Single-cell RNA sequencing analysis reveals the relationship of bone marrow and osteopenia in STZ-induced type 1 diabetic mice

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Highlights
- scRNA-seq analysis reveals the profiles of bone marrow cells in STZ-induced T1D mice.
- scRNA-seq analysis reveals the heterogeneity of bone marrow immune cells in STZ-induced T1D mice.
- The ratio of BM-neutrophils to B lymphocytes is increased in the bone marrow of STZ-induced T1D mice.
- Osteopenia is occurred in STZ-induced T1D mice.
- This increased ratio is negatively correlated with osteopenia in STZ-induced T1D mice.

Graphical Abstract

Introduction: Type 1 diabetes (T1D) is a multifactorial autoimmune disease. Broad knowledge about the genetics, epidemiology and clinical management of T1D has been achieved, but understandings about the cell varieties in the bone marrow during T1D remain limited.

Objectives: We aimed to present a profile of the bone marrow cells and reveal the relationship of bone marrow and osteopenia in streptozotocin (STZ)-induced T1D mice.

Methods: The whole bone marrow cells from the femurs and tibias of healthy (group C) and STZ-induced T1D mice (group D) were collected for single-cell RNA sequencing analysis. Single-cell flow cytometry...
Introduction

Type 1 diabetes (T1D) is characterized by chronic insulin deficiency and resultant hyperglycemia developed due to an autoimmune attack on pancreatic β cells [1,2]. The peak incidence of T1D onset is between 0 and 14 years old [3], but symptomatic onset has no age limitations [1]. Due to T1D’s severe symptoms and complications, increasing attention has been paid to its pathogenesis and related complication prevention. Increasing evidence has confirmed that T1D is associated with inflammation, and immune cells, such as neutrophils, macrophages, dendritic cells, and B lymphocytes, can initiate the diabetogenic T cell response and promote T1D [4]. In addition, changes in peripheral immune cell proportions are involved in aortic stiffness, a complication caused by T1D [5]. Moreover, T1D is complicated with mild to modest deficits in osteopenia and even osteopenic fracture [6–8]. However, there exists no definitive understanding of T1D-induced osteopenia’s mechanism nor standard guidelines for its prevention. Therefore, there is an urgent need to better reveal the mechanism of T1D-induced osteopenia to reduce the risk of fractures, eventually hoping to improve the quality of life of T1D patients.

Neutrophils and lymphocytes are indispensable participants involved in various immune responses, and their levels are readily available through routine blood count analyses [5]. In addition, the neutrophil-to-lymphocyte ratio (NLR), with its simple, non-invasive, and cost-effective characteristics, has been widely available as a marker for inflammation and diseases, including cancer [9–11], coronavirus pneumonia 2019 (COVID-19) [12], and acute coronary syndrome [13]. A multisite longitudinal cohort study showed that high bone mineral density (BMD) loss was associated with an increased NLR [14,15]. Furthermore, patients with T1D were confirmed to have significantly higher peripheral NLR [5], implying that an increased NLR in peripheral blood is closely associated with osteopenia and T1D.

Bone marrow, thymus, and lymphoid tissues are dedicated immune system organs for immune cell development [16]. The most important is bone marrow, serving as the initial organ that produces immune cells. However, the changes in immune cells in the bone marrow of T1D patients are still unclear. Despite our knowledge about osteopenia [17,18] and increased NLR [5] in T1D patients and in peripheral blood, a hitherto unacknowledged role for NLR in the bone marrow during T1D has not yet been revealed.

In this study, using single-cell RNA sequencing technology, we revealed the profile and characteristics of whole bone marrow cells and immunohistochemistry were performed to confirm the proportional changes among bone marrow neutrophils (BM-neutrophils) (Cxcr2+, Ly6g*) and B lymphocytes (Cd19*). X-ray and micro-CT were performed to detect bone mineral density. The correlation between the ratio of BM-neutrophils/B lymphocytes and osteopenia in STZ-induced T1D mice was analyzed by nonparametric Spearman correlation analysis.

Results: The bone marrow cells in groups C and D were divided into 12 clusters, and 249 differentially expressed genes were found. The diversity of CD45+ immune cells between groups C and D were greatly affected: the proportion of BM-neutrophils showed a significant increase while the proportion of B lymphocytes in group D showed a significant decrease. X-ray and micro-CT analyses confirmed that osteopenia occurred in group D mice. In addition, the results of single-cell flow cytometry and correlation analysis showed that the ratio of BM-neutrophils/B lymphocytes negatively correlated with osteopenia in STZ-induced T1D mice.

Conclusion: A single-cell RNA sequencing analysis revealed the profile and heterogeneity of bone marrow immune cells in STZ-induced T1D mice for the first time. The ratio of BM-neutrophils/B lymphocytes negatively correlated with osteopenia in STZ-induced T1D mice, which may enhance understanding for treating T1D and preventing T1D-induced osteopenia.

