Integrative analysis of genomic and epigenomic data reveal underlying superenhancer-mediated microRNA regulatory network for human bone mineral density

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Abstract
Bone mineral density (BMD) is a highly heritable complex trait and is a key indicator for diagnosis and treatment for osteoporosis. In the last decade, numerous susceptibility loci for BMD and fracture have been identified by genome-wide association studies (GWAS); however, fine mapping of these loci is challengeable. Here, we proposed a new long-range fine-mapping approach that combined superenhancers (SEs) and microRNAs (miRNAs) data, which were two important factors in control of cell identity and specific differentiation, with the GWAS summary datasets in cell-type-restricted way. Genome-wide SE-based analysis found that the BMD-related variants were significantly enriched in the osteoblast SE regions, indicative of potential long-range effects of such SNPs. With the SNP-mapped SEs (mSEs), 13 accessible long-range mSE-interacted miRNAs (mSE-miRNAs) were identified by integrating osteoblast Hi-C and ATAC-seq data, including three known bone-related miRNAs (miR-132-3p, miR-212-3p and miR-125b-5p). The putative targets of the two newly identified mSE-miRNAs (miR-548aj-3p and miR-190a-3p) were found largely enriched in osteogenic-related pathway and processes, suggesting that these mSE-miRNAs could be functional in the regulation of osteoblast differentiation. Furthermore, we identified 54 genes with the long-range ‘mSE-miRNA’ approach, and 24 of them were previously reported to be related to skeletal development. Besides, enrichment analysis found that these genes were specifically enriched in the post-transcriptional regulation and bone formation processes. This study provided a new insight into the approach of fine-mapping of GWAS loci. A tool was provided for the genome-wide SE-based analysis and the detection of long-range osteoblast-restricted mSE-miRNAs (https://github.com/Zheng-Lab-Westlake/Osteo-Fine-Mapp-SNP2SE2miRNA).

Introduction
Osteoporosis is a common skeletal disease characterized by reduced bone mass and deterioration in bone microarchitecture, leading to an increased propensity to fragility fracture (1). Bone mineral density (BMD), which is generally evaluated by dual energy X-ray absorptiometry scan (DXA), is a key indicator for diagnosis and treatment for osteoporosis (2). Bone mass could also be estimated by ultrasound approach by considering both the speed of sound and bone ultrasound attenuation. In the last
decade, genome-wide association studies (GWAS) have identified many susceptible loci for BMD at several anatomical sites such as lumbar spine (LS), femoral neck (FN), forearm (FA) and total body (3–6). In 2018, a GWAS with 426,824 individuals from UK biobank dataset identified 518 genome-wide significant loci associated with heel estimated BMD (eBMD), including 301 novel loci (7). However, it is challengeable to pinpoint susceptible genes from an associated locus, some studies have successfully identified the determinant genes of BMD and fracture, such as EN1 and DAAM2 (5,7). The commonly used approach for susceptible gene selection is to identify the lead SNP-overlapped protein coding gene directly or the nearest gene (8). However, the vast majority of associated variants are non-coding, residing at intronic and intergenic regions in genome. Besides that, some variants appear to be in one gene but influencing the expression of others due to the spatial state of chromatin (9). One example is that obesity-associated variants within FTO gene form long-range functional connections with IRX3 (10). Variants reside in some functionally important region, such as enhancers, may affect distant genes by long-range interactions, cell-type-specific Hi-C and ATAC-seq, which could show spatial interactions and the accessibility of chromatin (11,12), might detect these long-range interactions.

Superenhancers (SEs) are large clusters of regulatory regions that consist of multiple enhancers occupied by high density of mediator complexes that drive expression of cell-identity genes (13–15), previous studies found that enhancers within SEs possess stronger activating features than normal enhancers (16–18). SEs are generally enriched for disease-associated variants, especially for the SEs defined in the disease-relevant cell type or tissue (15,19). For example, SEs identified in human pancreatic islet cells are enriched for SNPs associated with type 2 diabetes (20), and Alzheimer’s disease-associated SNPs are found to reside in brain tissue SEs (15).

MicroRNAs (miRNAs) are a class of short non-coding RNAs that block protein translation or modulate mRNA stability by specifically binding the complementary sequences in the 3’ untranslated region (UTR) of mRNAs (21). Many miRNAs are found involved in multiple developmental osteogenic signaling pathways and osteoblast growth and differentiation pathways (22–25), such as miR-125b could downregulate cell proliferation and inhibit osteoblastic differentiation (26), and miR-212 and miR-384 could promote osteogenic differentiation via targeting RUNX2 (27). MiRNAs are transcribed by RNA polymerase (Pol) II/III into primary miRNA (pri-miRNA), which are processed by Drosha/Dgcr8 into precursor miRNA (pre-miRNA) (28).

