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Gene set control analysis predicts hematopoietic control mechanisms from genome-wide transcription factor binding data

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Transcription factors are key regulators of both normal and malignant hematopoiesis. Chromatin immunoprecipitation (ChIP) coupled with high-throughput sequencing (ChIP-Seq) has become the method of choice to interrogate the genome-wide effect of transcription factors. We have collected and integrated 142 publicly available ChIP-Seq datasets for both normal and leukemic murine blood cell types. In addition, we introduce the new bioinformatic tool Gene Set Control Analysis (GSCA). GSCA predicts likely upstream regulators for lists of genes based on statistical significance of binding event enrichment within the gene loci of a user-supplied gene set. We show that GSCA analysis of lineage-restricted gene sets reveals expected and previously unrecognized candidate upstream regulators. Moreover, application of GSCA to leukemic gene sets allowed us to predict the reactivation of blood stem cell control mechanisms as a likely contributor to LMO2 driven leukemia. It also allowed us to clarify the recent debate on the role of Myc in leukemia stem cell transcriptional programs. As a result, GSCA provides a valuable new addition to analyzing gene sets of interest, complementary to Gene Ontology and Gene Set Enrichment analyses. To facilitate access to the wider research community, we have implemented GSCA as a freely accessible web tool (http://bioinformatics. cscr.cam.ac.uk/GSCA/GSCA.html). © 2013 ISEH - Society for Hematology and Stem Cells. Published by Elsevier Inc.

Cell type–specific gene expression is an inherent property of all multicellular organisms and indeed represents a major determinant that underlies the generation of differentiated cell types with distinct functionality. Elucidating the molecular mechanisms controlling cell type–specific expression has the power to reveal fundamental insights into the regulatory circuitry controlling both human and model organism development. Moreover, identification of control mechanisms in normal cells provides potential avenues for manipulating cellular fates, as exemplified by the recent explosion in cellular reprogramming studies [1]. It also enables the rational design of new therapies aiming to revert abnormal pathological cellular states back to their normal condition [1].

The blood or hematopoietic system has long been recognized as a powerful model system for studying cell type– specific gene expression [2]. Within the blood system, more than 10 distinct mature hematopoietic lineages (e.g., red blood cells, T cells, B cells) are generated from pluripotent hematopoietic stem cells (HSCs) via a sequence of intermediate progenitors, often represented as a lineage differentiation tree. Both the mature lineages as well as the various immature blood stem and progenitor populations can be purified based on the expression of combinations of specific cell surface markers, thus enabling powerful studies of cellular differentiation.

Transcription factors have long been recognized as major regulators of hematopoietic cell type specification [3–6]. To understand the mechanisms underlying cell type specification by transcription factors, it will be essential to identify their transcriptional targets. An important advancement in this research area was provided by the introduction of chromatin immunoprecipitation (ChIP) coupled to massively parallel sequencing (ChIP-Seq),

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which allows genome scale identification of all DNA sequences (regions) bound by a given transcription factor (TF) in a given cell type [7]. The technique has been rapidly adopted with over 100 individual studies now deposited in public databases for the murine hematopoietic system alone. This wealth of new data represents unprecedented opportunities to unravel the transcriptional control mechanisms that mediate expression of specific sets of genes within the various hematopoietic cell lineages [8].

Gene ontology [9] overrepresentation analysis provides information on various types of functional categories enriched within a given gene set of interest [10] and GSEA determines whether a gene set of interest shows statistically significant expression differences between two or more cell types [11]. However, neither of these approaches explicitly links a gene set to transcriptional control mechanisms. In this study, we report a new computational framework for linking gene sets with transcriptional control, called Gene Set Control Analysis (GSCA). Unlike previous algorithms developed to provide functional enrichment [10], GSCA links gene sets to likely upstream regulators responsible for coordinated expression. By exploiting multiple transcription factor binding patterns from genome-wide ChIP-Seq studies, GSCA can provide previously unattainable insights into possible transcriptional control mechanisms operating in both normal and malignant cells. To gain insights into combinatorial control mechanisms (i.e. multiple transcription factors occupying the same binding site in a gene locus), we further developed a novel tool called combinatorial-GSCA (C-GSCA). Through integrated analysis of 142 blood-specific ChIP-Seq binding datasets, C-GSCA identifies likely combinatorial transcriptional control mechanisms by revealing TF cooccupancy patterns specifically associated with gene regulatory elements from a given gene set. A web-based implementation of GSCA and C-GSCA allows user-friendly access for the wider research community, and thus provides a substantial new addition to the bioinformatic toolbox for hematopoietic gene set analysis.

Methods

ChIP-seq compendium

Binding events for 35 transcription factors in seven major hematopoietic lineages were obtained from Hannah et al. [8]. Sixty new ChIP datasets from 18 publications and ENCODE murine datasets were analyzed, starting from the raw data set in each case, and peaks were identified in each sample using the protocol described previously [8]. A supplementary website (http://bioinformatics. cscr.cam.ac.uk/BLOOD_compendium_PUBLISHED.html) lists the number of peaks, reference, and peak calling method for each of the ChIP dataset. All binding events were mapped to genes using the same protocol described previously [12]. Binding events in the promoter and gene body were associated to the corresponding gene, whereas intergene peaks were associated to the nearest gene on either side within 50 kb, such that each peak is assigned to at most two genes.

Tissue-specific enhancer elements in mouse were downloaded from [13] and p value was calculated for overlap between each of the 61 tissue-specific enhancer regions and blood-specific regulatory regions [8] using a hyper-geometric test (Supplementary Table 1, online only, available at www.exphem).

GSCA method

Of 270,261 genomic regions bound by at least one TF (N), for a set of user-defined genes, we calculate the number of genomic regions mapped to the genes (n). For each ChIP-Seq ChIP dataset, the number of peaks (m) near user defined genes (k) is calculated. The p value is calculated using a hypergeometric test (Fischer exact test).

cGSCA method

A matrix of binding events with 270,261 genomic regions as rows and overrepresented ChIP-seq data sets (K) from GSCA step as columns is generated. The ChIP-seq data sets (K columns) are then clustered using a hierarchic clustering with Pearson's correlation coefficient as a distance measure.

Reference data set

Gene sets for 80 clusters of tightly coexpressed genes (their induction patterns) in 38 hematopoietic cell types were obtained from Novershtern et al. [14]. Human genes were mapped to orthologous mouse genes using MGI mammalian orthology (http://www.informatics.jax.org/orthology.shtml). We calculated the p value for each gene set with respect to each signature cluster using a hypergeometric test. We used the number of Novershtern clusters significantly overrepresented (Bonferroni corrected p < 0.001) for one or more transcription factor targets as a measure to evaluate performance while comparing different methods.

Gene expression datasets

Nine gene expression signatures (d-erythroid, differentiated, d-lymphoid, d-myeloid, r-myelolymphoid, s-erythroid, s-mpp, s-myelolymphoid, and stem) were obtained from [15]. Differentially expressed genes in various leukemia datasets were downloaded from their respective publications. Gene lists were then interrogated against the ChIP-seq compendium using both GSCA and C-GSCA.

GSCA web tool

The GSCA output was produced using R, and the web user interface of the application was done using Perl/CGI/HTML. R commands are executed through the perl–cgi script to produce the image. The web tool can be accessed at the following URL: http://bioinformatics.cscr.cam.ac.uk/GSCA/GSCA.html.

Results

Definition of a candidate regulatory genome in mouse hematopoiesis

We recently reported a compendium of more than 50 TF ChIP-Seq experiments in mouse blood cells collected from publicly available datasets [8]. We have doubled the compendium by adding 60 new ChIP datasets from 18 recently published studies [16–33] and ENCODE murine unpublished datasets to obtain genome-wide binding

patterns for 53 unique transcription factors in 15 major blood lineages and three types of leukemia (Table 1). TFbound peaks were determined for all new datasets using the same parameters as before [8], which resulted in a total of 270,261 genomic regions bound by at least one transcription factor. When added together, these 270,261 regions corresponded to 936 Mb, thus constituting 5.78% of the mouse genome. ChIP-Seq samples of the same transcription factor in related cell types were merged together to provide a consolidated set of 78 samples (Table 1).

