An experimentally validated network of nine haematopoietic transcription factors reveals mechanisms of cell state stability

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Abstract

Transcription factor (TF) networks determine cell type identity by establishing and maintaining lineage-specific expression profiles, yet reconstruction of mammalian regulatory network models has been hampered by a lack of comprehensive functional validation of regulatory interactions. Here, we report comprehensive ChIP-Seq, transgenic and reporter gene experimental data that have allowed us to construct an experimentally validated regulatory network model for haematopoietic stem/progenitor cells (HSPCs). Model simulation coupled with subsequent experimental validation using single cell expression profiling revealed potential mechanisms for cell state stabilisation, and also how a leukemogenic TF fusion protein perturbs key HSPC regulators. The approach presented here should help to improve our understanding of both normal physiological and disease processes.
Introduction

Tight regulation of gene expression is essential for both the establishment and maintenance of cellular phenotypes within metazoan organisms. The binding of transcription factor proteins (TFs) to specific DNA sequence motifs represents the primary step of decoding genetic information into specific gene expression patterns. TF binding sites (TFBSs) or motifs are usually short (6-10 bp), and therefore found just by chance throughout the genome. Functional TFBSs often occur as evolutionarily conserved clusters, which in the case of enhancers can act over long distances, thus necessitating comprehensive analysis of entire gene loci to understand the transcriptional control mechanisms acting at mammalian gene loci.

Given the complex regulatory circuitries that arise when control of multiple genes is considered, transcriptional control is often represented in the form of gene regulatory networks (GRNs), which carry most mechanistic information when constructed from detailed knowledge on the TFs and the cis-regulatory elements with which they interact (1-6). Importantly, regulatory network models can provide much more than a representation of existing knowledge, because network simulations can reveal possible molecular mechanisms that underlie highly complex biological processes. Boolean modelling approaches have been used to reconstruct core regulatory networks in blood stem cells (7) and myeloid progenitors (8), but neither of these studies took into account the underlying regulatory structure of the relevant gene regulatory elements. Full gene-regulatory information has been used for an ordinary differential equation-based model (9, 10), but was restricted to a small three-gene core circuit. Large consortia efforts such as ImmGen and FANTOM5 have created comprehensive networks of either regulatory elements or gene signatures important for multipotency and differentiation (11, 12). Furthermore, studies looking at gene regulation circuitry in embryonic stem (ES) cells have proposed regulatory networks important for ES cell identity (13, 14). While the complexities of transcriptional control demand approaches such as network modelling, no
single experimental method can provide the complex biological data required for the construction of accurate models. The previously mentioned studies focus their attention on one specific aspect of network modelling and importantly did not combine network analysis with comprehensive functional validation. Given that the key building blocks are gene regulatory sequences and the TFs bound to them, essential information for network reconstruction includes (i) comprehensive TF binding data, (ii) \textit{in vivo} validation of the functionality of TF-bound regions as bona fide regulatory elements, and (iii) molecular data on the functional consequences of specific TF binding events (e.g. activation vs. repression). The regulatory network model that we present in this study comprises all of the aforementioned components and is accompanied by functional validation of model predictions.

\section*{Results}

\textit{In vivo} validation of cis-elements as regulatory network nodes connecting 9 HSPC TFs

For the reconstruction of a core GRN model for HSPCs, we focussed on nine major HSPC regulators (ERG, FLI1, GATA2, GFI1B, LYL1, MEIS1, PU.1, RUNX1, TAL1), for which genome-wide binding patterns in the murine multipotent progenitor cell line HPC7 have previously been published (15). First, we searched the literature to summarise known \textit{cis}-regulatory regions for the nine TFs that possess haematopoietic activity in transgenic mouse embryos, which recovered a total of 14 regions: \textit{Erg}+85 (16), \textit{Fli1}-15 (17), \textit{Fli1}+12, \textit{Gata2}-3 (18), \textit{Gata2}+3 (= \textit{Gata2}+9.5) (19), \textit{Gfi1b}+13, \textit{Gfi1b}+16, \textit{Gfi1b}+17 (16, 20), \textit{Lyl1} promoter (21), \textit{Spi1}-14 (22), \textit{Runxl}+23 (23), \textit{Tal1}-4 (24), \textit{Tal1}+19 (25) and \textit{Tal1}+40 (26).

To extend this partial knowledge of relevant gene regulatory sequences to a comprehensive definition of how these nine TFs might cross-regulate each other, we made use of the genome-wide binding data for the nine TFs (15) as well as information on acetylation of histone H3 at
lysine 27 (H3K27ac) (27) in the HPC7 blood progenitor cell line. Additional candidate gene regulatory regions for all nine TFs were selected based on binding of at least three TFs and H3K27ac, since it has been shown previously that transcriptionally active regions are commonly bound by multiple TFs and display H3K27 acetylation (28). To assign putative candidate regions to a given TF, they had to be located between its respective upstream and downstream flanking genes, i.e. within the gene body itself or its 5’ and 3’ intergenic flanking regions. The Erg gene locus for example contains five candidate cis-regulatory regions based on these criteria, namely Erg+65, Erg+75, Erg+85, Erg+90 and Erg+149 (Fig. 1a), of which only the Erg+85 region had previously been tested in transgenic mice (16). Inspection of the gene loci of all nine TFs resulted in the identification of 35 candidate cis-regulatory regions (Fig. 1b, Fig. 1-figure supplements 1-8). In addition to the 14 haematopoietic enhancers previously published, eight of the 35 new candidate regulatory regions had previously been shown not to possess activity in tissues of the blood system of mouse embryos: Gata2-83 (Gata2-77), Gfi1b promoter (20), Spi1-18, Spi1 promoter (22), Runx1 P1 promoter (29), Tal1-9, Tal1 promoter (30) and Tal1+6 (31). Of the remaining 27 candidate cis-regulatory regions, two coincided with genomic repeat regions (Runx1-322 and Runx1+1) and were excluded from further analysis because mapping of ChIP-Seq reads to such regions is ambiguous. Since a comprehensive understanding of regulatory interactions among the nine HSPC TFs requires in vivo validation of candidate regulatory regions, we next tested the remaining 25 candidate cis-regulatory regions for their ability to mediate reporter gene expression in embryonic sites of definitive haematopoietic cell emergence and colonisation, namely the dorsal aorta and foetal liver of E10.5 to E11.5 transgenic LacZ-reporter mouse embryos. For the Erg locus, this analysis revealed that in addition to the previously known Erg+85 enhancer, the Erg+65 and Erg+75 regions also displayed activity in the dorsal aorta and/or the foetal liver, while the Erg+90 and Erg+149 regions did not (Fig. 1c). Careful
inspection of a total of 188 transgenic mouse embryos revealed that nine of the 25 identified regions showed \textit{LacZ} expression in the dorsal aorta and/or foetal liver (Fig. 1b, Fig. 1-figure supplements 1-9). This large scale transient transgenic screen therefore almost doubled the number of known \textit{in vivo} validated early haematopoietic regulatory elements for HSPC TFs.

\textbf{ChIP-Seq maps for a second progenitor cell line validate core regulatory interactions}

Although HPC7 cells are a useful model cell line for the prediction of genomic regions with haematopoietic activity in transgenic mouse assays (16), they are refractory to most gene transfer methods and therefore not suitable for functional characterisation of regulatory elements using standard transcriptional assays. By contrast, the 416b myeloid progenitor cell line can be readily transduced by electroporation and therefore represents a candidate cell line for functional dissection of individual regulatory elements. As ChIP-Seq profiles in 416b cells had not been reported previously, we performed ChIP-Seq for H3K27ac and the nine TFs in this cell line (Fig. 2a, Fig. 2-figure supplements 1-8). Alongside with our previously published HPC7 data, this new 416b dataset represents the most complete genome-scale TF-binding analysis in haematopoietic progenitor cell lines to date, with all new data being freely accessible under the following GEO accession number GSE69776 and also at http://codex.stemcells.cam.ac.uk/. Genome-wide TF binding patterns in 416b and HPC7 cells were closely related when compared with binding profiles for the same factors in other haematopoietic lineages (Fig. 2b, Fig. 2-figure supplement 9). Inspection of the gene loci for the nine HSPC TFs not only revealed many similarities between 416b and HPC7 cells, but also some differences in TF binding patterns. Overall, TF occupancy at the 23 regions with activity in haematopoietic tissues (14 previously published (16-26) and 9 newly identified) does not change between the two cell types in 71 \% of all cases (147 of 207 binding events), is gained in
416b cells in 16% (33 of 207) and lost in 13% (27 of 207) of cases compared to HPC7 cells (Fig. 2c). Next, all 23 elements were filtered to only retain those elements which were bound by at least 3 of the 9 TFs and displayed elevated H3K27ac in HPC7 and 416 cells. This led to the removal of the $Gata2$-3, which is not bound by any of the nine TFs in either cell type, $Gata2$-92 and $Gfi1b$+13, which are only bound by one or no TFs in 416b cells, and $Fli1$-15, which is not acetylated in 416b cells (Fig. 2c, Fig. 2-figure supplements 1-3). Overall, 19 cis-regulatory regions were therefore taken forward as a comprehensively validated set of regions for the reconstruction of an HSPC regulatory network model.

Comprehensive TFBS mutagenesis reveals enhancer-dependent effects of TF binding

The reconstruction of a core regulatory network model requires information about the effect of TF binding on gene expression, which can be activating, repressing or non-functional. In order to analyse the effects of all TF binding events at all 19 regulatory regions, we performed luciferase reporter assays in stably transfected 416b cells. Based on multiple species alignments between five species (mouse, human, dog, platypus, opossum), we identified conserved TFBSs for the nine TFs (Fig. 3a, Fig. 3-figure supplements 1-19), and generated mutant constructs for each of the 19 regulatory regions, resulting in 87 reporter constructs that were tested by luciferase assays (19 wild-type, 68 mutants). To ensure that DNA binding of the TFs was abrogated, the key DNA bases involved in DNA-protein interactions were mutated and the resulting sequences were scanned to ensure that no new binding sites were created (32). For each of the 19 regulatory regions the conserved TFBSs were mutated by family, for example all six Ets sites within the $Erg$+65 region were mutated simultaneously in one construct and this element was then treated as the $Erg$+65_Ets mutant. TFBS mutations reduced or increased activity compared to the wild-type enhancer, or indeed had no significant
effect (Fig. 3b, Fig. 3-figure supplements 1-18). For instance, at the Erg+65 region, mutation of
the six Ets binding sites or the three Gata binding sites reduced luciferase activity, whereas
mutation of the three Ebox or the three Gfi motifs increased luciferase activity (Fig. 3b).
Comparison of the luciferase assay results for all 19 cis-regulatory regions (Fig. 3c) reveals
that for each motif class mutation can result in activation, repression or no-change. This
observation even extends to single gene loci, where for example mutation of the Gata site
reduced activity of the Erg+65 region, but increased activity of the Erg+85 enhancer (Fig. 3c).
Taken together, this comprehensive mutagenesis screen highlights the dangers associated with
extrapolating TF function simply from ChIP-Seq binding events and thus underlines the
importance of functional studies for regulatory network reconstruction.