Interestingly, the tissue-specific miRNA expression atlas was found largely shaped by SEs and H3K4me3 domains, SEs could facilitate transcription, Drosha/Dgcr8 recruitment and pri-miRNA processing to boost cell-type-specific miRNA production (14). In addition, previous studies have found that SEs and miRNAs both play a key role in the control of cell-type-specific transcriptomes (14,29), in view of this, SE-interacted miRNAs (SE-miRNAs) may have some specific properties in the maintenance of cell identity and differentiation. We therefore applied a cell-type-restricted approach that combined SEs and miRNAs for the fine-mapping of GWAS SNPs. In this study, we mapped SEs with BMD-associated SNPs from 10 large-scale human GWAS datasets and identified long-range osteoblast SNP-mapped SE-miRNAs and genes for human BMD and fracture by integrating multiple genomic and epigenomic datasets. Our approach could also be generalized to the other cell-type-specific traits or diseases.

Results

BMD-associated SNP landscape in human osteoblast SEs and genome-wide SE-based analysis

The overall study design was shown in Figure 1. We employed histone modification marker of H3K27ac to shape the enhancer regions in genome in human osteoblast. A total of 72,936 and 116,190 peaks were initially called from two replicates H3K27ac ChIP-seq data, and a total of 1224 and 1149 SEs were detected from the peaks (Supplementary Material, Fig. S1, Supplementary Material, Tables S1 and S2). We next compared them with the human osteoblast SEs from SEA and dbSUPER database, the results showed a high degree of consistency between these SEs (Supplementary Material, Fig. S2). Besides, we also compared our SEs with the human fetal osteoblasts (undifferentiated and differentiated) SEs from SEdb, there were few differences were found, this could be caused by the different osteoblastic cell type used (Supplementary Material, Fig. S2). We extracted SNPs that resided in SEs from 10 large-scale BMD-related GWAS datasets, the landscapes of such SNPs in the genome were shown in Figure 2 and Supplementary Material, Figure S3. We found that many genome-wide significant BMD-associated SNPs were enriched in osteoblast SEs, particularly for heel eBMD and total body DXA BMD (Fig. 2). This result suggested a potential long-range effect of these SNPs via SEs. We then pruned out all non-independent SNPs in the 10 GWAS datasets. The SE-based mapping analysis for two replicates then respectively yielded 202 and 291 SNP-mapped SEs (mSEs) that passed Bonferroni multiple-test, these osteoblast mSEs could be functional important for BMD (Supplementary Material, Tables S3 and S4).

Identification of Long-range mSE-miRNA in human osteoblast

We first defined the promoter bait regions of 1914 pri-miRNAs (Supplementary Material, Table S5). After applying the promoter-focused Hi-C data and overlapping analysis, we found there were 74 pri-miRNAs that had DNA-looping interaction with mSEs (Supplementary Material, Table S6), most of them had at least two Hi-C interactions and a half of mSEs were mapped by 50–90 independent SNPs (Supplementary Material, Figure S4). The SNP signals in these mSEs were mainly from heel ultrasound eBMD GWAS, a few of signals were from total body and other anatomical sites DXA BMD GWAS. Among 74 mSE-miRNAs, 12 of them were previously found to be associated with osteogenic differentiation, and two pri-miRNAs were associated with muscle proliferation (Supplementary Material, Table S7).

We further investigated the chromatin accessibility of the 74 mSE-miRNAs by ATAC-seq data in primary human mesenchymal stem cells (MSCs)-derived osteoblasts (9). Thirteen pri-miRNAs were finally identified that had the open chromatin conservative region in the corresponding promoter or transcriptional start sites (TSS) region, Figure 3 demonstrated the detailed mapping and long-range interactions results. The distributions of H3K27ac peaks on the genome in two biological replicates showed the similar pattern, and the number of long-range interactions and distance between mSE and miRNAs were varied (Fig. 3). Among all mSE-miRNAs, mir-125b-1 (Fig. 3A), mir-548sj-1 (Fig. 3D), mir-614 (Fig. 3F) and mir-5702 (Fig. 3J) were found to have the most DNA-looping interactions with mSE, which respectively reached 0.11, 1.01, 0.56 and 0.3 per kb in mSE, reflecting the high spatial contact density in genome (Table 1), the mir-614 (Fig. 3F) and mir-3650 (Fig. 3H) showed the furthest
distance (>250 kb) with the corresponding mSE, whereas mir-6870 (Fig. 3 L) was found overlapped within mSE, the mir-132 (Fig. 3B), mir-212 (Fig. 3B) and mir-614 (Fig. 3F) were found interacted by two remote mSEs simultaneously. Besides, there were two pri-miRNAs, mir-125b-1 (Fig. 3A, mSE P-value: 1.9e−11, best SNP P-value: 3.6e−8) and mir-583 (Fig. 3E, mSE P-value: 1.7e−17, best SNP P-value: 8.6e−14), that were found interacted with the mSEs mapped by intergenic SNPs, indicative of the functional importance of GWAS signals in such regions (Table 1). These accessible mSE-miRNAs could be the vehicles for SNP loci to regulate the expression of genes in osteoblast.