Pennacchio et al. [13] developed a phylogenetic conservation and motif based approach to predict tissue specific enhancers, which allowed them to annotate ~5,500 high-confidence mouse tissue-specific enhancers for 61 murine tissue types by integrating tissue-specific expression data, conservation information, and cis-regulatory motifs. Only 4 of these 61 tissues corresponded to hematopoietic cells, and predicted only enhancers for those four tissues showed significant overlap with our ChIP-enriched regions (B220⁺ B cells, p = 1.9e-10; CD4⁺ T cells, p = 1.4e-4; CD8⁺ T cells, p = 7.0e-7; lymph node, p = 1.0e-4; see Supplementary table 1). This analysis therefore supports the validity of a compendium built on TF binding events in hematopoietic cells.

A new GSCA tool matches weighted TF-peak lists to gene sets

We next explored whether our blood-specific TF ChIP-Seq peak catalogue could be used to predict transcriptional control mechanisms that may regulate the coordinated expression of a given set of genes. Computational tools for the identification of statistically significant overlaps between a given gene set and peak regions from single ChIP-Seq experiments have been described previously [34,35]. However, these tools do not exploit the ever-increasing number of datasets for multiple TFs in the same or related cell types.

Novershtern et al. [14] reported gene expression profiles in 38 distinct purified populations of human hematopoietic cells ranging from hematopoietic stem cells, through multiple progenitor and intermediate maturation states, to 12 terminally differentiated cell types. Using the Module Networks algorithm [36], they identified 80 modules or gene sets of tightly coexpressed genes with distinct expression patterns and enrichment for specific biological functions, which they termed induction patterns. When we used the 80 Novershtern modules as gene sets, 37 of 80 gene sets (Supplementary Table 2, online only, available at www.exphem) showed a statistically significant correlation with one or more TF peak files from our compendium when using the previously described ChIP Enrichment Analysis (ChEA) [34] and Csan [35] tools. Of note, there was a good overlap between the cell type used for ChIP-Seq and the expression/induction patterns as annotated by Novershtern et al. (Supplementary Table 2). For example, gene set 727 with induction pattern "Late Erythroid" was associated with Eto2 in Erythroid, Scl, and Ldb1 in HSCs and Scl in MELs, and gene set 979 overrepresented for "immune response" genes with induction pattern "Late MYE" was associated with Cebpa, Cebpβ, P65, Pparg, and Stat1 in macrophages.

Because the ChEA [34] and Csan [35] tools could associate candidate upstream regulators to less than half of the 80 Novershtern gene sets, we set out to develop an alternative approach by incorporating the concept of weighted

Table 1. Seventy-eight ChIP-Seq binding peak files covering 53 unique transcription factors in 15 major blood lineages

Cell type	Transcription factors
Lymphocytes	
B cells	E2A, Ebf, Foxo1, Oct2, Pax5, Pu.1
T cells	Gata3, Fli1, Pu.1, Stat3, Stat4, Stat5, Stat5a, Stat5b, Stat6, Tbet
Thymocytes	Cbfb, Rag2, Ring1b, Runx1
Progenitors	
HPC	Gata2, Ldb1, Scl
HPC7	Erg, Fli1, Gata2, Gfi1b, Lmo2, Meis1, Pu.1, Lyl1, Runx1, Scl
EML	Runx1, Tcf7
Erythroid progenitors	Gata1, Gata2, Smad1
MK progenitors	Cbfb, Ring1b, Runx1
Myeloid progenitors	Myb
Pro B cells	Ebf1, Smad1
Myeloerythroid	
MK (megakaryocytes)	Gata1
Macrophages	Cebpα, Cebpβ, P65, Pparg, Pu.1, Stat1
Erythroid	Eto2, Gata1, Ldb1, Mtgr1, Pu.1, Scl
Leukemias	
Leukemia	Notch1
MLL leukemia	Af9
T cell leukemia	RbpJ
T-ALL	Notch1
MEL	Cmyb, Cmyc, Chd2, Gata1, JunD, MafK, Max, Mxi1, NelfE, Scl, Smc3, Tbp, Usf2

peak-to-gene mapping recently reported as part of the Genomic Regions Enrichment of Annotations Tool (GREAT) [37]. GREAT links a list of ChIP-Seq peak regions to gene lists with particular functional significance and unlike previous approaches incorporates binding sites not only in the promoter region of a gene. Taking inspiration from this approach, we developed a new tool by mapping each peak to its nearest gene within 50 kb and then considering the number of binding events in each gene locus to calculate the significance of association between a gene locus and a given upstream regulator. (Essentially this is the reverse of GREAT, which associates peaks with genes, whereas our new procedure associates genes with peaks). Specifically, our new tool determines the number of binding events in the loci of genes of interest for each ChIP dataset (Fig. 1A, red arrows), and then calculates a p value using a simple hypergeometric test. Datasets with statistically significant overlaps (corrected p value cutoff < 0.001) are then selected by interrogating all ChIP datasets independently against the gene list (Fig. 1B). When applied to the 80 gene modules from Novershtern et al. [14], our new tool reported significant associations with ChIP-Seq peaks for 65 gene modules (Supplementary Table 2, online only, available at www.exphem), which corresponds to 81% of all gene sets compared with only 46% using the previously reported ChEA and Cscan tools. Incorporation of weighted gene lists therefore results in a significant increase in the percentage of gene modules that can be linked to candidate upstream regulators. We named this new approach Gene Set Control Analysis, or GSCA. Only 61% of all Novershtern gene sets (49 of 80 gene modules) were enriched when the binding events only in promoters were selected, thus highlighting the likely importance of binding to nonpromoter regions, which compose 57% of all binding events in our datasets.

GSCA correlates relevant combinations of transcription factors with hematopoietic gene sets

To investigate the potential biological relevance of the candidate upstream regulatory transcription factors matched with the 65 Novershtern gene sets by GSCA, we again used the induction patterns defined by Novershtern et al. as a measure of lineage-specific expression. The majority of gene sets (97%) showed good correspondence between the induction patterns and the cell types in which the TFs had been chipped (Supplementary Table 3, online only, available at www.exphem).

For example, gene sets 667 and 829 (enriched for T cell receptor activity) were associated by GSCA with Stats and Gata3 in T cells, whereas gene sets 649 and 961 (enriched for B cell receptor activity) were associated with Pu.1, E2A, and Pax5 in B cells. Gene set 721 (involved in inflammatory and antibacterial response) was linked by GSCA with Cebp α , Cebp β , P65, Pu.1, and Stat1 in macrophages. Gene sets 727 and 889 with Late Ery induction pattern (en-

riched for protein amino acid glycosylation and blood group antigen functional annotations) significantly overlapped only with targets of Eto2, Gata2, Ldb1, Mtgr1, and Scl in erythroid cells. Taken together therefore, there is good concordance between the induction patterns of Novershtern gene sets and the matching ChIP-Sequencing TF datasets identified by GSCA.

Combinatorial regulatory pattern discovery from multi factor ChIP-Seq data

Compared with previous tools, our new GSCA tool performs better by associating gene lists with ChIP-Seq peaks by calculating weighted associations between factors and genes based on the number of binding events within a gene locus. However, all individual ChIP-Seq datasets are treated independently, thus making it difficult to infer whether two overrepresented transcription factors work combinatorially (e.g. whether they show statistically significant co-occupancy of the same regulatory regions), rather than binding to overlapping sets of gene loci, but using distinct *cis*-regulatory regions. To address this issue of combinatorial binding, we developed a new tool called combinatorial GSCA (C-GSCA), and then applied this new tool to our hematopoietic ChIP-Seq compendium. For a given gene list, we first run GSCA to select the TFs showing overrepresented binding. Assuming that m TFs are selected out of 78 ChIP-seq datasets, we generate a binary matrix $(n \times m)$ of m columns representing the m ChIP datasets and n rows representing the genomic regions occupied by two or more of the *m* TFs, with 1s and 0s indicating the presence or absence of binding, respectively. We filter genomic regions bound by only one factor ($\sim 16\%$ of genomic regions; Supplementary Table 2, online only, available at www.exphem) because they are not informative in terms of combinatorial control mechanisms. We then perform hierarchical clustering of *n* overrepresented ChIP datasets using Pearson's correlation coefficient as a distance measure. Unlike GSCA, all overrepresented ChIP datasets are considered together, making the prediction of combinatorial control feasible (Fig. 2).

