBMC	NS (not entered into stepwise model)	ns
	79.6+/-17 (M)		and young		TBBMD	NS (not entered into stepwise model)	ns

			adulthood (20- 25 y)		TBBMC net gain	0.33 +/- 0.042	<0.001
Wang et al, 1997, USA	0.97	DXA	Older post F	125	TBBMC TBBMC net gain TBBMD LSBMD FNBMD	Unstandardized Beta+/-SE (adjusted for sex, current height and weight, physical activity level, and other dietary nutrients) Females only with adequate calcium at pert-adolescence/early adulthood 0.21+/- 0.095 0.21+/-0.080 0.32+/-0.32 0.04 -0.01 Spearmans correlations	<0.05 <0.05 <0.05 Ns Ns
Wang et al. 1999, USA	1.05 g/Kg/d	QUS	18-18 year old women	63	Bone indices at18-19 years	Protein intake when aged 9-11 years:	
						Spearmans Rho	
					BUA	0.16	ns
					BV	0.27	<0.05
					SOS	0.25 Pearson's:	Ns
					BUA	0.11	Ns
					BV	0.21	Ns
					SOS	0.17	Ns
Weikert et al, 2005, Germany	67.9g/d	QUS/BUA	F 35-67y	8178	Os calcis TP AP VP AP:VP ratio TP	Coefficients not shown for multiple regression as ns for protein (adjusted for calcium, magnesium, vitamin C, race, height and weight)- no effect size Beta (Standardized) coefficient -0.03 (0.013) -0.03 (0.012) (controlling for VP) 0.11 (0.042) (controlling for AP) -1.12 (0.31) (controlling for TP) 0.014 (0.017) (controlling for AP:VP ratio)	0.017 0.010 0.007 <0.001 0.41
						Pearson's Correlations: r	
						rearson s Correlations: r	

					VP	0.03	0.009
					AP	0.02	0.015
					TP	0.03	0.002
Whiting et al,	1.15	DXA	М	57		Pearson's correlations:	
2002, Canada			39-42y		TBBMD	0.383(adj)	<0.01
,			5		LSBMD	0.419 (adj)	< 0.01
					THBMD	0.322 (adj)* chosen for pooling men as closest to Jaime	<0.05
						-controlling for anthropometry and energy intake	
					TBBMD	Linear regression: (non-standardized B) 0.00193 (0.00065)	<0.01
						Adjusted for lean body mass, height, fat mass, energy intake	
Yazdanpanah et al, 2007, The	81.3g/d , 1.1g/Kg/d	DXA	Men and Women aged	5304		Standardized Beta coefficient (adjusted for age, BMI, other dietary nutrients, sex)	
Netherlands			55 years and			Protein intake:	
			over		FNBMD	-0.03	0.29
					LSBMD	-0.03	0.27
Zhang et al. 2010,	1.7 g/Kg/d	DXA	Girls (Mean	757		Beta(adjusted for baseline bone mass,	
China			age 10 years)			tanner stage, age, physical activity). Beta	
						represents % change with doubling of protein intake: All participants:	
					DRBMC	-4.82	
					DRBMD	-3.18	< 0.01
					DRBA	~	<0.01
					PRBMC	-10.2	-
					PRBMD	~	<0.01
					PRBA	-9.11	- <0.01
					TBBMC	-1.92	NU.U1
							0.00
					TREMD	~	0.02
					TBBMD TBBA	~	0.02

* simple r coefficients unless otherwise stated; for r2 the brackets indicate if corresponding regression coefficient + or - ** total protein in g/Kg/d unless otherwise stated. ALP= Alkaline Phosphatase; AP=animal protein; BMC=Bone Mineral Content; BMD=Bone Mineral Density; BUA-Broadband Ultrasound Attenuation; BV=Bone Volume; Calc=Calcaneus; Cr=Creatinine; Deoxypyd=Deoxypyridinoline; DRBA=Distal Radial Bone Area; DRBD=Distal Radial Bone Density; DTB=Distal Tibial; DXA=Dual Xray Absorptiometry; FN=Femoral Neck; FNBA=Femoral Neck Bone Area; FNBMD=Femoral Neck Bone Mineral Density; FNvBMD=Femoral Neck volumetric Bone Mineral Density; FSBMD=Femoral Shaft Bone Mineral Density; HBMD=Humerus Bone Mineral Density; HPO=Hydroxyproline; HumBMC=Humerus Bone Mineral Content ;IntertrochBMD=Intertrochanter Bone Mineral Density; LSBMC=Lumbar Spine Bone Mineral Content; LSBMD=Lumbar Spine Bone Mineral Density; MRBMC Midradial Bone Mineral Content; MRBMD Midradial Bone Mineral Density; OC=Osteocalcin; P1NP= Procollagen type 1 N-terminal propeptide; PC=Periosteal Circumference; pQCT=Peripheral Quantitative Computed Tomography; PRBMC=Proximal Radial Bone Mineral Content; PYD=Pyridinoline; QUS=Quantitative Ultrasound; R=Radial RBMC=Radial Bone Mineral Content; sBAP=serum Bone Alkaline Phosphatase; sCTX serum C-telopeptide of collagen; sOC serum Osteocalcin; SP=Soy Protein; SSI=Stength Strain Index; Stand.=Standardised; TBBA=Total Body Bone Area; TBBMD Total Body Bone Mineral Density; VP=Vegetable Protein; WBBMC=Whole Body Bone Mineral Content; wBBMD=Whole Body Bone Mineral Density

Table S2 Pooled r values for protein intake and bone health for gender and age subgroups (non-adjusted data)

Parameter	Model	r	R ²	Lower	Upper	р	I ²	Total n	Included Studies
				limit	limit				
MEN BMD	Fixed	0.1201	0.01	0.0291	0.2091	0.010	44%	470	Coin, Jaime(Black), Jaime(White), Whiting
	Random	0.1549	0.02	0.0184	0.2858	0.026			
MEN BMC	Fixed	0.2881	0.08	0.1346	0.4281	0.0003	0%	154	Orwoll(group 1), Orwoll (group 2)
	Random	0.2881	0.08	0.1346	0.4281	0.0003			
POST F BMD	Fixed	0.1148	0.01	0.0791	0.1502	< 0.001	1%	2987	Alissa 2014, Cooper, Chiu, Devine, Gunn, Horiuchi,
									Lau, Rapuri, Thorpe 2008, Wang 1997
	Random	0.1147	0.01	0.0787	0.1503	< 0.001			
POST F BMC	Fixed	0.181	0.03	0.0618	0.2941	0.003	0%	267	Freudenheim, Lacey, Rubinacci (Early Post),
									Rubinnacci (Late Post)
	Random	0.181	0.03	0.0618	0.2941	0.003			
PRE F BMD	Fixed	0.0748	0.01	0.0384	0.1111	<0.001	74%	2896	Chan 2009, Chiu, Cooper, Gregg, Henderson, Hirota,
									Lau, New, Orozco, Quintas
	Random	0.1158	0.01	0.0376	0.1925	0.004			
PRE F BMC	Fixed	0.2834	0.08	0.1986	0.3640	<0.001	47%	485	Freudenheim, Lacey, Teegarden, Quintas
	Random	0.2748	0.08	0.1442	0.3959	<0.001			
OLDER ADULT	Fixed	0.1131	0.01	0.0736	0.1522	<0.001	0%	2448	Chiu, Coin, Devine, Gunn, Lau, Rapuri, Thorpe2008,
(OVER 60 YEARS)									Wang1997
M/F BMD									
	Random	0.1131	0.01	0.0736	0.1522				
CHILD M/F BMC*	Fixed	0.3154	0.10	0.2251	0.4003	< 0.001	0%	416	Bounds, Chevalley 2008, Ekbote, Hoppe

Random	0.3154	0.10	0.2251	0.4003	< 0.001
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BMD=Bone Mineral Density BMC=Bone Mineral Content n=number of partipenats in analysis *only radius BMC and total body BMC available for pooling. Where studies have multiple outcomes eligible for inclusion, choice of measures for pooling was as follows: Hip indices (first choice), Spine indices (2nd choice), Radial indices (3rd choice).