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teum on the outer surfaces of the femur and tibia, three femurs and tibias of group C were combined into one sequencing sample and three femurs and tibias of group D were combined into one sequencing sample. The bones were cut into approximately 1 mm long fragments and digested with 1 mg/ml collagenase I/dispase II (Roche, 46793022). After incubating for 40 min at 37 °C, the cell suspension was filtered through a 40-µm cell strainer (Corning, 431751) to obtain a single-cell suspension and centrifuged at 1500g for 5 min [23]. Next, we resuspended the cells in 1x RBC lysis buffer (BioLegend, 420301) on ice for 5 min to remove the red blood cells. The solution was then centrifuged at 500g for 5 min and resuspended in PBS. Finally, the sample was stained with trypan blue (Sigma), and cell survival was evaluated microscopically.

Single-cell RNA sequencing

Single-cell suspensions were prepared at a concentration of 1 × 10^6 cells/ml in PBS then loaded onto microfluidic devices, where scRNA-seq libraries were constructed according to the Single-Cell GE揽COPE® protocol by the GE揽COPE® Single-Cell RNA Library Kit (Singleion Biotechnologies) [24]. Individual libraries were diluted to 4 nM and pooled for sequencing. Pools were sequenced on an Illumina HiSeq X with 150 bp paired-end reads.

scRNA-seq quantifications and statistical analysis

Raw reads were processed to generate gene expression profiles using an internal pipeline. Briefly, after filtering out read one without poly T tails, the cell barcode and unique molecular identifier (UMI) were extracted. Adapters and poly A tails were trimmed (fastp 2.5.3a and featureCounts 1.6.2) [25]. Reads with the same cell barcode, UMI, and gene were grouped to calculate the number of UMIs per gene per cell. Each cellular barcode’s UMI count table was stored for further analysis. Cell type identification and clustering analysis were performed using the Seurat program [26,27] (http://satijalab.org/seurat/, R package, v.3.0.1), which was also applied for the RNA sequencing data analysis. UMI count tables were loaded into R using the read.table function. Then, we set the parameter resolution to 0.6 for the FindClusters function for clustering analyses. Next, differentially expressed genes (DEGs) between different samples or consecutive clusters were identified with the function FindMarkers. In addition, Gene Ontology (GO) function enrichment analysis and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway analysis were performed on the gene set using clusterProfiler software to identify biological functions or pathways that were significantly associated with the specific genes expressed [28].

RNA isolation and quantitative real-time PCR

Total RNA was extracted from the bone marrow form healthy and STZ-induced T1D mice with TRIzol reagent (Takara, China) according to the manufacturer’s instructions. For mRNAs quantification, and reversely transcribed into cDNAs with the PrimeScript RT reagent Kit (Takara, China). The resulting cDNAs were quantified with SYBR Green reagent (Takara, China) by using the LightCycler 480II (Roche, Germany). The relative expression levels and quantification of mRNAs (Asprv1, Pim1, H110, Prok2, H2-Aa, H2-Eb1, Vpreb3 and Akap12) were calculated with the 2^(-ΔΔCt) method. Beta-actin was used as an internal control. All the primers were provided by Sunya (Zhejiang, China) listed in Table S2.

Immunohistochemistry

The femurs from the two groups at aged 22 weeks were fixed in 4% paraformaldehyde for 24 h. Then, the femur sections (5 µm) were decalcified, dehydrated, and paraffin-embedded using standard histologic techniques. Endogeneous peroxidases were blocked with 3% H2O2. Next, antigen retrieval was completed with a complex digestive solution (AR0022; Boster), blocked with goat serum (AR0009; Boster), and then stained with CXCR2 (1:500; PAB18401; Abnova) and CD79a (1:500; RAB00679; Abnova) at 4 °C overnight. The sections were incubated with enhanced enzyme-labeled goat anti-rabbit IgG polymer (PV-9001; ZSGB-BIO), and the color reaction was conducted using DAB (ZLI-9018; ZSGB-BIO). The sections were scanned by a slide scanner using specific software (3DHISTECH, Ltd. Budapest, Hungary). The experiment was repeated more than three times.

Flow cytometry analysis

For single-cell flow cytometry, a single-cell suspension of bone marrow was collected as described in single-cell RNA sequencing [23]. After lysing and removing the red blood cells, we washed and resuspended the samples in staining buffer (BioLegend, 420201). Then, the samples were incubated with Fc block (BioLegend, 101302) for 10 min on ice to reduce non-specific immunofluorescent staining. For cell surface marker detection (CD45, Cxcr2, Ly-6g, CD19), cells were resuspended in 100 µl staining buffer, stained with primary antibody against surface markers for 30 min at 4 °C in the dark, and washed twice with staining buffer. For intracellular staining (Ki67), the cell pellet was resuspended in 250 µl Fix/Perm solution for 20 min at 4 °C, then washed twice in 1 × BD Perm/Wash buffer (BD Bioscience, 554714). Cells were then stained for intracellular molecules for 30 min at 4 °C in the dark and washed twice with 1 × BD Perm/Wash buffer. Finally, cell pellets were resuspended in staining buffer for flow cytometric analysis (Cytoflex Lx, Beckman Coulter), where gates and compensations were set before every experiment. All the antibodies mentioned above are listed in Table S1. The experiment was repeated more than three times.