Using ChIP-Seq analysis of 10 transcription factors in the hematopoietic progenitor cell line HPC7, we have shown previously that combinatorial interactions between a heptad of TFs (SCL, LYL1, LMO2, GATA2, RUNX1, ERG, and FLI-1) were overrepresented in the loci of genes specifically expressed in HSPCs and therefore associated with gene sets specifically expressed in HSCs [12]. When the heptad-bound genes were interrogated using GSCA, 49 of 78 ChIP-Seq datasets were enriched, thus identifying multiple new transcription factors as candidate upstream regulators in addition to the seven factors (Supplementary Figure 1, online only, available at www.exphem). Using C-GSCA, these 49 datasets could be split into four cell type-specific groups of T cells, macrophages, HSCs, and erythroid (Supplementary Figure 1, online only, available at www.exphem). This observation suggests that gene loci bound by the heptad in blood



Figure 1. Schematic representation of the Gene Set Control Analysis (GSCA) protocol. For a given gene set of interest (red arrows), the number of peaks in gene loci is determined and a p value is calculated using a hypergeometric test. The TFs from overrepresented ChIP datasets (corrected p < 0.001, yellow bars in the figure) are then reported as candidate upstream transcriptional regulators. (For interpretation of the reference to color in this figure legend, the reader is referred to the web version of this article.)

stem and progenitor cells not only include genes specifically expressed in HSCs, but could also include a subset of genes affiliated with various different hematopoietic differentiation programs—an observation that would be consistent with the concept of lineage priming developed in the 1990s [38]. These results suggested that the C-GSCA procedure outlined here may be useful more generally to associate hematopoietic gene sets to upstream regulators and thus able to predict combinatorial control mechanisms driving the expression of a given gene set. We next applied the new C-GSCA tool to all 80 hematopoietic gene sets from the Novershtern et al. study [14], which allowed us to associate 65 of the 80 Novershtern gene sets overrepresented for ChIP datasets using GSCA for combinatorial TF signatures. For example, Novershtern gene set 583 with induction pattern "Late Ery + T/B cell + GRAN" is associated with entirely different sets of transcription factors in two different cell types, because it was linked with Gata1, Gata2, Scl, and Smad1 in erythroid progenitors, and Rag2 in thymocytes, Max,



Figure 2. (A) Schematic representation of combinatorial Gene Set Control Analysis (cGSCA). A binary matrix of combinatorial binding patterns is generated using the overrepresented ChIP datasets from GSCA. (B) A hierarchical tree is then generated by clustering similar patterns. Color figure online.

Mxi1, and Tbp in mouse erythroleukemia (MEL) (Fig. 3A). Similarly, gene set 745 with induction pattern "NK + T cell" is linked with Myb in myeloid progenitors and Stat3, Stat4, and Stat5 in T cells (Fig. 3B). Indeed, more than 60% (40 of 65) of the overrepresented Novershtern gene sets with matched upstream regulators were linked with more than one combinatorial pattern (Supplementary Table 3, online only, available at www.exphem). Therefore, unlike the GSCA approach (Fig. 1), C-GSCA has the potential to identify distinct subsets of candidate upstream regulators for a given gene set (Fig. 2).

GSCA web tool

As GSCA and C-GSCA provide potentially powerful ways of predicting candidate upstream regulators for a given list, we

developed a web tool to facilitate gene set control analysis for the wider community (http://bioinformatics.cscr.cam.ac.uk/ GSCA/GSCA.html). In this section we provide a brief explanation of the functionality of the GSCA web tool using a recent transcriptome analysis of murine HSCs and early multipotent, bipotent, and unipotent progenitors [15], which reported nine gene expression signatures ranging from those characteristic for the most immature HSCs to those affiliated with differentiation into the individual hematopoietic lineages. We interrogated these nine experimentally obtained gene expression signatures using the GSCA web tool. Eight of these nine mouse stem–progenitor gene signatures showed significant overlap with multiple ChIP-Seq data sets, thus providing an independent test case to examine the biological relevance of predicted combinatorial regulatory signatures in



Figure 3. (A) Overrepresented regulators determined using GSCA (left) and C-GSCA (right) for gene module 583 from Novershtern et al. [14], with "Late Ery + T/B cells + GRAN" induction pattern. Unlike GSCA, C-GSCA can separate overrepresented independent binding patterns in different cell types (Gata1, Gata2, and Smad1 Erythroid progenitors and Max, Mxi1, and Tbp in MELs in this case).

addition to testing the functionality of the web tool (Supplementary Figure 3, online only, available at www. exphem). Figure 4A shows a screenshot of the web tool in which users can paste a query gene list or upload it from a file (human or mouse).

Upon choosing GSCA, a gene list of interest is interrogated against 78 ChIP-Seq datasets across 15 blood cell types. GSCA calculates the significance of overlap between each ChIP-Seq dataset and the gene set of interest and displays all ChIP-Seq datasets, with those showing enrichment in yellow color. For example, the self-renewing signature (stem signature from Ng et al. [15]) is provided as a test dataset for the users and shows statistically significant overlap with multiple transcription factors in HPC7 and progenitors. When the same stem signature gene list is analyzed using C-GSCA, the overrepresented ChIP datasets are clustered into two distinct cell type specific clusters HPC7 and MK progenitors (Fig. 4B). Six of the seven transcription factors in the HPC7 cluster overlap with the heptad signature-a binding pattern that we have previously shown is overrepresented in the loci of genes specifically expressed in HSPCs and therefore associated with gene sets specifically expressed in HSCs [12]. Similarly, the gene signature associated with the third wave of the myeloid lineage program (d-my signatures) from Ng et al. [15] shows statistically

significant overlap with two combinatorial binding events, Cebp α , Cebp β , Stat1, P65, and Pu.1 in macrophages and Myb in myeloid progenitors. In addition to showing the functionality of the web tool, these results suggest that combinatorial control signatures generated by C-GSCA have the potential to provide insights into combinatorial transcriptional control mechanisms, and that the GSCA web tool provides access to this type of analysis to the wider community.

GSCA analysis of gene sets associated with hematologic malignancies

We have shown that GSCA can be used to link lineagespecific gene sets to combinations of candidate upstream regulatory TFs, and these associations are consistent with expectations based on current knowledge of regulatory control within hematopoiesis. This consistency attests to the potential robustness of the GSCA approach and suggests that it may also be useful to reveal biological insights into transcriptional programs operating in malignant hematopoietic cells, where diagnostic or prognostic gene sets have been derived for many types of leukemia, yet the combinations of TFs driving expression of these gene sets remain largely unknown. We therefore explored



Figure 3. (continued). (B) Overrepresented regulators determined using GSCA (left) and C-GSCA (right) for gene module 745 from Novershtern et al. [14] with "NK + T cell" induction pattern. C-GSCA is able to separate combinatorial patterns in T cells and myeloid progenitors.

the utility of GSCA for linking leukemic gene sets with candidate upstream regulators.

We first analyzed a gene set recently reported by McCormack et al. [39], in which the investigators showed that overexpression of Lmo2 in T-lymphoid progenitors induced a preleukemic state characterized by extensive self-renewal capacity. When the authors performed comparative gene expression profiling of normal and LMO2 expressing thymocytes, they noted upregulation of several HSC specific genes and suggested that ectopic expression of Lmo2 might activate an HSC specific transcription program. To test this hypothesis further, we analyzed the list of genes upregulated in Lmo2 transgenic DN thymocytes [39] by GSCA. This analysis suggested that the LMO2 overexpression gene set was under the transcriptional control of stem cell transcription factors such as Scl, Gata2, Runx1, Fli1 and Erg and also showed a strong overlap with LMO2 binding itself in nonleukemic progenitor cells.

We next analyzed gene expression profiling data generated as part of a recent study investigating transcriptional programs downstream of mixed lineage leukemia (MLL) transformation in mouse models of acute myeloid leukemia (AML) [40]. Expression analyses following MLL-AF9 withdrawal had prompted the authors to propose a model whereby MLL-AF9 enforces a Myb-coordinated program of aberrant self-renewal that involves genes linked to leukemia stem cell potential and poor prognosis in human AML patients. Of note, when we analyzed the genes downregulated following MLL-AF9 withdrawal by GSCA, we observed statistically significant overlaps with the two Myb ChIP-Seq datasets in our compendium (Fig. 5A). In addition, GSCA also recovered associations with MAX and the MAX interacting protein MXI1, both of which have also been linked to a range of human cancers [41]. GSCA analysis therefore not only corroborated the findings by Zuber et al. [40]; it also provided additional hypotheses on likely mechanisms that might control transcriptional programs downstream of MLL-AF9 in AML.