Dynamic Bayesian network modelling can incorporate complex regulatory information
and shows stabilization of the HSPC expression state

We next set out to construct a regulatory network model that incorporates the detailed
regulatory information obtained for potential cross-regulation of the nine HSPC TFs obtained
in the previous sections (summarised in Fig. 4a). We focussed on three categories of causal
relationships: (i) one or several TFs can bind to a certain type of motif at a given regulatory
region, and the probability of a motif being bound depends on the expression levels of the
relevant TFs; (ii) TFBS mutations at a given regulatory region altered luciferase activities
compared to the wild-type, thus capturing the impact of TF binding on activity of the given
regulatory region; (iii) individual regulatory regions show varying degrees of activation over
baseline controls, which translate into different relative strengths of individual cis-regulatory
regions. To incorporate this multi-layered experimental information, we constructed a three-tier
dynamic Bayesian network (DBN) to jointly represent all those causal relationships (see
Material and Methods and Fig. 4b). The reconstructed DBN represents a first-order time-homogeneous Markov process, which is a stochastic process where the transition functions are the same throughout all time points and the conditional probability distribution of future states depends only on the present state (see Material and Methods). The model is calculated so that the expression at t+1 is influenced by the expression at t0; analogously, the expression at t0 is influenced by the expression at t-1, and so on. Therefore, though the model does not incorporate “epigenetic memory”, past expression levels directly influence current expression levels. Model execution therefore permits the simulation of gene expression states in single cells over time, as well as the calculation of gene expression distributions for each gene across a population of simulated single cells.

Having generated a DBN model incorporating extensive experimental information, we next investigated the expression states following model execution. First, we investigated whether the network model was compatible with the HSPC expression profile from which all the experimental data are derived, namely co-expression of all nine TFs. To this end, model execution was initiated with expression levels for all nine TFs set at the midpoint level of 0.5. A representative single cell modelled over time rapidly adopts characteristic levels of expression for each of the nine genes, with some genes showing perpetual fluctuations (Fig. 4c). The same expression levels were reached when the model was initiated with expression starting at 0.2, 0.8 or with initially only FLI1, RUNX1 and TAL1 being expressed at 0.5 (Fig. 4-figure supplement 1). We next modelled the overall distribution of the nine TFs as might be seen in a cell population by running 1000 model simulations (Fig. 4d). This analysis demonstrated that our model is compatible with co-expression of all nine genes within the same single cell. Moreover, stable expression over time for some genes as well as oscillations around a characteristic mean expression level for other genes suggests that our model may have
captured those aspects of HSPC regulatory networks that ensure maintenance of stem/progenitor cells.

Relative stability to experimental perturbation is recapitulated by the model

The TFs TAL1 and LYL1 are important regulators of adult haematopoiesis, but the deletion of each factor individually has only minor effects on adult HSC function (33-35). Combined deletion in adult HSCs however causes a severe phenotype with rapid loss of HSPCs (36). We wanted to investigate to what extent our computer model could recapitulate these known phenotypes through in silico perturbation simulations. To quantify if a change in the expression profile of a given TF was significant we performed a Wilcoxon rank-sum test. Interestingly, this significance calculation demonstrated that both large and small fold-changes can be significant. Simulated perturbation of just LYL1 caused significant alterations to the expression profiles of Gfi1b, Tal1, Fli1 and Gata2, but none of these were associated with a substantial shift in mean expression levels (Fig. 5a, Fig. 5-figure supplement 1). Perturbation of just TAL1 caused significant changes to the expression profiles of Runx1, Gfi1b and Gata2, and again none of these were associated with a substantial shift in expression levels (Fig. 5b, Fig. 5-figure supplement 1). Simultaneous deletion of both factors caused significant changes in gene expression profiles in all TFs except for Fli1. Unlike for the single TF perturbations, Gata2 and Runx1 showed substantial shifts in expression levels when both LYL1 and TAL1 were simulated to be knocked down (Fig. 5c, Fig. 5-figure supplement 1). Of note, the significance calculations highlight that there may be no one perfect way to visualize these small fold-change alterations. We therefore also generated histogram plots as an alternative visualization (Fig. 5-figure supplement 2).
We next wanted to compare model predictions with actual experimental data in the 416b cell line, from which the information for model construction had been derived. Because our DBN model is particularly suited to model the expression states in single cells, we compared predicted and experimentally observed effects of knockdown or overexpression in single cells. To this end we knocked down the expression of TAL1 in 416b cells by transfecting the cells with siRNA against *Tal1* (siTal1) or control siRNA (siCtrl). Forty-eight hours after transfection, gene expression for the nine network genes was analysed in 44 siTal1 treated cells and 41 siCtrl treated cells. Importantly, 29 of 44 cells (66%) transfected with siTal1 showed no expression of *Tal1* anymore, demonstrating the successful knockdown (Fig. 5d, Fig. 5-source data). Down-regulation of TAL1 caused a significant change in the expression profiles of *Tal1, Fli1* and *Gfi1b*, but a substantial shift of median expression was only observed for *Tal1* (Fig. 5-figure supplement 1). Experimental validation therefore confirmed the occurrence of statistically significant small-fold changes in expression profiles following single TF knockdown, although there was no perfect match between the genes affected in the model and experiment. To extend comparisons between model predictions and experimental validation, we investigated the consequences of knocking down the expression of PU.1 and overexpressing GFI1B. Complete removal of PU.1 *in silico* after the model had reached its initial steady state had no effect on the expression levels of the other TFs (Fig. 6a). To investigate whether the model prediction is comparable to experimental data obtained from single cells, single cell gene expression analysis using the Fluidigm Biomark HD platform was performed using 416b cells transduced with shRNA against PU.1 (shPU.1) or luciferase (shluc). Three days after transduction, 121 shPU.1 and 123 shluc transduced single cells were analysed for their expression of *Spy1* and the other eight TFs of the network. 18 shPU.1-transduced cells (15%) showed a complete loss of *Spy1*, and expression of *Spy1* in the remaining cells was markedly reduced compared to the control cells (shluc) (Fig. 6a, Fig. 5-
source data), highlighting the efficiency of the PU.1 knockdown. Spi1, Runx1, Erg and Fli1 showed a significant change in expression profiles after the depletion of PU.1, but this involved a substantial shift in median expression levels only for Spi1 and Runx1 (Fig. 5-figure supplement 1). Expression profiles of the remaining five TFs did not change as a result of reduced PU.1 levels (Fig. 6a, Fig. 5-source data), therefore mostly confirming the model prediction.

Next, we modelled GFI1B overexpression in silico by increasing the expression level of Gfi1b to the maximum value after the model had reached its initial steady state which led to a significant change in the expression profiles of Gfi1b, Meis1, Erg and Runx1, although only Gfi1b and Meis1 showed a substantial shift in median expression levels (Fig. 6b, Fig. 5-figure supplement 1, Fig. 5-source data). Expression profiles of the other five TFs were unaltered. Single cell gene expression analysis of 90 single 416b cells transduced with a Gfi1b-expressing vector and 104 single 416b cells transduced with an empty control vector showed a significant increase in the expression of Gfi1b and a significant alteration to the expression profile of Erg, but only the changes to Gfi1b involved a substantial shift in median expression levels. No significant expression changes were seen in any of the other seven network genes (Fig. 6b). Both PU.1 and GFI1B perturbation studies therefore emphasize the resilience of the HSPC TF network to single TF perturbation. Moreover, our in silico model reflects this, thus suggesting that the comprehensive experimental information used to construct the network model has allowed us to capture key mechanistic aspects of HSPC regulation. Of note, there were no short-term major expression changes immediately after the perturbation in the silico simulations for the three single TF perturbation described above. For completeness we performed in silico modelling for all permutations of single TF knockdown / overexpression as well as all pairwise combinations of all 9 TFs analysed (a total of 162 simulations, Fig. 6-figure supplement 1).
Major perturbations by the AML-ETO oncoprotein are captured by the network model

As the TF network described above is relatively stable to single TF perturbations, we set out to test whether a simulation that mimics the situation present in leukemic cells can influence the expression states of the nine TFs in our network. The *Aml-Eto9a* translocation is amongst the most frequent mutations in AML (reviewed in (37)). The resulting fusion protein is thought to act in a dominant negative manner to repress RUNX1 target genes. To simulate the leukemic scenario caused by AML-ETO expression, we fixed the level of *Runx1* to be the maximum value 1 and at the same time converted all activating inputs of RUNX1 to inhibiting inputs in our DBN model. Interestingly, this simulation of a “leukemic” perturbation caused significant expression changes to all eight of the core HSPC TFs (Fig. 6c). To compare the AML-ETO simulation results with experimental data, we utilised a doxycycline-inducible expression system to generate 416b cells with inducible expression of AML1-ETO fused to a mCherry reporter via a self-cleaving 2A peptide spacer. Following doxycycline induction, 56 single mCherry positive and 122 single mCherry negative 416b cells were analysed by single cell gene expression. Significant gene expression changes can be seen in six of the nine core HSPC TFs (all except *Tal1*, *Erg* and *Gata2*) thus highlighting significant overlap between predictions and experimental validation, although there are also notable differences between model predictions and the experimental data (see for example Gata2; Fig. 6c, Fig. 5-figure supplement 1, Fig. 5-source data). These results demonstrate that our new HSPC network model can capture many gene expression changes caused by ectopic expression of a leukemia oncogene as well as providing a useful model for normal HSPC transcriptional regulation. The inability of any model to completely recapitulate experimental data is not unexpected. Possible reasons in our case may include more complex activities of the onco-fusion protein than would be captured by our assumption that its “only” function is as a straightforward dominant-negative
Discussion

Transcription factor networks are widely recognised as key determinants of cell type identity. Since the functionality of such regulatory networks is ultimately encoded in the genome, the logic that governs interactions between network components should be identifiable, and in due course allow for the construction of network models that are capable of capturing the behaviour of complex biological processes. However, the construction of such network models has so far been severely restricted because the identification and subsequent functional characterisation of mammalian regulatory sequences represent major challenges, and the connectivity and interaction rules within regulatory networks can be highly complex. Here we report a comprehensive mammalian transcriptional network model that is fully grounded in experimental data. Model simulation coupled with subsequent experimental validation using sophisticated single cell transcriptional assays revealed the mechanistic basis for cell state stability within a haematopoietic progenitor model cell line, and also how a leukemogenic TF fusion protein can perturb the expression of a subset of key blood stem cell regulators.