Parameter	Model	r	\mathbb{R}^2	Lower limit	Upper limit	р	I ²	Total n	Included Studies
ADULTS									
TBBMC	Fixed	0.12	0.01	0.0662	0.1683	<0.001	73%	580	Ho, Loenekke, Meng, Teegarden,
	Random	0.14	0.02	0.0133	0.2622	0.0304			
DEPYD	Fixed	-0.23	0.05	-0.3859	-0.052	0.01	0%	128	Horiuchi, Nakamura
	Random	-0.23	0.05	-0.3859	-0.052	0.01			
FNBMD	Fixed	0.07	0.00	0.0374	0.0942	< 0.001	26%	4786	Alissa 2014, Chan 2009, Chiu, Coin, Cooper (post), Cooper (pre),
	Random	0.07	0.00	0.0391	0.1090	<0.001			Gunn, Henderson, Ho, Jaime(Black), Jaime (White), Kumar, Lau, Michaelsson, New, Orozco, Rapuri, Teegarden, Wang
FEMORAL SHAFT BMD	Fixed	0.06	0.00	-0.0394	0.1563	0.240	0%	405	Cooper(post), Cooper(pre), Henderson
	Random	0.06	0.00	-0.0394	0.1563	0.240			
TROCH/INTTRO CH BMD	Fixed	0.09	0.008	0.0528	0.1330	<0.001	68%	2375	Coin, Cooper (post), Cooper (pre), Henderson, Lau, New, Orozco, Rapuri, Teegarden
	Random	0.12	0.014	0.0401	0.2027	0.004			<u>r</u> , <u>8</u>
TOTAL HIP BMD	Fixed	0.09	0.008	0.0389	0.1491	0.001	86%	1259	Alissa 2014, Chan 2009, Coin, Quintas, Thorpe M, Whiting
- •	Random	0.14	0.02	-0.0118	0.2919	0.07			
WARDS BMD	Fixed	0.02	0.0004	-0.0325	0.0654	0.51	0%	1616	Kumar, Lau, New, Orozco, Teegarden
	Random	0.02	0.0004	-0.0325	0.0654	0.51			
HUMERUS BMC	Fixed	0.16	0.03	-0.0613	0.3648	0.16	0%	84	Freudeneheim (Post), Freudenheim (Pre)
	Random	0.16	0.03	-0.0613	0.3648	0.16			
HYDROXYPROL INE	Fixed	-0.07	0.00	-0.1838	0.0466	0.24	68%	290	Cooper (post), Cooper (pre)
	Random	-0.11	0.01	-0.3363	0.1240	0.35			
LSBMD	Fixed	0.07	0.005	0.0410	0.1012	0.0001	58%	4257	Chiu, Cooper (post), Cooper (pre), Henderson, Ho, Horiuchi, Kumar,
	Random	0.09	0.008	0.0373	0.1385	0.0007			Lau, Michaelsson, Quintas, Rapuri, Thorpe M, Teegarden, Wang, Whiting
LSBMC	Fixed	0.31	0.10	0.2329	0.3876	<0.001	41%	533	Orwoll (group 1), Orwoll (group 2), Teegarden, Quintas
	Random	0.31	0.10	0.2057	0.4146	<0.001			
RADBMD	Fixed	0.07	<0.01	0.0180	0.1267	0.009	53%	795	Cooper (post), Cooper (pre), Hirota, Quintas, Rapuri, Teegarden
IN IDDINID	Random	0.07	<0.01	-0.0101	0.1207	0.084	5570	175	cooper (post), cooper (pro), rinou, Quinus, Rupun, reegaluen
OSTEOCALCIN	Fixed	0.00	0.00	-0.0101	0.0809	0.004	40%	593	Cooper (post), Cooper (pre), Horiuchi, Nakamura, Michaelsson
of House of House	Random	-0.01	0.00	-0.1175	0.1039	0.90	1070	575	cooper (post), cooper (pro), nonuem, numeration
RADBMC	Fixed	0.16	0.026	0.0987	0.2268	<0.001	0%	915	Freudeneheim (Post), Freudenheim (Pre), Lacey (Pre), Lacey (Post),
	Random	0.16	0.026	0.0987	0.2268	<0.001	0.10	210	Orwoll (group 1), Orwoll (group 2), Quintas, Rubinacci (early post), Rubinacci (late post), Teegarden
TOTAL BODY BMD	Fixed	0.17	0.03	0.114	0.2334	<0.001	59%	1028	Knurick, Loenekke, Michaelsson, Rapuru, Teegarden, Whiting

 Table S3: Pooled r values for protein intake and bone health by outcome (non-adjusted data)

	Random	0.22	0.05	0.0114	0.3263	<0.001			
ULNABMC	Fixed	-0.02	0.00	-0.2395	0.197	0.84	0%	84	Freudeneheim (Post), Freudenheim (Pre)
	Random	-0.02	0.00	-0.2395	0.197	0.84			
TOTALHIP BMC	Fixed	0.16	0.026	0.0766	0.2330	0.001	94%	602	Ho, Quintas
	Random	0.24	0.06	-0.1358	0.5494	0.211			
	Random	0.13	0.02	0.0771	0.1913	< 0.001			
CHILDREN									
ALL BMC*	Fixed	0.32	0.10	0.2251	0.4003	<0.001	0%	416	Bounds, Chevalley 2008, Ekbote, Hoppe
	Random	0.32	0.10	0.2251	0.4003	<0.001			
TBBMC	Fixed	0.37	0.14	0.2386	0.4927	<0.001	0%	184	Bounds, Ekbote, Hoppe
	Random	0.37	0.14	0.2386	0.4927	<0.001			
TBBA	Fixed	0.48	0.23	0.3591	0.5892	<0.001	79%	176	Ekbote, Hoppe
	Random	0.46	0.21	0.1641	0.6821	0.003			
	Random	0.40	0.21	0.1041	0.0021	0.005			
TDDMD	Einad	0.02	0.0004	0 1222	0.0001	0.71	87%	314	Dounda Janas
TBBMD	Fixed	-0.02	0.0004	-0.1322	0.0901	0.71	01%	514	Bounds, Jones
	Random	0.11	0.01	-0.3055	0.4853	0.62			

All pooled effects calculated in R using 'meta' and 'metacor' packages, which use the inverse variance method, DerSimonian and Laird (random effects models) and Fisher's Z Transformation. BMD=Bone Mineral Density DEPYD=Deoxypyridinoline FNBMD= Femoral Neck Bone Mineral Density INTTROCH=Intertrochanter LSBMC=Lumbar Spine Bone Mineral Content LSBMD=Lumbar Spine Bone Mineral Density RADBMC= Radial Bone Mineral Content RADBMD= Radial Bone Mineral Density TBBA=Total Body Bone Area, TBBMC=Total Body Bone Mineral Content, TBBMD=Total Body Bone Mineral Density TROCH=Trochanter ULNABMC=Ulna Bone Mineral Content

Table S4 Associations between protein dose, calcium dose and calcium:protein ratio and FNBMD and LSBMD (non-adjusted for confounders)

