

Review Article

The genetic architecture of osteoporosis and fracture risk

Katerina Trajanoska, Fernando Rivadeneira*

Department of Internal Medicine, Erasmus MC University Medical Center, Rotterdam, Netherlands



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ABSTRACT

Osteoporosis and fracture risk are common complex diseases, caused by an interaction of numerous disease susceptibility genes and environmental factors. With the advances in genomic technologies, large-scale genome-wide association studies (GWAS) have been performed which have broadened our understanding of the genetic architecture and biological mechanisms of complex disease. Currently, more than ~90 loci have been found associated with DXA derived bone mineral density (BMD), over ~500 loci with heel estimated BMD and several others with other less widely available bone parameters such as bone geometry, shape, and microarchitecture. Notably, several of the pathways identified by the GWAS efforts correspond to pathways that are currently targeted for the treatment of osteoporosis. Overall, tremendous progress in the field of the genetics of osteoporosis has been achieved with the discovery of *WNT16*, *EN1*, *DAAM2*, and *GPC6* among others. Assessment of the function and biological mechanisms of the remaining genes may further untangle the complex genetic landscape of osteoporosis and fracture risk. With this review we aimed to provide a general overview of the existing GWAS studies on osteoporosis traits and fracture risk.

1. Introduction

Osteoporosis is a progressive silent disease with devastating clinical and economic consequences. Approximately 1/3 of postmenopausal women suffer osteoporosis worldwide and at least half of these will experience a fragility fracture during their lifetime. Fragility fractures are often associated with increased morbidity and mortality, dramatically decreasing quality of life [1–3]. As population age, the prevalence of osteoporosis and its sequelae will increase substantially, becoming one of the largest global healthcare burdens.

Osteoporosis and fracture risk are determined by a complex interplay of genetic and environmental factors. Positive family history of osteoporosis is an important risk factor for fracture, which underscores the pivotal relationship between an individual's genetic makeup and disease susceptibility. Many monogenic forms of bone fragility have been identified, which are caused by a single mutation in a gene that has a major role in skeletal biology such as observed in X-linked osteoporosis, osteogenesis imperfecta and Paget disease among many others [4]. However, these monogenic mutations explain a very small fraction of the variation in bone mineral density (BMD) and osteoporosis risk in the general population. Just as in other complex diseases, advances in high-throughput genomic technologies, increasing insight on how genetic variation is organized in the genome (i.e., HapMap project) and the availability of large biobank studies have led to the

advent of genome-wide association studies (GWAS) in the osteoporosis field. Given the enormous progress in the genomics of osteoporosis, the aim of this review is to provide a general overview of the existing GWAS studies on osteoporosis and fracture risk.

2. Heritability of bone properties

In order to evaluate the genetic architecture of any trait it is important first to establish if that trait is heritable. DXA derived BMD, used for the diagnosis of osteoporosis, is a highly heritable trait ($h^2 = 50\text{--}80\%$) and an excellent biomarker capturing intrinsic properties of bone biology that have led to the identification of hundreds of associated loci [5]. Twin and family studies have also shown that other bone parameters like geometry ($h^2 = 30\text{--}70\%$) [6], bone ultrasound measures ($h^2 = 40\text{--}50\%$) [5] and high resolution peripheral quantitative computed tomography (HR-pQCT) measures of bone microarchitecture ($h^2 = 20\text{--}80\%$) [7] are also highly heritable. Such other determinants of bone strength like geometry, cortical thickness and porosity, trabecular bone morphology and intrinsic properties (quality) of bone tissue contribute to the genetic predisposition to fragility fractures but have been less well-studied due to limitations in sample size. Large samples are needed since all these bone parameters are expected to be highly polygenic (i.e., determined by many variants with small effects).

* Corresponding author.

E-mail address: f.rivadeneira@erasmusmc.nl (F. Rivadeneira).<https://doi.org/10.1016/j.bone.2019.04.005>