Pictorial representations of putative network models are commonly shown in publications reporting ChIP-Seq TF binding datasets (38). However, due to the lack of experimental underpinning, such representations are simple images that do not encode any of the underlying gene regulatory logic, and importantly therefore cannot provide executable computational models that can be used to simulate biological systems. Although the experimentally-grounded network model shows good agreement with the relative expression states of the nine TFs for
Apart from the obvious caveat that any computer model is an abstraction of reality and therefore will not be correct in every detail, it also needs to be stressed that we treat the nine TFs as an isolated module for the computer simulations, and therefore could not account for possible influences by additional genes that may affect single cell gene expression measurements in the perturbation experiments.

Statistical significance calculations demonstrated that both the computer model and the experimental data showed significant changes in gene expression profiles that were associated with minimal fold-change alterations to median expression levels. Such alterations to expression profiles were prevalent in both single and double-gene perturbations, whereas substantial shifts in median expression were mostly restricted to the double perturbations (and also the AML-ETO oncogene overexpression). This observation suggests that (i) our approach has the capacity to reveal aspects of the fine-grained nature of biological networks, and (ii) the network presented in this study is largely resistant to perturbations of individual TFs in terms of substantial fold-change alterations in median expression levels. We believe that it may well be possible that the statistically significant small-fold changes in HSPC network genes may be responsible for the mild phenotypes seen when major HSPC regulators are deleted in adult HSPCs. *Tal1*^-/-^ mice for example are not viable because TAL1 is absolutely required for embryonic blood development (39), yet deletion of TAL1 in adult HSCs only causes minor phenotypes (33). Another noteworthy observation is that it would have been impossible to detect the statistically significant yet small fold-changes using conventional expression profiling, because they only become apparent following the statistical analysis of expression distributions generated by assaying lots of single cells. More generally it is important to acknowledge that the question of how close the present model comes to capturing the
underlying biological processes can only be revealed through much more exhaustive experimental validation studies.

A potential caveat for network reconstruction based on identification of regulatory elements comes from the difficulties associated with capturing negative regulatory elements. As shown elegantly for CD4 and CD8 gene silencers in the lymphoid lineage, TFs involved in the early repression of a locus are not required for maintenance of the silenced state (40, 41). Identification of negative regulatory inputs may therefore require an expansion of datasets to look across sequential developmental stages. It will therefore be important in the future to extend the work presented here to include additional HSPC regulators as well as additional stages along the haematopoietic differentiation hierarchy. Of note, TF-mediated cellular programming experiments have demonstrated that modules of 3-4 TFs are able to confer cell-type specific transcriptional programmes (42-45), consistent with the notion that a network composed of nine key HSPC regulators is able to capture useful information about HSPC regulatory programmes.

One of the most striking observations of the regulatory network defined here is the high degree to which the HSPC expression state is stabilised. As such, this model is different from previous experimentally-grounded transcriptional regulatory network models (46). These earlier model organism networks have inherent forward momentum, where the model captures the progression through successive embryonic developmental stages characterised by distinct expression states.

The model reported here is based on and validated with data from haematopoietic progenitor cell lines, which can differentiate (47, 48), but can also be maintained in stable self-renewing conditions. A recent study by Busch and colleagues tracked labelled Tie2+ HSCs in the bone marrow, and showed that haematopoietic progenitors \textit{in vivo} are also characterised by a
substantial self-renewal capability, therefore highlighting the stable state in which they can reside for several months (49). The observed stability of the HSPC expression state presented here is therefore likely to capture aspects of the regulatory mechanisms maintaining the steady state of primary haematopoietic progenitor cells, a notion reinforced further by the fact that our model is based on \textit{in vivo} validated regulatory elements.

The two types of models therefore accurately capture the properties of the distinct biological processes, e.g. driving developmental progression on the one hand, and maintaining a given cellular state on the other. Different design principles are likely to be at play, with feed-forward loops representing key building blocks of early developmental GRNs, while the network described here shows an abundance of auto-regulatory feedback loops and partially redundant enhancer elements, both of which may serve to stabilise a given cellular state.

Of particular interest may be the organisation of the \textit{Runx1} gene locus, where RUNX1 protein provides positive feedback at some, and negative feedback at other HSPC enhancers. Given that these different enhancers employ overlapping yet distinct sets of upstream regulators, it is tempting to speculate that such an arrangement not only stabilises a given expression level, but also provides the means to either up- or down-regulate RUNX1 expression in response to diverse external stimuli that may act on specific RUNX1 co-factors at either the repressing or activating RUNX1 binding events. Taken together, we report widely applicable experimental and computational strategies for generating fully validated regulatory network models in complex mammalian systems. We furthermore demonstrate how such a model derived for blood stem/progenitor cells reveals mechanisms for stabilisation of the progenitor cell state, and can be utilised to analyse core network perturbations caused by leukemic oncogenes.

\textbf{Materials and Methods}
ChIP-Sequencing and data processing

The mouse myeloid progenitor 416b cell line (48) was received from Chester Beatty lab and confirmed to be mycoplasma free. The cells were cultured in RPMI with 10 % FCS and 1 % Penicillin/Streptomycin.

ChIP assays were performed as previously described (16, 27), amplified using the Illumina TruSeq ChIP Sample Prep Kit and sequenced using the Illumina HiSeq 2500 System following the manufacturer’s instructions. Sequencing reads were mapped to the mm10 mouse reference genome using Bowtie2 (50), converted to a density plot and displayed as UCSC genome browser custom tracks. Peaks were called using MACS2 software (51). Mapped reads were converted to density plots and displayed as UCSC genome browser custom tracks. The raw and processed ChIP-Seq data have been submitted to the NCBI Gene Expression Omnibus (www.ncbi.nlm.nih.gov/geo) and assigned the identifier GSE69776. A binary binding matrix was created using in-house scripts, clustered using the dice coefficient and a heatmap was plotted using gplots in R in order to compare newly generated ChIP-Seq data with previously published data (52).

Analysis of enhancer activity in transient transgenic mouse embryos

Genomic fragments spanning the candidate cis-regulatory regions were generated by PCR or ordered as gBlocks (Life Technologies GmbH) and cloned downstream of the LacZ gene in an hsp68LacZ (Runx1 constructs) or SVLacZ (all other constructs) reporter vector. Coordinates of candidate chromosomal regions and corresponding primer sequences are given in Fig. 3-figure supplement 20. For Runx1, E10 mouse transient transgenic embryos carrying LacZ enhancer-reporter constructs were generated by pronuclear injection of (C57BL/6 x CBA)/F2 zygotes following standard procedures. Transgenic embryos were identified by LacZ-specific PCR on
genomic DNA isolated from yolk sac (5'-GCAGATGCACGTTACGATG-3'; 5'-GTGGCAACATGGAAATCGCTG-3'). Xgal staining and cryostat sectioning were performed as previously described (23). Embryos were photographed using a Leica MZFLIII microscope, Leica DFC 300F digital camera (Leica Microsystems, Milton Keynes, UK) and Openlab software (Improvision, Coventry, UK) and sections were examined using a Nikon Eclipse E600 microscope (Nikon, Japan) equipped with 20x and 40x Nomarski objectives. Photographs were taken using a Nikon DXM 1200c Digital Camera (Nikon, Tokyo, Japan). E11.5 transient transgenic embryos of all other candidate cis-regulatory regions were generated by Cyagen Biosciences Inc (Guangzhou, China). Whole-mount embryos were stained with 5-bromo-4-chloro-3-indolyl-β-d-galactopyranoside (X-Gal) for β-galactosidase expression and photographed using a Nikon Digital Sight DS-FL1 camera attached to a Nikon SM7800 microscope (Nikon, Kingston-upon-Thames, UK). Candidate transgenic mouse embryos with LacZ staining in haematopoietic tissues were subsequently embedded in paraffin, stained with 0.1 % (w/v) Neutral Red and cut into 6 μm deep longitudinal sections. Images of sections were acquired with a Pixera Penguin 600CL camera attached to an Olympus BX51 microscope. All images were processed using Adobe Photoshop (Adobe systems Europe, Uxbridge, United Kingdom).

Luciferase reporter assays

Wild-type and mutant DNA fragments for candidate regulatory regions were either cloned using standard recombinant DNA techniques, ordered as gBlocks (Life Technologies) or obtained from GeneArt® by Life Technologies. DNA fragments were cloned into pGL2 basic or pGL2 promoter vectors from Promega using restriction enzymes or by Gibson Assembly. TFBSs for the nine TFs of interest (corresponding DNA sequences are listed in Fig. 3-figure supplement 19) were identified based on multiple species alignments between five species
mouse, human, dog, platypus, opossum). Where a region contained multiple instances of the same motif, a single mutant construct with all relevant motifs mutated simultaneously was generated (for generated point mutations check Fig. 3a and Fig. 3-figure supplements 1-18). Where TF binding was observed in ChIP-Seq experiments in 416b cells, but the TFBS was not conserved, the motifs present in the mouse sequence were mutated. Stable transfections of the 416b cell line were performed using 10 μg reporter construct, 2 μg neomycin resistance plasmid and 1x10^7 416b cells in 180 μl culture medium per pulse. The sample was electroporated at 220V and capacitance of 900 μF using the GenePulser Xcell Electroporation System (Bio-Rad). Immediately after transfection, the sample was split into four culture plates. Twenty-four hours after transfection Geneticin G418 (Gibco by Life Technologies) at a final concentration of 0.75 mg/ml was applied to the culture to select for transfected cells. The activity of the luciferase reporter constructs was measured 12-16 days after transfection by using a FLUOstar OPTIMA luminometer (BMG LABTECH). The luciferase activity was normalised to the cell number and presented as relative activity compared to the wild-type construct. All assays were performed at least three times in quadruplicates.

**Single cell gene expression and data analysis**

The TAL1 knockdown was performed using pools of siRNA against Tal1 (Dharmacon) which were transfected into 416b cells. Briefly, 1 x 10^6 cells were electroporated with either a control or Tal1 siRNA. Forty-eight hours after transfection, cells were sorted into 96 well PCR plates containing lysis buffer using the BD Influx Cell Sorter.

The PU.1 knockdown was performed as previously described (27).