Linear Model

X	r for FNBMD				r for LSBMD				Actual FNBMD				Actual LSBMD			
Model	Estimate*	SE	Model P	N studies	Estimate *	SE	Model P	N studies	Estimate *	SE	Model P	N studies	Estimate *	SE	Model P	N studies
Protein (g/kg/d)	-0.05	0.16	0.76	16	0.25	0.18	0.19	14	0.39	0.15	0.02	19	0.33	0.13	0.02	16
Calciu m (mg/kg/ d)	-0.01	0.008	0.17	16	0.007	0.01	0.57	17	0.02	0.010	0.06	19	0.02	0.010	0.06	17
Ca:Prot ratio (mg/g/d)	-0.01	0.01	0.20	18	-0.008	0.02	0.63	17	0.0005	0.013	0.97	19	0.005	0.01	0.74	17

*Intercept not shown for clarity. Equation: y=x+c (where c=intercept, y=dependent variable). Note: No results were statistically significant after Bonferroni correction (36 tests, 0.05/36=0.001) Note: some studies had Ca:Prot ratio but not protein or calcium. This is because for some studies the body weight was not given so protein in g/Kg/d was not calculated. However, if studies gave Ca mg/d and Protein g/d the Ca:Prot ratio could still be calculated for these studies. Ca=Calcium FNBMD=Femoral Neck Bone Mineral Density LSBMD= Lumbar Spine Bone Mineral Density N Studies =Number of studies Prot=Protein SDE=Standard Error

Quadratic model

X ²	r for FNBMD				r for LSBMD				Actual FNBMD				Actual LSBMD			
Model	Estimate*	SE	Model P	N studies	Estimate *	SE	Model P	N studies	Estimate *	SE	Model P	N studies	Estimate *	SE	Model P	N studies
Protein ² (g/kg/d)	-1.28	1.16	0.53	16	1.26	0.87	0.17	14	0.33	0.77	0.06	19	0.28	0.70	0.08	16
Calciu m ² (mg/kg/ d)	-0.0007	0.003	0.40	16	0.002	0.004	0.75	17	0.0003	0.003	0.18	19	-0.0004	0.003	0.19	17
Ca:Prot ratio ² (mg/g/d)	-0.002	0.003	0.38	18	-0.003	0.004	0.75	17	-0.002	0.004	0.89	19	1.3 x 10 ⁻⁵	4.4 x 10 ⁻³	0.95	17

*Other model estimates not shown for clarity. Ca=Calcium, Prot=Protein. Equation: $y=x+x^2+c$ (where c=intercept, y=dependent variable). Note: No results were statistically significant after Bonferroni correction (36 tests, 0.05/36=0.001). Note: some studies had Ca:Prot ratio but not protein. This is because for some studies the body weight was not given so protein in g/Kg/d was not calculated. However, if studies gave Ca mg/d and Protein g/d the Ca:Prot ratio could still be calculated for these studies. Ca=Calcium FNBMD=Femoral Neck Bone Mineral Density LSBMD= Lumbar Spine Bone Mineral Density N Studies =Number of studies Prot=Protein SDE=Standard Error

Cubic model

X ³	r for FNBMD				r for LSBMD				Actual FNBMD				Actual LSBMD			
Model	Estimate*	SE	Model P	N studies	Estimate *	SE	Model P	N studies	Estimate *	SE	Model P	N studies	Estimate *	SE	Model P	N studies
Protein ³ (g/kg/d)	0.06	0.40	0.74	16	-0.10	0.46	0.33	14	-0.90	0.36	0.01	19	-0.84	0.29	0.009	16
Calciu m ³ (mg/kg/ d)	-0.0006	0.001	0.56	16	-0.0008	0.002	0.86	17	4.9 x 10- 5	1.4 x 10 -3	0.35	19	-0.0003	0.001	0.35	17
Ca:Prot ratio ³ (mg/g/d)	-0.001	0.001	0.51	18	-0.0002	0.002	0.91	17	-0.001	0.002	0.84	19	-0.003	0.002	0.58	17

*Other model estimates not shown for clarity. Ca=Calcium, Prot=Protein. Equation: $y=x+x^2+x^3+c$ (where c=intercept, y=dependent variable) Note: No results were statistically significant after Bonferroni correction (36 tests, 0.05/36=0.001). Note: some studies had Ca:Prot ratio but not protein. This is because for some studies the body weight was not given so protein in g/Kg/d was not calculated. However, if studies gave Ca mg/d and Protein g/d the Ca:Prot ratio could still be calculated for these studies. Ca=Calcium FNBMD=Femoral Neck Bone Mineral Density LSBMD= Lumbar Spine Bone Mineral Density N Studies =Number of studies Prot=Protein SDE=Standard Error

Study	Mean Protein *	Populatio n	Lengt h	Total n	Fracture / BMD site	Protein type	RR **≠	95% CI	P value	Confounder Adjustments
Beasley et al.	<13.3%	Women	бу	144,58			HR:			Age, BMI, race-ethnicity, calibrated energy intake, general
2014, USA	to	aged 50-		0	Any	TP	0.99	(0.97,	-	health, physical activity, history of fracture at age 55 y,
	≥15.6%	79 y at			Hip	TP	0.91	1.02)	-	history of parental
	of	baseline			Spine	TP	1.05	(0.84,	-	fracture, current smoking, corticosteroid use, glucocorticoid
	energy				Forearm	TP	0.93	1.00)	-	use, treated diabetes,
	intake							(0.98,		rheumatoid arthritis, and hormone use
	from							1.13)		
	protein							(0.88,		
								0.98)		
Dargent- Molina et al,	TP: 46(7.5)g	Postmenop ausal	8.37 (1.73)	36217 (2408	Any low impact	Energy adjuste				(Also has calcium intake stratification data)
2008, France	/d	women	(1.7 <i>5</i>) y	with	fracture	d	1.06	0.94-1.19	-	Adjusted for BMI, physical activity, parity, maternal history
E3N study	AP:29	wonnen	5	inciden	mueture	TP	1.10	0.98-1.24	_	of hip fracture, HT use, smoking status,
Leitbeaug	(8.8) g/d			t		AP	0.95	0.85-1.06	-	and alcohol intake
	VP:			fractur		VP				
	12(3.0)g			e,						
	/d			33809						
				fractur						
				e free)						
Feskanich et	79.6g/d	Caucasian	12y	85,900						Adjusted for questionnaire time period; age (5-year intervals),
al, 1996, USA	median	F, 35-59y			FF	AP	1.25	1.07-1.46	0.004	BMI and hours of
						TP	1.22	1.04-1.43	0.01	vigorous activity per week (qulntjles); menopause) status and
						VP	0.9	0.77-1.06	0.17	use of postmenopausal hormones (premenopausal,
					HF	AP	0.98	0.65-1.47	0.7	postmenopausal-never
						TP	0.96	0.64-1.45	0.7	user, postmenopausal-past user, postmenopausal-current
						VP	1.11	0.75-1.66	0.58	user); cigarette smoking (never, past, current); use of thyroid hormone medication
										and thiazlde diuretics (yes or no); and alcohol and caffeine Intakes (quintiles).