**Putative targets of 13 accessible mSE-miRNAs and biological process enrichment**

We predicted the target genes of the 13 osteoblast-restricted mSE-miRNAs by a two-step approach. TargetScan (30) and
Table 1. Accessible long-range mSE-miRNAs and implicated hub-genes in human osteoblast mSE. GWAS SNP-mapped superenhancer. mSE-miRNA: mSE-interacted miRNA. Interaction density was the number of Hi-C interactions per Kb in mSE. Accessibility of miRNA was evaluated by open chromatin regions. There were a total of 54 unique hub-genes detected from protein–protein interaction (PPI) network, one hub-gene could be targeted by multiple miRNAs. The hub-gene marked by asterisk meant the gene has been reported by previous studies, the detailed information of relative studies was listed in Supplementary Material, Table S11. Note that for a SE mapped by multiple GWAS sets, only the best P-value was showed.

| GWAS Locus | SNP-mapped SE | Distant miRNA | P-value of mSE | Interactions density | Implicated hub-gene | GWAS Set |
|------------|---------------|---------------|---------------|---------------------|---------------------|-----------|
| JAG1, SLX4IP | chr20: 10536348, 10656253 | mir-6870 | 2.46E-244 | 0.03 | DCAF7, ZFP36, AGO1, PIK3R2, DDX6, RBFOX2, ATXN1L, PAK2 | Heel eBMD, LS BMD, TB BMD |
| SMG6 | chr17: 1962802, 2042483 | mir-212 | 6.73E-219 | 0.01 | DCAF7, YWHAG, MEF2A*, FOXO3*, AGO1, SOS1, PIK3R1, LYN*, PIK3CA*, FRS2, EIF4A2, UHMK1*, SMAD2*, ACVR2B*, SMAD5*, DAZAP2, ATXN1, DCUN1D1 | Heel eBMD |
| SMG6, LOC101927839 | chr17: 2115397, 2168091 | mir-212 | 1.77E-70 | 0.09 | – | Heel eBMD |
| EMP1 | chr12: 13324312, 13377792 | mir-614 | 2.33E-36 | 0.56 | DCAF7, YWHAG, MEF2A*, FOXO3*, AGO1, SOS1, PIK3R1, LYN*, PIK3CA*, FRS2, EIF4A2, UHMK1*, SMAD2*, ACVR2B*, SMAD5*, DAZAP2, ATXN1, DCUN1D1 | Heel eBMD |
| HIEBP1, LOC100506314 | chr12: 13128286, 13146200 | mir-614 | 2.33E-36 | 0.56 | – | Heel eBMD |
| SMG6, LOC101927839 | chr17: 2115397, 2168091 | mir-132 | 1.77E-70 | 0.02 | – | Heel eBMD |
| Intergenic | chr5: 95530421, 95595730 | mir-583 | 1.66E-17 | 0.11 | DYRK2*, YWHAG, MEF2A*, FOXO3*, AGO1, SOS1, PIK3R1, LYN*, PIK3CA*, FRS2, EIF4A2, UHMK1*, SMAD2*, ACVR2B*, SMAD5*, DAZAP2, ATXN1, DCUN1D1 | Heel eBMD |
| SERTAD2 | chr2: 64864825, 64889632 | mir-4434 | 1.19E-13 | 0.04 | FOXO3*, CBL*, PIK3R1, LYN*, PIK3CB, PRKAR1A*, CCNC, DDX6, RUNX2*, RBFOX2, ATXN1L, YWHAE, PTPN14, PAR2, SPRY2*, NECAPI | Heel eBMD |
| Intergenic | chr11: 121903479, 121921160 | mir-125b-1 | 1.95E-11 | 0.11 | – | Heel eBMD |
| KDM2A, ANKRDI3D, ADRB1K1 | chr11: 67006556, 67060753 | mir-6860 | 6.17E-08 | 0.02 | DYRK2*, YWHAG, MEF2A*, FOXO3*, AGO1, SOS1, PIK3R1, LYN*, PIK3CA*, FRS2, EIF4A2, UHMK1*, SMAD2*, ACVR2B*, SMAD5*, DAZAP2, ATXN1, DCUN1D1 | Heel eBMD |
| BCAR3 | chr1: 94018520, 94142585 | mir-760 | 2.45E-06 | 0.02 | CBL*, NECAPI | Heel eBMD |
| LACTB, TPM1, RPS27L | chr15: 63305237, 63446803 | mir-190a | 4.06E-06 | 0.03 | – | Heel eBMD |
| OSNRI, OSNRI-AS1, IRS1 | chr9: 38825860, 38877448 | mir-3650 | 2.81E-05 | 0.02 | AGO3, CCNC, SMAD1*, NCK2, PRKAA2, WASL | Heel eBMD |
| LINC01013 | chr6: 132451318, 132476057 | mir-548s-1 | 2.37E-05 | 1.01 | DCAF7, ZFP36, GSK3B*, MYCN, MEF2A*, ERBIN, AGO3, GAB1*, PIK3R1, PIK3CA*, PIK3CB, FGRF2*, PRKAR1A*, CCNC, DDX6, TGFBR1*, UHMK1*, SMAD2*, RUNX2*, SMAD6*, ACVR2B*, SMAD5*, BMP2*, YAP1*, DAZAP2, ATXN1, DCUN1D1, MOB1B, NCK2, PRKAA2, WNT5A*, WASL, NECAPI | Heel eBMD |
| OSMR, OSMR-AS1 | chr5: 38825860, 38877448 | mir-5702 | 3.69E-05 | 0.30 | DCAF7, ZFP36, GSK3B*, MYCN, MEF2A*, ERBIN, AGO3, GAB1*, PIK3R1, PIK3CA*, PIK3CB, FGRF2*, PRKAR1A*, CCNC, DDX6, TGFBR1*, UHMK1*, SMAD2*, RUNX2*, SMAD6*, ACVR2B*, SMAD5*, BMP2*, YAP1*, DAZAP2, ATXN1, DCUN1D1, MOB1B, PTPN14, PAR2, SPRY2*, WNT5A*, WASL, NECAPI | Heel eBMD |
miRDB (31) database with their respective prediction confidence score were jointly used to form the initial pool of putative targets, and the osteoblast Ago2 CLIP-seq data was then used to filter out the false positive targets. The results showed that the number of putative targets for each miRNA was varied, miR-614 had only 21 putative targets whereas miR-548aj-3p and miR-190a-3p had 503 and 527, respectively, most of the rest of miRNAs had 100–200 putative targets (Fig. 4A). Besides, we found that the targets of miRNAs were largely shared with each other, indicating the functional similarity between them in human osteoblast (Fig. 4A). Furthermore, our results showed that miR-132-3p and miR-212-3p, which belongs to the same cluster, had almost the same putative targets. The whole lists of putative target genes of mature miRNAs were detailed in Supplementary Material, Table S8.