The MigR1-Gfi1b retroviral expression vector and the corresponding empty vector control (53) were used for GFI1B overexpression. Two million 416b cells were transduced with the above
listed vectors by adding viral supernatant and 4 μg/ml polybrene to the cells, followed by centrifugation at 900 x g for 90 min at 32°C and incubation with 5% CO₂ at 32°C. Half of the media was then replaced with fresh culture media and cells were incubated at 37°C with 5% CO₂. Forty-eight hours after transduction, GFP⁺ cells for each cell population were sorted into 96 well PCR plates containing lysis buffer using the BD Influx Cell Sorter.

To induce AML1-ETO9a expression, the 416b cell line was co-transfected with: 1) a plasmid containing the tetracycline transcription silencer (tTS), the tetracycline transactivator (rtTA) and blasticidin resistance under the control of the \( EF1\alpha \) promoter; 2) a plasmid containing the entire \( Aml-Eto9a \) cDNA (obtained from vector MigR1-AE9a, Addgene no. 12433) in frame with a F2A element and the mCherry protein under the control of a tetracycline responsive element; and 3) transposase PL623 (54) (kindly donated by Pentao Liu, Sanger Institute, Cambridge) in order to promote simultaneous stable integration of the two constructs described above. After 6 days of culture without selection, cells were incubated with 1 μg/ml of Doxycycline for 24 hours and then stained with DAPI. mCherry positive and negative cells that did not stain with DAPI were sorted into 96 well PCR plates containing lysis buffer using the BD Influx Cell Sorter.

Single cell gene expression analysis was performed using the Fluidigm BioMark platform followed by bioinformatics analysis as previously described (20). All cells that express less than 48% of genes assayed were removed from the analysis for PU.1 knockdown and GFI1B over-expression, all cells expressing less than 56% of genes assayed were removed from the TAL1 knockdown and all cells that express less than 44% of genes assayed were removed from the analysis for the AML-ETO9a induction. Importantly, this thresholding resulted in the removal of similar numbers of cells in both the perturbation and control arms of the experiments. The raw data as well as the normalised data (normalised to Ubc and Polr2a) of the gene expression analysis are listed in Fig. 5-source data).
**Computational modelling**

The first-order DBN shown in Fig. 4b was established on the basis of regulatory information summarized in Fig. 4a. The DBN essentially presents a discrete-time stochastic process that has the Markov property, i.e. the state of the process at the next time point depends purely on its state at the current time point. Also note that this is a time-homogeneous (or time-invariant) DBN, where the transition functions/matrices are the same throughout all time points.

To specify parameters of the DBN, we defined a motif family at a specific regulatory region as a unique binary variable; with value “1” indicating that no motif of a motif family is bound at the specific region and value “2” indicating that at least one motif of the motif family is bound by a TF at this region. We assumed that any of the following three factors can lead to a higher probability of a motif being bound by a TF and therefore taking the value 2: (i) more motifs of the same type present within a regulatory region; (ii) multiple TFs that can bind to the same motif, such as TAL1 and LYL1 both binding to Eboxes; (iii) higher expression levels of the TFs. The probabilities were thus calculated based on these three sources of information (see below for an example). We next defined that every regulatory region was a continuous variable on the close interval [0, 1], and its value was determined by the accumulated effects of all motifs present with the regulatory region. Finally, the expression levels of the nine TFs were also defined as continuous variables ranging from 0 to 0.8, and their expression levels were determined by the accumulated activities of the relevant regulatory regions.

Considering that variables in the top tier of the DBN are binary whereas those in the middle and bottom tiers are continuous, we found conditional linear Gaussian distribution (55) to be an appropriate generic representation of the intra-slice conditional probability distributions. Specifically, the regression coefficient of a regulatory region on a motif family was estimated by normalizing the logarithmic deviation of luciferase activity, where deviation refers to the
change of luciferase activity between the wild-type and the mutated (one motif family at a
time) regulatory region (see below for a demonstration). Using the logarithmic deviation
allowed us to account for the differences in effect sizes of various motif families by rescaling
the differences to a comparable range. Similarly, for each of the nine genes, the regression
coefficient of its expression level on a relevant regulatory region was estimated by normalizing
the logarithmic deviation of luciferase activity, where deviation refers to the change of
luciferase activity compared to the empty vector controls. All Matlab source codes are
available at https://github.com/Huange and also http://burrn-sim.stemcells.cam.ac.uk/.

Detailed explanation of the modelling of each tier of the DBN:

a) Estimating the discrete probability distribution of a motif variable

The probability of a motif family at a given regulatory region taking value 1 or 2 (i.e., being
unbound or bound) was calculated based on: (i) the number of such motifs in that regulatory
region; (ii) the expression levels of the relevant TFs.

For example, three Ebox motifs were found at \( \text{Erg}+65 \) (Fig. 3a). They can be bound by either
TAL1 or LYL1. Thus, we assigned that \( P(\text{Ebox}@\text{Erg}+65=1) \) and \( P(\text{Ebox}@\text{Erg}+65=2) \) were
determined by \{3, TAL1, LYL1\}. We assumed that (i) the expression level of a TF is
proportional to the probability of that TF binding to a target motif; and (ii) the bindings of TFs
to multiple motifs are independent events. Gene expression levels were defined within the
closed interval \([0, 1]\), which is identical to the possible range of probabilities. For ease of
calculation, we took the expression level of a TF as its probability of binding to a motif.

Accordingly, we have

\[
P(\text{Ebox}@\text{Erg}+65=1) = (1 - p)^3 \times (1 - q)^3
\]

\[
P(\text{Ebox}@\text{Erg}+65=2) = \sum_{n=1}^{3} C(3, n) \times p^n \times (1 - p)^{3-n} \times (1 - q)^3
\]
\begin{align*}
+ \sum_{n=1}^{3} C(3,n) \times q^n \times (1-q)^{(3-n)} \times (1-p)^3 \\
+ \sum_{n=1}^{2} \sum_{m=1}^{3-n} C(3,n) \times p^n \times (1-p)^{(3-n)} \times C((3-n),m) \times q^m \times (1-q)^{(3-m)}
\end{align*}

where \( p \) and \( q \) represent the expression levels of TAL1 and LYL1, respectively.

However, to remove the bias introduced by simply taking the expression level of a TF as its probability of binding to a motif, we further normalized the resulting probabilities as below:

\( \tilde{Z} = \tilde{P}(\text{Ebox@Erg}+65=1) + \tilde{P}(\text{Ebox@Erg}+65=2) \)  

\( P(\text{Ebox@Erg}+65=1) = \tilde{P}(\text{Ebox@Erg}+65=1)/\tilde{Z} \)  

\( P(\text{Ebox@Erg}+65=2) = \tilde{P}(\text{Ebox@Erg}+65=2)/\tilde{Z} \)  

It should be mentioned that the number of the same motifs in a regulatory region was directly taken into account in the estimation of probabilities. One may raise the question of whether this number has such strong power. Specifically, should the exponents in equations (1) and (2) change linearly, or less than linearly, along with the increase in the number of Ebox motifs? To address this issue, we replaced all exponents with their square roots and rerun the whole set of simulations (data not shown). Results showed that using the square roots instead of the original numbers (i) caused a more evenly distributed expression of the nine TFs over the hypothetical interval \([0, 1]\), (ii) captured the same trend in gene expression changes in some perturbations (e.g. the AML-ETO simulation), but (iii) led to decreased expression levels of certain TFs in other perturbations (e.g. PU.1 knockdown and GFI1B over-expression), which therefore disagrees with the experimental data. In order to capture a better agreement of computational and experimental results, we directly used the number of motifs to estimate the discrete probability distributions.

\textit{b) Estimating the activity of a regulatory region}
The regression coefficient of a regulatory region on a motif family was estimated by normalizing the logarithmic deviation of luciferase activity, e.g. comparing the change of luciferase activity between the wild-type and mutated constructs. For example, when the luciferase activity for the wild-type Erg+65 region was set to 100 %, the simultaneous mutation of all Ebox or Gfi motifs at this region resulted in increased luciferase activity (181.2 % or 475.9 %, respectively) (Fig. 3b). In contrast, simultaneous mutation of all Ets or Gata motifs at this region led to reduced luciferase activity (1.3 % or 14.5 %, respectively). Based on this information, we estimated the regression coefficient of the Erg+65 region on a relevant motif family in the following way:

\[
\alpha_i = \log \left( \frac{100}{l_i} \right) \times \left( \sum_k \log \left( \frac{100}{l_i} \right) \right)^{-1}
\]

(6)

where \( k \in \{1, \ldots, 4\}, \ l_1 = 181.2, \ l_2 = 475.9, \ l_3 = 1.3, \ l_4 = 14.5 \); accordingly, \( \alpha_1 = -0.070, \ \alpha_2 = -0.185, \ \alpha_3 = 0.515, \ \alpha_4 = 0.230 \). We can then formulate a linear regression equation as:

\[
\tilde{y} = \alpha_1 x_1 + \alpha_2 x_2 + \alpha_3 x_3 + \alpha_4 x_4
\]

(7)

where \( \tilde{y} \) denote the estimated luciferase activity of Erg+65, and \( x_1, x_2, x_3 \) and \( x_4 \) represent the binding status of Ebox, Gfi, Ets and Gata motifs at Erg+65.

However, the minimum and maximum \( \tilde{y} \) obtained by the above formula are 0.235 (when \( x_1 = x_2 = 2 \) and \( x_3 = x_4 = 1 \)) and 1.235 (when \( x_1 = x_2 = 1 \) and \( x_3 = x_4 = 2 \)). To make the values of \( \tilde{y} \) fall in the desired closed interval \([0, 1]\), an intercept of -0.235 has to be introduced into the linear regression model. In addition, a disturbance term has been included in the model in order to satisfy the generic assumption of conditional linear Gaussian distribution. Finally, the fully defined linear regression model regarding Erg+65 is given as:

\[
\tilde{y} = c + \alpha_1 x_1 + \alpha_2 x_2 + \alpha_3 x_3 + \alpha_4 x_4 + \epsilon
\]

(8)
where \( c = -0.235 \), \( \epsilon \sim N(0, \sigma^2) \), and \( \sigma \) should be a very small value.

\[ c = -0.235, \quad \epsilon \sim N(0, \sigma^2), \quad \text{and} \quad \sigma \text{ should be a very small value.} \]

\[ \]  

**c) Estimating the expression level of a gene**

For each gene studied, the regression coefficient of its expression level on a relevant regulatory region was estimated by normalizing the logarithmic deviation of luciferase activity, where deviation refers to the change of luciferase activity compared to an empty vector control.

For example, when setting the luciferase activity of the wild-type constructs to 100 %, the luciferase activity of the empty vector controls relative to Erg+65, Erg+75 and Erg+85 wild-types are 1.9 %, 1.0 % and 15.2 %, respectively (Fig. 3b, Fig. 3-figure supplement 1 and 2).