Table S5: Characteristics and outcomes of the 29 studies reporting fracture or osteoporosis diagnosis data (6 of which also in Table 1)a) 14 Cohort studies

Study	Mean Protein *	Populatio n	Lengt h	Total n	Fracture / BMD site	Protein type	RR **≠	95% CI	P value	Confounder Adjustments
Gunn et al, 2014, New Zealand	79g/d	Bone markers, DXA	POM wome n, 60 years of age	142	Osteopo rosis diagnosi s)	TP Mean(SD) Protei n intake by catego ry:	BMD: Normal 79(21) Mild Osteope nia 83(18) Significa nt osteopen ia 77(22) Osteopor osis 76(21)	n 51 21 53	NS	Non confounder adjusted
Key et al, 2007, UK	Women: 73.1 (21.6) g/d 77.8(22. 6)g/d	26 749 women and 7947 men aged 20–89 years.	5.2y	26 749 women , 7947 men, aged 20–89 years	All sites, incident fractures (includin g high trauma fractures, but still 72% from a fall)	TP: Women n=362 fracture s Men n=76 fracture s	Incident Rate Ratio: 0.97 1.29	0.74-1.27 0.72-2.31	0.55 0.68	Confounder adjusted: Method of recruitment and adjusted for age, smoking, intakes of energy and each other nutrient, alcohol consumption, body mass index, walking, cycling, vigorous exercise, other exercise, physical activity at work, marital status and, for women, parity and use of hormone replacement therapy
Langsetmo et al, 2015, Canada	TP: 0.79(0.6 0-1.03) AP (Non- dairy): 17.6(12. 8-23)g/d	Men and Women, aged 25- 49 and ≥50 years	5y	6510	Fragility fracture: n=4543	s TP: Men Women :	HR= 0.66 0.85	0.35-1.24 0.67-1.09	-	Confounder adjusted: Age, height, TEI, center (women only), education, smoking, alcohol intake, physical activity, sedentary hours, calcium and vitamin D supplement use, hormone therapy (women only), bisphosphonate use (women only), and diagnosis of osteoporosis (women only);

Study	Mean Protein *	Populatio n	Lengt h	Total n	Fracture / BMD site	Protein type	RR **≠	95% CI	P value	Confounder Adjustments
	VP: 24.3(18. 8- 31.0)g/d				Main fracture: n=4570	TP: Men Women	0.55 0.90	0.28-1.09 0.69-1.19	-	
Meyer et al, 1997, Norway	0.8	M/F (mean age 47.1y)	11.4y	19752 F 20035 M	HF- F HF-M	AP AP	0.96 1.3	Q4= highest: 0.62-1.49 0.63-2.68 Q1=refer ence,	0.37 0.48	Adjusted for age at screening, body height, body mass index, serf-reported physical activity at work and during leisure time, diabetes mellltus, disability pension, marital status, and smoking
Misra et al, 2011, USA	64g/d (energy	Men and women	11.6y	946 (n=100	HF	TP: M/F	HR=	RR=1, lowest		Confounder adjusted: age, sex, weight, height and total energy intake
	adjusted)	mean age=75 years		had hip fractur e)		(n=100) F (n=80)	0.63 0.82	0.37-1.09 0.44-1.51	-	
Munger et al, 1999, USA	1.2	Postmenop ausal F (55-69y)	1-3y	32 050	HF	AP TP VP	RR= 0.31 0.44 1.92	0.10-0.93 0.16-1.22 0.72-5.11	0.037 0.049 0.11	Age, body mass index, number of pregnancies, smoking, alcohol use, estrogen use, and physical activity.
Mussolino et al, 1998, USA	<56g/d - >98g/d	Caucasian M (45- 74y)	22y	2879	HF	TP	RR: 0.55	0.20-1.55	-	BMI, previous fracture, smoker, physical activity, alcohol, chronic health condition, calcium intake, weight loss.
Sahni et al, 2010, USA	Men TP: 79.0(27)	Men and women aged	7 to 14 years	3656	HF	Low calcium intake	HR for highest tertile of			Adjusted for sex and menopause status (group 1: men; group 2: premenopausal women; group 3: postmenopausal women), age
Framingham Offspring Study	g/d AP: 54.3(22) VP: 24.6(9)	mean= 55 (9.9)years				(<800 mg/d) n=2124 :	protein intake:			(years), weight at baseline (kg), height at baseline (m), physical activity index, intake of energy (MJ/day) and total vitamin D (IU/day), and smoking status (current versus former/never) and calcium intake

Study	Mean Protein *	Populatio n	Lengt h	Total n	Fracture / BMD site	Protein type	RR **≠	95% CI	P value	Confounder Adjustments
	Women: TP:75.7(27) AP: 52.5(22) VP: 23.1(9)					Energy adjuste d TP: g/d AP:g/d VP:g/d AP:VP ratio	2.20 3.17 0.60 1.81	0.88-5.54 1.30-7.78 0.20-1.85 0.68-4.86	0.09 0.01 0.34 0.29	
						High calcium intake (≥800 mg/d) n=1532 Energy adjuste d TP: g/d AP:g/d VP:g/d AP:VP ratio	0.54 0.32 0.23 2.02	0.12-1.30 0.05-2.08 0.05-1.03 0.37- 11.05	0.38 0.33 0.06 0.32	
Sellmeyer et al, 2001, USA	49.8g/d	Caucasian F aged over 65y	7.0y +/- 1.5y	1035	Hip Fracture	VP Ratio AP:VP	0.3 3.7	-	0.03 0.04	Age and body weight
Zhong et al, 2009 USA	Mean(S E)=61+/ -0.8 g/d	Postmenop ausal women at least 50 y of age	<7y	2006	All fragility fractures (hip, wrist, spine)	AP TP	2.7 OR data	- Data in Figure Only	0.04	Age, race, body mass index (underweight/normal, overweight, obese), physical activity level, smoking status, alcohol use (heavy, moderate/none), hormone use, general health status, osteoporosis, arthritis, vision impairment, and stroke.

Study	Mean Protein *	Populatio n	Lengt h	Total n	Fracture / BMD site	Protein type	RR **≠	95% CI	P value	Confounder Adjustments
Zhang 2005	SP: 9.6g/d Non Soy: 134g/d	Women aged 40- 70 years old	4.5 y	24403	All fractures	SP	<4.98 g/d (Referen ce) ≥13.27 g/d 0.63	0.53-0.76	<0.001	Age, body mass index, hours of exercise per week, cigarette smoking, alcohol consumption, history of diabetes mellitus, level of education, family income, season of recruitment, and intakes of total calories, calcium, non-soy protein, fruits, and vegetables

AP, Animal Protein; BMD, Bone Mineral Density; DXA, Dual X-ray Absorptiometry; HF, Hip Fracture; HR, Hazard Ratio; POM, Postmenopausal; RR, Relative Risk; SP, Soy protein; TEI, Total Energy Intake; TP, Total Protein; VP, Vegetable Protein

4b.2 Cross cultural studies

Study	Mean Protein **	Method	Population	n	Outcome	Coefficient*	р	Confounders
Abelow et al, al1992, USA cross cultural	10.4g/d- 77.8g/d AP	Fracture	F over 50y	34 studies 16 countries	Hip fracture and animal protein	r2=0.66(+) (by study) r2=0.67 (+) (by country)	<0.001 <0.001	Age adjusted
Frassetto et al, 2000, USA Cross Cultural	48 to 110.9 g/d	Fracture	F aged over 50y	33 countries	Hip fracture TP AP VP	0.67; 0.82; -0.370;	p<0.001 p<0.001 p<0.04	Age. Also, for AP, TP and VP

