Functional footprinting of regulatory DNA

Jeff Vierstra1,7, Andreas Reik2,7, Kai-Hsin Chang3, Sandra Stehling-Sun1, Yuanyue Zhou2, Sarah J Hinkley2, David E Paschon2, Lei Zhang2, Nikoletta Psatha3, Yuri R Bendana2, Colleen M O’Neil2, Alexander H Song2, Andrea K Mich2, Pei-Qi Liu2, Gary Lee2, Daniel E Bauer4, Michael C Holmes2, Stuart H Orkin4, Thalia Papayannopoulou3, George Stamatoyanopoulos3, Edward J Rebar2, Philip D Gregory2, Fyodor D Urnov2 & John A Stamatoyanopoulos1,6

Regulatory regions harbor multiple transcription factor (TF) recognition sites; however, the contribution of individual sites to regulatory function remains challenging to define. We describe an approach that exploits the error-prone nature of genome editing–induced double-strand break repair to map functional elements within regulatory DNA at nucleotide resolution. We demonstrate the approach on a human erythroid enhancer, revealing single TF recognition sites that gate the majority of downstream regulatory function.

Transcription regulatory regions harbor the majority of human disease-associated sequence variants1, rendering them attractive targets for elucidating disease mechanisms via targeted genome engineering2. However, our understanding of the impact of regulatory DNA variation is severely limited by the difficulty of precisely assigning the functional contribution of individual nucleotides to phenotypic outcomes. The error-prone nature of double-strand break repair triggered during targeted genome editing typically yields small deletions—or, more rarely, insertions—of variable size (1 to >10 nt, typically 2–6 nt). We reasoned that this byproduct of targeted genome editing could be systematically exploited to create a broad spectrum of variant regulatory alleles within a single experimental cycle and by coupling these alleles to a functional readout (such as protein expression), we could pinpoint the contribution of specific nucleotides to regulatory activity (Fig. 1a).

To test this paradigm (Fig. 1a), we studied the erythroid enhancer region of BCL11A (Fig. 1b), a transcriptional repressor of fetal hemoglobin production in adult erythroblasts3. Naturally occurring variants within this region are associated with reduced BCL11A expression, with consequent elevation of fetal globin (γ-globin) to levels that are clinically ameliorative for sickle-cell disease and beta thalassemia4.

To assess enhancer function in a physiologically relevant context, we first developed an optimized approach for high-efficiency genomic editing in primary human (adult) mobilized CD34+ hematopoietic stem cells, whereby a single exposure to an engineered zinc finger5 or transcription activator-like (TAL) effector6 nuclease could produce 60–80% per-allele editing rates (Supplementary Fig. 1a, Supplementary Tables 1 and 2 and Online Methods) within the population of edited cells. High-efficiency (70% of alleles) disruption of the BCL11A open reading frame in CD34+ cells followed by erythroid differentiation7 yielded dramatic and highly reproducible elevation of γ-globin mRNA, providing a physiologically relevant readout for loss of BCL11A function (Supplementary Fig. 1a,b). By contrast, control nucleases that drove high-efficiency (>60%) targeted disruption of a neutral locus (AAVS1)8 in the same cell type, coupled with the same differentiation protocol, had no impact on globin expression profiles (Supplementary Fig. 1b).

The BCL11A erythroid enhancer region encompasses three DNase I–hypersensitive sites (DHSs) located at +55 kb, +58 kb and +62 kb relative to the transcriptional start site (Fig. 1b). TAL effector nucleases (TALENs) targeting each of these DHSs were delivered to human CD34+ cells, and TALEN delivery was followed by erythroid differentiation. Notably, nuclease-treated cells developed morphologically and physiologically indistinguishably from untreated cells, as reviewed by an expert hematologist (T.F.; Supplementary Fig. 2). Complete deletion of the +55 or +58 DHS reproducibly elevated γ-globin mRNA levels in mature CD34+–derived erythroblasts, whereas deletion of the +62 DHS did not (Supplementary Fig. 1c,d). Of note, single-nucleotide polymorphisms in the latter DHS are associated with lower BCL11A expression and elevated γ-globin3; however, it is unclear whether these variants are in fact causal.