Radiography

X-ray radiography was performed using a small-animal X-ray apparatus (UltraFocus, Faxitron) to detect the difference in BMD changes in group C(n = 6) and D(n = 5) mice’s left legs. The exposure time and kV were set to the auto setting [29]. The window level was set from 2058 to 7200.

Computed tomography

Computed tomographic images of the left femurs in groups C(n = 6) and D(n = 5) were acquired using a micro-CT scanner (VivaCT80, SCANCO Medical Company, Switzerland) at high resolution. After the femur was reconstructed, BMD (mg HA/ccm), trabecular bone volume/total volume (%), trabecular thickness (Th.Th, mm), trabecular number (Th.N, 1/mm), and trabecular space (Th.Sp, mm) [30] were assessed by scanner software (μCT Evaluation Program V6.5–3).

Statistical analysis

The data were presented as the means ± SEM and were statistically analyzed using GraphPad Prism (version 9.0; GraphPad Software, LLC). Student’s t-tests were used when comparing two groups of data. Nonparametric Spearman correlation analysis was
used for correlation analyses. P values < 0.05 were considered to be significant.

Results

Profiles of bone marrow cells in healthy and STZ-induced T1D mice

We collected the whole bone marrow cells from the femurs and tibias of healthy (group C) and STZ-induced T1D mice (group D). After digestion and erythrolysis, 12,279 cells from group C and 15,118 cells from group D were collected for single-cell RNA-sequencing analysis (Fig. 1A). Fig. 1B demonstrated the unbiased clustering of all cells in group C and group D, originating from their co-analysis. The result revealed 12 cell clusters: spanning neutrophils (mainly expressing Cxcr2, Ly6g, and Ltyf), monocytes (mainly expressing F11, Ccr2, and F13a1), neutrophil-myeloid progenitor (NMP, mainly expressing Elane, Mpo, and Ptnn3), T lymphocytes (mainly expressing Cd3d, Cdl5, and Ms4a4b), B lymphocytes (mainly expressing Cd79a, Cd19, and Igdm), dendritic cells (DCs, mainly expressing Siglech, Irf8, and Bst2), erythrocytes (mainly expressing Hba-a1, Hbb-bs, and Hbb-bt), hematopoietic stem cell (HSC, mainly expressing Cd34, Adgrg1, and Cdk6), macrophages (mainly expressing C1qa, Vcam1, and Mrc1), basophils (mainly expressing Cd200r3, Mcpt8, and Prss34), plasma cells (mainly expressing Jchain, Igglc2, and Mzb1), and mesenchymal stem cells (MSC, mainly expressing Cxcl12, Col1a2, and Lepr) (Fig. 1C, D, G, H).

The proportional changes of different cell clusters in the bone marrow were discussed here. CD45, known as the common leukocyte antigen, is expressed by all immune cells. CD45+ immune cells accounted for 95.44% of cells in group C and 96.33% of cells in group D (Fig. 1F); This result was consistent with that from the flow cytometry for CD45 in group C and group D (Fig. S1). In addition, these following cell clusters presented an increase in their proportion: BM-neutrophils increased from 46.94% to 64.04%; NMP increased from 3.47% to 6.48%; plasmas and T lymphocytes increased slightly. On the other hand, these following cell clusters presented a decrease in their proportion: B lymphocytes decreased sharply from 18.12% to 3.34%; DCs decreased significantly from 4.00% to 0.58%; macrophages, basophils, and monocytes decreased slightly. Among all immune cells mentioned above, the BM-neutrophil cell cluster had the most significant proportional increase, while the B lymphocytes cell cluster experienced the most significant proportional decrease (Fig. 1E, F).

There were 249 DEGs between groups C and D, including 69 upregulated and 180 downregulated genes. The top 20 most highly upregulated and downregulated DEGs are shown in Fig. S2. Besides, to confirm the results of single-cell RNA sequencing, the relative expression levels of eight DEGs selected randomly between groups C and D were validated by quantitative real-time PCR (Fig. S3). The results showed that there was the same DEGs expression trend between single-cell RNA sequencing and quantitative real-time PCR. GO enrichment analysis and KEGG pathway analysis were performed on the upregulated and downregulated DEGs. The most highly enriched biological process (BP), cellular component (CC), and molecular function (MF) terms among the upregulated DEGs were the response to interleukin-1, secretory granule, and antioxidant activity, respectively (Fig. S4). Conversely, the most enriched BP, CC, and MF terms among the downregulated DEGs were cytoktoplasmic translation, cytosolic ribosome, and structural constituent of ribosome, respectively (Fig. S5). In addition, the most enriched pathways among the upregulated DEGs were the IL-17 signaling pathway, AGE-RAGE signaling pathway in diabetic complications, TNF signaling pathway, and osteoclast differentiation (Fig. S5). In contrast, the most enriched pathways among the downregulated DEGs were those associated with ribosome, antigen processing and presentation, and B cell receptor signaling pathway (Fig. S7).