AP, Animal Protein; TP, Total Protein; VP, Vegetable Protein

4c 13 Case control studies

Study	Protein intake*	Population	n	Site	Group/outcome	OR**≠	р	Confounders
Alissa et al, 2011, Saudi Arabia Non- Prospective	77g/d	DXA	Postmenopausal women, aged 50-60 years	122	Normal BMD Osteopenic	Dietary protein intake g/d 77.5 76.6	ns	Non adjusted for confounders
Chevalley et al. 2011, Switzerlan d	Age 7.4 (0.4): 1.78 (0.46) Age 15.2(0.5): 1.08 (0.41)	DXA	Caucasian boys- measured during pre- puberty and adolescence	176	Age 7: Without Fracture: n=89 With Fracture: n=87	Dietary Intake, (g/d) 48.5 (13.3) 45.2(11.1)	0.08	Non adjusted for confounders
					Age 15: Without Fracture :n=89 With Fracture: n=87	65.4 (24.1) 61.2 (23.1)	0.24	
Chiu et al, 1997, Taiwan Non-	1.09	DPA (BMD)	Older POM F	258	Osteopenia of: Lumbar Spine Femoral Neck	Energy intake from protein (%) 0.51 (0.30-0.89) 0.71 (0.33-1.54)	- Significant NS	Adjusted for age, BMI, physical activity, calcium intake, non-protein energy intake, long term vegan/vegetarianism
Prospectiv e								
Coin et al, Italy, 2008	75.8+/-22.1 g/d Weight=74.2+/- 13.4	DXA	Males, mean age 73.9+/-5.6 years	136	Only data for men included protein in model:	OR (95% CI) of low total hip BMD<=0.83h/cm2)		Adjusted for BMI
Non- Prospective	So 1.02 g/Kg/d				MEN Protein<65.7g/d Protein>=65.7g/d	3.69 (1.40-9.70) 1.00	0.008	
Farrin et al. 2008, Iran	81.4g/d	DXA	Postmenopausal Women	58	LSBMD based diagnosis: Normal Osteopenic	Mean (SD) Protein intake:g/d 68.7 +/- 5.0 95.5+/- 67.6	<u>One way</u> <u>ANOVA</u> p<0.05	Unadjusted

Non- prospective					Osteoporotic	67.6+/-5.3	Post hoc tests: Normal- Osteopenia: P=0.009 Normal- Osteoporoti c P=0.75	
Kim et al, 2008, Korea Non- prospective	TP= 60g/d AP= 19g/d VP= 40g/d	DXA	Postmenopausal women, 134 osteoporotic cases and 137 non- osteoporotic controls	271	Osteoporotic (n=134) Non- Osteoporotic (n=137)	OR for Osteoporosis by protein intake: TP: g/d Lowest 1.0(reference) Middle 0.91 (0.68- 1.21) Highest 1.47 (1.03- 2.05)	P=0.004	Adjusted for age, smoking, alcohol drinking, BMI, exercise, family history of osteoporosis, and energy intakes
					Osteoporotic (n=134) Non- Osteoporotic (n=137)	AP: g/d Lowest=1.0 (reference) Middle= 1.21(0.58- 2.52) Highest= 1.62(1.03- 3.92)	P=0.03	
					Osteoporotic (n=134) Non- Osteoporotic (n=137)	VP: g/d Lowest=1.0(reference) Middle=0.62(0.31- 1.23) Highest=0.42(0.23- 0.83)	P=0.011	
Martinez- Ramirez et al, 2012, Spain	TP:105 (1.0) g/d AP:66-70 (1.3) g/d VP: 38 (0.63)g/d	Aged 65 years or over, cases from hospital record and	167 cases and 167 controls	All low energy fractures (e.g. from a fall)	TP AP VP AP:VP ratio	OR: 1.10 (0.18, 6.80) 0.38 (0.10-1.41) 0.52(0.16-1.65) 0.75(0.14-3.99)	0.291 0.115 0.460 0.121	Adjusted for age, sex, energy intake, vegetable protein intake or animal protein intake (according to the analysis), serum vitamin C, calcium intake, underlying

Non- Prospective	AP:VP ratio: 2 (0.1) g/d	controls drawn from local community population, 80% female						chronic disease, home access, Katz's index, physical activity (METS), HDL cholesterol, and MUFA/PUFA intake.
Nieves et al, 1992, USA	<24g/d to >55g/d	F 50 to 103y	329 (161 cases, 168 controls)	Hip (OR)	Hip fracture	1.04 (0.43, 2.55)	ns	Hospital, age, BMI, oestrogen use, chronic disease status
Non- prospective								
Park et al, 2014,	81.93+/-52.31 g/d	Z score from DXA	Young Women	1157		Protein Intake g/d:		Non-adjusted
Korea Non- prospective					Z-Score ≥0 (n=171) Z-score<0 (n=986)	85.96+/-55.81 81.23+/-51.67	0.276	
Perez- Durillo et al, 2011, Spain Non- prospective	Cases 60 (19)g/d; controls 94 (19) g/d	Women older than 65 y, medical outpatients	44 cases and 42 controls	HF	% energy TP	16.7 (4.7)% (cases (3.0)%) vs 18.3 (control) OR of being a case: (continuous protein	0.07	Non adjusted
					TP intake: (n=86)	intake) 0.96	0.92-1.00	BMI, carbohydrate intake and calcium intake
Preisinger et al, 1995, Austria	15 % total energy, 45-96 g/d	Osteoporosis diagnosis	Post F 50-70 years old	23	IF Intake. (II=60)	Protein intake % mean+/-SEM	0.92-1.00	Non adjusted
Non- prospective					Group 1- Osteoporotic n=12	<u>TP</u> 15.5+/-0.9	NS	
					Group 2 Non- osteoporotic n=11	15.4+/-0.9 AP (g/d) 46.9+/-4.1		

					Group 1- Osteoporotic n=12	42.8+/-3.3		
					Group 2 Non- osteoporotic n=11	VP (g/d) 25.0+/-4.1		
					Group 1- Osteoporotic n=12	25.4+/-2.3		
					Group 2 Non- osteoporotic n=11			
Samieri et 11, 2013, France	70-76 g/d	Men and women 65y and over	1482	Incident fracture of hip, spine	Cases (n=155) Controls(n=1327)	70.4 (26.3) g/d 75.8 (26.8) g/d	0.02	Not adjusted for confounders
Prospective				or wrist		Baseline protein intake		
Wengreen et al, 2004,	1.2g/Kg/d	50-89y M/F	2501 (1157 cases, 1334	Hip (OR)	50-69y (TP) 70-89y (TP)	0.35 1.28	<0.001 0.06	BMI, smoking, alcohol, physical activity, oestrogen use, gender, total
USA			controls		50-69y (AP) 70-89y (AP)	0.43 1.54	0.21 0.95	Calcium and Vitamin D intakes (diet and supplements), potassium intake,
Non- prospective					50-69y (VP) 70-89y (VP)	0.52 0.79	0.19 0.46	age. AP model also adjusted for VP intake, VP model also adjusted for AF intake.

*in g/Kg/d unless otherwise stated, * *(Highest Quartile/Quintile of intake, lowest quartile=1), #OR unless otherwise stated

AP, Animal Protein; BMD, Bone Mineral Density; DPA, Dual Photon Absorptiometry; DXA, Dual X-ray Absorptiometry; LSBMD, Lumbar Spine Bone Mineral Density; OR, Odds Ratio; POM, Postmenopausal; TP, Total Protein; VP, Vegetable Protein

Study, Country,	Design	Baseline protein intake	Supp. (g/d) vs control	Subject Total n	Outcomes Measured	n (I)	Mean, SD(I)	n(p)	Mean/SD(p)	p
Alekel et al, 2000, USA, 24wks	Parallel	No information in paper	Soy vs Whey	2002 PERI F	LSBMC LSBMD BAP	24 24 24	52.96+/-8.72 0.933+/-0.12 15.05+/-5.11	21 21 21	56.57+/-9.74 0.989+/-0.132 12.51+/-4.3	Ns Ns -
Aoe et al, 2001, Japan	Parallel	No information in paper	40mg/d MBP vs Placebo	PRE F	% change in Calcaneal BMD	17	3.42+/-2.05 %	16	2.01+/-1.75 %	0.042
Aoe et al 2005, Japan, 6mo	Parallel	No information in paper	MBP vs Inactive placebo	27 PERI F	NTX OC LSBMD	14 14 14	47.3+/-8.3 5.73+/-0.59 1.11+/-0.03	13 13 13	58.7+/-8.3 5.82+/-0.59 1.09+/-0.03	Ns Ns <0.05
Arjmandi et al, 2003, USA, 3mo	Parallel	Mean (SE)	40g/d Soy protein vs MBP	42 POM F	BAP DPYD	20 20	0.41+/-0.14 7.19+/-3.31	22 22	0.35+/-0.15 6.79+/-3.24	-
Cao et al, 2011, USA	Crossover- 7 weeks in each arm	Soy group – 60(6)g/d	61g/d ('lower protein control- US daily recommenda tion) vs. 118g/d ('higher protein' group)	N=16 40-75 year old postmen opausal women,	NTX Log DPYD	16 16	270 +/- 153 3.7+/-0.61	16 16	227+/-153 3.5+/-0.61	0.41 0.20