Based on these data, we estimated the expression level of Erg on a relevant regulatory region in the following way:

\[
\beta_i = \log \left( \frac{100}{l_k} \right) \times \left( \sum \log \left( \frac{100}{l_k} \right) \right)^{-1}
\]

(9)

where \( k \in \{1, 2, 3\} \), \( l_1 = 1.9 \), \( l_2 = 1.0 \), \( l_3 = 15.2 \); accordingly, \( \beta_1 = 0.379 \), \( \beta_2 = 0.441 \), \( \beta_3 = 0.180 \). We can then formulate a linear regression equation as below:

\[
\tilde{z} = \beta_1 \tilde{y}_1 + \beta_2 \tilde{y}_2 + \beta_3 \tilde{y}_3
\]

(10)

where \( \tilde{z} \) denote the estimated expression level of Erg; and \( \tilde{y}_1 \), \( \tilde{y}_2 \) and \( \tilde{y}_3 \) represent the estimated activities of Erg+65, Erg+75 and Erg+85. Again, a disturbance term has been introduced to the model in order to meet the generic assumption of conditional linear Gaussian distribution. Thus, the fully defined linear regression model regarding Erg is given as:

\[
\tilde{z} = \beta_1 \tilde{y}_1 + \beta_2 \tilde{y}_2 + \beta_3 \tilde{y}_3 + \epsilon
\]

(11)

where \( \epsilon \sim N(0, \sigma^2) \) and \( \sigma \) should be a very small value.

\[ \]

**Statistics**
Significance for the results of the luciferase reporter assays was calculated by combining the p-values of each experiment (generated by using the t-test function in Excel) using the Fisher’s method, followed by the calculation of Stouffer’s z trend if necessary. Significance tests for changes in TF expression levels caused by TF perturbations (both computational and experimental) were evaluated by Wilcoxon rank-sum tests.

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Competing Interests

The authors declare that no competing interests exist.

References


Figure legends

Figure 1: Identification of haematopoietic active cis-regulatory regions. (a) UCSC screenshot of the Erg gene locus for ChIP-sequencing data for nine haematopoietic TFs (ERG, FLI1, GATA2, GFI1B, LYL1, MEIS1, PU.1, RUNX1 and TAL1 (15)) and for H3K27ac (27) in HPC7 cells. Highlighted are all regions of the Erg gene locus that are acetylated at H3K27 and are bound by three or more TFs. Numbers indicate the distance (in kb) from the ATG start codon. (b) Summary of the identification of candidate cis-regulatory regions for all nine TFs and subsequent analysis in transgenic mouse assays. The inspection of the nine gene loci and the application of the selection criteria (≥ 3 TFs bound and H3K27ac) identified a total of 49 candidate cis-regulatory regions. The heatmap shows the binding pattern of the nine TFs to all candidate regulatory elements in HPC7 cells: green = bound, grey = unbound. Haematopoietic activity in E11.5 transgenic mice is indicated by the font color: black = active, red = not active. Grey indicates genomic repeat regions that were not tested in transgenic mice. Detailed experimental data corresponding to the summary heatmap can be found in Fig. 1-figure supplement 1-8. (c) Haematopoietic activity of the five candidate Erg cis-regulatory regions was determined in E11.5 transgenic mouse assays. Shown are X-Gal-stained whole-mount embryos and paraffin sections of the dorsal aorta (DA, ventral side on the left/top) and foetal liver (FL), sites of definitive haematopoiesis. Colour coding as in B.
Figure 1 – figure supplement 1: Identification of haematopoietic active cis-regulatory elements for *Fli1*. (a) The candidate cis-regulatory elements were identified by ChIP-Seq analysis of the TFs ERG, FLI1, GATA2, GFI1B, LYL1, MEIS1, PU.1, RUNX1 and TAL1 as well as H3K27 acetylation in the haematopoietic stem/progenitor cell line HPC7. Highlighted in pink are the candidate cis-regulatory regions which are bound by at least three of the nine TFs and showed H3K27 acetylation. The numbering represents the direction and distance in kilobases from the start codon ATG (pro = promoter). (b) Candidate regions were assayed for haematopoietic enhancer activity in mouse transient transgenic embryos. X-Gal stained whole-mount E11.5 embryos and paraffin sections of the dorsal aorta (DA; longitudinal section, ventral side on the left/top) and foetal liver (FL) are shown for the candidate cis-regulatory regions. Transgenic mouse data are not shown for previously published regions, but relevant publications are listed.

Figure 1 – figure supplement 2: Identification of haematopoietic active cis-regulatory elements for *Gata2*. (a) The candidate cis-regulatory elements were identified by ChIP-Seq analysis of the TFs ERG, FLI1, GATA2, GFI1B, LYL1, MEIS1, PU.1, RUNX1 and TAL1 as well as H3K27 acetylation in the haematopoietic stem/progenitor cell line HPC7. Highlighted in pink are the candidate cis-regulatory regions which are bound by at least three of the nine TFs and showed H3K27 acetylation. The numbering represents the direction and distance in kilobases from the start codon ATG (pro = promoter). (b) Candidate regions were assayed for haematopoietic enhancer activity in mouse transient transgenic embryos. X-Gal stained whole-mount E11.5 embryos and paraffin sections of the dorsal aorta (DA; longitudinal section, ventral side on the left/top) and foetal liver (FL) are shown for the candidate cis-regulatory
regions. Transgenic mouse data are not shown for previously published regions, but relevant publications are listed.

**Figure 1 – figure supplement 3: Identification of haematopoietic active *cis*-regulatory elements for *Gfi1b*.** (a) The candidate *cis*-regulatory elements were identified by ChIP-Seq analysis of the TFs ERG, FLI1, GATA2, GFI1B, LYL1, MEIS1, PU.1, RUNX1 and TAL1 as well as H3K27 acetylation in the haematopoietic stem/progenitor cell line HPC7. Highlighted in pink are the candidate *cis*-regulatory regions which are bound by at least three of the nine TFs and showed H3K27 acetylation. The numbering represents the direction and distance in kilobases from the start codon ATG (pro = promoter). (b) All candidate regions were previously published regions. Relevant publications are listed.

**Figure 1 – figure supplement 4: Identification of haematopoietic active *cis*-regulatory elements for *Lyl1*.** (a) The candidate *cis*-regulatory elements were identified by ChIP-Seq analysis of the TFs ERG, FLI1, GATA2, GFI1B, LYL1, MEIS1, PU.1, RUNX1 and TAL1 as well as H3K27 acetylation in the haematopoietic stem/progenitor cell line HPC7. Highlighted in pink are the candidate *cis*-regulatory regions which are bound by at least three of the nine TFs and showed H3K27 acetylation. The numbering represents the direction and distance in kilobases from the start codon ATG (pro = promoter). (b) Candidate regions were assayed for haematopoietic enhancer activity in mouse transient transgenic embryos. X-Gal stained whole-mount E11.5 embryos and paraffin sections of the dorsal aorta (DA; longitudinal section, ventral side on the left/top) and foetal liver (FL) are shown for the candidate *cis*-regulatory regions. Transgenic mouse data are not shown for previously published regions, but relevant publications are listed.
Figure 1 – figure supplement 5: Identification of haematopoietic active cis-regulatory elements for *Meis1*. (a) The candidate cis-regulatory elements were identified by ChIP-Seq analysis of the TFs ERG, FLI1, GATA2, GFI1B, LYL1, MEIS1, PU.1, RUNX1 and TAL1 as well as H3K27 acetylation in the haematopoietic stem/progenitor cell line HPC7. Highlighted in pink are the candidate cis-regulatory regions which are bound by at least three of the nine TFs and showed H3K27 acetylation. The numbering represents the direction and distance in kilobases from the start codon ATG (pro = promoter). (b) Candidate regions were assayed for haematopoietic enhancer activity in mouse transient transgenic embryos. X-Gal stained whole-mount E11.5 embryos and paraffin sections of the dorsal aorta (DA; longitudinal section, ventral side on the left/top) and foetal liver (FL) are shown for the candidate cis-regulatory regions.

Figure 1 – figure supplement 6: Identification of haematopoietic active cis-regulatory elements for *Runx1*. (a) The candidate cis-regulatory elements were identified by ChIP-Seq analysis of the TFs ERG, FLI1, GATA2, GFI1B, LYL1, MEIS1, PU.1, RUNX1 and TAL1 as well as H3K27 acetylation in the haematopoietic stem/progenitor cell line HPC7. Highlighted in pink are the candidate cis-regulatory regions which are bound by at least three of the nine TFs and showed H3K27 acetylation. The numbering represents the direction and distance in kilobases from the start codon ATG (pro = promoter). (b) E10 embryos and cryosections of the DA (transverse; ventral down) and FL are shown. For the *Runx1*+204 region, a larger 12 kb fragment (chr16:92,620,915-92,631,936, mm9) was used for transient transgenesis, but similar results were obtained with the +204 fragment alone (data not shown). The +24 element was tested in conjunction with the +23 and did not change its tissue specificity (Bee et al., 2010).
Preliminary data show that the +24 on its own does not mediate robust tissue specific expression of reporter genes. Transgenic mouse data are not shown for previously published regions, but relevant publications are listed.

Figure 1 – figure supplement 7: Identification of haematopoietic active cis-regulatory elements for Spi1. (a) The candidate cis-regulatory elements were identified by ChIP-Seq analysis of the TFs ERG, FLI1, GATA2, GFI1B, LYL1, MEIS1, PU.1, RUNX1 and TAL1 as well as H3K27 acetylation in the haematopoietic stem/progenitor cell line HPC7. Highlighted in pink are the candidate cis-regulatory regions which are bound by at least three of the nine TFs and showed H3K27 acetylation. The numbering represents the direction and distance in kilobases from the start codon ATG (pro = promoter). (b) All candidate regions were previously published regions. Relevant publications are listed.

Figure 1 – figure supplement 8: Identification of haematopoietic active cis-regulatory elements for Tal1. (a) The candidate cis-regulatory elements were identified by ChIP-Seq analysis of the TFs ERG, FLI1, GATA2, GFI1B, LYL1, MEIS1, PU.1, RUNX1 and TAL1 as well as H3K27 acetylation in the haematopoietic stem/progenitor cell line HPC7. Highlighted in pink are the candidate cis-regulatory regions which are bound by at least three of the nine TFs and showed H3K27 acetylation. The numbering is based on the distance (in kb) to promoter 1a. (b) All candidate regions were previously published regions. Relevant publications are listed.