Heterogeneity of BM-neutrophils in the bone marrow of STZ-induced T1D mice

As previously mentioned, BM-neutrophils presented the largest proportional increase among all the immune cells in group D. To explore the heterogeneity of BM-neutrophils, we analyzed the characteristics of the BM-neutrophil subsets. The results showed that the BM-neutrophils could be subdivided into 6 subsets (Fig. 2A–C): BM-neutrophil_1 (mainly expressing Asprv1, Prok2, and Fbxs5), BM-neutrophil_2 (mainly expressing Ltf, Zmpste24, and Cybb), BM-neutrophil_3 (mainly expressing Clec4d, Thbs1, and Egr1), BM-neutrophil_4 (mainly expressing Ube2c, Mk67, and Top2a), BM-neutrophil_5 (mainly expressing Pcnna, Rm2, and Pclaf), and BM-neutrophil_6 (mainly expressing Actb, Cto10467.1, and Tmsb4x) (Fig. 2B, F). The proportional changes of each subset were as the following: BM-neutrophil_1 increased the most, spiking from 5.97% to 50.14%; BM-neutrophil_2 decreased from 34.4% to 19.12%; BM-neutrophil_3 decreased from 38.53% to 3.13%; BM-neutrophil_4 decreased from 16.22% to 15.33%; BM-neutrophil_5 increased slightly from 4.2% to 5.67%; BM-neutrophil_6 increased slightly from 0.68% to 6.61%. Compared with group C, the proportion of BM-neutrophil_1 demonstrated an impressive increase. Conversely, the proportion of BM-neutrophil_3 presented a most significant decrease (Fig. 2D). Moreover, we performed immunohistochemistry staining to validate the increase in BM-neutrophils (marked by Cxcr2) in group D (Fig. 2E).

Among all DEGs in the 6 neutrophil subsets, Asprv1, Ltf, Clec4d, Ube2c, Pcnna, and Actb were DEGs with the most significant changes in proportion. Asprv1, a mammalian retroviral-like enzyme [31], and Clec4d, a trehalose-6,6'-dimycolate receptor [32], were both expressed in the BM-neutrophil_1 and BM-neutrophil_3 subsets (Fig. 2E). Asprv1 and Clec4d showed the most significant changes among BM-neutrophil_1 and BM-neutrophil_3, respectively. Ltf, an extracellular iron-binding glycoprotein [33], and Actb, were both expressed in whole bone marrow cells and the 6 BM-neutrophil subsets (Fig. 2E). Ube2c, a member of the ubiquitin modification system [34], and Pcnna, a gene associated with cell proliferation, were mainly expressed in BM-neutrophil_4 and BM-neutrophil_5, respectively (Fig. 2F).

Next, pseudotime analysis of the 6 subsets of BM-neutrophils was carried out using Monocle. The results suggested that there were two diverging cell fates, one starting with BM-neutrophil_4 and BM-neutrophil_5 and differentiating towards BM-neutrophil_1 or BM-neutrophil_3, and the other starting with BM-neutrophil_2 and BM-neutrophil_6 and differentiating towards BM-neutrophil_1 or BM-neutrophil_3 (Fig. 3A–D). Moreover, the ordering was captured by latent time and RNA velocity (Fig. 3E).

Furthermore, because BM-neutrophil_1 made up the highest proportion of cells in group D, GO enrichment analysis and KEGG pathway analysis were performed for the upregulated DEGs in BM-neutrophil_1. The results showed that most of the biological processes enriched among the DEGs in BM-neutrophil_1 were related to the hexose metabolic process, another glucose metabolism, anabolism, and cytokine–mediated signaling pathway. Extracellular vesicles, including exosomes and other organelles, were the most common cellular components (Fig. 3G). KEGG pathway analysis indicated that osteoclast differentiation, IL–17 signaling pathway, and TNF signaling pathway were the most highly enriched pathways among the DEGs (Fig. 3H). As listed in Fig. 3I–J, the top 5 most upregulated DEGs were Asprv1, Pim1, Cebpβ,
H1f0, and Prok2, and the top 5 most downregulated DEGs were Lyz2, Ifitm3, Thbs1, S100a8, and Lcn2. We mixed bone marrow cells from three mice into one sequencing sample in group C or in group D. The biological duplication was achieved. But as for single-cell sequencing sample analysis, the sample number is only one, so histograms in Fig. 3 did not present statistical analysis.

Heterogeneity of B lymphocytes in the bone marrow of STZ-induced T1D mice

We next examined the characteristics and differentiation of B lymphocytes. Common marker genes to annotate B lymphocytes are Bank1 [35], CD83 [36], and Ighd [37] (Fig. 4F). The top 5 most upregulated genes were H2-Aa, Ighd, Macf1, Ptpn22, and H2-Eb1. The top 5 most downregulated genes were Vpreb3, Top2a, Mki67, Akap12, and Myb. Both sets of DEGs were shown in Fig. 4J and K. Because the sample number was one in each group for single-cell sequencing sample analysis, so the histograms in Fig. 4 did not present statistical analysis. B lymphocytes could be separated into two subsets, precursor B cells (Pre_B) and Naïve_B cells (Naïve_B). Their unbiased clustering was shown in Fig. 4A–C. We observed that the Pre_B group was the dominant subset in group C, while only the Naïve_B subset was present in group D (Fig. 4D). Immunohistochemistry validated that the number of B lymphocytes (marked by CD79a) were decreased in group D (Fig. 4E). In addition, latent time and RNA velocity analysis showed that the Pre_B subset was produced earlier than the Naïve_B subset (Fig. 4G).

