Single-cell transcriptomic reconstruction reveals cell cycle and multi-lineage differentiation defects in Bcl11a-deficient hematopoietic stem cells

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Abstract

Background: Hematopoietic stem cells (HSCs) are a rare cell type with the ability of long-term self-renewal and multipotency to reconstitute all blood lineages. HSCs are typically purified from the bone marrow using cell surface markers. Recent studies have identified significant cellular heterogeneities in the HSC compartment with subsets of HSCs displaying lineage bias. We previously discovered that the transcription factor Bcl11a has critical functions in the lymphoid development of the HSC compartment.

Results: In this report, we employ single-cell transcriptomic analysis to dissect the molecular heterogeneities in HSCs. We profile the transcriptomes of 180 highly purified HSCs (Bcl11a+/+ and Bcl11a−/−). Detailed analysis of the RNA-seq data identifies cell cycle activity as the major source of transcriptomic variation in the HSC compartment, which allows reconstruction of HSC cell cycle progression in silico. Single-cell RNA-seq profiling of Bcl11a−/− HSCs reveals abnormal proliferative phenotypes. Analysis of lineage gene expression suggests that the Bcl11a−/− HSCs are constituted of two distinct myeloid-restricted subpopulations. Remarkably, similar myeloid-restricted cells could also be detected in the wild-type HSC compartment, suggesting selective elimination of lymphoid-competent HSCs after Bcl11a deletion. These defects are experimentally validated in serial transplantation experiments where Bcl11a−/− HSCs are myeloid-restricted and defective in self-renewal.

Conclusions: Our study demonstrates the power of single-cell transcriptomics in dissecting cellular process and lineage heterogeneities in stem cell compartments, and further reveals the molecular and cellular defects in the Bcl11a-deficient HSC compartment.

Background

Continuous and responsive hematopoiesis is essential for hematopoietic homeostasis throughout the life of a mammalian individual. The classical model of hematopoiesis described this process in a developmental hierarchy, where multipotent hematopoietic stem cells (HSCs) situate at the apex [1]. Purified HSCs are able to provide long-term reconstitution to all blood lineages in transplantation experiments [2–5]. In adult mice, HSCs reside in the bone marrow and remain predominantly mitotically quiescent. Only a small portion of HSCs are cycling to maintain hematopoietic homeostasis or in response to stress and injury [6]. Lymphoid and myeloid blood cells are derived from successive differentiation of respective lineage progenitors with the loss of reconstituting potential upon commitment. HSCs generate multipotent progenitors, which in turn give rise to lineage progenitors, i.e., the common myeloid progenitors (CMPs) and common lymphoid progenitors (CLPs). CLPs subsequently produce B/T lymphocytes, whereas CMPs further differentiate to megakaryocyte-erythroid progenitors (MEPs) and granulocyte-macrophage progenitors (GMPs) to reconstitute erythrocytes, platelets and myeloid cells. Transcription regulators play critical roles in the HSC compartment and direct stem cell differentiation [7, 8].
We have previously reported that the C2H2 zinc finger transcription factor B cell CLL/lymphoma 11A (Bcl11a) is essential for both fetal and adult lymphoid development [9, 10]. Bcl11a is also highly expressed in the adult HSC compartment, and Bcl11a-deficient HSCs fail to contribute to CLP development in vivo [9]. Increasing evidence suggests that the phenotypically defined HSC compartment is heterogeneous and a portion of HSCs display lineage bias or restriction [8, 11–15]. It is possible that the lymphoid defect in Bcl11a-deficient HSCs is dependent on its functions in the regulation of lineage-priming heterogeneities in the HSC compartment. However, unbiased transcriptomic dissection of the HSC compartment at the single-cell level was previously hampered by the rarity of HSCs. This technical obstacle has recently been overcome by the advance of RNA-seq technology [16, 17], which allows genome-wide transcriptome analysis to be conducted at the single-cell level.

In this report, we applied a semi-automated microfluidic system to dissect the transcriptomic heterogeneities and reconstruct the cell cycle progression in the Lin− Sca1+ Kit+ (LSK) CD150+48− and Lin− Sca1+ Kit+ (LSK) CD150+48−34−135− HSC compartments at the single-cell level. These exercises further revealed the molecular and cellular defects in the Bcl11a-deficient HSC compartment, which only contains myeloid-restricted progenitor-like cells with substantial self-renewal defects.

Results

Single-cell transcriptome profiling of adult HSCs using a microfluidic system

We employed the Fluidigm microfluidic system coupled with multiplex barcoding to streamline the workflow in cell sorting and cDNA generation. The use of the microfluidic system allowed us to visually confirm the presence of single cells in individual capture sites under the microscope and to filter out debris that may otherwise be mistaken as single cells in flow cytometry. A maximum of 96 cells can be captured and cDNA reverse transcription could be generated in situ in each integrated fluidic circuit. We analyzed the LSK CD150+48− HSC compartment (Bcl11a+/+) from eight Bcl11a-GFP reporter conditional knockout mice that we previously described [9] (Fig. 1a). An IRES-eGFP cassette was targeted at the 3′ untranslated region of the Bcl11a locus, which enabled tracking of Bcl11a expression by green fluorescent protein (GFP). The reporter mice have normal HSC development and functions [9]. Additionally, we also purified HSCs (sBcl11a+/+) with a more stringent sorting scheme (LSK CD150+48−34−135−) from ten wild-type C57BL/6 mice (Fig. 1a). Notably, the majority (81 %) of the LSK CD150+48−34−135−, in contrast to the progenitor cell compartment (14 % of LSK CD150+48−), indicating significant overlap between the sorting schemes. The single-cell capture rate by the microfluidic integrated fluidic circuit was 86.5 % (83/96) and 54.2 % (52/96) for Bcl11a+/+ and sBcl11a+/+ HSCs, respectively (Fig. 1b). To investigate the role of Bcl11a in the HSC compartment, we used Bcl11afllox/fllox; ROSA26-Cre-ERT2 mice to obtain LSK CD150+48− Bcl11a+/− HSCs (Bcl11a+/−) [9]. Bcl11a deletion was induced by treating the mice with tamoxifen [9]. One week after tamoxifen treatment, HSCs were purified using fluorescence-activated cell sorting (FACS) from five mice and the single-cell capture rate was 85.4 % (82/96). Cell lysis, cDNA reverse transcription and pre-amplification by PCR were performed and harvested by the C1 Single-Cell Auto Prep system. The sequencing libraries of individual cells of each experiment group (Bcl11a+/+, sBcl11a+/+ or Bcl11a+/−) were then pooled separately and sequenced. After quality control, cells with low numbers of reads (<500,000 in annotated genes), low numbers of detectable genes (<3000 annotated genes), high fractions of mitochondrial encoded transcripts (>10 %) or anomalies seen under microscope were filtered out. We also removed cells showing incomplete Bcl11a exon 4 deletion in the Bcl11a+/− dataset (Additional file 1). In total, 76 Bcl11a+/+ HSCs, 44 sBcl11a+/+ HSCs and 61 Bcl11a+/− HSCs were further analyzed. The average number of unique counts of genes was 3.16 million (range 1.43–4.52 million) per cell (Bcl11a+/+), 0.89 million (range 0.55–1.71 million) per cell (sBcl11a+/+) and 3.67 million (range 0.54–12.5 million) per cell (Bcl11a+/−). Standardized External RNA Controls Consortium (ERCC) RNA spike-ins were added to the sequencing library to account for the technical variability of the protocol [18]. One significant outlier cell from the Bcl11a+/+ dataset was removed from downstream analysis after principal component analysis (PCA; Fig. 1c). The lower count number from the sBcl11a+/+ dataset is not unexpected due to the lower sequencing depth (“Methods and materials”), but the numbers of genes detected (normalized count >1) between two wild-type datasets after size factor normalization are comparable (Wilcoxon rank sum test p = 0.362) (Fig. 1d).