Figure 1 – figure supplement 9: Number of PCR and LacZ positive transgenic embryos (E10.5-11.5) for each regulatory region.
Figure 2: Comparison of TF binding pattern at haematopoietic active cis-regulatory regions in two haematopoietic progenitor cell lines, HPC7 and 416b. (a) UCSC screenshot of the Erg gene locus for ChIP-Sequencing data for nine haematopoietic TFs (ERG, FLI1, GATA2, GFI1B, LYL1, MEIS1, PU.1, RUNX1 and TAL1) and for H3K27ac in 416b cells. Highlighted are those haematopoietic active Erg cis-regulatory regions that were identified based on acetylation of H3K27 and TF binding in HPC7 cells followed by transgenic mouse assays. Numbers indicate the distance (in kb) from the ATG start codon. (b) Hierarchical clustering of the binding profiles for HPC7, 416b and other published datasets. The heatmap shows the pairwise correlation coefficient of peak coverage data between pairs of samples in the row and column. The order of the samples is identical in columns and rows. Details about samples listed can be found in Fig. 2-figure supplement 9. (c) Pair-wise analysis of binding of the nine TFs to haematopoietic active cis-regulatory regions of the nine TFs in HPC7 versus 416b cells. Green = bound in both cells types, blue = only bound in 416b cells, orange = only bound in HPC7 cells, grey = not bound in either cell type.

Figure 2 – figure supplement 1: UCSC screenshot for the Fli1 gene locus demonstrating binding patterns for nine key haematopoietic TFs and H3K27ac in 416b cells. Highlighted in pink are cis-regulatory regions that were identified based on the selection criteria (≥ 3 TFs bound and H3K27ac) in HPC7 cells and were shown to possess haematopoietic activity. The numbering represents the distance (in kb) from the start codon ATG.

Figure 2 – figure supplement 2: UCSC screenshot for the Gata2 gene locus demonstrating binding patterns for nine key haematopoietic TFs and H3K27ac in 416b cells. Highlighted
in pink are *cis*-regulatory regions that were identified based on the selection criteria (≥ 3 TFs bound and H3K27ac) in HPC7 cells and were shown to possess haematopoietic activity. The numbering represents the distance (in kb) from the start codon ATG.

Figure 2 – figure supplement 3: UCSC screenshot for the *Gfi1b* gene locus demonstrating binding patterns for nine key haematopoietic TFs and H3K27ac in 416b cells. Highlighted in pink are *cis*-regulatory regions that were identified based on the selection criteria (≥ 3 TFs bound and H3K27ac) in HPC7 cells and were shown to possess haematopoietic activity. The numbering represents the distance (in kb) from the start codon ATG.

Figure 2 – figure supplement 4: UCSC screenshot for the *Lyl1* gene locus demonstrating binding patterns for nine key haematopoietic TFs and H3K27ac in 416b cells. Highlighted in pink is the promoter (“pro”) that was identified based on the selection criteria (≥ 3 TFs bound and H3K27ac) in HPC7 cells and was shown to possess haematopoietic activity. The promoter is labelled with “pro”.

Figure 2 – figure supplement 5: UCSC screenshot for the *Meis1* gene locus demonstrating binding patterns for nine key haematopoietic TFs and H3K27ac in 416b cells. Highlighted in pink is the *cis*-regulatory region that was identified based on the selection criteria (≥ 3 TFs bound and H3K27ac) in HPC7 cells and was shown to possess haematopoietic activity. The numbering represents the distance (in kb) from the start codon ATG.
Figure 2 – figure supplement 6: UCSC screenshot for the Runx1 gene locus demonstrating binding patterns for nine key haematopoietic TFs and H3K27ac in 416b cells. Highlighted in pink are cis-regulatory regions that were identified based on the selection criteria (≥ 3 TFs bound and H3K27ac) in HPC7 cells and were subsequently shown to possess haematopoietic activity. The numbering represents the distance (in kb) from the start codon ATG.

Figure 2 – figure supplement 7: UCSC screenshot for the Spi1 gene locus demonstrating binding patterns for nine key haematopoietic TFs and H3K27ac in 416b cells. Highlighted in pink is the cis-regulatory region that was identified based on the selection criteria (≥ 3 TFs bound and H3K27ac) in HPC7 cells and was shown to possess haematopoietic activity. The numbering represents the distance (in kb) from the start codon ATG.

Figure 2 – figure supplement 8: UCSC screenshot for the Tal1 gene locus demonstrating binding patterns for nine key haematopoietic TFs and H3K27ac in 416b cells. Highlighted in pink are cis-regulatory regions that were identified based on the selection criteria (≥ 3 TFs bound and H3K27ac) in HPC7 cells and were shown to possess haematopoietic activity. The numbering represents the distance (in kb) from promoter 1a.

Figure 2 – figure supplement 9: List of ChIP-Seq samples included in the heatmap in Figure 2b.
Figure 3: TFBS mutagenesis reveals enhancer-dependent effects of TF binding on gene expression. (a) Multiple species alignment of mouse (mm9), human (hg19), dog (canFam2), opossum (monDom5) and platypus (ornAna1) sequences for the Erg+65 region. Nucleotides highlighted in black are conserved between all species analysed, nucleotides highlighted in grey are conserved between four of five species. Transcription factor binding sites (TFBS) are highlighted in: blue = Ebox, purple = Ets, green = Gata, yellow = Gfi, red = Meis. The nucleotides changed to mutate the TFBSs are indicated below the alignment. All binding sites of one motif family (e.g. all Ebox motifs) were mutated simultaneously. (b) Luciferase assay for the Erg+65 wild-type and mutant enhancer in stably transfected 416b cells. Each bar represents the averages of at least three independent experiments with three to four replicates within each experiment. Results are shown relative to the wild-type enhancer activity, which is set to 100%. Error bars represent the standard error of the mean (SEM). Stars indicate significance: ** = p-value < 0.01, *** = p-value < 0.001. P-values were calculated using t-tests, followed by the Fisher's method. (c) Summary of luciferase assay results for all 19 high-confidence haematopoietic active regulatory regions. Relative luciferase activity is illustrated in shades of blue (down-regulation) and red (up-regulation). Crossed-out grey boxes indicate that there is no motif for the TF and/or the TF does not bind to the region. Detailed results and corresponding alignments with highlighted TFBSs and their mutations can be found in Figure 3-figure supplements 1-18.

Figure 3 – figure supplement 1: Multiple species alignment and luciferase assay results for Erg+75. (a) Multiple species alignment (MSA) with the following species: mouse (mm9), human (hg19), dog (canFam2), opossum (monDom5) and platypus (ornAna1). Nucleotides highlighted in black are conserved between all species analysed, nucleotides highlighted in grey are conserved between four of five species. Transcription factor binding sites (TFBS) are...
highlighted in: blue = Ebox, purple = Ets, yellow = Gfi. The nucleotides that were changed to mutate the TFBSs are indicated below the MSA. All conserved binding sites of one motif family (e.g. all Ebox motifs) were mutated simultaneously. Where TF binding was observed in ChIP-Seq experiments in 416b cells, but the TFBS was not conserved, the motifs present in the mouse sequence only were mutated. (b) For the luciferase reporter assays in stably transfected 416b cells the averages of at least three independent experiments with three to four replicates within each experiment are shown. Error bars represent the standard error of the mean (SEM). Stars indicate significance: ** = p-value < 0.01, *** = p-value < 0.001. P-values were generated using t-tests, followed by the Fisher’s method and if necessary Stouffer’s z trend.

Figure 3 – figure supplement 2: Multiple species alignment and luciferase assay results for Erg+85. (a) Multiple species alignment (MSA) with the following species: mouse (mm9), human (hg19), dog (canFam2), opossum (monDom5) and platypus (ornAna1). Nucleotides highlighted in black are conserved between all species analysed, nucleotides highlighted in grey are conserved between four of five species. Transcription factor binding sites (TFBS) are highlighted in: blue = Ebox, purple = Ets, green = Gata, yellow = Gfi. The nucleotides that were changed to mutate the TFBSs are indicated below the MSA. All conserved binding sites of one motif family (e.g. all Ebox motifs) were mutated simultaneously. (b) For the luciferase reporter assays in stably transfected 416b cells the averages of at least three independent experiments with three to four replicates within each experiment are shown. Error bars represent the standard error of the mean (SEM). Stars indicate significance: *** = p-value < 0.001. P-values were generated using t-tests, followed by the Fisher’s method and if necessary Stouffer’s z trend.
Figure 3 – figure supplement 3: Multiple species alignment and luciferase assay results for *Fli1*+12. (a) Multiple species alignment (MSA) with the following species: mouse (mm9), human (hg19), dog (canFam2), opossum (monDom5) and platypus (ornAna1). Nucleotides highlighted in black are conserved between all species analysed, nucleotides highlighted in grey are conserved between four of five species. Transcription factor binding sites (TFBS) are highlighted in: blue = Ebox, purple = Ets. The nucleotides that were changed to mutate the TFBSs are indicated below the MSA. All conserved binding sites of one motif family (e.g. all Ebox motifs) were mutated simultaneously. (b) For the luciferase reporter assays in stably transfected 416b cells the averages of at least three independent experiments with three to four replicates within each experiment are shown. Error bars represent the standard error of the mean (SEM). Stars indicate significance: *** = p-value < 0.001. P-values were generated using t-tests, followed by the Fisher’s method and if necessary Stouffer’s z trend.

Figure 3 – figure supplement 4: Multiple species alignment and luciferase assay results for *Gata2*-93. (a) Multiple species alignment (MSA) with the following species: mouse (mm9), human (hg19), dog (canFam2), opossum (monDom5) and platypus (ornAna1). Nucleotides highlighted in black are conserved between all species analysed, nucleotides highlighted in grey are conserved between four of five species. Transcription factor binding sites (TFBS) are highlighted in: blue = Ebox, purple = Ets, green = Gata, red = Meis, turquoise = Runt. The nucleotides that were changed to mutate the TFBSs are indicated below the MSA. All conserved binding sites of one motif family (e.g. all Ebox motifs) were mutated simultaneously. (b) For the luciferase reporter assays in stably transfected 416b cells the averages of at least three independent experiments with three to four replicates within each experiment are shown. Error bars represent the standard error of the mean (SEM). Stars
Figure 3 – figure supplement 5: Multiple species alignment and luciferase assay results for Gata2+3. (a) Multiple species alignment (MSA) with the following species: mouse (mm9), human (hg19), dog (canFam2), opossum (monDom5) and platypus (ornAna1). Nucleotides highlighted in black are conserved between all species analysed, nucleotides highlighted in grey are conserved between four of five species. Transcription factor binding sites (TFBS) are highlighted in: blue = Ebox, purple = Ets, green = Gata. The nucleotides that were changed to mutate the TFBSs are indicated below the MSA. All conserved binding sites of one motif family (e.g. all Ebox motifs) were mutated simultaneously. (b) For the luciferase reporter assays in stably transfected 416b cells the averages of at least three independent experiments with three to four replicates within each experiment are shown. Error bars represent the standard error of the mean (SEM). Stars indicate significance: *** = p-value < 0.001. P-values were generated using t-tests, followed by the Fisher’s method and if necessary Stouffer’s z trend.