GO enrichment analysis and KEGG pathway analysis were performed for the upregulated DEGs in B cells. GO enrichment analysis showed that most of the enriched biological processes were associated with antigen processing and presentation of peptide antigen via MHC class II, antigen processing and presentation of peptide or polysaccharide antigen via MHC class II, and lymphocyte proliferation. Enriched cellular components mainly included the MHC protein complex, late endosome, and lysosome. MHC protein complex binding was the major molecular function (Fig. 4H). KEGG pathway analysis indicated that DEGs were mainly enriched for immune-related diseases, including tuberculosis and asthma, antigen processing and presentation, and hematopoietic cell lineage (Fig. 4I).

Heterogeneity of monocytes, T lymphocytes, and DCs in the bone marrow of STZ-induced T1D mice

We analyzed monocytes, T lymphocytes, and DCs, respectively. The changes in the proportion of each subset were analyzed and shown in Fig. 5.
Monocytes were subdivided into 5 subsets, including classical monocytes (mainly expressing Fn1, Tmsb10, and Crip1), neutrophils (mainly expressing Retnlg, Ngp, and Camp), NMP (mainly expressing Mpo, Prtn3, and Ncl), nonclassical monocytes (mainly expressing Ace, Cd36, and Apoe), and macrophages (mainly expressing Cd74, H2-Aa, and H2-Ab1) (Fig. 5A, D). The unbiased clusters of these subsets were shown in Fig. 5A–B. The proportional changes of each subset were as follows: classical monocytes decreased from 71.09% to 60.75%; neutrophils increased from 15.39% to 27.85%; NMP increased from 5.46% to 7.66%; classical nonclassical monocytes increased slightly from 2.91% to 3.48%; macrophages decreased from 5.15% to 0.26%. Except for the macrophage subset that decreased noticeably, the proportions of the other subsets experienced no obvious changes compared to group C (Fig. 5C).

T lymphocytes play a critical role in the pathogenesis of T1D [38,39]. Therefore, we next analyzed the profile of T lymphocytes in the bone marrow of STZ-induced T1D mice. The T lymphocytes were divided into 4 subsets, including natural killer T cells (NKT, expressing Ly6c2, Cc15, and Cd69), naïve T cells/central memory T cells (Naïve T cells or Tcm, mainly expressing Ccr7, Hbb-bs, and Gm14085), natural killer cells (NK, mainly expressing Gzma, KlrA8, ...
and Klra4), and CD4+ T cells (CD4+ T, mainly expressing Rora, Ikd2, and Ttn) (Fig. 5E, H). The unbiased clusters of these subsets were shown in Fig. 5E–F. The proportional changes of each subset were as follows: NKT decreased from 49.32% to 36.0%; Naïve T cells or Tcm increased from 26.03% to 51.73%; NK decreased from 18.15% to 10.4%; CD4+ T decreased from 6.51% to 1.87%. Except for the obvious increase in the proportion of Naïve T cells or Tcm, the other T lymphocyte subsets had a decreasing trend in proportion in group D (Fig. 5G).

Fig. 3. Differential trajectory of BM-neutrophil subsets and enrichment of DEGs. (A–D) Differential directions of 6 BM-neutrophil subsets. Heatmap presenting relative expressions of markers of BM-neutrophils along inferred trajectories in 6 BM-neutrophil subsets. The red and blue branches correspond to the two differential directions (A). Monocle pseudotime trajectory expression pattern of 6 BM-neutrophil subsets (B). Pseudotime trajectories of the BM-neutrophil subsets (C–D). (E) Latent time and RNA velocity of 6 BM-neutrophil subsets. Latent time and RNA velocity showed the internal clock of BM-neutrophil subsets. Different colors of latent time represent different differentiation times, the colder the color, the earlier, the warmer the color, the later. The arrows of RNA velocity represent the different directions of differentiation. (F) Signature genes of BM-neutrophil subsets. Subset signature genes highlighted on left. Expression of signature genes (rows) across the cells (columns), the warmer the color, the higher the expression. (G) GO analysis of BM-neutrophil_1. Bubble diagram of upregulated DEGs of BM-neutrophil_1 enriched in GO analysis. (H) KEGG analysis of BM-neutrophil_1. Bubble diagram of upregulated DEGs of BM-neutrophil_1 enriched in KEGG analysis. (I–J) The top 5 DEGs of BM-neutrophils. The top 5 upregulated DEGs between groups C and D (I). The top 5 downregulated DEGs between groups C and D (J).