Cell cycle activity represents the dominant source of transcriptional heterogeneity in the HSC compartment

Single-cell transcriptomic analysis allows the detection of gene expression variability between individual cells and identification of cellular subpopulations. Expression variability of a particular gene could come from technical sources (e.g., stochasticity of reverse transcription reaction and library preparation) or genuine biological sources (e.g., differences in cellular states, distinct biological subpopulations and transcription kinetics). It is crucial, therefore, to properly account for the technical
Fig. 1 Single-cell transcriptome profiling of mouse HSCs by microfluidic system. 

a. Gating strategy for HSC purification. Bcl11a+/+ and Bcl11a−/− HSCs were isolated by sorting markers LSK CD150+48− and sBcl11a+/+. Lineage markers used for enrichment included B220, CD19, CD3, CD4, CD8, TCRγδ, TCRβ, NK1.1, CD11b, Gr-1, Ter119. FSC: Forward scatter, SSC: Side scatter.

b. Single-cell capturing efficiency by the C1 AutoPrep microfluidic system and representative microscopic images of individual capture sites. A representative high-quality single HSC at an individual capture site is indicated by the red arrow. Representative pictures of poor quality cells, an empty capture site and a multiplet capture site are framed in colors as indicated in the key.

c. Principal component analysis of the transcriptome of all 181 HSCs passed initial computational quality control. One significant outlier from the Bcl11a+/+ dataset was identified (marked by red arrow). It was removed from subsequent analysis.

d. Boxplot comparing the number of genes detected (normalized count >1) in the sBcl11a+/+ and Bcl11a−/− datasets. The two datasets were comparable, despite low sequencing depth of the sBcl11a+/+ dataset.
variability in single-cell transcriptomic data interpretation. Brennecke et al. [18] recently described a statistical approach to address this problem by the addition of standardized external RNA spike-ins to the sequencing library. The null hypothesis is that the expression variability detected in a particular gene is not different from the technical variability measured by the external RNA spike-ins; thus, genes that display higher than expected variability imply genuine biological fluctuation from possible cellular subgroups. We identified 6,597, 7,716 and 5503 highly variable genes in the Bcl11a+/−, sBcl11a+/− and Bcl11a−/− datasets, respectively (Fig. 2a; Additional file 2). Remarkably, gene ontology (GO) term enrichment analysis showed that terms related to cell cycle were significantly over-represented in all three datasets (p < 0.0001; Fig. 2b; Additional file 2). This result suggested that cell cycle activity is the dominant source of transcriptomic heterogeneity among HSCs.

Transcriptional reconstruction of cell cycle progression in the HSC compartment

To resolve the cell cycle heterogeneities in the HSC compartment, we focused on the expression pattern of cell cycle-associated genes in the merged Bcl11a+/− HSC dataset (Bcl11a+/− and sBcl11a+/−). We compiled a list of 2212 cell cycle-associated genes based on the annotations in the GO term “cell cycle”, the Cyclebase database and a recent study on G0/G1 transition by Oki et al. [19] (Additional file 3). PCA was performed to transform the high dimensional expression data into individual linearly uncorrelated principal components. Interestingly, we observed that HSCs displayed a characteristic distribution pattern in the first two principal components (PC1/PC2; Fig. 3a). To aid visualization, we grouped HSCs into five smaller clusters (C1–5) (Fig. 3a) based on their proximity in the PCA plot (Fig. 3b). The grouping is largely consistent with hierarchical clustering (Fig. 3c). To understand if the distribution of the HSCs depicted a cell cycle progression trajectory, we inspected the expression pattern of stage-specific genes based on their annotations in the Cyclebase database and from literature [20]. It appeared that genes situated at different spatial domains of the plot showed clustered expression of stage-specific genes. For instance, the C1 clusters showed high expression of the quiescence regulator of Cdkn1c (p57) and G1 arrest factor Txnip [21, 22] and lack of the methylation and G1/S entry regulator Uhrf1 [23] (Fig. 3d). C3 was dominated by S phase-specific genes such as Rrm2 [24] and DNA replication-licensing factors (e.g., Mcm2, Mcm5, Mcm7), while the C4 cluster expressed high levels of genes involved in cell division (e.g., Prc1, Mki67 (Ki-67)) (Fig. 3d). Correspondence of individual clusters to specific cell cycle stages is further supported by the distinct pattern of cyclin and cyclin-dependent kinase expression in specific clusters (Fig. 3d). Cyclins are known to oscillate at specific cell cycle stages when they form complexes with partnering cyclin-dependent kinases to drive cell cycle progression [25]. Cyclin E (Ccene2) regulates G1/S transition with Cdk2 (Cdk2), cyclin A (Ccn2a) is specifically active in S phase, and cyclin B (Ccnb2) functions at G2/M phase transition with CDK1 (Cdk1). Cyclin F expression closely followed that of cyclins A and B [26]. Cyclin D (Cnd2) is first expressed in G1 phase and involved in G1/S transition, and its expression is sustained in proliferative cells. The expression levels of specific cyclin genes therefore provided a good molecular marker to define the cell cycle stages of different cell clusters. Concordant to the clustering pattern in PCA, the cyclin expression patterns across clusters C1–4 closely recapitulated the known cyclin oscillation during cell cycle progression (Fig. 3d). This correlation combined with the expression patterns of Cdkn1c and Txnip allowed us to reconstruct a transcriptionally defined cell cycle and assign the cell cycle stage of C1 (72/119 cells) as G0/early G1 phase, C2 (14/119) as late G1 phase, C3 (21/119) as S phase and C4 (10/119) as G2/M phase (Fig. 3b, d). The status of the C5 clusters is undetermined due to the low number of cells. Interestingly, the fraction of HSCs classified as G0/early G1 phase in both the Bcl11a+/− (28 %, 54/75) and sBcl11a+/− (27.3 %, 32/44) datasets are similar, consistent with our finding that the sorting schemes of LSK CD150−/48 and LSK CD150−/48 34 135 are mostly overlapping (Fig. 1a).