We performed the functional pathway and process enrichment analysis for the putative targets of each miSE-miRNA. The results of top 20 enriched Gene Ontology (GO) biological processes showed that the putative targets of miR-548aj-3p, miR-190a-3p, miR-132-3p, miR-212-3p, miR-125b-5p and miR-3650 were significantly enriched in the muscle structure development (GO Term ID: 0061061), Wnt signaling pathway (GO Term ID: 0016055), skeletal system development (GO Term ID: 0015051) and mesenchyme development (GO Term ID: 0060485), implying that these miRNAs were likely to be involved in biological processes of bone formation in human osteoblast (Fig. 4B and Supplementary Material, Table S9). Besides, we found putative targets of these miRNAs were also significantly enriched in some growth and cell differentiation pathways, such as response to growth factor (GO Term ID: 0070848) and tissue morphogenesis (GO Term ID: 0048729) (Fig. 4B), indicative of the potential effects of these miRNAs in growth and differentiation of osteoblast.

To further reveal the functional enrichment of 13 osteoblast-restricted accessible miSE-miRNAs in bone-related biological process, we extracted enrichment results of 18 GO terms (Fig. 4C) based on several key words, including ‘osteoblast’, ‘osteoclast’, ‘skeletal system’, ‘skeletal muscle’, ‘bone’, ‘ossification’, ‘BMP’ and ‘wnt signaling pathway’ from all enriched GO biological process results (Supplementary Material, Table S10). Our results showed that the skeletal system development (GO Term ID:
Figure 3. The regional association plots for 13 accessible mSE-miRNAs. mSE: GWAS SNP-mapped superenhancer. mSE-miRNA: mSE-interacted miRNA. The peak tracks in blue showed H3K27ac modifications in two biological replicates in osteoblast and green track showed the open chromatin regions that detected by ATAC-seq. The miRNA was considered to be accessible when the open chromatin regions found in its promoter or transcriptional start sites. The yellow bar represented SEs that were detected by ROSE and pooled in two H3K27ac datasets, and red curves represented DNA-looping interactions. Note that the genome scale for each panel was different and only most significant mapping results were plotted across all GWAS datasets. The regional associations were drawn by LocusZoom, the peak tracks and Hi-C interactions were drawn by WashU Epigenome Browser.

0001501) was enriched by the majority of the miRNAs with a high target gene ratio (Fig. 4C). Besides, the putative targets of miR-548aj-3p and miR-190a-3p were found respectively enriched in 16 and 14 out of a total of 18 bone-related processes (Fig. 4C). Two previously reported osteogenic differentiation regulator miRNAs, miR-212-3p (27) and miR-132-3p (32), were also found largely enriched in such processes (Fig. 4C). These results suggested that the osteoblast-restricted mSE-miRNAs could potentially maintain the cell identify and regulate osteogenic differentiation. However, no osteogenic process was found enriched for miR-614. Moreover, we found that the putative target genes of miR-125b-5p were largely enriched in multiple miRNA production and metabolic pathways, such as the production of miRNAs involved in gene silencing by miRNA (GO Term ID: 2000637) (Fig. 4D), indicating that miR-125b-5p may act as a master miRNA to dominate the miRNA pool via SEs in osteoblast (14).