DCs also experienced one of the most significant decreases in proportion among cells and were subdivided into 4 subsets, described as DCs_1 (mainly expressing Klk1, Cd74, and H2-Aa), DCs_2 (mainly expressing Stat1, Runx2, and Ly6d), DCs_3 (mainly expressing Top2, Mki67, and Hist1h1b) and DCs_4 (mainly expressing Prtn3, F13a1, and Mcm3) (Fig. 5I, L). The unbiased clustering of these subsets was shown in Fig. 5I–J. The proportional changes of each subset were as follows: DCs_1 decreased from 39.89% to 7.94%; DCs_2 increased from 26.59% to 39.68%; DCs_3 increased from 18.28% to 25.40%; DCs_4 increased from 15.24% to 26.98% (Fig. 5K). Except for the significant decrease in the proportion of DCs_1, the other DC subsets experienced slight increases in cell proportion in group D.
Fig. 4. The characteristics of B lymphocytes between groups C and D. (A) UMAP of the cluster of B lymphocytes among all cells. (B-C) Two B lymphocyte subsets. UMAP of unbiased clustering and cell annotation of the subsets of B lymphocytes in groups C and D (B). UMAP of unbiased clustering of the subsets of B lymphocytes in groups C and D (C). (D) The proportion of the subsets of B lymphocytes in groups C and D. (E) B lymphocytes were labeled by immunohistochemistry of the sections of femur in groups C and D (CD79a; Scale bar, 40 μm or 20 μm). (F) Key marker genes of B lymphocyte subsets. UMAP of key marker genes of the subsets of B lymphocytes, along with the corresponding distribution of expression levels among 12 clusters. Key marker genes included Bank1, CD83 and Ighd. (G) Latent time and RNA velocity of 3B lymphocyte subsets. Latent time and RNA velocity showed the internal clock of B lymphocyte subsets. Different colors of latent time represent different differentiation times, the colder the color, the earlier, the warmer the color, the later; The arrows of RNA velocity represent the different directions of differentiation. (H) GO analysis of Naïve_B. Bubble diagram of upregulated DEGs of Naïve_B enriched in GO analysis. (I) KEGG analysis of Naïve_B. Bubble diagram of upregulated DEGs of Naïve_B enriched in KEGG analysis. (J-K) The top 5 DEGs of B lymphocytes. The top 5 upregulated DEGs between groups C and D (J). The top 5 downregulated DEGs between groups C and D (K).
Osteopenia in STZ-induced T1D mice

The weight and blood sugar levels of all groups of mice were measured at 22 W and 28 W of age. The results showed that our T1D model was successful (Fig. 7A). To identify whether osteopenia occurred in STZ-induced T1D mice, we used X-ray and micro-CT to analyze the changes in bone mass in groups C (28w, n = 6) and D (28w, n = 5). Fig. 6A showed the X-ray images of femurs and tibias, where it was clear that BMD had decreased in group D compared with group C. Moreover, the micro-CT images of the same part of the femur showed that the cortical bone was obviously thinner in group D. BMD ($P < 0.001$), trabecular bone volume/total volume (BV/TV) ($P < 0.001$), and trabecular thickness (Tb.Th) ($P < 0.001$) also experienced significant decreases in the femurs of group D (Fig. 6C). Meanwhile, trabecular number (Tb.N) ($P = 0.098$) and trabecular space (Tb.Sp) ($P = 0.085$) experienced decreased and increased trend, respectively (Fig. 6C). In summary, it was confirmed that osteopenia could occur in STZ-induced T1D mice.

Ratios of immune cells in the bone marrow of STZ-induced T1D mice

Increased NLR in the blood is correlated with bone loss and has been proposed as an index of related diseases, including osteoporosis in postmenopausal women and elders [15,40]. This critical correlation prompted us to explore if the change in NLR in peripheral blood originates from the change in ratios of immune cells in the bone marrow in T1D-induced osteopenia. Compared with group C, group D demonstrated an increased proportion of particular CD45+ immune cells, mainly BM-neutrophils, monocytes (including macrophage-like monocytes), and T lymphocytes. Meanwhile, group D also demonstrated a decreased proportion of other particular CD45+ immune cells, mainly B lymphocytes and DCs. The specific proportions of immune cells came from the
results of the single-cell RNA sequencing analysis. The ratios of the above immune cells between groups C and D are shown in Fig. 7C. RD/RC (the ratio of cells in group D to cells in group C) was used to illustrate the differences in the ratios between groups C and D. We can see from Fig. 7C that the RD/RC of BM-neutrophil_1/Naive_B was the highest. Moreover, there was an increasing trend in the RD/RC of BM-neutrophils/DCs, BM-neutrophils/B lymphocytes, monocytes/DCs, and monocytes/B lymphocytes. The above result found that the ratio of BM-neutrophils/B lymphocytes increased in bone marrow, similar to the increase of NLR in peripheral blood.