Figure 3 – figure supplement 6: Multiple species alignment and luciferase assay results for Gfi1b+16. (a) Multiple species alignment (MSA) with the following species: mouse (mm9), human (hg19), dog (canFam2), opossum (monDom5) and platypus (ornAna1). Nucleotides highlighted in black are conserved between all species analysed, nucleotides highlighted in grey are conserved between four of five species. Transcription factor binding sites (TFBS) are highlighted in: blue = Ebox, purple = Ets, green = Gata, yellow = Gfi, red = Meis, turquoise = Runt. The nucleotides that were changed to mutate the TFBSs are indicated.
below the MSA. All conserved binding sites of one motif family (e.g. all Ebox motifs) were 
mutated simultaneously. Where TF binding was observed in ChIP-Seq experiments in 416b 
cells, but the TFBS was not conserved, the motifs present in the mouse sequence only were 
mutated. (b) For the luciferase reporter assays in stably transfected 416b cells the averages of 
at least three independent experiments with three to four replicates within each experiment are 
shown. Error bars represent the standard error of the mean (SEM). Stars indicate significance: 
** = p-value < 0.01, *** = p-value < 0.001. P-values were generated using t-tests, followed by 
the Fisher’s method and if necessary Stouffer’s z trend.

Figure 3 – figure supplement 7: Multiple species alignment and luciferase assay results 
for Gfi1b+17. (a) Multiple species alignment (MSA) with the following species: mouse 
(mm9), human (hg19), dog (canFam2), opossum (monDom5) and platypus (ornAna1). 
Nucleotides highlighted in black are conserved between all species analysed, nucleotides 
highlighted in grey are conserved between four of five species. Transcription factor binding 
sites (TFBS) are highlighted in: blue = Ebox, purple = Ets, green = Gata, yellow = Gfi, red = 
Meis. The nucleotides that were changed to mutate the TFBSs are indicated below the MSA. 
All conserved binding sites of one motif family (e.g. all Ebox motifs) were mutated 
simultaneously. (b) For the luciferase reporter assays in stably transfected 416b cells the 
averages of at least three independent experiments with three to four replicates within each 
experiment are shown. Error bars represent the standard error of the mean (SEM). Stars 
indicate significance: ** = p-value < 0.01, *** = p-value < 0.001. P-values were generated 
using t-tests, followed by the Fisher’s method and if necessary Stouffer’s z trend.
Figure 3 – figure supplement 8: Multiple species alignment and luciferase assay results for Lyl1 promoter. (a) Multiple species alignment (MSA) with the following species: mouse (mm9), human (hg19), dog (canFam2) and opossum (monDom5). Nucleotides highlighted in black are conserved between all species analysed, nucleotides highlighted in grey are conserved between three of four species. Transcription factor binding sites (TFBS) are highlighted in: purple = Ets, green = Gata. The nucleotides that were changed to mutate the TFBSs are indicated below the MSA. All conserved binding sites of one motif family (e.g. all Ets motifs) were mutated simultaneously. (b) For the luciferase reporter assays in stably transfected 416b cells the averages of at least three independent experiments with three to four replicates within each experiment are shown. Error bars represent the standard error of the mean (SEM). Stars indicate significance: *** = p-value < 0.001. P-values were generated using t-tests, followed by the Fisher’s method and if necessary Stouffer’s z trend.

Figure 3 – figure supplement 9: Multiple species alignment and luciferase assay results for Meis1+48. (a) Multiple species alignment (MSA) with the following species: mouse (mm9), human (hg19), dog (canFam2), opossum (monDom5) and platypus (ornAna1). Nucleotides highlighted in black are conserved between all species analysed, nucleotides highlighted in grey are conserved between four of five species. Transcription factor binding sites (TFBS) are highlighted in: purple = Ets, green = Gata, yellow = Gfi, red = Meis. The nucleotides that were changed to mutate the TFBSs are indicated below the MSA. All conserved binding sites of one motif family (e.g. all Ets motifs) were mutated simultaneously. (b) For the luciferase reporter assays in stably transfected 416b cells the averages of at least three independent experiments with three to four replicates within each experiment are shown. Error bars represent the standard error of the mean (SEM). Stars indicate significance: *** = p-value < 0.001.
p-value < 0.001. P-values were generated using t-tests, followed by the Fisher’s method and if necessary Stouffer’s z trend.

Figure 3 – figure supplement 10: Multiple species alignment and luciferase assay results for Spi1-14. (a) Multiple species alignment (MSA) with the following species: mouse (mm9), human (hg19), dog (canFam2), opossum (monDom5) and platypus (ornAna1). Nucleotides highlighted in black are conserved between all species analysed, nucleotides highlighted in grey are conserved between four of five species. Transcription factor binding sites (TFBS) are highlighted in: blue = Ebox, purple = Ets, turquoise = Runt. The nucleotides that were changed to mutate the TFBSs are indicated below the MSA. All conserved binding sites of one motif family (e.g. all Ebox motifs) were mutated simultaneously. (b) For the luciferase reporter assays in stably transfected 416b cells the averages of at least three independent experiments with three to four replicates within each experiment are shown. Error bars represent the standard error of the mean (SEM). Stars indicate significance: ** = p-value < 0.01, *** = p-value < 0.001. P-values were generated using t-tests, followed by the Fisher’s method and if necessary Stouffer’s z trend.

Figure 3 – figure supplement 11: Multiple species alignment and luciferase assay results for Runx1-59. (a) Multiple species alignment (MSA) with the following species: mouse (mm9), human (hg19) and dog (canFam2). Nucleotides highlighted in black are conserved between all species analysed, nucleotides highlighted in grey are conserved between two of three species. Transcription factor binding sites (TFBS) are highlighted in: blue = Ebox, purple = Ets, green = Gata, red = Meis. The nucleotides that were changed to mutate the TFBSs are indicated below the MSA. All conserved binding sites of one motif family (e.g. all Ebox
motifs) were mutated simultaneously. (b) For the luciferase reporter assays in stably transfected 416b cells the averages of at least three independent experiments with three to four replicates within each experiment are shown. Error bars represent the standard error of the mean (SEM). Stars indicate significance: *** = p-value < 0.001. P-values were generated using t-tests, followed by the Fisher’s method and if necessary Stouffer’s z trend.

Figure 3 – figure supplement 12: Multiple species alignment and luciferase assay results for Runx1+3. (a) Multiple species alignment (MSA) with the following species: mouse (mm9), human (hg19), dog (canFam2), opossum (monDom5) and platypus (ornAna1). Nucleotides highlighted in black are conserved between all species analysed, nucleotides highlighted in grey are conserved between four of five species. Transcription factor binding sites (TFBS) are highlighted in: blue = Ebox, purple = Ets, green = Gata, yellow = Gfi, red = Meis, turquoise = Runt. The nucleotides that were changed to mutate the TFBSs are indicated below the MSA. All conserved binding sites of one motif family (e.g. all Ets motifs) were mutated simultaneously. Where TF binding was observed in ChIP-Seq experiments in 416b cells, but the TFBS was not conserved, the motifs present in the mouse sequence only were mutated. (b) For the luciferase reporter assays in stably transfected 416b cells the averages of at least three independent experiments with three to four replicates within each experiment are shown. Error bars represent the standard error of the mean (SEM). Stars indicate significance: * = p-value < 0.05, ** = p-value < 0.01, *** = p-value < 0.001. P-values were generated using t-tests, followed by the Fisher’s method and if necessary Stouffer’s z trend.

Figure 3 – figure supplement 13: Multiple species alignment and luciferase assay results for Runx1+23. (a) Multiple species alignment (MSA) with the following species: mouse
(mm9), human (hg19), dog (canFam2) and opossum (monDom5). Nucleotides highlighted in black are conserved between all species analysed, nucleotides highlighted in grey are conserved between three to four species. Transcription factor binding sites (TFBS) are highlighted in: blue = Ebox, purple = Ets, green = Gata, red = Meis, turquoise = Runt. The nucleotides that were changed to mutate the TFBSs are indicated below the MSA. All conserved binding sites of one motif family (e.g. all Ebox motifs) were mutated simultaneously. ** For the luciferase reporter assays in stably transfected 416b cells the averages of at least three independent experiments with three to four replicates within each experiment are shown. Error bars represent the standard error of the mean (SEM). Stars indicate significance: * = p-value < 0.05, *** = p-value < 0.001. P-values were generated using t-tests, followed by the Fisher’s method and if necessary Stouffer’s z trend.

**Figure 3 – figure supplement 14: Multiple species alignment and luciferase assay results for Runx1+110. (a) Multiple species alignment (MSA) with the following species: mouse (mm9), human (hg19), dog (canFam2), opossum (monDom5) and platypus (ornAna1). Nucleotides highlighted in black are conserved between all species analysed, nucleotides highlighted in grey are conserved between four of five species. Transcription factor binding sites (TFBS) are highlighted in: blue = Ebox, purple = Ets, green = Gata. The nucleotides that were changed to mutate the TFBSs are indicated below the MSA. All conserved binding sites of one motif family (e.g. all Ets motifs) were mutated simultaneously. Where TF binding was observed in ChIP-Seq experiments in 416b cells, but the TFBS was not conserved, the motifs present in the mouse sequence only were mutated. (b) For the luciferase reporter assays in stably transfected 416b cells the averages of at least three independent experiments with three to four replicates within each experiment are shown. Error bars represent the standard error of the mean (SEM). Stars indicate significance: ** = p-value < 0.01, *** = p-value < 0.001. P-
values were generated using t-tests, followed by the Fisher’s method and if necessary Stouffer’s z trend.

Figure 3 – figure supplement 15: Multiple species alignment and luciferase assay results for Runx1+204. (a) Multiple species alignment (MSA) with the following species: mouse (mm9), human (hg19), dog (canFam2), opossum (monDom5) and platypus (ornAna1). Nucleotides highlighted in black are conserved between all species analysed, nucleotides highlighted in grey are conserved between four of five species. Transcription factor binding sites (TFBS) are highlighted in: blue = Ebox, purple = Ets, yellow = Gfi, turquoise = Runt. The nucleotides that were changed to mutate the TFBSs are indicated below the MSA. All conserved binding sites of one motif family (e.g. all Ets motifs) were mutated simultaneously. Where TF binding was observed in ChIP-Seq experiments in 416b cells, but the TFBS was not conserved, the motifs present in the mouse sequence only were mutated. (b) For the luciferase reporter assays in stably transfected 416b cells the averages of at least three independent experiments with three to four replicates within each experiment are shown. Error bars represent the standard error of the mean (SEM). Stars indicate significance: *** = p-value < 0.001. P-values were generated using t-tests, followed by the Fisher’s method and if necessary Stouffer’s z trend.