Protein–protein interaction network and hub-gene group

We next performed protein–protein interaction (PPI) network analysis for the 1279 unique putative targets of 13 osteoblast-restricted mSE-miRNAs (Supplementary Material, Table S8). A total of 54 hub-genes were finally identified as densely connected network components and constituted seven hub-gene groups (Molecular Complex Detection (MCODE1–7)) (Fig. 5A), and each hub-gene could be targeted by multiple mSE-miRNAs, indicative of a complementary regulation patterns in
osteoblast (Fig. 5B). We found that there was no overlap between the genes directly resided in or near GWAS loci and the hub-genes identified in the long-range fine-mapping way by the corresponding GWAS loci (Table 1). Besides, the genes fine-mapped in this way were more than twice the corresponding GWAS loci genes (Table 1). Moreover, we investigated all 54 hub-genes identified in the long-range ‘mSE-miRNA’ fine-mapping approach with previous functional experiments studies, and a total of 24 of them have been found to be directly related to the bone mass and skeletal development (Supplementary Material, Table S11), whereas rest 30 have not been reported before.

We further conducted the biological pathway and process enrichment analysis for the hub-genes. Our results showed that MCODE1 group, which contained 12 hub-genes was mainly functional in mRNA cleavage involved in gene silencing processes (Table 2), suggesting an important function in the post-transcriptional regulation network; MCODE2 group genes were significantly enriched in some kinase signaling pathways, including phosphatidylinositol 3-kinase signaling (PI3K) which was implicated in osteoblast proliferation and differentiation; MCODE3 group genes were involved in cell proliferation and tissue growth of cardiac and striated muscle; besides, MCODE4 group genes we identified, including SMAD2, RUNX2, SMAD6, SMURF1, ACVR2B, SMAD5, BMPR2 and YAP1, were significantly enriched in bone morphogenetic proteins (BMP) signaling pathway (Table 2). These results suggested that these hub-genes, which were predicted to be targeted by osteoblast-restricted mSE-miRNAs, could play a critical role in the regulation network in bone formation. The complete pathway and process enrichment results were detailed in Supplementary Material, Table S12.

**Discussion**

In this study, we aimed to uncover the miRNA regulatory network that was long-range mediated by SEs which were mapped by BMD-associated SNPs in human osteoblast. We presented the landscape of BMD-related SNPs in human osteoblast SEs, and
identified 13 mSE-miRNAs and 54 genes that may maintain the osteoblast differentiation. We provided a new insight into the fine-mapping approach of GWAS loci and a tool for the genome-wide SE-based analysis and the detection of long-range osteoblast-restricted mSE-miRNAs. This approach could also be generalized to the other cell-type-specific traits or diseases.

We collected BMD-associated SNPs from large-scale human heel eBMD, DXA BMD and fracture GWAS datasets. Rather than the gene-overlapped/nearest SNPs, we focused on the long-range effective SNPs which are usually neglected in disease-associated gene selection, we assumed that a plenty of SNPs in genome are effective via affecting the SE they reside in. We plotted the SNP landscapes for 10 traits of GWAS datasets in osteoblast SEs. The genome-wide significant BMD SNP signals ($P$-value < 5e$^{-8}$) residing in SEs were mostly from heel eBMD, followed by total body DXA BMD, while only few were from other DXA BMD, such as FN and LS (Fig. 2 and Supplementary Material, Fig. S3). Despite the BMD measurements, this disequilibrium could be mainly caused by two reasons: (i) sample size of GWAS. Heel eBMD and total body BMD GWAS was conducted in 426,824 samples and 66,628 samples, respectively. However, the samples in other datasets were <30,000, low sample size...


Table 2. Seven hub-gene groups in protein-protein interaction network and the Top 3 enriched GO biological processes. For each hub-gene group, three best-enriched GO biological process terms by P-value were shown in the table. The enrichment was performed by MetaScape, the complete pathway and process enrichment results for hub-gene groups were detailed in Supplementary Material, Table S11