We then applied this classification to explore the molecular differences between the “G0/early G1 phase” C1 and “S phase” C3 clusters. We performed gene set enrichment analysis (GSEA) on gene sets curated in the Kyoto Encyclopedia of Genes and Genomes (KEGG) database. Gene sets related to “cell cycle” and “DNA replication” are significantly enriched as expected (Fig. 3e). In addition, we also detected significant enrichment of gene sets involved in DNA damage repair such as “mismatch repair” and “nucleotide/base excision repair”. This finding is in accordance with recent reports that quiescent adult HSCs accumulate DNA damage and this damage is repaired upon entry into the cell cycle [27–29]. Similarly, enrichment of “p53 signaling pathway” is also detected in the proliferative C3 cluster, consistent with its role in the orchestration of DNA damage repair [30]. Furthermore, gene set enrichment of “oxidative phosphorylation” and “citrate cycle” in the C3 cluster is supported by the observation that quiescent HSCs undergo a glycolytic to aerobic metabolic transition upon activation [31]. Interestingly, gene sets associated with “ubiquitin-mediated proteolysis” is also significantly enriched, affirming the role of the ubiquitin proteasome system in the regulation of HSC cell cycle regulation [32]. These results suggest that
Fig. 2 (See legend on next page.)
transcriptomic cell cycle reconstruction by single-cell RNA-seq data permits direct comparison of the biological characteristics of HSCs at different cell cycle stages in silico.

**Acute Bcl11a deletion in the HSC compartment alters cell cycle progression**

We next examined the HSC compartment when Bcl11a was deleted. Notably, the distribution of Bcl11a−/− HSCs in PCA closely mimicked that of merged Bcl11a+/+ HSCs in terms of cell cycle-related genes (Fig. 4a). Nevertheless, there were more Bcl11a−/− cells (59.0%, 36/61 cells) in the space occupied by the proliferative C3 and C4 clusters of the control Bcl11a+/+ HSCs (26.1%, 31/119) (Fig. 4a), suggesting that Bcl11a deficiency is associated with increased proliferation in the HSC compartment. Moreover, we observed a significant increase of transcription activity in the proliferating Bcl11a+/− HSCs compared with controls, estimated by the total number of ERCC-normalized counts per cell (Fig. 4b). This is consistent with a previous report that transcription rate periodically increases from G1 to S/G2/M phase [33] and suggests that Bcl11a−/− HSCs are more transcriptionally active. There was also evident general up-regulation of cyclin genes and down-regulation of the quiescence regulator Cdkn1c (p57) and G2/M markers such as Prc1, Plk1 and Mki67 (Ki-67) [20] (Fig. 4c). Notably, genes encoding HSC activation and progenitor markers such as Cd34 and Cd244 were also elevated [34, 35]. Moreover, examination of the gene expression correlation of transcription factors revealed that there were two anti-correlative gene clusters (Fig. 4d, cluster I in red and II in blue) in the HSC compartment. Cluster I genes were enriched for G0 cell cycle terms like “interphase”, suggesting their actions in cell cycle regulation (Fig. 4e). Meanwhile, the expression of Bcl11a in cluster II correlated with multiple known HSC quiescence regulators such as Egr1[36], Fos [37], Mecom [38], Meis1 [39], Hif1α [40], and Nrk4l1 [41] (Fig. 4d). Signaling mediators important for HSC quiescence, such as Notch signaling (Hes1) [42], Wnt signaling (Tcf7/ly) [42] and TGFβ signaling (Id1 and Id3) [43] can also be found in cluster II. These suggest that Bcl11a may cooperate with these regulators in maintaining HSC quiescence. To confirm the prediction that Bcl11a−/− HSCs have reduced quiescence, we performed in vivo 5-bromodeoxyuridine (BrdU) staining for cell cycle profiling. As predicted, significantly higher numbers of BrdU− cells were found in either the Bcl11a+/− or Bcl11a−/− HSC compartment compared with the control (Fig. 4f, g). Furthermore, Bcl11a+/− HSC compartment had significantly more proliferative cells compared with haploinsufficient Bcl11a−/− HSCs, indicating a dose-dependent requirement for Bcl11a. These transcriptomic and experimental results demonstrate a cell cycle defect in the Bcl11a−/− HSC compartment.

**Bcl11a-deficient HSCs have defects in long-term self-renewal potential**

The cell cycle defect in Bcl11a-deficient HSCs prompted us to examine their self-renewal potential. We first examined the expression of the self-renewal gene signature by GSEA. We retrieved the signature gene set described by Krivtsov et al. [44] as associated with HSC and leukemic stem cell self-renewal (Additional file 3). We detected a highly significant reduction of expression in Bcl11a+/− HSCs (Fig. 5a). To experimentally test this observation, we transplanted purified Bcl11a+/− or Bcl11a−/− LSKs (CD45.1+) to sublethally irradiated CD45.1+ recipient mice. Donor contribution in the secondary recipient mice had drastically decreased Bcl11a+/− HSCs, both percentage-wise and total cell numbers (Fig. 5c, d), which was further confirmed by using an alternative HSC sorting scheme (LSK CD34+ 135+) (Fig. 5d).