TableS6: Characteristics and outcomes of the 30 intervention studies
Ceglia et	Cross-over	MBP group	0.5g/Kg/d	M/F 54-	OC (ng/mL)	10	6.2+/-2.6	10	6.9+/-4.3	Ns
al, 2009	study	(75(9) g/d	(low) vs.	82 years	Urinary NTX/Cr	10	41.0+/-15.2	10	40.4+/-19.1	ns
41 d			1.5g/Kg/d (high)	old N=10	(nmol/mmol)					
			(ingii)	(placebo						
				group						
				used)						
Cuneo et	Parallel	No	Hydrolysed	N=36	BAP	36	26.2(7.2)	35	32.0(10.6)	-
al, 2010,		information	collagen	collagen	CTX	36	0.48(0.1)	35	0.57(0.2)	-
Brazil		in paper	(10g/d	, N=35	OC	36	29.0(8.5)	35	31.8(10.5)	-
			protein) vs. maltodextrin	placebo 45-65						
			placebo	year old						
			practice	post						
				women						
Dalais et	Parallel	Mean (SD):	40g	106	PYD	38	70+/-24.97	40	72.72+/-21.31	Ns
al, 2003,		69.1 (22.1)	Soy protein	POM F	DPYD	38	14.48+/-8.15	40	14.19+/-6.58	ns
AUS, 3mo		g/d	vs casein placebo	50-75 y						
Dawson-	Parallel	Mean (SD)	High	32			High protein		Low protein	
Hughes et		((0.75g/Kg/d)	Elderly	NTX	16	102.3+/-34.5	16	170+/-118.4	0.038
al			vs low (0.04	M/F	OC	16	3.4+/-0.9	16	3.2+/-1.5	0.795
2004,USA,			g/kg/d)							
63d			protein							
Evans et	Cross-over	67(18.8) g/d	Soy protein	Postmen	Change in:					
al, 2007,		(placebo)	isolate (I) vs.	opausal	TBBMD;	21	-0.009 ± 0.013	22	-0.011 ± 0.018	0.72
USA			Milk protein	women	LSBMD	21	-0.011 ± 0.028	22	-0.014 ± 0.022	0.65
9 mo			isolate (p),	N=22,	ProximalFemur	21	0.002 ± 0.016	22	-0.003 ± 0.015	0.29
			exercise	Mean	BMD	21	0.002 + 0.022	22	0.006 + 0.025	0.20
			counterbalan ced across	age 63 years	FNBMD TrochBMD	21 21	$\begin{array}{c} 0.003 \pm 0.022 \\ 0.004 \pm 0.013 \end{array}$	22 22	-0.006 ± 0.025 -0.002 ± 0.018	0.20
			groups (1/2	years	IntertrochBMD	$\frac{21}{21}$	0.004 ± 0.013 0.000 ± 0.025	$\begin{vmatrix} 22\\22 \end{vmatrix}$	-0.002 ± 0.013 -0.002 ± 0.023	0.23
			in each							
			group							
			exercise, ¹ / ₂		BAP	21	-2.1 ± 4.0	22	1.2 ± 4.7	0.02
			in each		CTX	21	-0.08 ± 0.09	22	-0.02 ± 0.11	0.02
			group no exercise)							
			(ACICISE)							

Hunt et al, 2009, USA	Cross-over	61.9(24) g/d (collagen)	Low Calcium	N=13 in two LC	Group 1 LC: LCLP (p) vs.					
		_	(LC)(670mg/	arms,	LCHP (I)					
7wk each			d) Low	n=14 in						-
arm			protein	two HC	Log DPYD	13	2.3 (0.2)	13	2.4 (0.2)	-
			(10% of total	arms	Log OC	13	1.74(0.74)	13	1.94(0.74)	-
			energy,		BAP	13	0.55(0.08)	13	0.52(0.08)	-
			0.8g/Kg/d) vs. High	POM F	TRAP	13	52.5(8.0)	13	55.1(8.0)	
			protein (20%		Group 2 HC:					
			of total		HCLP (p) vs.					
			energy,		HCHP(I)					
			1.6g/Kg/d).							
					Log DPYD					-
			HighCalcium		Log OC	14	2.2(0.2)	14	2.3(0.2)	-
			(HC)		BAP	14	1.90(0.74)	14	1.73(0.74)	-
			(1500mg/d)		TRAP	14	0.58(0.02)	14	0.53(0.02)	-
			Low protein			14	57.6(8.0)	14	55.7(8.0)	
			(10% of total							
			energy, 0.9g/Kg/d) vs. High protein (20% of total energy1.7g/ Kg/d).							
Ince et al	Cross-over	109(7) g/d	High	39 Pre			High protein		Low protein	
2004,		Soy group	(1.1g/Kg/d)	F, 22-	NTX	39	442+/-124.9	39	360+/-99.9	<0.001
USA,		boy group	vs low	39y	OC	39	15.8+/-8.74	39	13.4+/-8.1	0.166
2wks			(0.8g/Kg/d) protein diet							
Kenny et	Parallel	112(6)g/d	Soy protein	Women			Mean(SEM)		Mean(SEM)	
al, 2009,		Placebo	(I) vs. Mixed	over 60	Change in	24	0.001 + / -0.005	22	-0.003(0.005)	0.317
USA, 1y			control	years	FNBMD	- '				0.017
			protein	old	Change in					
			(Casein,	(mean=71y)	LSBMD	24	0.001+/-0.008	22	0.010+/-0.007	0.181