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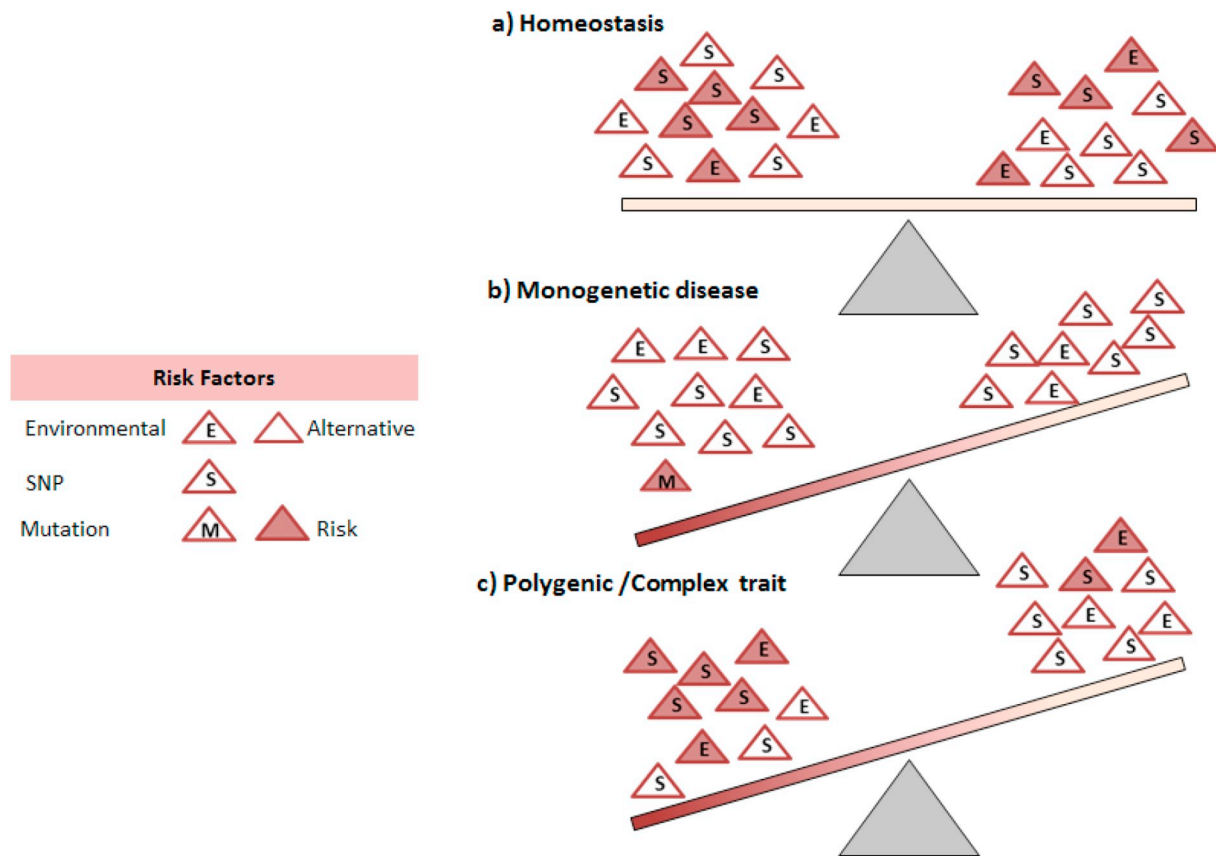


Fig. 1. Genetic and environmental factors both act on the musculoskeletal system and increase the risk of fracture. Complex traits influenced by environmental factors and/or genetic variants with weak effects need to synergize their effects in order to increase fracture risk. By contrast, monogenetic conditions are sufficient on its own to cause fracture.

3. Fracture risk: an extremely complex trait

Numerous genetic and environmental factors, individually or combined, can cause fracture. These risk factors can act through one or multiple systems, triggering different processes across hormonal/metabolic, biomechanical and material property pathways, which eventually in combination will lead to fracture. Importantly, certain non-genetic factors can exert their influences during different life stages where distinct processes have greater influence like bone accrual in children or bone loss with aging. Genes can act alone or in combination with environmental factors (gene x environment interactions) to have an effect on fracture risk (Fig. 1). Sometimes it may be difficult to distinguish fractures with high environmental influence from fractures having considerable influence from genetic variants. In addition, the musculoskeletal system undergoes adaptation, in principle directed to repair frequent and repetitive micro-damage and to preserve bone-strength to the needs set by strain and stresses. This adaption can also occur in presence of genetic susceptibility to fracture, making the search for fracture genes even more difficult.

Phenotype definition is also an important determinant of the success of GWAS, frequently confronted with a trade-off between accurate trait definition and sample size. Ideally, one would choose to study fractures with a strong hereditary component i.e., low-trauma fractures occurring after falls from a standing height or less which are the consequence of osteoporosis (BMD T-score < −2.5). Nevertheless, it is well established that the majority of fractures occur above the T-Score osteoporosis threshold [8] in individuals with osteopenia (low BMD with $-2.5 < \text{T-score} < -1.0$). Moreover, low-trauma fractures may also occur in individuals with normal BMD (T-score > −1.0), who are for example carriers of rare mutations which affect distinct bone properties such as

in the monogenetic forms of hypophosphatemic rickets or osteogenesis imperfecta. These fractures occur beyond what is expected from a low BMD level and are typically attributed to impaired “bone quality” – an ambiguous term for bone properties influencing mechanical performance but not well-characterized by BMD. The mechanistic pathways affecting bone quality have been set to arise from three distinct tissue material properties that describe mechanical failure of bone: including tissue strength, fracture toughness and fatigue strength [9]. Lacking to date adequate, affordable and wide-spread methods to assess bone quality, it is no wonder that the majority (if not all) current GWAS have focused on BMD-based and fracture risk traits. Interestingly, as described below all fracture risk loci identified to GWAS are BMD loci, further supporting the mediating role of BMD on fracture. Overall, the GWAS field has prioritized sample size (high powered setting) over the use of accurate phenotype definition as the most cost-effective strategy to reduce the noise introduced by phenotypic heterogeneity and identify real genetic signals.

4. Genetic studies of bone mineral density

The early genetic studies in the osteoporosis field were confined to linkage and candidate gene association studies, which have been particularly focused on BMD. However, these studies have turned out to be typically underpowered leading to negative or irreproducible findings. With the advent of GWAS a new era in bone genomics has begun. The findings from the first GWA study from the Framingham study were initially underwhelming as a result of poor marker density (100,000 SNPs) and inadequate statistical power ($N_{\text{total}} = 1141$) [10]. Thanks to the advances in genomic technologies and statistical genetic methodologies it became clear that the success of GWAS depends on sample

size, effect size of causal genetic variants, the frequency of those variants and the LD between observed genotyped variants and the unknown causal variants [11]. This knowledge has yielded a dramatic change shaping the success of subsequent GWAS, where with the rise of sample size the loci associated with BMD started to increase dramatically.