**Bcl11a deletion eliminated lymphoid-competent HSCs and resulted in expansion of two myeloerythroid-restricted subpopulations in the HSC compartment**

We have previously demonstrated experimentally that Bcl11a is essential in the generation of lymphoid progenitors, including lymphoid-primed multipotent progenitors and CLPs, but its effects on HSCs at the single-cell level were unclear [9]. We thus attempted to interrogate the effect of Bcl11a deficiency on the lineage multipotency of HSCs with our single-cell transcriptomic dataset.
Fig. 3 (See legend on next page.)
We compiled a list of genes from the annotation “hematopoiesis and lymphoid organ development” in the GO database and the literature (Additional file 3) and studied their expression in HSCs with PCA. Annotated hematopoietic genes also involved in cell cycle activity were removed from the compilation to reduce their effect on heterogeneity dissection. Notably, significant overlap of Bcl11a+/− and sBcl11a+/− HSCs can be observed in the PCA (Fig. 6a, upper panel). The Bcl11a−/− HSCs are segregated into two uneven subpopulations, with the larger subpopulation partially overlapping the merged Bcl11a+/− (Bcl11a+/− and sBcl11a+/−) HSCs. The smaller and more scattered Bcl11a−/− HSC subpopulation clusters with few sBcl11a+/− HSCs at the right lower quadrant of the loading plot of the first two principal components (Fig. 6a, upper panel).

To aid visualization and downstream comparison, we again grouped the HSCs shown in Fig. 6a into four subgroups (H1–4) based on their locations on the loading plot (Fig. 6a, middle panel). Intriguingly, the expression pattern of hematopoietic genes pointed to a clustered expression of markers associated with specific lineages (Fig. 6b). For instance, expression of mega-erythroid genes such as Epor, Selp, Gp9, and Gfi1b [45, 46] were clustered in subgroups H2 and H3, while myelopoietic genes such as Mpo, Cebpα [47], Gfi1 [48], and Csf1r were concentrated at the H4 subgroup. Interestingly, the known antagonistic regulation between Gfi1 and Gfi1b is recapitulated by the reciprocal exclusion of the two regulators in the H3 and H4 subgroups [49, 50]. Compared with the H1 subgroup, the H3 and H4 subgroups showed reduced expression of lymphoid and HSC activation regulators such as Egr1, Bcl11a, and Jun, but prevalent expression of progenitor markers like Cd34 and Cd48 (Fig. 6b).

Overlaying the cell cycle staging information on the PCA plot of hematopoietic lineage genes (Fig. 6a, lower panel), we found that most of the cells in the H1 subgroup belong to the C1 (G0/early G1 phase) and C2 (late G1 phase) clusters. Wild-type HSCs which clustered with Bcl11a+/− HSCs in the H4 subgroup belong to either C3 (S phase) or C4 (G2/M phase) (Fig. 6a, lower panel).

We then performed GSEA to further compare lineage progenitor signature enrichment between the different subgroups (Fig. 6c, d). The signature gene sets were retrieved from Sanjuan-Pla et al. [13] (Additional file 3) and comparison between Bcl11a+/− and Bcl11a−/− HSCs showed an expected depletion of the lymphoid signature, consistent with our previous report [9] (Fig. 6c). In addition, the H1 subgroup is enriched with the lymphoid and self-renewal signature, while the H3 and H4 subgroups are enriched with the erythroid and granulocyte-macrophage signatures, respectively (Fig. 6d). Further HSC transplantation experiments also confirmed that Bcl11a−/− HSCs, which dominated the H3 and H4 subgroups, are myeloid-restricted, and that the myeloid progenitor compartments (CMPs, GMPs and MEPs) were not affected in both acute Bcl11a deletion mice and primary recipients after Bcl11a deletion (Fig. 6e–h).

Collectively, a trajectory of gradual loss of lymphoid and self-renewal potential can be discerned as wild-type HSCs become activated and proliferative when transiting from the lymphoid-competent H1 subgroup to the mega-erythroid-restricted H3 subgroup or the granulocyte-macrophage-restricted progenitor-like H4 subgroup. More importantly, the existence of sBcl11a+/− HSCs in the H4 subgroup suggest that myeloid-restricted cells similar to Bcl11a−/− HSCs have already existed in the wild-type HSC compartment. The Bcl11a−/− HSCs likely come from expansion or activation of existing myeloid-restricted HSCs in the wild-type compartment after the elimination of lymphoid-competent HSCs by Bcl11a deletion.

Discussion

The hematopoietic system has been an area of intense research because of its paramount clinical potential. The discovery of specific surface markers for purification of HSCs in the adult mouse model allowed us to study the key biological properties of self-renewal and lineage commitment. Recently, increasing evidence from clonal transplantation assays supports the phenotypically defined HSC compartment as being functionally heterogeneous, with subpopulations of platelet-primed or myeloid-restricted HSCs [13, 14]. In this study, we have profiled the transcriptomes of 180 HSCs in the mouse adult HSC compartment and studied the transcriptome structure and the role of Bcl11a in the HSC compartment. We show that single-cell transcriptomics could be used to reconstruct biological processes, dissect...
Fig. 4 (See legend on next page.)
cellular subpopulations in the HSC compartment, and predict stem cell behavior, such as self-renewal capacity and differentiation potential in silico. Reconstruction of cell cycle progression at the single-cell level provides a new avenue to study the molecular characteristics of quiescent and proliferative HSCs without additional staining or sorting. This advantage is illustrated by our demonstration of the direct comparison of quiescent and proliferative HSCs defined in the reconstruction by GSEA. Reported features of proliferative HSCs, such as activation of the DNA damage repair pathway and transition to oxidative metabolism, can be clearly identified from such an exercise. We noted that the fraction of proliferative cells classified by the transcriptomic approach is higher than the estimation by conventional BrdU staining in this study (~27 % versus 12 %). Others have reported an even lower fraction (~2 %) of cycling cells with alternative DNA content staining and Ki67 expression [35]. The discrepancy is not, in fact, unexpected; instead, the transcriptomic approach may provide a more sensitive staging as cell cycle progression requires further translational and post-translational regulation. Changes in the transcriptome are therefore one of the earliest events in entry to the cell cycle. We further extended the use of the reconstruction in predicting cell cycle phenotype in the Bcl11a<sup>−/−</sup> HSC compartment. Comparison of Bcl11a<sup>−/−</sup> and the control datasets predicted reduction of quiescence in the Bcl11a<sup>−/−</sup> HSC compartment, which was experimentally confirmed by BrdU staining. Gene expression correlation analysis revealed a correlative expression pattern of Bcl11a with multiple known quiescence regulators and signaling mediators. The analysis provided a basis for future mechanistic dissection of Bcl11a in the regulation of HSC self-renewal and quiescence.