Figure 3 – figure supplement 16: Multiple species alignment and luciferase assay results for Tal1-4. (a) Multiple species alignment (MSA) with the following species: mouse (mm9), human (hg19) and dog (canFam2). Nucleotides highlighted in black are conserved between all species analysed, nucleotides highlighted in grey are conserved between two of three species. Transcription factor binding sites (TFBS) are highlighted in: purple = Ets. The nucleotides that
were changed to mutate the TFBSs are indicated below the MSA. All conserved binding sites of the Ets motif family were mutated simultaneously. (b) For the luciferase reporter assays in stably transfected 416b cells the averages of at least three independent experiments with three to four replicates within each experiment are shown. Error bars represent the standard error of the mean (SEM). Stars indicate significance: ** = p-value < 0.01. P-values were generated using t-tests, followed by the Fisher’s method and if necessary Stouffer’s z trend.

Figure 3 – figure supplement 17: Multiple species alignment and luciferase assay results for Tal1+19. (a) Multiple species alignment (MSA) with the following species: mouse (mm9), human (hg19), dog (canFam2) and opossum (monDom5). Nucleotides highlighted in black are conserved between all species analysed, nucleotides highlighted in grey are conserved between three of four species. Transcription factor binding sites (TFBS) are highlighted in: purple = Ets. The nucleotides that were changed to mutate the TFBSs are indicated below the MSA. All conserved binding sites of the Ets motif family were mutated simultaneously. (b) For the luciferase reporter assays in stably transfected 416b cells the averages of at least three independent experiments with three to four replicates within each experiment are shown. Error bars represent the standard error of the mean (SEM). Stars indicate significance: *** = p-value < 0.001. P-values were generated using t-tests, followed by the Fisher’s method and if necessary Stouffer’s z trend.

Figure 3 – figure supplement 18: Multiple species alignment and luciferase assay results for Tal1+40. (a) Multiple species alignment (MSA) with the following species: mouse (mm9), human (hg19) and dog (canFam2). Nucleotides highlighted in black are conserved between all species analysed, nucleotides highlighted in grey are conserved between two of three species.
Transcription factor binding sites (TFBS) are highlighted in: blue = Ebox, purple = Ets, green = Gata. The nucleotides that were changed to mutate the TFBSs are indicated below the MSA. All conserved binding sites of one motif family (e.g. all Ebox motifs) were mutated simultaneously. (b) For the luciferase reporter assays in stably transfected 416b cells the averages of at least three independent experiments with three to four replicates within each experiment are shown. Error bars represent the standard error of the mean (SEM). Stars indicate significance: * = p-value < 0.05. P-values were generated using t-tests, followed by the Fisher’s method and if necessary Stouffer’s z trend.

Figure 3 – figure supplement 19: List of TF binding sites and the TFs that bind to them.

Figure 3 – figure supplement 20: List of co-ordinates and primer sequences for the regulatory regions analysed in this study.

Figure 4: A three-tier DBN incorporating transcriptional regulatory information can recapitulate the HSPC expression state. (a) Representation of the complete network diagram generated using the Biotapestry software (56). (b) Schematic diagram describing the DBN which contains three tiers: I. TF binding motifs within regulatory regions, II. cis-regulatory regions influencing the expression levels of the various TFs, and III. genes encoding the TFs. The output of tier III, namely the expression levels of the TF, feed back into the TF binding at the various motifs of tier I. The model therefore is comprised of successive time slices (t). (c) Simulation of a single cell over time. The expression levels of all 9 TFs are the same at the beginning (0.5). The simulation rapidly stabilizes with characteristic TF expression levels. (d)
Simulation of a cell population by running the model 1000 times. The scale of the x-axis is linear. Each simulation was run as described in (c).

**Figure 4 - figure supplement 1:** Simulation of a single cell over time with different expression levels at the beginning. The simulation rapidly stabilizes with characteristic TF expression levels irrespective of the starting conditions. (a) The expression levels of all 9 TFs are 0.2 at the start of the simulation. (b) The expression levels of all 9 TFs are 0.8 at the start of the simulation. (c) The expression levels for FLI1, RUNX1 and TAL1 are set to be 0.5 at the beginning, with all other TFs not being expressed (value of 0).

**Figure 5:** The DBN recapitulates the consequences of TAL1 and LYL1 single and double perturbations as seen *in vivo* and *in vitro*. Computational prediction of gene expression patterns for the nine TFs of interest after perturbation of TAL1 (a), LYL1 (b) or both (c). Deletion of TAL1 or LYL1 on their own has no major consequences on the expression levels of the other eight TFs of the gene regulatory network, but simultaneous deletion of both TAL1 and LYL1 caused changes in expression of several genes, mainly a decrease in *Gata2* and *Runx1*. This major disruption of the core GRN for blood stem/progenitor cells is therefore consistent with TAL1/LYL1 double knockout HSCs showing a much more severe phenotype than the respective single knock-outs. One thousand simulations were run for each perturbation to determine the TFs expression levels in a “cell population” by selecting expression levels at random time points after reaching its initial steady state. Expression levels of 0 resemble no expression, whereas expression levels of 1 stand for highest expression level that is possible in this system. The scale of the x-axis is linear. (d) Gene expression levels measured in single 416b cells transfected with siRNA constructs against Tal1 or a control. The density plots of
gene expression levels after perturbation of TAL1 indicate the relative number of cells (y-axes) at each expression level (x-axes). The scale of the x-axis is linear. The values indicate the results of the Wilcoxon rank-sum test: alterations to the expression profiles are indicated by the p-value (statistical significance: p < 0.001 for computational data and p < 0.05 for experimental data); substantial shifts in median expression level are indicated by the shift of median (SOM) (SOM >0.1 for computational data and >1 for experimental data). For details, see Fig. 5 – figure supplement 1; for full expression data, see figure 5 – source data.

Fig. 5 - figure supplement 1: Significance tests for the computational and experimental data after TF perturbations. To determine statistical significance the Wilcoxon rank-sum test was used. Alterations to the expression profiles are indicated by the p-value; with statistically significance defined as follows: p < 0.001 for computational data and p < 0.05 for experimental data. Significance of a substantial shift in median expression levels are as follows: shift of median >0.1 for computational data and >1 for experimental data (because of different scales). If the number for the shift of median is negative, the median of the perturbation data is smaller than that of the wild-type control; if the number is positive, the median of the perturbation is larger than that of the control. For simplicity, all significant changes are highlighted in red (p-value) and blue (shift of median).

Figure 5 - figure supplement 2: Histogram plots showing the gene expression distributions of all nine genes of the network for the following perturbations: (a) LYL1 down-regulation; (b) TAL1/SCL down-regulation; (c) LYL1 and TAL1/SCL down-regulation; (d) PU.1 down-regulation; (e) GFI1B up-regulation; and (f) AML-ETO9a simulation.
Figure 5 - source data: Raw and normalised data for the single cell gene expression experiments presented in this study: 1) TAL1 down-regulation (related to Fig. 5 d), 2) PU.1 down-regulation (related to Fig. 6 a), 3) GFI1B up-regulation (related to Fig. 6b) and 4) AML-ETO9a perturbation (related to Fig. 6 c)

Figure 6: The DBN captures the transcriptional consequences of network perturbations. Left panel: Computational prediction of gene expression after perturbation of specific TFs. 1000 simulations were run for each perturbation to determine expression levels in a “cell population” (expression at 0 resembles no expression, whereas expression of 1 represents the highest possible expression level). The scale of the x-axis is linear. Right panel: Density plots of gene expression levels in single 416b cells after perturbation of specific TFs indicating the relative number of cells at each expression level. The scale of the x-axis is linear. The values indicate the results of the Wilcoxon rank-sum test: alterations to the expression profiles are indicated by the p-value (statistical significance: p < 0.001 for computational data and p < 0.05 for experimental data); substantial shifts in median expression level are indicated by the shift of median (SOM) (SOM >0.1 for computational data and >1 for experimental data). For details, see Fig. 5 – figure supplement 1. (a) PU.1 down-regulation: (Left) Computational prediction of gene expression after PU.1 knockdown (Spi1 was set to 0 after reaching its initial steady state). (Right) Gene expression levels measured in single 416b cells transduced with shRNA constructs against shluc (wild-type) or shPU.1 (PU.1 knockdown). (b) GFI1B over-expression: (Left) Computational prediction of gene expression after over-expression of GFI1B (Gfi1b was set to 1 after reaching its initial steady state). (Right) Gene expression levels in single 416b cells transduced with a Gfi1b-expressing vector compared to an empty vector control (wild-type). (c) Consequences of the AML-ETO9a oncogene: (Left) Computational prediction of gene expression patterns after introducing the dominant-negative effect of the AML-ETO9a
oncogene (Runx1 was fixed at the maximum value of 1 after reaching its initial steady state and in addition all Runt binding sites were set to have a repressive effect). (Right) Gene expression levels measured in single 416b cells transduced with an AML-ETO9a expressing vector fused to mCherry. mCherry positive cells were compared to mCherry negative cells (wild-type).

**Figure 6 – figure supplement 1: Summary of all computational simulations for perturbations of one or two TFs.** The results for a total of 162 simulations are shown. The data can be accessed using the embedded hyperlinks. The y-axes show the number of cells and the x-axes the relative expression level. Blue curves represent wild-type data and red curves represent perturbation data.
Figure 1
Figure 3
Figure 4
Figure 5

a) LYL1 down
   - wild-type
   - Ly1 knockdown

b) TAL1 down
   - wild-type
   - Scl knockdown

c) LYL1 and TAL1 down
   - wild-type
   - Ly1 and Scl double knockdown

d) TAL1 down (experimental)
   - wild-type
   - Scl knockdown

Expression levels and number of cells (x1000) are shown for various genes, including Erg, Fli1, Gata2, Gfi1b, Lyl1, Meis1, and Tal1, under different conditions.
**Computational**

**a** PU.1 down  
- - wild-type  
- - PU.1 knockdown

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**b** GFI1B up  
- - wild-type  
- - GFI1B over-expression

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<th>Over-expression</th>
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<tr>
<td>Tal1</td>
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**c** AML1-ETO9a  
- - wild-type  
- - AML-ETO9a simulation/expression

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**Figure 6**