In early 2008 two GWAS simultaneously identified five common variants associated with BMD variation in the general population. Richards et al. [12] identified two variants associated with lumbar spine and femoral neck BMD mapping to *LRP5* and *TNFRSF11B* (OPG) in 8557 individuals. The *LRP5* variants were also associated with osteoporotic fractures, reproducing the findings of the largest candidate gene study drawn by the GENOMOS consortium [13]. Almost simultaneously, Styrkársdóttir et al. [14] identified in addition to *TNFRSF11B* (OPG) variants mapping to *ESR1*, *ZBTB40* and the major histocompatibility complex (MHC) loci in 13,786 Icelandic individuals, associated with BMD and osteoporotic fractures. Subsequent work by the same group with expanded sample size ($N_{\text{total}} = 15,375$) identified additional novel variants mapping to/near *TNFRSF11A* (RANK), *SOST*, *MARK3*, and *SP7* (osterix) [15]. The latter gene (*SP7*) was also identified in the first BMD GWAS of children [16] ($N_{\text{total}} = 5275$). Medina-Gomez et al. [17] identified variants in *WNT16* associated with skull and total body BMD in children, accompanied by an effort identifying variants in the same locus associated with pQCT and wrist fracture outcomes [18]. The largest yield in discoveries has been facilitated by the rise of collaborative networks giving way to large-scale GWAS meta-analyses that identified novel bone regions and pathways. The first meta-analysis of the GEFOS consortium ($N_{\text{total}} = 19,195$) identified 13 novel loci associated with BMD [19] followed by a second GEFOS meta-analysis ($N_{\text{discovery}} = 32,961$) which replicated the majority of known BMD loci and identified additional 32 novel loci ($N_{\text{total}} = 83,894$) [20]. Fourteen of the BMD-associated loci were also associated with osteoporotic fractures, with those mapping to *FAM210A*, *SLC25A13*, *LRP5*, *MEPE*, *SPTBN1* and *DKK1* showing strongest association. This study was also the first one to identify sex-specific effects by examining the X chromosome. The variant (rs5934507), associated with BMD mapping to Xp22.31, has been previously associated with male serum testosterone levels [21]. The latter study was followed by additional large meta-analysis by Zheng et al. [22] (described below) which have provided evidence that low-frequency non-coding variants have large effects on BMD and fracture. Next, in 2018 Medina et al. [23] in a meta-analysis of 30 GWASs ($N_{\text{total}} = 66,628$) identified 80 loci associated with total body BMD, of which 36 had not been previously identified. Moreover, in the age-stratified analyses only two loci displayed evidence for age-specific effects, including variants in *ESR1* and in close proximity to RANKL. These findings suggest that most of the genes identified throughout the life-course, exert an effect on peak BMD acquisition and this effect can still be observed decades later [23]. Three recent studies have used bone mineral density estimated from heel ultrasound (eBMD), in the UK Biobank Study. In the first effort ($N_{\text{total}} = 142,487$) Kemp et al. [24] identified 203 loci associated with eBMD and in the following effort ($N_{\text{total}} = 426,924$) this number was increased to 518 (301 novel) [25]. In the latter UK biobank setting, Kim identified 613 novel loci using less stringent conditional analysis [77]. These studies highlight the value of expanding the sample size for GWAS and the amazing opportunities to unravel novel biology provided by the approach. To date, >20 GWAS have been published for different bone parameters from which three are large meta-analysis and three are based on the UKBiobank study (Table 1).

The majority of the GWA studies have scrutinized common variants ($\text{MAF} > 5\%$.) All these efforts have identified variants together explaining 10–20% of the variance in bone phenotypes. It is well established that less-common (rare) variants can have bigger effects than those from common ones. An alternative approach is to focus on individuals with extremely low or high BMD in order to identify rare variants with relatively large effect. In line with this contention, whole-

genome sequencing (WGS) efforts have been also successful in mapping rare variants associated with different monogenetic conditions. Using WGS and imputation to larger population sets, a rare novel variant ($\text{MAF} = 0.17\%$) was associated with low BMD (4931 low BMD cases and 69,034 controls) and fracture risk as a result of rare nonsense mutation within *LGR4* (c.376C > T) [26]. A few years later, also using WGS and imputation the same group discovered two rare mutations in *COL1A2* associated with low BMD (2984 cases and 206,675 controls) in participants without signs of osteogenesis imperfecta. In 2018 Duncan et al. [27] performed the most comprehensive extreme phenotype study in 240 individuals from UK with extreme high BMD ($Z\text{-scores} \geq +3.2$) and 1955 women with high ($N = 1055$) or low ($N = 900$) BMD. The analyses yielded two novel loci mapping near *NPR3* (rs9292469; $\text{MAF} = 0.33\%$) associated with lumbar spine BMD and *SPON1* (rs2697825; $\text{MAF} = 0.17\%$) associated with total hip BMD. Finally, Zheng et al. [22] using an extremely powerful WGS design identified novel rare variants associated with BMD variation in the general population. The rare noncoding variant mapped to *EN1* and showed large effects on BMD ($N_{\text{total}} = 53,236$, effect size = +0.20 standard deviations [SD]) and fracture risk ($N_{\text{total}} = 508,253$, $\text{OR} = 0.85$).