We have previously shown that Bcl11a<sup>−/−</sup> HSCs are deficient in lymphoid reconstitution [9]. This defect could be caused by blockage of lymphoid commitment at the HSC level or selective elimination of lymphoid-comm petent HSCs. The possibility to dissect subpopulations using single-cell RNA-seq data provides a new avenue to test these hypotheses.

Segregation of HSCs on the basis of hematopoietic gene expression by PCA revealed an interesting separation of Bcl11a<sup>−/−</sup> and Bcl11a<sup>+/+</sup> HSC expression patterns with partial overlap (Fig. 6a, upper panel). To better dissect their hematopoietic characteristics, we subdivided all the HSCs sequenced in this study into four groups (H1–4) based on their location proximity. The H1 subgroup overwhelmingly consisted of Bcl11a<sup>+/+</sup> HSCs. GSEA of the lineage signature showed that they are enriched with the self-renewal signature and lymphoid potential. Overlaying the PCA plot with cell cycle stage information, we found that the lymphoid-competent H1 HSCs contained all the G0/G1 Bcl11a<sup>+/+</sup> HSCs (Fig. 6a, lower panel). In contrast, the H2 subgroups contained both Bcl11a<sup>−/−</sup> and Bcl11a<sup>+/+</sup> HSCs and the Bcl11a<sup>−/−</sup> HSCs in H2 are predominantly proliferative. Few proliferative Bcl11a<sup>+/+</sup> HSCs are also present in the H4 subgroup clustering with Bcl11a<sup>−/−</sup> HSCs. Compared with the H1 lymphoid-competent subgroup, the H3 and H4 subgroups showed marked enrichment of MEP and GMP progenitor signatures (Fig. 6d) and high expression of activation markers such as Cdl48 and Cdl34. The computational exercises and experimental data indicate that, firstly, the lymphoid-competent H1 subgroup is selectively depleted after Bcl11a deletion. Secondly, the dominance of G0/G1 phase HSCs and the self-renewal signature in the H1 subgroup compared with the H2 subgroup is associated with lymphoid competency and HSC self-renewal. Thirdly, the presence of mixed myeloid-restricted Bcl11a<sup>−/−</sup> and Bcl11a<sup>+/+</sup> HSCs in the H2 and H4 subgroups suggests the pre-existence of myeloid-restricted Bcl11a<sup>−/−</sup>-like HSCs in the wild-type compartment. Fourthly, the clear MEP- and GMP-like signature distinction in the H3 and H4 subgroups suggests that the myeloid-restricted Bcl11a<sup>−/−</sup> HSC compartment is constituted of two distinct progenitor-like subpopulations. Finally, the close relationship of the H3 and H4 subgroups with the H2 subgroup suggests that they may arise from activation and expansion of the latter subpopulations after Bcl11a deletion. Based on this evidence, we propose that Bcl11a serves two roles in the HSC compartment: in the

Fig. 4 Bcl11a-deficient HSCs showed significant proliferative changes in the HSC compartment. a Distribution of Bcl11a<sup>−/−</sup> and Bcl11a<sup>+/+</sup> HSCs (purple) and cell cycle reconstructed-Bcl11a<sup>−/−</sup> HSC clusters (C1–C5) in a PCA loading plot of the first two principal components. b Boxplots comparing the transcriptional activity of G0/G1 stage (C1 and C2) and S/G2/M stage (C3 and C4) HSCs in the Bcl11a<sup>−/−</sup> and Bcl11a<sup>+/+</sup> datasets, estimated by the total number of read counts normalized by ERCC size factor per cell. c Violin plots of gene expression of selected cyclin genes, progenitor markers and cell cycle stage-associated genes in Bcl11a<sup>−/−</sup> and Bcl11a<sup>+/+</sup> HSCs. The black dots represent the mean expression for each gene. d Heatmap showing expression correlations of selected transcription regulators in the HSC compartment. Correlation coefficient was calculated by Spearman correlation coefficient and clustering was performed by complete linkage. Gene correlation cluster II (blue) is magnified. e Enriched GO terms in gene correlation cluster I (red in d). f Validation of cell cycle changes in Bcl11a<sup>−/−</sup> HSCs by 5-bromodeoxyuridine (BrdU) staining. The purple box marks the BrdU<sup>−</sup> fraction in the HSC compartment of different genotypes. FSC-A: Forward scatter area. g The dose-dependent changes in BrdU<sup>−</sup> cell number in the HSC compartment in different Bcl11a genotypes. HSCs were harvested and sorted from adult mouse bone marrow 5 days after tamoxifen-induced Bcl11a ablation. *<i>p</i> < 0.05, **<i>p</i> < 0.01, n = 3 mice for each group. Bcl11a<sup>−/−</sup>, CreERT2; Bcl11a<sup>−/−</sup/><sup>lox/lox</sup> (treated with tamoxifen); Bcl11a<sup>−/−</sup>, CreERT2; Bcl11a<sup>−/−</sup/><sup>lox/lox</sup> (treated with tamoxifen). The error bar represented mean ± 1 standard deviation.
maintenance of lymphoid-competent HSCs and in the regulation of HSC self-renewal. Deletion of Bcl11a selectively eliminated lymphoid-competent HSCs, causing the lymphoid deficiency in the Bcl11a−/− compartment. The myeloerythroid restrictive phenotype of Bcl11a−/− HSCs is contributed by the residual surviving myelo-restricted subpopulations in the HSC compartment, which may have limited self-renewal potential.

The detection of Bcl11a−/−-like wild-type HSCs (H2 and H4 subgroups) in the wild-type compartment in our dataset raises interesting questions about their resemblance to the recently reported myeloid-restricted repopulating progenitors in the myeloid bypass model proposed by Yamamoto et al. [14]. In that study, HSCs were purified with a variant sorting scheme (LSK CD150+ CD34−), which selectively enriched myeloid-biased HSCs [5, 11, 12, 14]. Notably, Itga2b (CD41) was also found to be expressed predominantly in the H2 and H3 clusters in our dataset, supporting the mega-erythroid signature enrichment in these cells (Fig. 6b). Nonetheless, we did not observe
Fig. 6 (See legend on next page.)
clustered expression of Vwf in our dataset, which is suggested to mark platelet-primed primitive HSCs in mice [13]. The transcriptomic relationship of these variant HSC compartments with the Bcl11a<sup>−/−</sup> HSCs thus warrants further investigation.