| Hub-gene group | GO terms ID | Description | P-value (log10) | MCODE Genes |
|----------------|-------------|-------------|----------------|-------------|
| MCODE1         | GO:0090625  | mRNA cleavage involved in gene silencing by siRNA | -9.4          | DCAF7, DYRK2, YWHAH, ZFP36, GSK3B, MYCN, MEF2A, FOXO3, ERB1N, AGO1, AGO2, AGO3 |
|                 | GO:0098795  | mRNA cleavage involved in gene silencing | -9             |             |
|                 | GO:0035279  | mRNA cleavage involved in gene silencing by miRNA | -9             |             |
| MCODE2         | GO:0007169  | Transmembrane receptor protein tyrosine kinase signaling pathway | -15.2         | SOS1, CBL, GAB1, PIK3R1, PIK3R2, LYN, PIK3CA, PIK3CB, FGFR2, FRS2 |
|                 | GO:0014065  | Phosphatidylinositol 3-kinase (PI3K) signaling | -13.4         |             |
|                 | GO:0051897  | Positive regulation of protein kinase B signaling | -13           |             |
| MCODE3         | GO:0060038  | Cardiac muscle cell proliferation | -5.8          | EIF4A2, PRKAR1A, CCNC, CNOT7, DDX6, SMAD1, TGFBR1, ZFYVE16, UHMK1 |
|                 | GO:0014855  | Striated muscle cell proliferation | -5.5          |             |
|                 | GO:0055017  | Cardiac muscle tissue growth | -5.2          |             |
|                 | GO:003509   | BMP signaling pathway | -14.5         | SMAD2, RUNX2, SMAD6, SMURF1, ACVR2B, SMAD5, BMP2R, YAP1 |
| MCODE4         | GO:0071773  | Cellular response to BMP stimulus | -14.2         | RBFOX2, DAZAP2, ATXN1, ATXN1L, YWHAE, DCUN1D1, MOB1B, PTPN14 |
|                 | GO:0071772  | Response to BMP | -14.2         |             |
| MCODE5         | GO:0051168  | Nuclear export | -4.5           |             |
|                 | GO:0007420  | Brain development | -4.3          |             |
|                 | GO:0060322  | Head development | -4.2          |             |
| MCODE6         | GO:0018105  | Peptidyl-serine phosphorylation | -7.6          | PAK2, NCK2, PRKAA2, SPRY2 |
|                 | GO:0018209  | Peptidyl-serine modification | -7.5          |             |
|                 | GO:0031400  | Negative regulation of protein modification process | -6.4          | WNT5A, WASL, NECAP1 |

could lead to a low statistical power (33). (ii) Weight-bearing condition of different anatomical sites. The eBMD was measured at heel, which is the usually weight-bearing site of human body. Mechanical loadings and some mechanically sensitive protein, such as Piezo1, could significantly influent BMD (34); hence some SNPs for heel may be more sensitive than for other less weight-bearing anatomical sites (35).

SEs is closely related to cell-identity and fate-determined processes (15). To avoid potential biases from cell type, all data we used in this study were specific to human osteoblast, including H3K27ac ChIP-seq for SE detection, Hi-C interactions, ATAC-seq peaks for chromatin accessibility checking and CLIP-seq data for false positive target genes filtering. A total of 74 miRNAs were identified to be interacted by SNP-mapped SEs (Supplementary Material, Table S6), 12 of them have been previously found to be associated with osteogenic-related pathway, such as miR-100-5p and miR-143-3p could increase osteogenic differentiation potential by modulation of mTOR signaling (36), miR-181a could promotes osteoblastic differentiation through repression of TGF-β signaling molecules (37), and miR-217 in MSCs could promote cell proliferation and osteogenic differentiation by targeting DKK1 (38) (Supplementary Material, Table S7). For the 13 accessible mSE-miRNAs, previous studies found that miR-125b (lead SNP in SE: rs725670 [G > A], BETA = 0.01 per G allele, P-value = 5.90E-08) could inhibit osteogenic differentiation whereas the interference of miR-212 (lead SNP in SE: rs11657636 [C > T], BETA = -0.05 per C allele, P-value = 4.60E-116) could promote it (26,27). In our study, the putative targets of miR-212 was largely enrich in osteogenic-related pathway and processes (Fig. 4C). In addition, we noticed that no accessible mSE-miRNA except miR-4434 was enriched in osteoclast differentiation process (Fig. 4D). This could be caused by the cell type specificity since our data were all specific to osteoblast. So far, no study reported the function of miR-4434 in bone-related tissue. Although we identified 13 mSE-miRNAs that could be potentially affected by the BMD-associated SNPs, there was a limitation in our study, we did not verify the association between such SNPs and the expression level of mSE-miRNAs. Actually, the expression quantitative trait loci (eQTL) analysis could be an approach to provide some clues. We retrieved the public eQTL databases, such as GTEx (https://www.gtexnodeal.org/home/), however, no bone-related tissues or osteoblastic cell-types were found.

Among the 54 hub-genes we identified, some of them were classic and well-studied osteogenic-related gene, such as RUNX2 (Runx family transcription factor 2), SMAD1/5 (Smad family...
member 1 and 5), BMFR2 (bone morphogenetic protein receptor type 2), TGFBR1 (transforming growth factor beta receptor 1) and WNT5A (Wnt family member 5A) (39–44). In addition, the expression of some hub-genes, such as YAP1, ZFP36 and UHMK1, were found to be up-regulated in high BMD samples (45–47). Out of a total of 54 hub-genes, 24 have been studied and found to be involved in bone formation (Supplementary Material, Table S11).