5. Genetic studies of fracture risk

Fracture is the most important clinical outcome of osteoporosis. In the past, most of the genes shown to be associated with fracture risk have been discovered by testing known GWAS BMD loci for association with fracture as described above. To date two GWAS have been performed using vertebral fractures as an endpoint. In the first meta-analysis one locus on chromosome 16q24 (rs11645938) was associated with the risk of radiographic vertebral fractures, which failed to replicate across 5720 cases and 21,791 controls [28]. A recent meta-analysis reported a locus mapping on chromosome 2q13 to be significantly associated with clinical vertebral fractures [29]. The first GWAS study on non-vertebral osteoporotic fractures ($N = 700$) was performed in elderly Chinese individuals and identified one fracture-associated locus within the *ALDH7A1* gene [30]. However, this gene failed to replicate in any of the larger European meta-analyses. In 2018, Trajanoska et al. [31] conducted the largest GWAS on osteoporotic fractures to date comprising 37,857 cases and 227,116 non-cases with replication in up to 300,000 individuals (147,200 cases). Altogether, the effort identified 15 fracture loci with modest effects. Interestingly, all identified loci were known BMD loci. Overall, the effect of these SNPs on fracture was smaller than the effect on BMD (Fig. 2). Thus, the genetics of any-type of fractures in the general population is mediated through the genetic influence on BMD. This is well characterized by the genetic correlations of fracture risk with BMD. Further, among 15 tested clinical factors (including vitamin D levels and milk calcium intake) only BMD had a major causal effect on fracture [31].

6. Genetic studies of other bone parameters

Studies have been performed on other bone parameters. Loci mapping near RANK/OPG have been associated with cortical volumetric BMD [32,33]. While genetic variants in the *FMN2/GREM2* locus were associated with trabecular volumetric BMD and fracture risk [33]. Moreover, five loci have been reported to be associated with lumbar spine volumetric BMD ($N_{\text{total}} = 15,275$) mapping near *WNT4* and *ZBTB40*, *TNFRSF11B*, *AKAP11*, and *TNFSF11*; from which two loci (5p13 and 1p36.12) were associated with vertebral fractures [34]. Several GWA studies have identified *RAP1A*, *TBC1D8*, and *OSBPL1A* to be associated with hip structure analysis (HSA) parameters [35]. Finally, Baird et al. [36] identified nine loci associated with hip shape. Seven SNPs were within 200 kb of genes involved in endochondral bone formation, namely *SOX9*, *PTHrP*, *RUNX1*, *NKX3-2*, *FGFR4*, *DICER1*, and *HHIP* [36].

Table 1
GWASs for different bone parameters and osteoporotic fractures.

Study	Trait	Sample size	Total number of GWs loci	Total number of GWs novel loci	Genes
Richards et al. (2008)	Lumbar spine and femoral neck BMD	8557	2	2	<i>TNFRSF11B, LRP5</i>
Szykarski et al. (2008)	Lumbar spine and femoral neck BMD	13,786	4	3	<i>RANKL, OPG, ESRI, ZBTB40, MHC</i>
Szykarski et al. (2008)	Lumbar spine and femoral neck BMD	15,375	7	4	<i>SOST, MARK3, SP7, TNFRSF11A</i>
Timpson et al. (2009)	Lumbar spine and femoral neck BMD	5275	1	1	<i>SP6</i>
Xiong et al. (2009)	Lumbar spine and femoral neck BMD	9109	2	2	<i>ADAMTS18, TGFBR3</i>
Rivadeneira et al. (2009)	Lumbar spine and femoral neck BMD	19,195	20	13	<i>GPR177, SPTBN1, CTNBN1, MEPE, MEF2C, STARD3NL, FLJ42280, LRP4, ARHGAP1, F2, CDC5, SOX6, FOXL1, HDAC5, CRHR1</i>
Zhao LJ et al. (2009)	Femoral neck bone geometry	5676	1	1	<i>RTP3</i>
Guo et al. (2010)	Hip BMD and osteoporotic fractures	11,568	1	1	<i>ALDH7A1</i>
Kung et al. (2010)	Lumbar spine and femoral neck BMD	18,898	1	1	<i>JAG1</i>
Hsu et al. (2010)	Lumbar spine and femoral neck BMD	11,290	4	0	
Koller et al. (2010)	Lumbar spine and femoral neck BMD	2193	0	0	
Paternoster L et al. (2010)	Cortical vBMD	5789	1	0	<i>RANKL</i>
Duncan et al. (2011)	Extreme high or low hip BMD	21,798	2	2	<i>GALNT3, RSP03</i>
Estrada et al. (2012)	Lumbar spine and femoral neck BMD	83,894	56	32	<i>*DNM3, ANAPC1, LEKRI, IDUA, WNT16, FUBP3, MPP7, MBL/DKK1, AXIN1, SOX9, FAM9B/KAL1</i>
Zheng et al. (2012)	Cortical bone thickness, forearm BMD and fracture	5878	1	0	<i>WNT16</i>
Medina-Gomez et al. (2012)	Total Body BMD	13,712	1	0	<i>WNT16</i>
Paternoster L et al. (2013)	Cortical and trabecular vBMD	6930	5	1	<i>RANKL, OPG, ESRR, LOC285735, FMN2/GREM2</i>
Zhang et al. (2013)	Lumbar spine, hip and femoral neck BMD	27,061	15	2	<i>SMOC1, CLDN14</i>
Moayyeri A et al. (2014)	BUA, VOS, BMD	15,514	9	1	<i>TMEM135</i>
Oei et al. (2014)	Radiological vertebral fractures	2995	1	1	<i>16q24</i>
Zheng et al. (2015)	Lumbar spine, femoral neck and forearm BMD	53,236	1	1	<i>EN1</i>
Pei et al. (2016)	Hip trochanter and intertrochanteric BMD	9174	3	1	<i>RP11-384F7.1</i>
Mullin et al. (2017)	BUA and VOS	16,627	8	3	<i>PPP1R3B, LOC387810, SEPT5</i>
Kemp et al. (2017)	heel BMD	142,487	203	153	<i>*GPC6</i>
Medina-Gomez et al. (2018)	Total Body BMD	66,628	80	36	<i>*RERE, CSF1, SLC8A1, PLCL1, AQP1, SMAD9, TOM1L2, ADAMTS5, ETS2</i>
Duncan et al. (2018)	Extreme high or low BMD	2195	4	2	<i>NPR3, SPON1</i>
Braid et al. (2018)	(DXA)-derived hip shape	15,934	8	8	<i>FGFR4, ASTN2, 17q42.3, 14q32.13, 12p11.22, 21q21.1, 4p15.33, 4q31.21, 17q24.3</i>
Kim (2018)	heel BMD	394,929	899	613	<i>*WNT1, RSP03, ESR, SPTBN1</i>
Morris et al. (2018)	heel BMD	426,824	518	301	<i>*DAAM2</i>
Nerea et al. (2018)	Clinical vertebral fractures	10,683	1	1	<i>2q13</i>
Trajanoska et al. (2018)	Any type of fractures	562,258	15	9**	<i>SPTBN1, CTNBN1, RSP03, ESRI, WNT16, STARD3NL, GRR10, FUBP3, MBL2/DKK1, LRP5, SOST, FAM210A, ETS2</i>