**Conclusions**

We have successfully applied single-cell transcriptomic analysis in reconstructing the cell cycle process and dissecting the molecular heterogeneities of the rare HSC population. Our data reveal the diversity of lineage-biased subpopulations and the effect of Bcl11a in the maintenance of the HSC compartment. Bcl11a is essential in the maintenance of lymphoid-competent HSC survival and self-renewal.

**Materials and methods**

**Mice**

Bcl11a<sup>lox/lox</sup>Rosa26<i>CreERT2</i>/<i>CreERT2</i> and Bcl11a<sup>GFP/+</sup>/<i>GFP</i> mice were previously reported [9]. F1 hybrid wild-type mice from C57BL/6 CD45.1<sup>+</sup> crossed with 129S5 mice were used as recipients for transplantation of bone marrow or LSK cells. All mice used were from colonies maintained at the Research Support Facility of the Wellcome Trust Sanger Institute. Housing and breeding of mice and experimental procedures were according to the UK 1986 Animals Scientific Procedure Act and the Animal Welfare and Ethical Review Body of the Wellcome Trust Sanger Institute.

**Flow cytometry**

Single-cell suspensions were incubated with purified anti–CD16/32 (clone 93) for 10 min on ice to block Fc receptors. Fluorochrome- or biotin-labeled monoclonal antibodies (clones denoted in parentheses) against B220 (RA3-6B2), CD19 (1D3), CD3e (145-2C11), CD4 (RM4-5), CD8α (53–6.7), TCRβ (B20.6), NK1.1 (PK136), CD11b (M1/70), Gr1 (RB6-8C5), Ter119 (TER-119), Sca1 (D7), c-kit (2B8), CD48 (HM48-1), CD150 (TC15-12 F12.2), CD34 (RAM34), and Flt3 (A2F10) were purchased from BD Biosciences, BioLegend or eBioscience. Cells were stained with antibodies on ice for 20 min before washing. Cells were run on a Fortessa (BD Biosciences) or MoFlo XDP (BD Biosciences) and analyzed by Flowjo (Tree Star). For HSC sorting, Lin<sup>−</sup> cells were enriched by Lineage Cell Depletion Kit (Miltenyi Biotec) before antibody staining.

**Cell-cycle analysis**

Mice were given a single intraperitoneal injection of (BrDU (100 μg g<sup>−1</sup> mouse weight) and then maintained on 0.8 mg/ml BrDU in the drinking water for 14 or 20 h before being euthanized. HSCs were double FACS-sorted from mouse bone marrow. BrDU incorporation was measured by flow cytometry using an APC BrDU Flow Kit (BD Pharmingen) according to the manufacturer’s instructions.

**In vivo transplantation assay**

Transplantation assays were performed using the CD45.1/CD45.2 congenic mouse system. The total nucleated bone marrow cells or LSK cells were purified from bone marrow of the flox/flox mice and the control mice (+/+flox) treated with tamoxifen for 4–5 days. The bone marrow cells (1–5 × 10<sup>5</sup>) or sorted LSK cells (2000 cells) were injected with helper CD45.1<sup>+</sup> bone marrow cells (2 × 10<sup>3</sup> cells) into lethally irradiated (2 × 500 cGy) recipient mice (CD45.1<sup>+</sup>) via the tail vein. For the second bone marrow transplantation, bone marrow cells (0.2–1 × 10<sup>6</sup>) were obtained from recipient mice 16 weeks after first transplantation, and transplanted into a second set of lethally irradiated (2 × 500 cGy) CD45.1<sup>+</sup> recipient mice.

**Single-cell RNA-seq library preparation**

Two thousands FACS-sorted Bcl11a<sup>+/+</sup> and Bcl11a<sup>−/−</sup> HSCs pooled from eight and five mice, respectively, were loaded onto two separate 5–10 μm C1 Single-Cell Auto Prep integrated fluidic circuits (Fluidigm) and cell
Read alignment and quality control of dataset

Paired-end reads were aligned to the Mus musculus genome (Ensembl GRCm38.75 assembly) by GSNAP (version 2013-08-14) with the following parameter settings: ‘gsnap -A sam -B 5 -t 8 -n 1 -Q -nofails’ [51]. The quantification of expression was performed by the htseq-count module from the HTSeq package [52] with gene annotation from GTF files (Ensembl GRCm38.75 assembly) with parameter ‘--s no’ in intersection-nonempty mode [53]. Cells with less than 500,000 counts in annotated features, expression of less than 3000 genes or a high percentage (>10 %) of counts mapping to mitochondrial genes were removed from subsequent analysis. Cells with reads detected on the Bcl11a locus chr11:24163146–24165166 were removed from the Bcl11a−/− dataset as they represent incomplete exon 4 deletion after tamoxifen induction. Significant outliers in PCA based on the whole transcriptome were removed from the merged dataset of Bcl11a+/+, sBcl11a+/+ and Bcl11a−/− HSCs (Fig. 1d).

Detection of highly variable genes and statistical analysis

The details of the statistical model for testing highly variable genes have been described by Brennecke et al. [18]. We set the minimal biological dispersion parameter at 0.5. PCA was performed with function prcomp() in the stats package of R (version 3.0.2 — “Frisbee Sailing”) based on log2 transformed count matrix by log2(n + 1).

GO enrichment analysis

GO term enrichment analysis was performed by DAVID Bioinformatics Resources 6.7 [54] with the default mouse genome as background [55].

Gene set enrichment analysis

GSEA was performed with the javaGSEA application (version 2.0.14) available online [56] with default settings [57]. The lineage-specific gene sets (MkP, PreCFU-E, PreGM, PreMEGE, CLP) were retrieved from a previously published dataset by Sanjuan-Pla et al. [13]. The self-renewal signature gene set was retrieved from Krivtsov et al. [44]. The gene identifiers were remapped by BioMart [58] to official Ensembl gene symbols. A normalized count matrix from DESeq2 was supplied as the input expression dataset. Enrichment is considered significant if the false discovery rate is below 0.1 and the nominal p value is below 0.05.

Statistical analysis

Experimental data are shown as mean and standard deviation. All statistical analyses were either conducted with Prism (GraphPad) or as specified in relevant sections.