**Ratio of BM-neutrophils and B lymphocytes in bone marrow was related to osteopenia in STZ-induced T1D mice**

Next, we performed single-cell flow cytometry to validate the changes in proportion for BM-neutrophil (Cxcr2+, Ly6g+) [41,42] and B lymphocyte (Cd19+) [43] in groups C (right, n = 4) and D (right, n = 4) (Fig. 8A). The ratio of BM-neutrophils/B lymphocytes (P = 0.040) was shown in Fig. 8A. We performed a correlation analysis (GraphPad Prism 9.0) between the ratio of BM-neutrophils/B lymphocytes and the ratio of BV/TV (corresponding left, n = 4). The results demonstrated that there was a negative correlation between the ratio of BM-neutrophils/B lymphocytes and the ratio of BV/TV (r = -0.7619, P = 0.0368) (Fig. 8C).

**Discussion**

T1D is largely portrayed as an autoimmune disease. However, research has yet given the role of the bone marrow, the initial organ producing immune cells, in T1D sufficient attention. To understand the varieties of bone marrow cells of T1D, we conducted a single-cell RNA sequencing analysis of whole bone marrow cells from the femurs and tibias of STZ-induced T1D mice for the first time. STZ-induced T1D was used as the model since it has been considered a representative model of T1D [21]. The result presented an unbiased clustering of all cells, and we focused on the changes in immune cells in the bone marrow between healthy and STZ-induced T1D mice. Furthermore, the characteristics of BM-neutrophils, B lymphocytes, monocytes, T lymphocytes, and DCs were identified.

As a result of the increased incidence and prevalence of T1D, the number of patients at risk for T1D-induced osteopenia is expected to rise [6,44]. Thus, we also explored the relationship between immune cells in the bone marrow and osteopenia in STZ-induced T1D mice. We found that the proportion of BM-neutrophils increased most significantly, and the proportion of B lymphocytes decreased most significantly. The increased ratio of BM-neutrophils/B lymphocytes was negatively correlated with T1D-induced osteopenia. Our results revealed the profiles of bone marrow cells in STZ-induced T1D mice and the correlation between immune cells and osteopenia, which may contribute to a better understanding of the pathogenesis of STZ-induced T1D and osteopenia in STZ-induced T1D mice.

As universally acknowledged, bone marrow neutrophils (immature neutrophils) are usually divided into 5 subsets according to the morphology of the nucleus, including myeloblast, promyelocyte, myelocyte, metamyelocyte, and stab granulocyte. Our discovery could provide a new classification for BM-neutrophils,
subdividing them into 6 subsets according to their DEGs for the first time. We speculate that there may be an intersection among the two classifications of the BM-neutrophils subset. More importantly, the DEG-based classification may be more helpful for studying specific functions of neutrophil subsets. The proportion of BM-neutrophil_1 was highest, while the proportion of other subsets all tended to differentiate into BM-neutrophil_1. GO analysis of the upregulated DEGs between BM-neutrophil_1 and the other subsets indicated that extracellular vesicles, including exosomes and other organelles, were the most enriched cellular component term, consistent with the GO analysis of the upregulated DEGs between healthy and STZ-induced T1D mice. KEGG pathway analysis of BM-neutrophil_1 indicated that it was mainly enriched for osteoclast differentiation, IL-17 signaling pathway, and TNF signaling pathway. This result was also consistent with the KEGG analysis of the upregulated DEGs between healthy and STZ-induced T1D mice.

KEGG pathway analysis of BM-neutrophil_1 indicated that it was mainly enriched for osteoclast differentiation, IL-17 signaling pathway, and TNF signaling pathway. This result was also consistent with the KEGG analysis of the upregulated DEGs between healthy and STZ-induced T1D mice. These results demonstrated that the BM-neutrophil_1 subset might be closely involved in the pathological processes of T1D and T1D-induced osteopenia. Furthermore, it has been confirmed that there is a reduction in circulating neutrophils in T1D patients [45–48]. Several explanations have been proposed for the decrease in circulating neutrophils, such as an impairment in neutrophil output from the bone marrow, the differentiation of neutrophils, an increase in peripheral consumption, destruction of neutrophils, or tissue sequestration [45]. However, the exact mechanisms responsible for the reduction in circulating neutrophils are still unclear. According to our results, flow cytometry of Ki67 showed no significant difference in cell proliferation (Fig. S8). Meanwhile, the proportion of neutrophils increased notably, suggesting that a limited neutrophil outflow from the bone marrow seems to be involved. However, this potential mechanism requires further in-depth study.

Additionally, evidence has shown that B lymphocytes also play a key role in the pathogenesis of T1D [43,49]. Among bone marrow cells, B lymphocytes experienced the most significant decrease in proportion. B lymphocytes could be subdivided into Pre_B and Naïve_B. Only Naïve_B was presented in STZ-induced T1D mice, and the enrichment analysis mainly reflected terms related to promoting the immune response by antigen processing and presentation. In addition, enhanced lymphocyte proliferation was highly expressed in the bone marrow of STZ-induced T1D mice, which is consistent with the critical role of B cells in the pathogenesis of T1D [49] and their therapeutic potential for T1D [43]. Other immune cells with significant proportional changes mainly included monocytes, T lymphocytes, and DCs. Excluding the proportion change of B lymphocytes, we found that DCs experienced the subsequent largest decrease in proportion. The increase in monocytes in STZ-induced T1D mice was mirrored by an increase in the neutrophil subset, which suggested that there may be a transformation of monocytes to neutrophils. The increase of the proportion of T cells in STZ-induced T1D mice mainly comprised an increase in Naïve T cells or Tcm. This increase in Naïve T cells could prevent chronic infection and control systemic secondary infections with their stem cell-like capacity to expand, differentiate, and self-renew [50].