BMD = Bone mineral density, BUA broadband ultrasound attenuation, VOS = velocity of sound, DXA = Dual energy x ray absorptiometry, v = volumetric. * due to the large number of genes the functional validated genes or a smaller curated list is presented. ** Novel in relation with fracture all of the loci are known BMD loci. † closest genes.

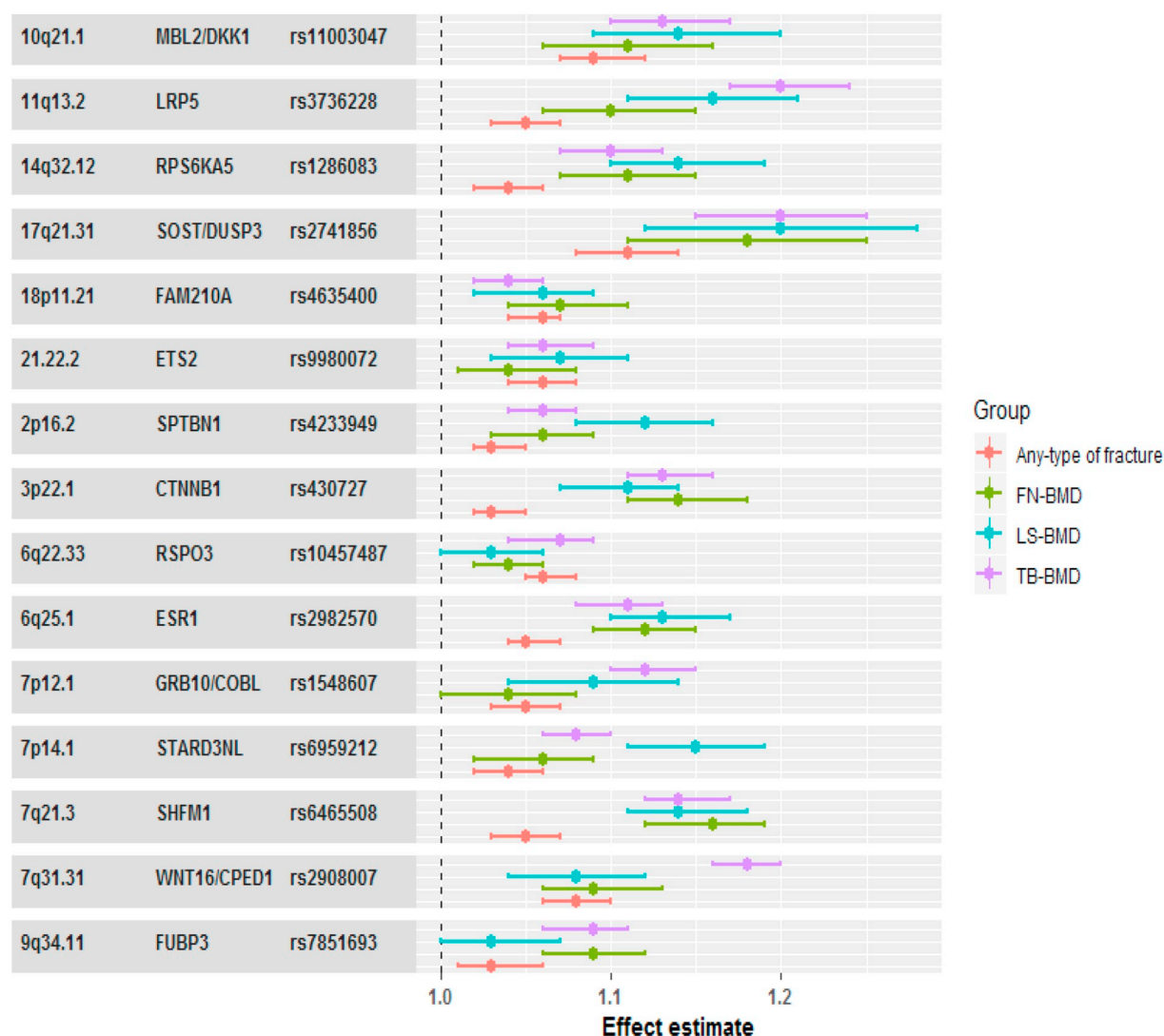


Fig. 2. Phenotype-wide effect for the fracture loci associated with femoral neck, lumbar spine and total body bone mineral density.