Additional files

| Additional file 1: Table S1. Quality control summary of the Bcl11a+/-, sBcl11a+/- and Bcl11a-/- HSC dataset. (XLSX 64 kb) |
| Additional file 2: Table S2. The list of highly variable genes in the Bcl11a+/-, sBcl11a+/- and Bcl11a-/- HSC dataset and the gene ontology term (biological processes) enrichment analysis results. (XLSX 3257 kb) |
| Additional file 3: Table S3. The list of annotated cell cycle genes and hematopoietic regulators used for PCA and the gene sets used for GSEA in this study. (XLSX 126 kb) |

Abbreviations
BrdU: 5-bromo-2′-deoxyuridine; CLP: common lymphoid progenitor; CMP: common myeloid progenitor; ERCC: External RNA Controls Consortium; FACS: fluorescence-activated cell sorting; GFP: green fluorescent protein; GMP: granulocyte–macrophage progenitor; GO: gene ontology; GSEA: gene set enrichment analysis; HSC: hematopoietic stem cell; LSK: Lin- Sca1+ Kit+; MEP: megakaryocyte–erythroid progenitor; PNA: principal component analysis.
Competing interests
The authors declare that they have no competing interests.

Authors' contributions
JT performed single-cell experiments, constructed sequencing libraries, performed bioinformatic analyses, flow cytometry experiments, and wrote the manuscript. YY performed flow cytometry experiments, serial transplantation experiments, cell cycle analysis and wrote the manuscript. SB performed serial transplantation experiments. FB assisted in computational analysis. CW provided assistance in experiments. AK advised on bioinformatics. ST provided reagents and intellectual inputs. LL provided intellectual input. PL conceived and supervised the study and wrote the manuscript. All authors read and approved the final manuscript.

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References
1. Orkin SH, Zon Li. Hematopoiesis: an evolving paradigm for stem cell biology. Cell. 2008;132:631–44.
2. Spangrude GJ, Heimfeld S, Weissman IL. Purification and characterization of mouse hematopoietic stem cells. Science. 1988;241:58–62.
3. Ogawa M, Matsuoka Y, Nishikawa S, Hayashi S, Kuniyoshi T, Sudo T, et al. Expression and function of c-kit in hematopoietic progenitor cells. J Exp Med. 1991;174:63–71.
4. Christensen JL, Weissman IL. Flk-2 is a marker in hematopoietic stem cell differentiation: a simple method to isolate long-term stem cells. Proc Natl Acad Sci U S A. 2001;98:14541–6.
5. Kiel MJ, Milham OH, Iwashita T, Yilmaz OH, Terhorst C, Morrison SJ. SLAM family receptors distinguish hematopoietic stem and progenitor cells and reveal endothelial niches for stem cells. Cell. 2005;121:1109–21.
6. Kiel MJ, He S, Ashkenazi R, Gentry SN, Teta M, Kucher JA, et al. Haematopoietic stem cells do not asymmetrically segregate chromosomes or retain BrdU. Nature. 2007;449:238–42.
7. Gottgens B. Regulatory network control of blood stem cells. Blood. 2015;125:2614–20.
8. Eaves CJ. Hematopoietic stem cells: concepts, definitions, and the new reality. Blood. 2015;125:2605–13.
9. Yu Y, Wang J, Khaled W, Burke S, Li P, Chen X, et al. Bcl11a is essential for lymphoid development and negatively regulates p53. J Exp Med. 2012;209:2467–83.
10. Liu L, Liu P, Kellermans T, Tressauro L, Rachel RA, Nakamura T, et al. Bcl11a is essential for normal lymphoid development. Nat Immunol. 2003;4:525–32.
11. Morita Y, Ema H, Nakauchi H. Heterogeneity and hierarchy within the most primitive hematopoietic stem cell compartment. J Exp Med. 2010;207:1173–82.
12. Gekas C, Graf T. CD41 expression marks myeloid-biased adult hematopoietic stem cells and increases with age. Blood. 2013;121:4643–72.
13. Sanjuan-Pla A, Macaulay IC, Jensen CT, Wol P, Luis TC, Mead A, et al. Platelet-biased stem cells reside at the apex of the haematopoietic stem-cell hierarchy. Nature. 2013;502:232–6.
14. Yamanoto R, Morita Y, Ooebara H, Hamanaka S, Onodera M, Rudolph KL, et al. Clonal analysis unveils self-renewing lineage-restricted progenitors generated directly from hematopoietic stem cells. Cell. 2013;154:1112–26.
15. Benz C, Copley MR, Kent DG, Wohrer S, Cortes A, Aghaeepour N, et al. Hematopoietic stem cell subtypes expand differentially during development and display distinct lymphopoietic programs. Cell Stem Cell. 2012;10:273–83.
16. S Milo, Westermann M, Gorka SA, Vogel J. Single-cell RNA-seq advances and future challenges. Nucleic Acids Res. 2014;42:8845–60.
17. Kolodziejczyk AA, Kim JK, Svensson V, Marioni JC, Teichmann SA. The technology and biology of single-cell RNA sequencing. Mol Cell. 2015;58:610–20.
18. Brennecke P, Anders S, kim JK, Kolodziejczyk AA, Zhang X, Prospero V, et al. Accounting for technical noise in single-cell RNA-seq experiments. Nat Methods. 2013;10:1093–5.
19. Oki T, Nishimura K, Kitaura J, Togami K, Maehara A, Iizawa K, et al. A novel cell-cycle-indicator, mVenus-p27KIC, identifies quiescent cells and visualizes G0-G1 transition. Sci Rep. 2014;4:4012.
20. Santos A, Wernersson R, Jensen LJ. Cyclebase 3.0: a multi-organism database on cell-cycle regulation and phenotypes. Nucleic Acids Res. 2015;43:D1140–4.
21. Matsumoto A, Takeishi S, Kanie T, Suzuki E, Onoyama L, Tateshi Y, et al. p57 is required for quiescence and maintenance of adult hematopoietic stem cells. Cell Stem Cell. 2019;12:262–71.
22. Zou P, Yoshihara H, Hosokawa K, Tai I, Shinmyozu K, Tsukahara F, et al. p57(Kip2) and p27(Kip1) cooperate to maintain hematopoietic stem cell quiescence through interactions with Hsc70. Cell Stem Cell. 2011;9:247–61.
23. Bonapace IM, Latella L, Papait R, Nacciso F, Sacco A, Muto M, et al. Np95 is regulated by E1A during mitotic reactivation of terminally differentiated cells and is essential for S phase entry. J Cell Biol. 2002;157:909–14.
24. Engstrom Y, Eriksson S, Jildevik I, Skog S, Thelander L, Tribukait B. Cell cycle-dependen
type of mammalian ribonucleotide reductase. Differential regulation of the two subunits. J Biol Chem. 1985;260:9114–6.
25. Satyanarayana A, Kaldis P. Mammalian cell-cycle regulation: several Cdkds, numerous cyclins and diverse compensatory mechanisms. Oncogene. 2009;28:2925–39.
26. Bai C, Richman R, Elledge SJ. Human cyclin F. EMBO J. 1994;13:6087–98.
27. Beerman I, Seita J, Inlay MA, Weissman IL, Rossi DJ. Quiescent hematopoietic stem cells accumulate DNA damage during aging that is repaired upon entry into cell cycle. Cell Stem Cell. 2014;5:357–50.
28. Flach J, Bakker ST, Mohrin M, Corvec PC, Petras EM, Reynaud D, et al. Replication stress is a potent driver of functional decline in ageing haematopoietic stem cells. Nature. 2014;512:198–202.
29. Walter D, Lier A, Geiselhart A, Thalheimer FB, Hunschta S, Sobotta MC, et al. Exit from dormancy provokes DNA-damage-induced attrition in haematopoietic stem cells. Nature. 2015;520:549–52.
30. Asai T, Liu Y, Bae N, Nimer SD. The p53 tumor suppressor protein regulates hematopoietic stem cell fate. J Cell Physiol. 2011;226:215–21.
31. Suda T, Takubo K, Semenza GL. Metabolic regulation of hematopoietic stem cells in the hypoxic niche. Cell Stem Cell. 2011;9:298–310.
32. Moran-Crusio K, Reavie LB, Aifantis I. Regulation of hematopoietic stem cell fate by the ubiquitin proteasome system. Trends Immunol. 2012;33:357–63.
33. Zopf CJ, Quinn K, Zeidman J, Maheshri N. Cell-cycle dependence of transcription dominates noise in gene expression. PLoS Comput Biol. 2013;9:e1003161.
34. Oguro H, Ding L, Morrison SJ. SLAM family markers resolve functionally distinct subpopulations of hematopoietic stem cells and multipotent progenitors. Cell Stem Cell. 2013;13:102–16.
35. Wilson A, Laurenti E, Oser G, van der Wath RC, Blanco-Bose W, Jaworski M, et al. Hematopoietic stem cells reversibly switch from dormancy to self-renewal during homeostasis and repair. Cell. 2008;135:1118–29.
36. Min IM, Pietramaggiori G, Kim FS, Passegue E, Stevenson KE, Wagers AJ. The transcription factor EGR1 controls both the proliferation and localization of hematopoietic stem cells. Cell Stem Cell. 2008;2:380–91.
37. Venezia TA, Merchant AA, Ramos CA, Whitehouse NL, Young AS, Shaw CA, et al. Molecular signatures of proliferation and quiescence in hematopoietic stem cells. PLoS Biol. 2004;2:e301.
38. Zhang Y, Stehling-Sun S, Lezon-Geyda K, Juneja SC, Coillard L, Chatterjee G, et al. PR-domain-containing Mds1-Evi1 is critical for long-term hematopoietic stem cell function. Blood. 2011;118:3853–61.