The final leukemic gene set analyzed by GSCA was taken from a 2009 study of the transcriptional programs in leukemic stem cells [42]. Comprehensive gene expression profiling analysis had lead the authors to speculate that leukemia stem cells in an MLL-driven mouse model of AML are characterized by a transcriptional program shared with embryonic rather than adult stem cells. This conclusion was subsequently challenged when it was suggested that the overlap with embryonic stem cell transcriptional programs was the reflection for a shared dependence on c-MYC activity rather than related to the stemness phenotype of ES cells [43]. Analysis of the leukemia stem cell associated gene set from



Figure 4. (A) Screen shot of Gene Set Control Analysis (GSCA) web tool with an option to either paste user defined gene list or upload from file, and to select method (GSCA or C-GSCA). (B) GSCA and C-GSCA output for stem signature dataset from Ng et al. [15] showing two cell type–specific distinct combinatorial patterns.





B Somervaille et. al., Cell Stem Cell. 2009 Feb 6;4(2):129-40. (Probe sets positively correlated with LSC frequency)



Figure 5. (A) Overrepresented regulators determined by C-GSCA for genes down regulated after MLL-AF9 withdrawal from Zuber et al. [40]. C-GSCA supports the notion that AF9 induces an Myb coordinated response. (B) Overrepresented regulators determined by C-GSCA for genes positively correlated with LSC frequency from Somervaille et al. [42]. C-GSCA identified cMyc and several other transcription factors to be overrepresented.

the Somervaille et al. [42] study by GSCA revealed a strong association with c-MYC ChIP-Seq datasets (Fig. 5B). However, there were also statistically significant associations with many additional ChIP-Seq datasets. GSCA analysis was therefore supportive of a role for c-MYC in the similarity between leukemic and embryonic stem cell expression signatures, but suggested that TFs more specifically expressed within blood cells also make important contributions to the leukemia stem cell transcriptional program. Of note, genes negatively associated with the leukemia stem cell phenotype in the study by Somervaille et al. [42] did not show the overlap with c-MYC, but it showed a distinct pattern of correlated ChIP-Seq datasets for the hematopoietic TFS, which interestingly contained several datasets for mature macrophages and was thus consistent with a relatively immature differentiation stage for the leukemia stem cells (Supplementary Figure 2, online only, available at www.exphem).

The application of GSCA to leukemic expression datasets supports the notion that integrated analysis of genome-wide transcription factor binding maps has significant potential as a new addition to the toolbox used by experimentalists to derive new hypothesis for experimental validation, which in the case of our current implementation of GSCA analysis would be geared specifically toward the identification of transcriptional mechanisms that control the behavior of normal and leukemic blood cells.

Discussion

Gene expression arrays have been used widely to characterize genes responsible for a particular cellular phenotype. The differentially expressed genes thus obtained can then be used for functional enrichment analysis. However, the important question of "What upstream regulatory mechanisms are responsible for the differential expression?" is not specifically addressed when using current approaches for gene set analysis, such as Gene Ontology or Gene Se Enrichment analysis tools.

As a result of the rapid progress in next-generation sequencing technology, ChIP-Seq analysis has become a favorite tool to investigate in vivo binding events because it offers higher resolution, less noise, and greater coverage compared with other techniques [44]. Nevertheless, the generation of genome-wide binding maps for multiple transcription factors across different cell types remains a formidable challenge for individual labs [45]. ChIP-Seq datasets from different labs can, however, be integrated at the computational level, which we recently demonstrated using 53 mouse ChIP-Seq experiments from different laboratories across the hematopoietic differentiation tree [8]. Since then, we have added 60 new ChIP datasets, thus more than doubling the size of the original compendium. In addition to highlighting a potentially major portion of the total regulatory genome involved in hematopoietic gene expression, a data compendium of this scale should have the potential

to provide new insights into regulatory mechanisms governing gene sets of interest.

To explore this further, we developed GSCA to identify enriched combinatorial binding patterns of transcription factors regulating a given gene set. This method uses experimental binding evidence, keeping the cell type specific context, unlike prediction methods based on overrepresentation of cis-regulatory sequence motifs in the promoters [46]. Using 80 clusters of tightly coexpressed genes in 38 hematopoietic cell types [14], we demonstrated that the transcriptional control mechanisms predicted are biologically coherent, and that GSCA performs better than current methods. Of note, this analysis also demonstrated that GSCA can be used in a cross-species fashion, with human gene sets analyzed using a murine ChIP-Seq compendium in this particular instance. The rationale for this crossspecies capability is provided by recent observations from ChIP-Seq data for the same transcription factor in multiple species where it was shown that, although a significant proportion of binding locations (peaks) are not conserved, there tends to be what was termed binding site turnover for these sites where loss of binding in one species is accompanied by gains elsewhere in the same gene locus in the other species [47]. The conserved and many of the nonconserved binding sites therefore map to the same gene loci, such as in human-mouse comparisons. Just as for many other gene set analysis tools, cross-species capability in GSCA is facilitated by the use of standard gene symbols that are standardized across mammals.

We further illustrated the utility of the GSCA tool to unravel potential regulatory mechanisms underlying a range of leukemia gene sets, thus suggesting potential future application of GSCA to build hypotheses to investigate transcriptional control mechanisms responsible for the expression of gene sets with diagnostic, prognostic, or therapeutic relevance. Finally, we built a web tool to facilitate similar analysis for the wider scientific community. Complementary to gene ontology functional overrepresentation analysis, GSCA calculates overrepresentation of binding events for a gene list of interest, thus predicting possible transcriptional control mechanisms.

Given the significant investment into several collaborative projects such as the ENCODE (Encyclopaedia of DNA Elements) and modENCODE (model organism ENCODE) initiatives [48,49], we are likely to witness a near exponential increase in ChIP-Seq datasets over the coming years. Although our current implementation of the GSCA web tool is geared toward predicting candidate upstream regulators within hematopoietic cells, the approach can be applied easily to other tissues when sufficient ChIP-Seq data become available.

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Author contributions: A.J. developed the GSCA software and performed analysis; R.L.H. collected and processed publicly available ChIP-Seq data; E.D. developed the web tool; B.G. designed the study and supervised the work; A.J. and B.G. wrote the manuscript.

Conflict of interest disclosure

No financial interest/relationships with financial interest relating to the topic of this article have been declared.

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GSCA

Supplementary Figure 1. GSCA analysis of Heptad signature identified by Wilson et al. [12].



Supplementary Figure 2. Hierarchical maintenance of MLL myeloid leukemia stem cells uses a transcriptional program shared with embryonic rather than adult stem cells [42]. (A) Probe sets positively correlated with LSC frequency. (B) Probe sets negatively correlated with leukemia stem cell (LSC) frequency.



Supplementary Figure 3. GSCA analysis of eight of nine gene expression signatures identified by Ng et al. [15].