In summary, we mapped genome-wide osteoblast SEs with BMD-associated SNPs from large-scale human GWAS summary datasets. Based on that, we revealed the mSE-miRNAs regulatory network and identified 54 key genes by integrating the Hi-C, ATAC-seq and Ago2 CLIP-seq data. Our study presented the BMD-related SNP loci landscape in SEs, and provided a new approach and insight for the fine-mapping of GWAS loci. We also provided a tool for the genome-wide SE-based analysis and the detection of long-range osteoblast-restricted mSE-miRNAs (https://github.com/Zheng-Lab-Westlake/Osteo-Fine-Mapp-SNP2SE2miRNA).

Materials and Methods

GWAS summary-statistic data

The overall study design was shown in Figure 1. We included 10 GWAS summary-statistic datasets of different BMD-related traits from Genetic Factors for Osteoporosis (GEFOS) Consortium, which is a large international collaborative effort we were involved in. The fracture summary statistics data were obtained from a study with ∼2.5 million SNPs in 20,439 fracture cases and 78,843 controls (1), and from UK biobank with ∼14 million SNPs in 53,184 fracture cases and 373,611 controls (7). The eBMD GWAS dataset was from UK biobank which contained ∼13.8 million SNPs in 426,824 samples (7). The DXA BMD at distal radius/FA, FN and LS were from UK10K project which included ∼10 million SNPs in 8143 samples, 29,188 samples and 25,225 samples, respectively (5). In addition, we included FN and LS DXA BMD GWAS data from another study which included ∼2.5 million SNPs in 32,961 and 31,800 samples, respectively (3). We also included GWAS summary-statistic data of total body DXA BMD (∼18.3 million SNPs in 66,628 samples) (6) and total body less head DXA BMD (∼2.3 million SNPs in 10,414 samples) (48). All datasets which were not based on hg19 were lifted over to hg19 since the miRBase v22 pri-miRNAs-self (Supplementary Material, Table S5).

Detection of SE in human osteoblast

SEs are usually modified by H3K27ac marks (18), hence we used the H3K27ac ChIP-seq data to detect SEs. The BAM data of H3K27ac ChIP-seq (ENCODE accession: ENCSR000APH) with two biological replications in human primary osteoblast and the control data (ENCODE accession: ENCSR000APU) were downloaded from the ENCODE Project (49). We performed quality control for the alignment datasets by SAMtools (50), PCR duplicate reads and any reads with Q-value < 20 were removed. The MACS2 was employed for peak calling with default settings (51), resulting 72,936 and 78,936 narrow peaks for two replications, respectively. Next, we employed the ROSE algorithm to identify the SE regions (17), in brief, H3K27ac peaks within 2.5 kb of TSS on genome were subtracted and the enhancers within a distance of 12.5 kb were stitched according to H3K27ac peaks occupancy rates (52). Finally, we plotted the ranked stitched enhancers and individual enhancers by H3K27ac enrichment, and using a line with a slope of one tangent to the curve as a cutoff to distinguish SEs above the point and typical enhancers below the point of tangency.

We downloaded the human osteoblast SEs from SEA and dbSUPER database (53,54), and the human fetal osteoblasts (hFOB, undifferentiated and differentiated) SEs from SEDb (55). The Wash U Epigenome Browser (http://epigenomewidget.wustl.edu/browser/) was employed (56) to compare the SEs from the databases with the SEs identified in our study by visualizing the results.

Genome-wide SE-based analysis in human osteoblast

For the SEs we detected in human osteoblast, we first mapped them with SNPs (P < 0.05) in 10 BMD-related GWAS datasets to investigate the landscape of SNP resided in SEs. And then we performed the SE-based mapping analysis by the independent SNPs. We extracted the EUR group genotypes (European ancestry) in 1000 Genome Project Phase 3 (1000GP, v5a) (57) as the SNPs reference and calculated the r² by PLINK v1.9 with setting window size at 50 SNP counts, step size at five counts (58). The non-independent SNPs with r² > 0.1 were then pruned out. We next mapped SE by the GLOSSI methods (59), the statistic for each SE was defined as below:

\[ S_k = -2 \sum_{j=1}^{J} g_k \ln (p_j) \]  

Where \( g_k \) is an indicator variable for each SE and each SNP, \( g_k = 1 \) if SNP, \( g_k = 0 \) otherwise; \( p_j \) is the P-value of SNP, in GWAS summary data. Based on Fisher’s transformation, we calculated the P-value for SE using corresponding \( \chi^2 \) distribution. The Bonferroni correction was then employed for the multitest, SNP-mapped SE (mSE) with P-value < 4.08e−5 (0.05/1224 for biological replicate 1) or 4.35e−5 (0.05/1149 for biological replicate 2) were regarded to be significant.