			Whey and		BAP	24	18.8+/-1.07	22	25.2+/-2.03	0.050
			Egg) (p). No		NTX	24	30.2+/-2.74	22	35.0+/-3.21	0.50
			isoflavones			2.	30.217 2.71		55.017 5.21	0.50
			in these two							
			study arms							
Kerstsetter	Parallel	No	45g Whey	Men			Mean(SEM)		Mean(SEM)	
et al, 2015,		information	protein (I) or	over 70	LSBMD	106	1.05(1.10+/-0.01)	102	1.02(1.11+/-	-
USA,		in g	isocaloric	y and	TotalHipBMD	106	1.06(0.88+/-0.01)	102	0.02)	-
18mo			maltodextrin	women	FNBMD	106	1.06(0.80+/-0.01)	102	1.02(0.89+/-	-
			(p)	over 60		100	45(00.2.4.4.20)	100	0.01)	
				years,	LSBMD P1NP nmol/L	106	45(99.3+/-4.29)	102	1.02(0.82+/-	-
				n=121	CTX ng/L	61 61	1.32+/-0.06 480+/-30	60 60	0.01)	0.395 0.041
					OC nmol/L	61	1.12+/-0.05	60	44(106+/-4.07)	0.041
						01	1.12+7-0.03	00	1.35+/-0.07	0.775
									440+/-30	
									1.18+/-0.06	
Kerstetter	Cross-over	(17-18% of	High	16 Pre			Mean+/-SEM		Mean+/-SEM	
et al, 1999,		total energy)	(2.1g/kg/d)vs	F, 20-	OC	-	5.7+/-0.8	-	7.6+/-1.4	Ns
USA, 4d			low	40g	BAP	-	57.2+/-7.8	-	69.4+/-8.8	Ns
			(0.7g/kg/d)		NTX	-	48.2+/-7.2se	-	32.7+/-5.3	<0.05
			protein							
Khalil et	Parallel	No	Soy vs Milk	64 M,	BAP	24	-	22	-	Ns
al, 2002,		information	protein (40g)	59.2+/-	DPYD	24	-	22	-	ns
US, 3mo		in paper		17.6y						
Jenkins et	Cross-over	63(15) g/d	Vegetable	20	NTX	20	584+/-340	20	461+/-259	
al, 2003,	Cross-over	05(15) g/u	diet (27%	Middle	BAP	$\frac{20}{20}$	20+/-4.5	$\begin{vmatrix} 20\\20 \end{vmatrix}$	401+/-239	-
USA, 2mo			protein) vs	aged	DAI	20	20+7-4.5	20	197/4.5	-
Crossover			Control	M/F						
010550701			diet(16%	141/1						
			protein)							
Lampl et	Parallel	69(17) g/d	Normal diet	7-13	Periosteal	26	5.9+/-0.1	30	5.7+/-0.1	< 0.05
al. 1978,			(11g/d)(p)	year old	breadth (mm)					
New			vs. normal	children	Endosteal	26	2.8+/-0.1	30	2.8+/-0.1	ns
Guinea, 8			diet plus	with	breadth (mm)					
mo			20g/d milk	low	Compact bone	26	3.1+/-0.1	30	2.8+/-0.1	ns
			protein		breadth (mm)					

Martin- Bautista, 2011, Spain 4 mo	Parallel	1.1 kg/d	supplement(I) Collagen (without calcium) group vs. Placebo	protein intakes 38	BAP OC TRAP CTX	20 20 20 20 20	GP 2 2.35+/-42.6 -4.0+/-8.1 -1.2+/-4.0 0.03+/-0.44	18 18 18 18	Mean+/-SEM GP 1 -28.6+/-29.9 -2.1+/-14.3 1.6+/-4.2 0.07+/-0.43	NS NS NS <0.05
Roughead et al, 2003, USA, 8wk	Cross-over	No information in g	High meat (20% of energy) versus low meat(12% of energy) diet	15 POM F	HPO OC NTX BAP	15 15 15 15	High 71.5 4.01 3.79 18.1	15 15 15 15	Low 64.5 3.94 3.83 18.3	0.001 Ns Ns Ns
Schurch et al, 1998 Switzerlan d, 6mo	Parallel	(18% of total energy)	Total protein (20g/d) vs placebo	82 Elderly M/F 80.7y+/- 7.4	%change DPYD FSBMD LSBMD OC PFBMD PYD TrochBMD TBBMD	- - - - - -	-9.2 -1.61 -3.05 7.9 -2.95 6.6 -3.02 -3.77		1.4 -1.23 -6.11 6.9 -3.37 17 -3.65 -3.1	>0.2 >0.2 >0.2 >0.2 >0.2 >0.2 >0.2 >0.2
Shapses et al, 1995, USA, 5d	Cross-over	Mean +/-SD	LPHC(0.44g/ Kg/D protein, p) vs. HPHC (2.71g/kg/d, I) Calcium in both groups=1600 mg/d	21-42 year old males and females	HPO (mol/mol)	15	0.011+/-0.008	13	0.010 +/-0.007	-
Spence et al, 2005, USA, 28d per phase	Cross-over	Soy group- 62.5 (13.7) g/d	Soy protein isolate without isoflavones (I) vs.	N=15 POM F	BAP ng/mL OC ng/mL NTX nmolBCE/mmol Cr	15 15 15	14.8+/-4.5 10.2+/-3.9 48.0+/-22.6	15 15 15	14.3+/-4.0 8.1+/-3.8 55.6+/-29.0	<0.05 <0.05 ns

Tkatch et al, 1992, Switzerlan d, 38days	Parallel	Mixed control group- 57.0(21.9)	casein-whey protein (p) 20.4g/d Protein in nutritional supplement vs. the same nutritional supplement without protein	62 M/F elderly, mean age 82y	Change: FNBMD FSBMD LSBMD OC	25 24 25 24	0.569+/-0.105 0.24+/-0.049 0.88+/-0.18 6.94+/-2.45	23 22 23 18	0.579+/-0.12 1.257+/-0.3 0.81+/-0.17 4.96+/-2.93	<0.05
Toba et al 2001, Japan, 16d	Parallel	1.0g/Kg/d	MBP (30mg/d) vs inactive	30 M, 36.2y+/- 8.5	NTX OC	30 30	26.8+/-9.6 5.4+/-1.8	30 30	31.5+/-10.2 3.7+/-1.8	<0.001 <0.001
Uenishi et al, 2007, Japan, 6mo	Parallel	Mean (SEM):	placebo 40mg/d MBP vs inactive placebo	35 Pre F	LSBMD %change in LSBMD	17 17	1.16+/-0.14 +1.75%	18 18	1.13+/-0.16 +0.13%	- 0.042
Vupadhya hula et al, 2009, USA	Parallel	72.9(1.8) Maltodextrin Group	25g soy protein (no isoflavones), 25g milk (casein, whey) protein	203 POM F Mean (SE) age 64 0.6)y	SpineBMD FNBMD TrochBMD TotalFemoralB MD TBBMD	48 48 48 48 48	Mean+/-SE 1.068+/- 0.02 0.845+/-0.01 0.741+/-0.01 0.892+/-0.02 1.078+/-0.01	52 52 52 52 52 52	Mean+/-SE 1.082+/-0.02 0.869+/-0.01 0.747+/-0.01 0.897+/-0.01 1.094+/-0.01	- - - -
					%change from baseline: NTX:Cr	22	2.27+/-2.1	30	-1.86+/-2.3	-
Yamamura et al, 2002, Japan	Parallel	73.9(1.9) Whey Group	MBP(40mg) vs inactive placebo	33 Pre F	RBMD	17	-Missing data	16	-Missing data	-
Zhu et al, 2011, AUS, 2y	Parallel	No information in paper	High protein drink (I) vs.	219 70- 80 year	Total Hip vBMD	67	Mean(SEM) -3.63+/-1.10	66	Mean(SEM) -3.82+/-1.43	-

			low protein	old	FN vBMD	67	-2.39+/-1.25	66	-0.24+/-1.19	
			drink (p)	women						
					Baseline FN	91	0.70+/-0.010	88	0.71+/-0.012	0.35
					aBMD					
					2 yr FN aBMD	91	0.69+/-0.010	88	0.70+/-0.012	0.33
Zou et al	Parallel	11g/d	Milk with	57	TBBMD	29	0.946+/-0.064	28	0.913+/-0.053	-
2009,			40mg MBP	women,	LSBMD	29	1.041+/-0.118	28	0.995+/-0.068	-
China, 8			(I) vs. Milk	20 years	DistalRadius/Ul	29	0.351+/-0.041	28	0.341+/-0.036	-
mo			without MBP	old	na BMD					
			(p)							

aBMD, areal Bone Mineral Density; BAP, Bone Specific Alkaline Phosphatase; BCE, Bovine Collagen Equivalents; BMD, Bone Mineral Density; Cr, Creatinine; CTX, Cterminal telopeptide of collagen; DPYD, Deoxypyridinoline; FNBMD, Femoral Neck Bone Mineral Density; FSBMD, Femoral Shaft Bone Mineral Density; GP, Group; HCHP, High Calcium High Protein; HCLP, High Calcium Low Protein; HPO, Hydroxyproline; IntertrochBMD, Intertrochanter Bone Mineral Density; LCHP, Low Calcium High Protein; LCLP, Low Calcium Low Protein; LSBMC, Lumbar Spine Bone Mineral Content; LSBMD, Lumbar Spine Bone Mineral Density; NTX, N-terminal telopeptide of collagen; OC, Osteocalcin; P1NP, Procollagen type 1 N-terminal propeptide; PERI, Perimenopausal; PFBMD, Proximal Femur Bone Mineral Density; POM, Postmenopausal; PRE, Premenopausal; RBMD, Radial Bone Mineral Density; TBBMD, Total Body Bone Mineral Density; TRAP, Tartrate Resistant Alkaline Phosphatase; TrochBMD, Trochanter Bone Mineral Density; vBMD, volumetric Bone Mineral Density