Supplementary Table 1. The overlap between tissue specific enhancers identified by Pennacchio et al. [13] and the blood compendium showing that the enhancers in the compendium are highly blood specific

Tissue type	Number of enhancers	Overlap	<i>n</i> value
	212	0.000 map	
adipose tissue	213	86	0.99995
Adrenal gland	1/0	47	1
Allyguala P220 ⁺ P collo	218	24 159	1 1 01E 10
Bladder	212	138	1.91E-10
Blastocysts	191	63	1
Bone	200	87	0.99795
Bone marrow	224	101	0.99466
Brown fat	224	47	1
CD4 ⁺ T cells	226	148	0.000149
CD8 ⁺ T cells	194	137	7.00E-07
Cerebellum	180	24	1
Cerebral cortex	190	29	1
Digits	263	56	1
Dorsal root ganglia	193	45	1
Dorsal striatum	193	30	1
Embryo day 10	171	58	1
Embryo day 6	167	68	0.99961
Embryo day 7	163	51	1
Embryo day 8	170	44	1
Embryo day 9	174	63	1
Epidermis	292	58 22	1
Eye Fortilized egg	233	32	1
Frontal cortex	107	21	1
Heart	227	62	1
Hippocampus	201	30	1
Hypothalamus	183	25	1
Kidney	230	43	1
Large intestine	208	62	1
Liver	267	27	1
Lung	241	75	1
Lymph node	245	160	0.000102
Mammary gland	198	33	1
Med	192	40	1
Olfactory bulb	194	32	1
Oocyte	173	37	1
Ovary	192	46	1
Pancreas	211	45	1
Pituitary	187	34	1
Placenta	202	59	1
Preopuc	170	29	1
Saliyary gland	221	49	1
Skeletal muscle	215	40	1
Small intestine	259	65	1
Snout epidermis	275	51	1
Spinal cord lower	197	39	1
Spinal cord upper	196	26	1
Spleen	228	108	0.97033
Stomach	206	46	1
Substantia nigra	183	29	1
Testis	197	31	1
Thymus	194	111	0.15904
Thyroid	239	47	1
Tongue	289	51	1
Trachea	250	72	1
Trigeminal	193	30	1
			(continued)

Supplementary	Table	1.	(continued)
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Tissue type	Number of enhancers	Overlap	p value
Umbilical cord	223	44	1
Uterus	181	58	1
Vomeralnasal organ	252	66	1

	Novershtern et al. clusters	Candidate upstream regulators		
#	Induction pattern	Transcription factor	Cell type	
583	Late $Ery + T/B cell + GRAN$	TCF7	EML	
	-	GFI1B	Erythroid	
		SCL	HPC	
		MAX, MXI1, NELFE, TBP	MEL	
		ETS1	MK progenitors	
		FLI1	T cells	
		RAG2	Thymocytes	
607		TCF7	EML	
007		GF11B. SCL	HPC7	
		P65 PPARG	Macrophages	
		MXI1 NELEE	MEL	
		MYB	Myeloid progenitors	
		FI I	T cells	
		PAG2 PING1R	Thymocytes	
640	P coll	$E_{2A} = E_{2A} = E$	P collo	
049	Всеп	EZA, EDFI, OCIZ, FAAJ, FUI DUNV1	E CEIIS	
		KUNAI CEUR LDRI MTCRI RUI SCI	ENIL	
		GFIIB, LDB1, MIGKI, PUI, SCL	Erythroid	
		SUL	HPC	
		FLII, GATA2, MEISI, PUI, SCL	HPC/	
		CEBPA, CEBPB, P65, STATT	Macrophages	
		CMYB, CHD2, JUND, MAFK, MAX, MXI1, NELFE, SCL	MEL	
		GATA1	MK cells	
		CBFB, RING1B, RUNX1	MK progenitors	
		AF9	MLL leukemia	
		MYB	Myeloid progenitors	
		EBF1, SMAD1	Pro B cells	
		RBPJ	T cell leukemia	
		FLI1, GATA3, PU1, STAT3, STAT5A, STAT5B, STAT5, TBET	T cells	
		CBFB, RAG2, RUNX1	Thymocytes	
655	Муе	LDB1, SCL	HPC	
		NELFE, SCL	MEL	
661	Late $Ery + T/B - cell + GRAN$	TCF7	EML	
		NELFE	MEL	
		FLI1	T Cells	
		RAG2	Thymocytes	
667	T cell + NK	RUNX1	EML	
		GATA2. RUNX1	HPC	
		RUNX1	HPC7	
		GATA3 STAT5A STAT5B	T Cells	
		CBEB_RUNX1	Thymocytes	
673	T/B cell	E2A EBEL OCT2	B cells	
075	i/b cell	GATA2 PUNY1 SCI	HPC7	
		CEDDA CEDDD D65 DDADC	Magnamhagaa	
		CEDIA, CEDID, F03, FFARO	Pro D collo	
		EBF1, SMADI	Pro B cells	
		FLII, GAIA3, PUI, SIAI3, SIAI4, SIAI5A, SIAI5B, SIAI5, SIAI6	I cells	
<0 .		CBFB, KINGIB, KUNAI	Inymocytes	
685	Early Mye $+$ T/B cell $+$ GRAN	RUNX1, TCF7	EML	
		GFIIB, PUI	Erythroid	
		SCL	HPC7	
		CEBPA, CEBPB, P65	Macrophages	
		CMYC, MAX, MXI1, NELFE, TBP	MEL	
		CBFB, ETS1, RUNX1	MK progenitors	
		MYB	Myeloid progenitors	
		FLI1, STAT4, STAT5B, STAT6, TBET	T cells	
		CBFB, RAG2, RING1B, RUNX1	Thymocytes	
703	T/B cell	RAG2	Thymocytes	
715	Early Mye + T/B cell + GRAN	RAG2	Thymocytes	
721	Late $MYE + DCs$	CEBPA, CEBPB, P65	Macrophages	

Supplementary Table 2. Thirty-seven gene sets of 80 with respective induction patterns from Novershtern et al. [14] found enriched using the method of Lachmann et al. [34] and Zambelli et al. [35]

Supplementary Table 2. (continued)

	Novershtern et al. clusters	Candidate upstream regulators		
#	Induction pattern	Transcription factor	Cell type	
727	Late Ery	ETO2	Erythroid	
		LDB1, SCL	HPC	
		SCL	MEL	
733	HSE + Early Mye	RUNX1, TCF7	EML	
		GATA1, GATA2	Erythroid progenitors	
		GFI1B, MTGR1, SCL	Erythroid	
		GATA2, GFI1B, LMO2, MEIS1, PU1, SCL	HPC7	
		CEBPA, CEBPB, P65, STAT1	Macrophages	
		GATA1, MAFK, MXI1, NELFE, TBP	MEL	
		GATA1	MK cells	
		GATA1, GATA2, RING1B	MK progenitors	
		MYB	Myeloid progenitors	
		GATA3, STAT3, STAT5A, STAT5B, STAT5, STAT6, TBET	T cells	
		RAG2, RING1B, RUNX1	Thymocytes	
739	Late $Ery + T/B cell + GRAN$	TCF7	EML	
		MXI1, NELFE	MEL	
		FLI1	T cells	
		RAG2	Thymocytes	
763	Late MYE	EBF1	B cells	
		RUNX1, TCF7	EML	
		PU1	Erythroid	
		FLI1, GFI1B, RUNX1, SCL	HPC7	
		CEBPA, CEBPB, P65, PPARG, STAT1	Macrophages	
		NELFE	MEL	
		CBFB	MK progenitors	
		GATA3, STAT4, TBET	T cells	
		RAG2	Thymocytes	
793	Late $Erv + T/B - cell + GRAN$	TCF7	EML	
170		SCL	HPC	
		NEL FE	MEL	
		ETS1 RUNX1	MK progenitors	
		FI II	T cells	
		CBEB RAG2	Thymocytes	
799	NK + T cells (2)	$F_{2A} = F_{2A} = F$	B cells	
177		RUNY1	EMI	
		FTO2 PU1	Erythroid	
		GATA2 LDB1 SCI		
		EDG ELLI GATA2 GELLE I MO2 I VI 1 MEIS1 DUI DUNVI SCI		
		CERDA CERDA DES DDADG DUI STATI	Macrophages	
		CMVP CHD2 IIND	MEI	
		GATA1	MEL MK cells	
		CDED GATA1 GATA2 DING1D	MK cells	
		MVD	Mulaid progenitors	
		MID SMAD1	Dro D collo	
		SMADI El 11 CATA 2 DU1 STAT2 STATA STATEA STATED STATE STATE STATE	Pro B cells	
		FLII, GAIA3, PUI, SIAI3, SIAI4, SIAI3A, SIAI3B, SIAI3, SIAI6, IBEI	I cells	
011		CBFB, KAG2, KUNAI	Thymocytes	
811	Early Mye $+ 1/B$ cell $+ GRAN$		EML UDC7	
		SUL	HPC/	
		CMYC, MXII, NELFE, TBP, USF2	MEL	
		ETS1, RUNX1	MK progenitors	
		FLII	T cells	
		RAG2, RUNXI	Thymocytes	

Supplementary Table 2. (continued)