Identification of long-range interactions between pri-miRNA and SEs in human osteoblast

The human miRNA data were downloaded from miRBase v22 (60), a total of 1914 pri-miRNAs transcripts and 2880 mature miRNAs were included. Note that the genome positions of miRNAs were lifted over from GRCh38 to hg19 since the miRBase v22 only provided miRNA in GRCh38 version. For each pri-miRNA, we designed its promoter baits region. Since only a part of TSS of pri-miRNAs were available, we employed two strategies for that here: (i) using the miRNA gene promoter baits which were designed by SureSelect library (Agilent) targeted TSSs in Alessandra Chesi et al. study (9); (ii) we obtained the TSS of pri-miRNAs from mirGen v3 database (61), and defined the promoter region of a particular pri-miRNA as the 2.5 kb upstream region before the TSS or itself. Finally, there were 1549 promoter baits region were obtained from Agilent SureSelect library, and 365 putative promoter baits regions were designed using miRGen v3 TSS or the pri-miRNAs-self (Supplementary Material, Table S5).

We obtained the high-resolution genome-scale, promoter-focused Capture C data and ATAC-seq data in primary MSCs-derived osteoblasts from ArrayExpress (E-MTAB-6862, E-MTAB-6834) (9), including 295,422 DNA-looping interactions (1-fragment and 4-fragment resolutions) and 156,406 ATAC-seq peaks. Note that these data we downloaded were not raw but had been processed using CHICAGO and ENCODE ATAC-seq pipeline (https://github.com/kundsjelab/atac_dnase_pipelines).
the processing steps, including quality control, were detailed in the original study (9). Then, we overlapped the genome-wide long-range interactions regions with the promoter of pri-miRNAs and significant mSEs we detected, respectively. For the long-range mSEs-interacted miRNAs (mSE-miRNAs), we further investigated the chromatin accessibility by overlapping them with ATAC-seq peaks. The mSE-miRNAs with open chromatin region in the promoter or TSS was considered to be accessible. The LocusZoom and WashU Epigenome Browser were employed for results plotting (62,63). Here, we developed a tool for the genome-wide SE-based analysis and the detection of long-range osteoblast-restricted mSE-miRNAs (https://github.com/Zheng-Lab-Westlake/Osteo-Fine-Mapp-SNP2SE2miRNA).

Target gene prediction of accessible mSE-miRNAs

We used a two-steps approach to predicted target genes of accessible mSE-miRNAs. First step, we accessed TargetScan v7.2 and miRDB v6.0 (50,51), the two most commonly used miRNA targets database, to initially conduct the prediction. For TargetScan results, any targets predicted with the absolute value of cumulative weighted context++ scores (CWCS) >0.2 will be filtered out, CWCS was a specific value that provided by TargetScan algorithm for the measurement of prediction confidence (30). For miRDB results, a prediction score ranged from 50 to 100 was used to measure the confidence (31). And as the database developer recommended, any target with the score <60 will be excluded. The union of results of both databases was then formed as the initial target genes pool. Second step, we downloaded the osteoblasts-specific Ago2 CLIP-seq data from GEO (GSE114432) (64), and downloaded the 3′ UTR RefSeq (hg19) of human genes from UCSC (http://genome.ucsc.edu/). We called the peaks from the bedGraph data of Ago2 CLIP-seq by using MACS2 bdgpeakcall utility (51). The 3′ UTR of predicted target genes were then overlapped with the Ago2 CLIP-seq peaks. Any targets failed the overlapping will be considered as false positive target and filtered out from prediction pool. Beside, for the pairwise mature miRNAs (i.e. -5p/-3p) that were derived from the same pri-miRNA, we found that the quantity of final targets was greatly different from each other, including miR-125b-5p/-3p (69 versus 527), miR-190a-5p/-3p (169 versus 37), miR-190a-5p/-3p (69 versus 527), miR-122-5p/-3p (45 versus 139), miR-212-5p/-3p (66 versus 136) and miR-6370-5p/-3p (143 versus 44) (Supplementary Material, Table S8). Here, only the one with more targets in each mature miRNA pair was kept for next analysis because of the quantity difference and overlapping of targets.

Biological pathway and process enrichment analysis

For the putative target genes of 13 accessible mSE-miRNAs, we performed biological pathway and process enrichment analysis using the web-based tool metaScape (65) and R package clusterProfiler (66). Enriched terms with a P-value < 0.01, a minimum miRNA target genes count of 3, and a ratio between the observed counts and the counts expected by chance > 1.5 (enrichment factor) were collected and grouped into clusters based on their membership similarities (65). The Benjamini–Hochberg procedure was used here for multiple testing (67). For enriched pathway and process clusters across the 13 miRNAs, hierarchical clustering was then performed by metaScape pipeline.

Hub-gene detection and enrichment analysis

We next performed PPI network analysis with the following database BioGrid (68), InWebIM and OmniPath (69) on the metaScape platform. The PPI network contains the subset of proteins that form physical interactions with at least one other member in the putative target genes of 13 mSE-miRNAs. MCODE algorithm (70) was then applied to identify hub-gene groups from the PPI network with default settings. Next, for each hub-gene group identified, the GO biological process enrichment analysis was applied independently using metaScape pipeline (65), and the three best-enriched terms by P-value were used as the functional description of the corresponding components.

Supplementary Material

Supplementary Material is available at HMG online.

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