Furthermore, the X-ray and micro-CT imaging analysis results showed a significant reduction in the BMD of STZ-induced T1D mice, consistent with previously published results [7]. We then determined the ratio between two sets of immune cells among BM-neutrophils (set monocytes and T lymphocytes, set B lymphocytes, set T lymphocytes, set monocytes/DCs, set BM-neutrophil_1/Naïve_B).
Fig. 8. The correlation between immune cells and osteopenia. (A) Flow cytometry of BM-neutrophils and B lymphocytes from bone marrow. The proportion of BM-neutrophils increased notably in group D and the proportion of B lymphocytes decreased notably in group D. (B) The ratios of BM-neutrophils/B lymphocytes from the results of flow cytometry. The ratio of BM-neutrophils/B lymphocytes was notably higher in group D ($p = 0.040$). (C) The correlation analysis between the ratio of BM-neutrophils/B lymphocytes and the ratio of BV/TV. There was a negative correlation between the ratio of BV/TV and the ratio of BM-neutrophils/B lymphocytes. ($r = -0.7619$, $p = 0.0368$). (*$p < 0.05$).
cytes and DCs) in healthy and STZ-induced T1D mice. The RD/RC of BM-neutrophils/B lymphocytes was increased in the bone marrow. Next, flow cytometry verified that the ratio of BM-neutrophils/B lymphocytes might be more meaningful between healthy and STZ-induced T1D mice. Moreover, as it is known that both osteopenia and increased NLR could occur in T1D patients [5,17,18], it is worth discussing whether the increased ratios of BM-neutrophils/B lymphocytes in the bone marrow was associated with osteopenia in STZ-induced T1D mice. We then conducted a correlation analysis, which indicated that there was a negative correlation between the ratio of BM-neutrophils/B lymphocytes and the ratio of BV/TV, consistent with increased peripheral NLR in T1D patients. Thus, we speculated that the increased peripheral NLR derived from increased NLR in the bone marrow. Moreover, the KEGG pathway analysis of upregulated DEGs in BM-neutrophil_1 was mainly enriched in osteoclast differentiation, which suggested that BM-neutrophil_1 may be closely involved in the mechanisms of osteopenia. Furthermore, the fact that only Naïve_B was left in STZ-induced T1D mice and that the RD/RC of BM-neutrophil_1/Naïve_B was the highest both suggested that BM-neutrophil_1 and Naïve_B were key cell subsets responsible for the negative correlation between the NLR and BMD. In addition, there was also an increasing trend in the RD/RC of BM-neutrophil_1/DCs, monocytes/DCs, and monocytes/B lymphocytes, suggesting that DCs and monocytes may become diagnostic indicators for T1D and T1D-induced complications.

Inevitably, there were still few limitations in our studies. In addition to the STZ-induced T1D model, nonobese diabetic (NOD) mouse, BioBreeding/Worcester rat, and lymphocytic choriomeningitis virus-induced model could also represent T1D animal models [51]. We limited our model in this research to STZ-induced T1D mice only. This limitation may result in an incomplete representation of the T1D model, but the STZ-induced T1D model was chosen as it could provide a relatively better understanding of the pathogenesis of T1D and T1D-induced osteopenia. We also acknowledged that NOD mice had become the preferred model for preclinical studies of T1D [51]. Thus, a complementary single-cell RNA sequencing analysis of the bone marrow in NOD mice might provide another comprehensive view.

Conclusion

Overall, our data objectively revealed the profiles of bone marrow cells in STZ-induced T1D mice for the first time through single-cell RNA sequencing analysis. In particular, the proportions of immune cells were chosen to analyze further since most of the bone marrow cells were CD45+ cells. The occurrence of osteopenia in STZ-induced T1D mice was also confirmed. Furthermore, through single-cell flow cytometry and correlation analyses, we found that the increased ratio of BM-neutrophil/B lymphocytes negatively correlates with osteopenia in STZ-induced T1D mice, which may be a marker of increased peripheral NLR in blood. Meanwhile, GO and KEGG analyses showed that BM-neutrophils and B lymphocytes were involved in the pathogenesis of T1D and T1D-induced osteopenia. Thereafter, in-depth studies on BM-neutrophils and B lymphocytes may be conducted to enhance understanding of T1D and T1D-induced osteopenia mechanisms. Studies of such may suggest that precise regulation of immune cells in the bone marrow could potentially become a novel strategy for treating T1D and preventing T1D-induced osteopenia.

Compliance with Ethics Requirements

All Institutional and National Guidelines for the care and use of animals (fisheries) were followed.

Data availability

All relevant data are available from the authors upon request. The raw data files of scRNA-seq have been deposited in the Gene Expression Omnibus with following accession number: GSE182986.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary material

Supplementary data to this article can be found online at https://doi.org/10.1016/j.jare.2022.01.006.

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