Novershtern et al. clusters		Candidate upstream regulators	
#	Induction pattern	Transcription factor	Cell type
817	T/B cell	E2A, EBF1, OCT2, PAX5	B cells
		RUNX1, TCF7	EML
		ETO2, GFI1B, PU1, SCL	Erythroid
		GAIA2, SCL	HPC UDC7
		EKG, FLH, GFHB, LMO2, MEISI, PUI, KUNAI, SCL CERDA CERDR D65 DDADG STAT1	HPC / Macrophages
		CEDFA, CEDFB, F05, FFARO, STATT CMVR, CHD2, MVI1, NELEE, SCI	MEI
		CRFR GATA2 RING1B RUNX1	MEL MK progenitors
		EBF1_SMAD1	Pro B cells
		FLI1, GATA3, STAT3, STAT4, STAT5A, STAT5B	T cells
		CBFB, RAG2, RING1B, RUNX1	Thymocytes
823	Early Mye + T/B cell + GRAN	MXI1, NELFE	MEL
		FLI1	T cells
		RAG2	Thymocytes
835	Early Mye + T/B - cell + GRAN	GFI1B	Erythroid
		ERG	HPC7
		CMYC, CHD2, MAX, NELFE, TBP	MEL
		ETS1, RUNX1	MK progenitors
		FLI1, STAT3, STAT6	T cells
		RAG2	Thymocytes
841	Early Mye + T/B cell + GRAN	TCF7	EML
		SCL	HPC7
		NOTCH1	Leukemia
		CMYC, MAX, MXI1, NELFE, TBP	MEL
		ETS1	MK progenitors
		FLI1	T cells
		RAG2	Thymocytes
859	T cell + NK	RINGIB	Thymocytes
8/1	HSC + Early MYE	MXII, NELFE	MEL
		FLII	I cells
007	Let $E_{T} = T/D$ coll $+ CDAN$	KAG2	I nymocytes
005	Late $EIy + I/B$ cell + $ORAIN$	PUI TCE7	E Cells
			ENIL Erythroid progenitors
		GATA1 GEUR	Erythroid
		FRG SCI	HPC7
		CFBPB	Macrophages
		CMYC CHD2 MXI1 NELFE TBP	MEL
		CBFB, RING1B, RUNX1	MK progenitors
		FLI1, STAT3, STAT4, STAT5, STAT6, TBET	T cells
		RAG2, RING1B, RUNX1	Thymocytes
889	Late Ery	GATA1, GATA2, SMAD1	Erythroid progenitors
	-	ETO2, GATA1, LDB1, MTGR1, SCL	Erythroid
		GATA2, LDB1, SCL	HPC
		LMO2, RUNX1	HPC7
		CEBPA, CEBPB	Macrophages
		CMYB, GATA1, MAFK, MAX, SCL, USF2	MEL
		GATA1	MK cells
		GATA1, GATA2, RING1B, RUNX1	MK progenitors
901	Early Mye + T/B cell + GRAN	TCF7	EML
		GFI1B	Erythroid
		NOTCH1	Leukemia
		CMYC, MXI1, NELFE, TBP	MEL
		ETS1	MK progenitors
		FLI1, STAT5B	T cells
		RAG2, RUNX1	Thymocytes

Supplementary Table 2. (continued)

Novershtern et al. clusters		Candidate upstream regulators	
#	Induction pattern	Transcription factor	Cell type
907	Late Ery + T/B cell + GRAN	TCF7 GATA1, GATA2 ETO2, GATA1, GFI1B, MTGR1 LDB1, SCL NOTCH1 CMYC, GATA1, MAX, MXI1, NELFE, SCL, TBP	EML Erythroid progenitors Erythroid HPC Leukemia MEL
		CBFB, ETS1, GATA1, RING1B FL11, STAT3, STAT5B, STAT6, TBET RAG2	MK progenitors T cells Thymocytes
925	Early Mye + T/B cell + GRAN	TCF7 CMYC, MXII, NELFE, TBP FLI1, STAT6 PAG2	EML MEL T cells Thymocytes
943	T/B cell	TCF7 NELFE ETS1 FLI1 RAG2	EML MEL MK progenitors T cells Thymocytes
961 973 979	B cell HSE + Early Mye Late MYE	E2A, EBF1, OCT2 NELFE CEBPA, CEBPB, P65, PPARG, STAT1 MYB	B cells MEL Macrophages Myeloid progenitors
985 991	Early Mye + T/B – cell + GRAN T/B cell	CHD2 E2A, OCT2, PAX5, PU1 RUNX1, TCF7 GATA1, GATA2 GATA1, GF11B, PU1 ERG, FL11, GF11B, MEIS1, PU1 CEBPA, CEBPB, P65, PPARG, STAT1 CMYC, CHD2, MAX, MX11, NELFE, TBP CBFB, ETS1, RING1B, RUNX1 MYB EBF1 FL11, PU1, STAT3, STAT4, STAT5A, STAT5B, STAT5, STAT6, TBET	MEL B cells EML Erythroid progenitors Erythroid HPC7 Macrophages MEL MK progenitors Myeloid progenitors Pro B cells T cells Thymocytes
1003	Late Ery + T/B - cell + GRAN	NELFE PAG2	MEL Thymocytes
1021	Early Mye + T/B cell + GRAN	TCF7 GFI1B NELFE ETS1 FLI1 RAG2, RING1B, RUNX1	EML Erythroid MEL MK progenitors T cells Thymocytes

Supplementary Table 3. Sixty-five gene sets of 80 with respective induction patterns from Novershtern et al. [14] enriched for transcription factor binding regions across multiple blood tissues using GSCA: 63 of 65 show cell type and induction pattern matching

Novershtern et al. clusters		Combinatorial control signature	
No.	Induction pattern	Transcription factor	Cell type
399	None	STAT4, STAT5	T cells
559	NK + T cell (2)	STAT3, STAT4, STAT5A, STAT5B, STAT5, STAT6, TBET	T cells
571	Late MYE	CEBPA, CEBPB, P65, PU1, STAT1	Macrophages
583	Late ERY $+$ T/B cell $+$ Gran	GATA1, GATA2, SMAD1	Erythroid progenitors
		SCL	Erythroid
		SCL	HPC
		MAX, MXI1, TBP	MEL
		RAG2	Thymocytes
607	Early MYE + T/B cell + Gran	PU1	B cells
		ERG, FLI1, GFI1B, MEIS1, PU1, SCL	HPC7
		CEBPA, CEBPB, P65, PPARG, PU1, STAT1	Macrophages
		MYB	Myeloid progenitors
		GATA3, PU1, STAT3, STAT4, STAT5A, STAT5B, STAT5, STAT6, TBET	T cells
613	T/B – cell	PU1	B cells
619	Late MYE	CEBPA, CEBPB, PU1, STAT1	Macrophages
637	Late Ery	GATA1, GATA2, SMAD1	Erythroid progenitors
		GATA1, LDB1, MTGR1, SCL	Erythroid
		ERG, LDB1	HPC
		ERG	HPC7
		GATA1, SCL	MEL
		GATA1, RING1B, RUNX1	MK progenitors
643	HSE + Early Mye	GATA2	HPC7
649	B cells	E2A, PAX5, PU1	B cells
		CEBPA, CEBPB, P65, PU1, STAT1	Macrophages
655	Mye	GATA1, GATA2, SMAD1	Erythroid progenitors
		GATA1, LDB1, MTGR1, SCL	Erythroid
		LDB1, SCL	HPC
		CEBPB, P65, PU1, STAT1	Macrophages
		CMYB, CMYC, GATA1, MAFK, MXI1, SCL, TBP	MEL
		CBFB, GATA1, RING1B	MK progenitors
661	Late $Ery + T/B cell + GRAN$	PU1	B cells
		GAIAI, GAIA2	Erythroid progenitors
		CMYB, CHD2, GAIAI, MXII, NELFE, TBP	MEL
		EISI	MK progenitors
		FLII	I cells
((7	$T \rightarrow 11 + NIZ$	ΚΑΟΖ Ο ΑΤΑ 2. ΟΠΑΤ2. ΟΠΑΤ4. ΟΠΑΤ5 Α. ΟΠΑΤ5 Ο ΟΠΑΤ5. ΟΠΑΤζ. ΠΡΕΤ	T nymocytes
00/	$1 \operatorname{cell} + \operatorname{NK}$	GAIAS, STATS, STAT4, STATSA, STATSB, STATS, STATO, TBET	There are the a
672	T/P. 2011	CDFD, KAO2, KINO1D, KUNAI	P collo
075	I/B cell	CEDDD D65 DU1 STAT1	D cells Maarambaaaa
		CEDED, FUJ, FUI, STATI CATA3 STAT3 STAT4 STAT5A STAT5B STAT5 STAT6 TBET	
670	HSE Forly Mye	GATA2	Fruthroid progenitors
685	I ate MVE \pm T/B cell \pm GRAN	RUNX1_TCF7	Englifold progenitors
005	Eace WITE $+$ 17D cell $+$ OKAN	GEI1B PU1	Enthroid
		FRG PUL SCI	HPC7
		CEBPA CEBPB P65 PU1 STAT1	Macronhages
		CMYC GATA1 MXI1 NELFE TBP	MEL
		CBFB. ETS1	MK progenitors
		FLI1. PU1	T cells
		RAG2, RUNX1	Thymocytes
703	T/B cell	TCF7	EML
		GFI1B	Erythroid
		ERG, PU1	HPC7
		NOTCH1	Leukemia
		CMYC, MAX, MXI1, NELFE, TBP	MEL
		CBFB, ETS1	MK progenitors
		FLI1, STAT3, STAT4, STAT5B, STAT6, TBET	T cells
		CBFB, RAG2	Thymocytes