7. Variance explained in bone traits by bone-associated variants

Harnessing the information from GWAS can help improve risk prediction of a particular disease; ultimately, this information can then be used for the prevention, diagnosis, prognosis, and treatment of a particular disease [37]. However, clinically-relevant prediction was not achieved by earlier genomic studies in the osteoporosis field; i.e., genetic risk scores did not substantially increase osteoporosis or fracture risk discrimination above the use of traditional clinical risk factors, probably as a result of low variance explained by the genetic variants [20]. The discovery of new loci using more powerful settings have led to substantial leaps in the variance explained (Fig. 3) which will substantially improve risk prediction models in the osteoporosis field. So far, common to less-frequent genetic variants explain around 20% of the BMD variation, which is up to two-fold higher of what can be achieved through the use of other traditional risk factors for osteoporosis such as age or weight (8–9%) [38]. Moreover, the combination of genetic factors, height, weight, age, and sex explains around 25% of the variance in eBMD [77]. This implies that the use of genetic markers has started to materialize in the clinical setting.

8. Biological pathways underlying common bone conditions

Impressively, a large proportion of the genes discovered by GWAS

are located in well-known bone-active pathways. Overall, there are five main pathways crucial for bone metabolism i) Mesenchymal cell differentiation ii) WNT, iii) NOTCH, iv) Hedgehog and v) OPG-RANK-RANKL signalling pathways.

8.1. Mesenchymal cell differentiation pathways

Mesenchymal stem cells (MSc) have the capacity to differentiate into several cell lineages including osteogenic and chondrogenic. Numerous transcription factors (TFs) such as RUNX2, Osterix (Ox), SOX9, can induce the osteogenic differentiation of MSc [39–41]. These factors have been also identified in various BMD GWAS. *RUNX2* is involved in both chondrocyte osteoblast differentiation [42] and is crucial for the early stages of osteoblast development whereas the *Ox* effects are more pronounced in the later stages (i.e. pre-osteoblast differentiation into functional osteoblast) [43].

8.2. WNT signalling

WNT signalling plays a pivotal role in bone development during embryogenesis and bone formation, resorption and accrual during postnatal growth. WNTs are proteins secreted from the cells and regulate the proliferation, differentiation, and apoptosis of bone cells [44]. There are several pathways that can be activated by Wnt proteins

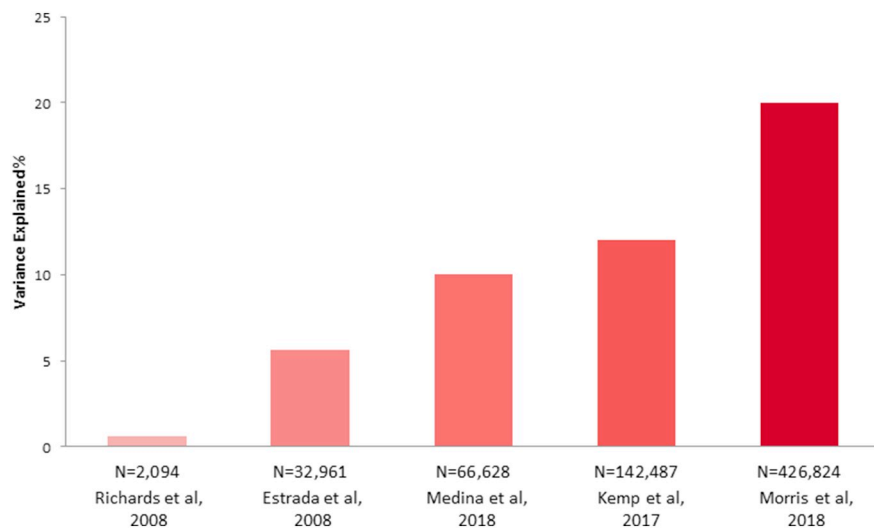


Fig. 3. Variance explained by GWAS on bone traits. With the increase of the sample size the variance explained increased accordingly.

among which the Wnt/ β -catenin (canonical) pathway is the most important one for bone biology. This pathway is activated when WNTs proteins bind to the Frizzled membrane receptors and the low-density lipoprotein (LDL) receptor-related protein 5/6 (LRP5/6). Thereupon the stabilized β -catenin is translocated to the nucleus where it binds to the TFs LEF1/TCF and initiates the transcription of the target genes. Many genes within the WNT pathway have been identified by GWAS: LRP5, WNT16, AXIN1, CTNNB1, DKK1, WLS, LPR4, MEF2C, RSPO3, SERP4, SNT16, SOST, WNT4, WNT5B, and EN1. LRP5 (encoding LDL receptor-related protein 5) is one of the first genes discovered in the osteoporosis field long before the GWAS era [45–47]. LRP5 is a key component of the Wnt signalling since it acts as a co-receptor that binds Wnt proteins with Frizzled-receptors. Functional studies have shown that LRP5 can lead to low bone mass (loss-of-function) [48] and high bone mass (gain-of-function mutation) [49]. LRP5 can be inhibited by several factors such as sclerostin and the dickkopf (DKK) proteins, thus, inhibit the formation of new bone. WNT16 biology has been confirmed by many GWAS studies in relation with different bone parameters such as areal BMD, ultrasound BMD, cortical thickness and fracture, both in adults and children [17,18,20,24,31] and in perimenopausal women [50]. Functional studies initially showed that the Wnt16^{-/-} mouse model has reduced cortical, but not trabecular, bone mass. This effect mediated by WNT signalling, was shown to inhibit human and mouse osteoclastogenesis through direct effects on osteoclast progenitors and indirect effects resulting in increased Opg expression in osteoblasts [51]. Subsequent work showed that overexpression of WNT16 increases mainly trabecular bone mass; and postulated that WNT16-targeted therapies might be useful for the treatment of postmenopausal trabecular bone loss as these effects were independent of oestrogen action [52]. One of the strongest associations implicates MEF2C (Myocyte enhancer factor-2), which regulates a group of transcriptional regulatory proteins relevant for skeletal muscle development. Recent studies have found that a super activating form of MEF2C causes precocious chondrocyte hypertrophy, ossification of growth plates, and dwarfism [53]. Studies have reported that loss-of-function mutations in WNT1 can cause dominantly inherited early-onset osteoporosis [54]. In addition, WNT1 deletion in osteocytes has been related with low bone mass and spontaneous fractures [55]. In addition, the same study has also shown that it increases the number of osteoblasts and regulates their activity. EN1 is another important novel gene involved in murine calvarial osteoblast differentiation and proliferation [56] and large effect on BMD and fracture risk in humans [22].