39. Unnisa Z, Clark JP, Roychoudhury J, Thomas E, Tessarollo L, Copeland NG, et al. Meis1 preserves hematopoietic stem cells in mice by limiting oxidative stress. Blood. 2012;120:4973–81.

40. Gazit R, Garrison BS, Rao TN, Shy T, Costello J, Ericson J, et al. Transcriptome analysis identifies regulators of hematopoietic stem and progenitor cells. Stem Cell Rep. 2013;1:266–80.

41. Land RH, Rayne AK, Vanderbeck AN, Barlowe TS, Manjunath S, Gross M, et al. The orphan nuclear receptor NRAA1 specifies a distinct subpopulation of quiescent myeloid-biased long-term HSCs. Stem Cells. 2015;33:278–88.

42. Duncan AW, Rattis FM, DiMascio LN, Congdon KL, Pazianos G, Zhao C, et al. Integration of Notch and Wnt signaling in hematopoietic stem cell maintenance. Nat Immunol. 2005;6:314–22.

43. Perry SS, Zhao Y, Nie L, Cochrane SW, Huang Z, Sun XH. Id1, but not Id3, directs long-term repopulating hematopoietic stem-cell maintenance. Blood. 2007;110:2351–60.

44. Kvitko AV, Twomey D, Feng Z, Stubbs MC, Wang Y, Faber J, et al. Transformation from committed progenitor to quiescent stem cell initiated by MLL-AF9. Nature. 2006;442:818–22.

45. Osawa M, Yarnaguchi T, Nakamura Y, Kaneko S, Onodera M, Sawada K, et al. Erythroid expansion mediated by the Gfi-1B zinc finger protein: role in normal hematopoiesis. Blood. 2002;100:2769–77.

46. Saleque S, Cameron S, Orkin SH. The zinc-finger proto-oncogene Gfi-1b is essential for development of the erythroid and megakaryocytic lineages. Genes Dev. 2002;16:301–6.

47. Zhang DE, Zhang P, Wang ND, Hetherington CJ, Darlington GJ, Tenen DG. Absence of granulocyte colony-stimulating factor signaling and neutrophil development in CCAAT enhancer binding protein alpha-deficient mice. Proc Natl Acad Sci U S A. 1997;94:5697–71.

48. Moignard V, Macaulay IC, Swiers G, Buettner F, Schutte J, Calero-Nieto FJ, et al. Characterization of transcriptional networks in blood stem and progenitor cells using high-throughput single-cell gene expression analysis. Nat Cell Biol. 2013;15:363–72.

49. Wu TD, Nacu S. Fast and SNP-tolerant detection of complex variants and splicing in short reads. Bioinformatics. 2010;26:873–81.

50. javaGSEA. http://www.broadinstitute.org/gsea/downloads.jsp.

51. Subramanian A, Tamayo P, Mootha VK, Mukherjee S, Ebert BL, Gillette MA, et al. Gene set enrichment analysis: a knowledge-based approach for interpreting genome-wide expression profiles. Proc Natl Acad Sci U S A. 2005;102:15545–50.