Supplementary Table 3. (continued)

Novershtern et al. clusters		Combinatorial control signature	
No.	Induction pattern	Transcription factor	Cell type
709	General mild induction	ETO2	Erythroid
		AF9	MLL leukemia
715	Early MYE + T/B cell + GRAN	PU1	B cells
		TCF7	EML
		GFILB	Erythroid
		CEBPA, CEBPB, PUI, SIAII	Macrophages
		CMYC, GAIAI, MXII, NELFE, IBP	MEL MK and a solitons
		E151 E11	T colle
			T cells
721	Late MVE \pm DCs	CERPA CERPR P65 PU1 STAT1	Macrophages
721	Late MTE + Des	MVB	Myeloid progenitors
727	Late Frv	GATA1 GATA2 SMAD1	Frythroid progenitors
121	Late Liy	ETO2 GATA1 GEIIB LDB1 MTGR1 SCL	Erythroid
		LDB1 SCL	HPC
		CMYC, GATA1, MAFK, MAX, MXI1, SCL, TBP	MEL
		CBFB. GATA1. GATA2. RING1B. RUNX1	MK progenitors
733	HSC + Early MYE	CEBPA, CEBPB, P65, PU1, STAT1	Macrophages
		МҮВ	Myeloid progenitors
739	Late ERY $+$ T/B cell $+$ Gran	PU1	B cells
		TCF7	EML
		GATA1, GATA2	Erythroid progenitors
		GFI1B, PU1	Erythroid
		ERG, PU1	HPC7
		NOTCH1	Leukemia
		CMYC, CHD2, GATA1, MAX, MXI1, NELFE, TBP	MEL
		CBFB, ETS1, RUNX1	MK progenitors
		FLI1	T cells
		RAG2, RUNX1	Thymocytes
745	General mild induction	MYB	Myeloid progenitors
		STAT3, STAT4, STAT5	T cells
757	T cell + NK	TCF7	EML
		CMYC, MAX, MXI1, NELFE, TBP	MEL
		FLI1, STAT3, STAT5, TBET	T cells
		RAG2	Thymocytes
763	Late MYE	ERG, FLI1	HPC7
		CEBPA, CEBPB, P65, PPARG, PU1, STAT1	Macrophages
		PU1, STAT3, STAT4, STAT5, TBET	T cells
769	T/B cell	PU1	B cells
		CEBPA, CEBPB	Macrophages
		GATA3, STAT3, STAT4, STAT5A, STAT5B, STAT5, STAT6, TBET	T cells
775	Mye	GATA1	MK cells
781	General mild induction	NOICHI	TALL
/8/	MYE	SMADI	Erythroid progenitors
		CEBPA, CEBPB, STATT	Macrophages
702	Lata EDV T/D call Crar	GAIAI, MAFK	MEL D selle
193	Late $ERY + 1/B$ cell + Gran	PAA5, PUI TCE7	B cells
		IUF7 CATA 1, CATA 2, SMA D1	ENIL Emuthacid macconitors
		CATA 1 DUL SCI	Erythroid progenitors
		DUI SCI	HPC
		DI1	HPC7
		CERPA CERPR PUL STATI	Macrophages
		CEDIA, CEDID, IUI, SIAII CMVR CMVC GATAI MAY MYII NELEE SCL TDD	MEI
		GATA1	
		CRER ETS1 RING1R RUNY1	MK progenitors
		ΕΠΙ GATA3 STAT3 STAT4 STAT5A STAT5R STAT5 STAT6 TRET	T cells
		CRER RAG2 RUNX1	Thymocytes
			inymocytes

Supplementary Table 3. (continued)

Novershtern et al. clusters		Combinatorial control signature	
No.	Induction pattern	Transcription factor	Cell type
799	NK + T cell (2)	E2A	B cells
		CEBPA, CEBPB, P65, PU1, STAT1	Macrophages
		GATA3, PU1, STAT3, STAT4, STAT5A, STAT5B, STAT5, STAT6, TBET	T cells
805	HSE + Early Mye	ETO2	Erythroid
811	Late MYE + T/B cell + GRAN	TCF7	EML
		GFI1B	Erythroid
		ERG, SCL	HPC7
		CEBPA, CEBPB, PUI	Macrophages
		CMYC, CHD2, MAX, MXII, NELFE, TBP, USF2	MEL
		CBFB, ETSI, RUNXI	MK progenitors
		FLII, SIAI4, SIAI5B, SIAI5, SIAI6, IBEI	I cells
017	T/D coll	CBFB, KAG2, KING1B, KUNAI E2A EDE1 DAV5 DI1	I nymocytes
817	I/B cell	EZA, EBFI, PAA5, PUI EDC, MEIS1, DUI	B cells
		ERG, MEISI, PUI CEDDA CEDDD D65 DDADC DUI STATI	HPC/ Maananhaasa
		DING1R	MK progenitors
			Bro D collo
		EDF1, SMADI ELLI DUI STAT2 STAT4 STAT5A STAT5B STAT5 STAT6 TRET	T cells
		CREP DAG2 DUNY1	Thymocytes
823	Early MYE + T/B cell + GRAN	TCF7	FMI
025		GATA?	Ervthroid progenitors
		GFUB	Erythroid
		SCL	HPC7
		NOTCH1	Leukemia
		CMYC CHD2 MAX MXI1 NELFE TBP	MEL
		CBFB_FTS1	MK progenitors
		RBPI	T cell leukemia
		FLI1. STAT3. STAT6	T cells
		RAG2	Thymocytes
829	T cell + NK	E2A	B cells
		GATA3, STAT3, STAT4, STAT5A, STAT5B, STAT5, STAT6, TBET	T cells
		CBFB, RUNX1	Thymocytes
835	Early MYE + T/B cell + GRAN	PAX5	B cells
		TCF7	EML
		GFI1B	Erythroid
		ERG	HPC7
		NOTCH1	Leukemia
		CMYC, CHD2, MAX, MXI1, NELFE, TBP	MEL
		CBFB, ETS1, RUNX1	MK progenitors
		RBPJ	T cell leukemia
		FLI1	T cells
		RAG2	Thymocytes
841	Early MYE + T/B cell + GRAN	PU1	B cells
		TCF7	EML
		GFI1B, PU1	Erythroid
		ERG, GFI1B, PU1, SCL	HPC7
		NOTCH1	Leukemia
		PU1	Macrophages
		CMYC, CHD2, MAX, MXI1, NELFE, TBP	MEL
		ETS1, RUNX1	MK progenitors
		AF9	MLL leukemia
			T cell leukemia
		FLII, FUI, SIAI3, SIAI4, SIAI3B, SIAI6, TBET	I Cells
		KAUZ	Inymocytes

Supplementary Table 3. (continued)