8.3. NOTCH signalling

In the skeleton, both osteoblasts and osteoclasts require NOTCH signalling for proper differentiation and function, and the specific roles of NOTCH are dependent on the differentiation status of the cell [57]. NOTCH is a family of four transmembrane proteins (NOTCH1–4) that are expressed on the cell surface and require cell-to-cell contact for activation [58] through several ligands (Dll1, Dll3, Dll4, JAG1, JAG2) binding to receptors expressed on the surface of neighboring cells. The ligand binding induces proteolytic cleavage and releases the NOTCH intracellular domain (NICD), which enters the cell nucleus and the transcription of NOTCH targeted genes starts. There is also crosstalk between the NOTCH and WNT signalling pathways which precise mechanisms are still not clear. Several BMD genes have been related to this pathway such as JAG1, MAPT, and NOTCH2.

8.4. Indian Hedgehog (IHH) signalling

The IHH signalling pathway consists of essential signalling molecules crucial for intra-membranous ossification of cranial bones and endochondral ossification in other parts of the skeletal system. During endochondral ossification, chondrocytes differentiate and go through a tightly regulated developmental program of proliferation, hypertrophy, and apoptosis to be eventually replaced by osteoblasts in the ossification centres [59]. IHH signalling may also regulate osteoblast differentiation during endochondral bone development in interaction with WNT/ β -catenin [60]. Within this cross-pathway signalling there are also several GWAS-identified genes such as WNT1, WNT4, WNT5b, WNT16, DHH, and PTCH1. The latter gene encodes the patched –1 receptor (Ptch1) which is essential for many developmental processes such as osteoblastogenesis and chondrocyte differentiation [61]. Patched1 haploinsufficiency (Ptch1 +/–) is characterized by increased adult bone mass in mice; while in culture cells exhibited accelerated osteoblast differentiation [62].

8.5. OPG–RANK–RANKL signalling

The OPG–RANK–RANKL signalling pathway predominantly regulates the coupling between osteoblasts and osteoclast activity [63]. Osteoblasts secrete the receptor activator of NF- κ B ligand (RANKL) which binds to its receptor activator of NF- κ B (RANK) on monocytes resulting in osteoclast differentiation and activation the presence of monocyte colony-stimulating factor (M-CSF). OPG also secreted by the osteoblast is a decoy receptor of RANKL and blocks osteoclast induction

by competing with RANK to bind RANKL. These factors are part of the tumour necrosis factor (TNF) superfamily of ligands and receptors and have been shown to have other functions beyond bone remodelling, including potential roles in other disease processes (i.e., vascular calcification, diabetes and cancer). *TNFRSF11* (RANKL), *TNFRSF11A* (RANK) and *TNFRSF11B* (OPG) were also one of the first genes associated with osteoporosis found and replicated by several studies. Functional studies have shown that in *RANK*^{-/-} mice the generation of osteoclasts from their myeloid progenitors is blocked resulting in absence of bone resorption and severe osteopetrosis [64]. Although this pathway has not been associated with fracture risk it holds an important role in increasing BMD and decreasing fracture risk as due to its antiresorptive effects discussed below.

9. The fulfilled and unfulfilled promises of genomics

The ultimate goal of osteoporosis genomics after GWAS is to perform functional validation, that will allow translating the discoveries into clinical practice. It has been shown that genetic information may significantly improve the search of drug targets [65] and may increase the success rate of preclinical and clinical trials. Nowadays, most of the osteoporosis agents in use (or undergoing trials) target pathways related to the discovered BMD genes. **Denosumab** is a human monoclonal antibody which binds to RANKL and inhibits bone resorption by preventing RANKL from activating RANK, its receptor on the osteoclast surface. Such mimicking of the decoy action of OPG reduces the formation, activation, and survival of osteoclasts [66]; This RANKL inhibitor is approved for use in postmenopausal women with risk of osteoporosis. Since its approval, it has shown sustained efficacy in increasing BMD and decreasing vertebral fracture risk [67]. Anabolic agents hold promising potential, constituting bone-building drugs with proven success, such as **PTH 1–34 (teriparatide)** and **PTHrP 1–36 (abaloparatide)**, which stimulate osteoblasts to make new bone. GWAS have identified genes in the PTH pathway including *PTH1LH* [20]. Teriparatide, is the first anabolic agent approved for the treatment of osteoporosis. In clinical trials to date, abaloparatide has shown promising results in a reduction of new-onset vertebral (approximately 86% reduction) and nonvertebral fractures (approximately 43% reduction) [68]. Finally, several GWAS on BMD and fracture risk have identified variants in several genes (*SOST*-sclerostin, *DKK1*, *LRP5*, *LRP4*, *AXIN1*, and *CTNBN1*- β -catenin) in pathways with anabolic potential [20]. **Sclerostin** (which is produced by osteocytes) inhibits the proliferation, differentiation, and survival of osteoblasts, leading to reduced bone formation. Sclerostin also stimulates (in neighboring osteocytes) the production of RANKL, leading to bone resorption. Anti-sclerostin antibodies, Romosozumab and Blosozumab, have been developed to counterattack the sclerostin effects and have undergone phase II clinical trials [69]. Only Romosozumab was followed to phase III where it showed a 73% reduction in vertebral fracture risk and a 36% reduction in clinical fractures at 1 year (reviewed by Reid et al) [70]. Overall the current agents have been mainly successful in reducing the risk of vertebral fractures (up to 70%), whereas reduction rates for non-vertebral fractures and hip fractures were of smaller magnitude (20% and 40%, respectively) [71]; indicating that trabecular and cortical bone might be regulated by different biological pathways or respond differently to signals involved in the regulation of skeletal homeostasis, as recently shown by the *WNT16* contrasting mechanisms of action [52,72].