		Correlation				
Study	Total		COR	95%-CI	W(fixed)	W(random)
Alissa2014	300			[0.07; 0.29]	6.3%	6.7%
Chan2009*	441			[-0.12; 0.07]	9.3%	8.7%
Chiu1997	258	+		[-0.04; 0.20]	5.4%	6.0%
Coin2008	136			[-0.07; 0.26]	2.8%	3.6%
Cooper1995* Postmenopausal	218		0.02	[-0.11; 0.15]	4.5%	5.3%
Cooper1995* Premenopausal	72	\ {	0.27	[0.04; 0.47]	1.5%	2.0%
Gunn2014	142		0.19	[0.03; 0.34]	2.9%	3.7%
Henderson1995	115		0.22	[0.04; 0.39]	2.4%	3.1%
Ho2003*	438		0.04	[-0.06; 0.13]	9.2%	8.6%
Jaime2006 Black	31	+	- 0.36	[0.01; 0.63]	0.6%	0.9%
Jaime2006 White	246		0.05	[-0.07; 0.18]	5.1%	5.8%
Kumar2010	255		0.04	[-0.08; 0.16]	5.3%	6.0%
Lau1998	76		0.13	[-0.10; 0.35]	1.5%	2.1%
Michealsson1995	175		0.12	[-0.03; 0.26]	3.6%	4.4%
New1997	994		0.02	[-0.04; 0.08]	21.0%	13.2%
Orozco1998	76		-0.03	[-0.25; 0.20]	1.5%	2.1%
Rapuri2003	473	- <u>-</u>	0.09	[0.00; 0.18]	9.9%	9.1%
Teegarden1998	215		0.08	[-0.05; 0.21]	4.5%	5.2%
Wang1997	125		-0.01	[-0.19; 0.17]	2.6%	3.3%
Fixed effect model	4786	†	0.07	[0.04; 0.09]	100%	
Random effects model		🔶	0.07	[0.04; 0.11]		100%
Heterogeneity: I-squared=25.5%, tai	u-squared <u>=0</u> .	0014, ρ=0.1498		_		
	-0.6	-0.4 -0.2 0 0.2 0.4 0.	6			

Figure S1 Femoral Neck Bone Mineral Density- correlation coefficients for association with dietary protein intake*=multivariate adjusted data

		Correlation				
Study	Total	1. 11	COR	95%-CI	W(fixed)	W(random)
Alissa2014	300			[-0.13; 0.09]	7.1%	7.0%
Chiu1997 Cooper1995* Postmenopausa	258 1 218			[-0.02; 0.23] [-0.18; 0.08]	6.1% 5.1%	6.7% 6.2%
Cooper1995* Premenopausal				[-0.16; 0.08]	1.6%	3.3%
Henderson1995	115			[-0.13; 0.23]	2.7%	4.5%
Ho2003*	438			[-0.08; 0.11]	10.3%	7.9%
Horiuchi2000	85			[0.01; 0.42]	2.0%	3.7%
Kumar2010	255		0.22	[0.10; 0.34]	6.0%	6.6%
Lau1998	76		0.09	[-0.14; 0.31]	1.7%	3.4%
Michealsson1995	175	- <u> </u>		[-0.09; 0.20]	4.1%	5.6%
New1997	994			[-0.03; 0.09]	23.6%	9.3%
Orozco1998	76			[-0.25; 0.20]	1.7%	3.4%
Quintas2003	164			[0.17; 0.45]	3.8%	5.4%
Rapuri2003	473			[-0.03; 0.15]	11.2%	8.1%
Teegarden1998	215			[0.06; 0.32]	5.0%	6.2%
Thorpe2008	161			[-0.16; 0.14]	3.8%	5.4%
Wang1997	125			[-0.14; 0.21]	2.9%	4.7%
Whiting2002	57		- 0.42	[0.18; 0.61]	1.3%	2.7%
Fixed effect model	4257	\$		[0.04; 0.10]	100%	
Random effects model	,		0.09	[0.04; 0.14]		100%
Heterogeneity: I-squared=58.1%, ta	u-squared=	-υ.υυσ3, ρ=υ.υυ11	1			
	-0.0	6 -0.4 -0.2 0 0.2 0.4 0	.6			

Figure S2 Lumbar Spine Bone Mineral Density- correlation coefficients with dietary protein intake *=multivariate adjusted data

Study	TE seTE	Hazard	Ratio	HR	95%-Cl	W(fixed)	W(random)
Langsetmo2015Men Langsetmo2015Women Misra2011 Sahni2010HighCa Sahni2010LowCa	-0.42 0.320 -0.16 0.125 -0.46 0.280 -0.62 0.740 - 0.79 0.470			0.85 [0 0.63 [0 0.54 [0	0.35; 1.23] 0.67; 1.09] 0.36; 1.09] 0.13; 2.29] 0.88; 5.54]	10.5% 68.9% 13.7% 2.0% 4.9%	18.6% 44.1% 22.2% 4.7% 10.4%
Fixed effect model Random effects model Heterogeneity: I-squared=35		=0.0479 , p=0.1855 0.2 0.5 1	2 5	-	.67; 1.01] .59; 1.14]	100% 	 100%

Figure S3 Total Protein intake and Hazard Ratio for Fracture (cohort studies) Lowest intake category=reference (OR=1)



Figure S4 Protein intake and Odds Ratio of Fracture (case control studies) Lowest intake category=reference (OR=1)

	E	perim	ental		Co	ntrol	Mean difference				
Study	Total	Mean	SD	Total	Mean	SD	1.1	MD	95%-CI	W(fixed)	W(random)
Tkatch1992 Kerstetter2015	25 105	- 17 TE-21	0.18 0.10	23 102		0.17 0.20			[-0.03; 0.17] [-0.01; 0.07]		16% 84%
Fixed effect model Random effects model Heterogeneity: I-squared=0%		uared=(), ρ=0.	125 4682					[0.00; 0.08] [0.00; 0.08]		 100%
							0.15-0.1-0.05 0 0.05 0.1 0.15	5			

Figure S5 Effects of Total Protein intake on areal Lumbar Spine Bone Mineral Density in randomized controlled trials

	Experimental	Control	Mean difference			
Study	Total Mean SD Total	Mean SD		MD 9	5%-CI W(fixed) W(random)
Tkatch1992 Kerstetter2015	25 0.57 0.105 23 106 1.06 0.100 102	0.58 0.14 -		Contraction of the second s	3; 0.06] 7.79 1; 0.07] 51.99	
Zhu2011	91 0.69 0.100 88	0.70 0.11		-0.01 [-0.04	2011년 2011년 - 11월 2011년 201 1월 2011년 2	
Fixed effect model Random effects model	222 213 %, tau-squared=0.0008, p=0.044	r		0.02 [0.00 0.01 [-0.03		 100%
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Figure S6: Effects of Total Protein intake on areal Femoral Neck Bone Mineral Density in randomized controlled trials



Figure S7: Milk Basic Protein supplementation: Effects on Lumbar Spine Bone Mineral Density

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