Novershtern et al. clusters		Combinatorial control signature	
No.	Induction pattern	Transcription factor	Cell type
847	Late Ery + T/B cell + GRAN	PAX5	B cells
		TCF7	EML
		GATA1, GFI1B	Erythroid
		GFI1B	HPC7
		NOTCH1	Leukemia
		CMYC, CHD2, MAX, MXI1, NELFE, TBP	MEL
		CBFB, ETS1, RUNX1	MK progenitors
		RBPJ	T cell leukemia
		FLI1, STAT3, STAT4, STAT5B, STAT6, TBET	T cells
		RAG2	Thymocytes
853	Late MYE	PU1	B cells
		ERG. PU1	HPC7
		CEBPA, CEBPB, P65, PU1, STAT1	Macrophages
		PUL STAT3 STAT4 STAT5A STAT5 STAT6	T cells
859	T cell + NK	GATA3 STAT3 STAT5 TBET	T cells
865	HSE \perp early Mye	GATA2 SMAD1	Erythroid progenitors
005	HSL + carry wryc	GATA1	MEI
971	USE Forly MVE	TCE7	
0/1	HSE + Earry WITE	CELLB	EWIL
		UFIID EDC MEIS1 DUI	Elyilloid
		EKG, MEISI, PUI	HPC/
		NOICHI	Leukemia
		CEBPA, CEBPB, PUI, STATT	Macrophages
		CMYC, MAX, MXII, NELFE, TBP	MEL
		CBFB	MK progenitors
		FLI1, STAT3, STAT6, TBET	T cells
		RAG2	Thymocytes
883	Late MYE + T/B cell + Gran	TCF7	EML
		GATA1, GFI1B	Erythroid
		SCL, SCL	HPC
		SCL	HPC7
		CMYC, CHD2, MXI1, NELFE, TBP	MEL
		RING1B, RUNX1	MK progenitors
		FLI1, STAT4, STAT6, TBET	T cells
		RAG2, RUNX1	Thymocytes
889	Late ERY	GATA1, GATA2, SMAD1	Erythroid progenitors
		ETO2, GATA1, LDB1, MTGR1, SCL	Erythroid
		LDB1, SCL	HPC
		GATA1, SCL	MEL
		GATA1, GATA2, RING1B	MK progenitors
		STAT4	T cells
895	Late ERY	GATA2	Erythroid progenitors
070		GATA1 LDB1	Erythroid
		CEBPA CEBPB PUI STATI	Macrophages
		MXI1	MEI
001	Late MVE T/B cell Gran	TCE7	FMI
901		GATA1 GATA2 SMAD1	Eruthroid progenitors
		CELLE DILL	Environd progenitors
		CELLB	
			Leukemia
		CMYC, GATAI, MAX, MXII, NELFE, TBP	MEL
		CBFB, ETSI	MK progenitors
		FLI1, STAT3, STAT5B, STAT5	T cells
		CBFB, RAG2, RING1B, RUNX1	Thymocytes

Supplementary Table 3. (continued)

Novershtern et al. clusters		Combinatorial control signature	
No.	Induction pattern	Transcription factor	Cell type
907	Late ERY + T/B cell + Gran	PAX5, PU1	B cells
		TCF7	EML
		GATA1, GATA2, SMAD1	Erythroid progenitors
		ETO2, GATA1, GFI1B, LDB1, MTGR1, SCL	Erythroid
		ERG, LDB1, SCL	HPC
		ERG	HPC7
		NOTCH1	Leukemia
		CMYB, CMYC, CHD2, GATA1, MAX, MXI1, NELFE, SCL, TBP	MEL
		CBFB ETS1 GATA1 RING1B RUNX1	MK progenitors
		FLI1	T cells
		RAG2	Thymocytes
919	HSE + Farly Mye	FOX01	B cells
	HSE + Earry Wyc	FRG	HPC7
		CMVC MVI1 NELEE TDD	MEI
025	Early MVE $+$ T/P coll $+$ GPAN	DAV5 DU1	MIEL B. collo
923	Early $MTE + T/B$ cell + $ORAN$	TALS, FUI	D Cells
		GATA1, GATA2, SMADI	Erythroid progenitors
		GAIAI, GHIB, PUI	Erythroid
		ERG, SCL, SCL	HPC
		ERG, SCL	HPC/
		CMYC, CHD2, GATA1, MAX, MXI1, NELFE, SCL, TBP	MEL
		CBFB, ETS1, RING1B, RUNX1	MK progenitors
		FLI1, STAT3, STAT4, STAT5B, STAT5, STAT6, TBET	T cells
		RAG2, RING1B	Thymocytes
931	None	GATA1	MEL
		STAT4, STAT5B, STAT5	T cells
943	T/B cell	TCF7	EML
		GATA2	Erythroid progenitors
		PU1	Erythroid
		CMYC, MXI1, NELFE, TBP	MEL
		ETS1	MK progenitors
		FL11, GATA3, STAT3, STAT4, STAT5B, STAT5, STAT6, TBET	T cells
		CBFB, RAG2, RUNX1	Thymocytes
949	T/B cell	GATA2	Erythroid progenitors
		RAG2	Thymocytes
955	T cell + NK	GATA3, STAT3, STAT4, STAT5A, STAT5B, STAT5, STAT6, TBET	T cells
961	B cell	E2A, EBF1, OCT2, PAX5, PU1	B cells
		CEBPA, CEBPB, P65, PU1, STAT1	Macrophages
967	Late ERY $+$ T/B cell $+$ Gran	PAX5	B cells
		TCF7	EML
		GATA1, GFI1B, LDB1	Ervthroid
		ERG. FLI1. MEIS1. PU1. SCL	HPC7
		NOTCH1	Leukemia
		CMYC CHD2 MAX MXI1 NELFE SMC3 TBP	MEL
		CBFB FTS1 RING1B RUNX1	MK progenitors
		FLI1 STATA STATE TBET	T cells
		PAG2	Thymocytes
073	HSE \perp Early Mya	αποί αποί ματά μαχ μχιι	MEI
913	Loto MVE	DIT	MEL P. colle
979			D CEIIS
			HPC/
		CEBPA, CEBPB, POJ, PPAKG, PUI, STATI	Macrophages
			Pro B cells
		STAT3, STAT4, STAT5, STAT6	T cells

Supplementary Table 3. (continued)

Novershtern et al. clusters		Combinatorial control signature		
No.	Induction pattern	Transcription factor	Cell type	
985	Early MYE + T/B cell + Gran	TCF7	EML	
		GFI1B	Erythroid	
		ERG, SCL	HPC7	
		CMYC, CHD2, MAX, MXI1, NELFE, TBP	MEL	
		ETS1, RUNX1	MK progenitors	
		FLI1, STAT3, STAT4, STAT6, TBET	T cells	
		CBFB, RAG2, RUNX1	Thymocytes	
991	T/B cell	PU1	B cells	
		GATA2	Erythroid progenitors	
		PU1	Erythroid	
		ERG	HPC7	
		CEBPA, CEBPB, PU1, STAT1	Macrophages	
		MXI1, TBP	MEL	
		FLI1, STAT3, STAT4, STAT5B, STAT5, STAT6, TBET	T cells	
		RAG2	Thymocytes	
997	NK + T cell (2)	STAT4	T cells	
1003	Late ERY + T/B cell + Gran	TCF7	EML	
		GATA1, GFI1B, PU1, SCL	Ervthroid	
		ERG, PU1, SCL	HPC	
		ERG, PU1	HPC7	
		NOTCH1	Leukemia	
		CMYC, CHD2, MAX, MXI1, NELFE, TBP	MEL	
		CBFB, ETS1, RING1B, RUNX1	MK progenitors	
		RBPI	T cell leukemia	
		FLI1, GATA3, STAT3, STAT6	T cells	
		CBFB, RAG2, RING1B, RUNX1	Thymocytes	
1009	HSE + Early Mye	CEBPA, CEBPB, P65, PU1, STAT1	Macrophages	
1021	Early MYE + T/B cell + GRAN	PAX5	B cells	
1021		TCF7	EML	
		GATA1	Ervthroid progenitors	
		GEI1B PU1	Erythroid	
		SCL	HPC7	
		NOTCH1	Leukemia	
		CMYC CHD2 MXI1 NELEE TBP	MEL	
		ETS1 RUNX1	MK progenitors	
		FLI1	T cells	
		RAG2 RING1B RUNX1	Thymocytes	
		KIG2, KINGID, KUIMI	i ily illocy tes	