9.1. Identification of causal genes for bone traits

It is a common approach to annotate the closest gene to a SNP with the lowest p-value as the most likely causal gene. However, it is important to note that the physical distance of a variant to a gene is not substantive evidence of causality [73]. Multitude of statistical and functional fine-mapping methods have been developed in order to

prioritize causal variants (reviewed by Spain et al) [73]. These methods are essential for highly polygenic traits such as bone density and can drastically improve the mapping of associated loci to their causal genes. For example, in the latest and largest GWAS to date in the field of osteoporosis, Morris et al. [25] have used UK Biobank data to build a robust analysis pipeline to prioritize eBMD-associated SNPs by combining several distinctive approaches. Firstly, two statistical fine-mapping methods were used to refine associated SNPs at each locus. These methods identify SNPs based on their conditional independence (GCTA-COJO) and posterior probability (log10 Bayes factor > 3) for causality (FINEMAP); yielding two conditionally independent SNPs and five SNPs with a log10 Bayes factor > 3 per locus. Second, all fine-mapped SNPs were then tested for enrichment for missense protein coding SNPs, DHSs (DNase I hypersensitive sites) from primary osteoblasts, and ATAC-seq (Assay for Transposase-Accessible Chromatin using sequencing) peaks from SaOS-2 osteosarcoma cells. Notably, the fine-mapped SNPs showed strong enrichment for both missense variants in protein-coding regions and osteoblast open chromatin sites. Finally, a list of positive control genes identified through pharmacotherapy or Mendelian disease which are critical for bone biology were curated. Six distinctive genomic features that link a SNP to a gene such as cell-specific 3-dimensional (3D) contact domains, cell-specific open chromatin states, physical proximity, and the presence of associated coding variation were used to construct *Target Gene* sets which were tested for enrichment for positive control bone genes. All six methods for linking fine-mapped SNPs to Target Genes yielded strong enrichment for positive control genes known to be central to bone biology [25]. Notably, *Target Genes* closest to fine-mapped SNPs in osteoblast-derived ATAC-seq peaks were seen to be ~60-fold enriched for positive control genes. This is quite an insightful finding, as transcription factor binding occurs mainly in regions of open chromatin (ATAC-seq peaks), thus, this implies that the mapped *Target Genes* may have substantial impact on bone biology.

10. Post-GWAS analyses and concluding remarks

In the fast-moving world of genomics, including the field of osteoporosis, hundreds of genetic markers have been identified as associated with complex traits. As discussed above the GWAS findings have aided the discovery of several novel osteoporosis drug targets. However, there is still an overwhelming amount of significant associations, for which the underlying biological mechanisms remain unknown; as the functional characterization of the discovered genetic variants have lagged far behind [74]. In order to understand the functional consequences of these loci future post-GWAS methods should be focused on bridging the gap between disease-associated loci and underlying disease biology [74]. An essential step in the translation studies, from gene discovery to biological mechanisms, is the identification of causal variants and genes. It is well known that the most strongly associated variants with a specific trait or diseases are likely to be in linkage disequilibrium (LD) with the causal variant, rather than have a biological function themselves [73]. In the last few years large numbers of studies have been devoted to pinpoint causal variants using both statistical evidence (e.g. reference panels, targeted resequencing, Bayesian methods, machine learning methods) from large association data sets and functional annotations of genetic variants (e.g. enrichment analysis, pathway prioritization, DNase sensitivity) [73,75]. Moreover, the emergence of expression quantitative trait loci (eQTLs), which characterize associations between genetic variants and gene expression at the cellular level, have provided a better biological context in disease studies [76]. Although a widely used approach, eQTL data from primary bone cells is limited. Establishing bone specific eQTLs may significantly improve the search of causal variants and provide valuable target genes. Finally, the identification of causal variants can further facilitate the interpretation of the GWAS findings and opens opportunities for more detailed downstream functional investigations such as in human and/or animal

cell and tissues models.

GWAS have provided us with a comprehensive understanding of the genetic architecture of osteoporosis and fracture risk. Moreover, key bone genes and pathways have been identified which have promoted novel drug targets and treatments. In the decade to come, the advances of GWAS and post-GWAS techniques and methods will enable fruitful incorporation of genetics in clinical practice that will ensure better disease prediction and risk stratification, leading to overall improvement in disease prevention or intervention.

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Resources

Fig. 2 was modified from Estrada et al.

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