We next asked whether it would be possible to identify the specific sequence elements within a DHS that underlie the functional effects of the entire element. For this purpose, we focused on the +58 DHS that showed the most potency in elevating γ-globin mRNA levels when deleted (Supplementary Fig. 1d). To map in vivo TF occupancy sites, we performed genomic footprinting9 on human erythroblasts (Online Methods) and delineated eight TF footprints (FPs) covering 162 bp (53%) of the +58 DHS (Fig. 1c). To perform an initial functional scan, we used zinc-finger nucleases (ZFNs) to disrupt five of the FPs, after which we

1Department of Genome Sciences, University of Washington, Seattle, Washington, USA. 2Sangamo BioSciences, Pt. Richmond, California, USA. 3Department of Medicine, Division of Hematology, University of Washington, Seattle, Washington, USA. 4Boston Children’s Hospital, Division of Hematology/Oncology, Boston, Massachusetts, USA. 5Department of Medicine, Division of Medical Genetics, University of Washington, Seattle, Washington, USA. 6Department of Medicine, Division of Oncology, University of Washington, Seattle, Washington, USA. 7These authors contributed equally to this work. Correspondence should be addressed to F.D.U. (turnov@sangamo.com) or J.A.S. (jstam@uw.edu).

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performed erythroid differentiation and measured γ-globin mRNA levels. ZFN-induced lesions within footprint 5 (FP5) exhibited marked depletion of BCL11A mRNA and corresponding elevation of γ-globin levels irrespective of β-globin expression or normalization strategy (Fig. 1d and Supplementary Fig. 3), indicating that a focused lesion in this regulatory element was sufficient to disrupt BCL11A enhancer function. Of note, FP5 overlaps a recognition sequence for the master regulator of erythropoiesis GATA1 (ref. 10; Fig. 1d), and Encyclopedia of DNA Elements (ENCODE) data show GATA1 occupying the γ-globin gene and corresponding elevation of γ-globin mRNA and corresponding elevation of γ-globin mRNA levels (Supplementary Fig. 4a).
Figure 2 | Functional footprinting via genome editing and cell phenotyping reveals the precise boundaries of a GATA1 binding site in the BCL11A erythroid enhancer. (a) Genotypes derived from the alleles present in cells with low and high γ-globin protein expression edited with ZFN Z5. (b) Proportion of total alleles genotyped that retain a match to the GATA1 consensus sequence in cells with low and high γ-globin protein expression. (c) Meta-analysis of deep-sequencing data on cells containing high vs. low levels of γ-globin reveals the precise boundaries of a GATA1 binding site. The two bar charts plot the frequency for which a given base pair is deleted in erythroblasts with high and low levels of γ-globin (top and bottom, respectively). The WebLogos (bottom) show the normalized (high γ-globin – low γ-globin) nucleotide deletion frequency compared to the idealized GATA1 consensus sequence.

coding sequence (Supplementary Fig. 7 and Supplementary Table 5). These results indicate that the function of the erythroid-specific BCL11A enhancer is gated on the GATA1 recognition site in the +58 DHS.

The distinct spectra of allele frequencies observed in cells with high and low γ-globin expression indicated that γ-globin expression itself could be used as a quantitative molecular sensor for the function of the GATA1 binding site at single-base-pair resolution. To explore this further, we computed the per-nucleotide editing rate for each position surrounding the GATA1 consensus motif considering the frequencies of all genotypes observed. Edits associated with high γ-globin expression were markedly enriched for positions that covered the core GATA1 consensus motif, and the normalized editing rate observed in the cells with high γ-globin expression strikingly recapitulated an idealized GATA1 consensus sequence (Fig. 2c). We additionally found increased rates of editing in both the upstream and downstream flanking sequences, indicating the presence of additional binding sites (for example, the GATA1 partner TAL1) that may serve to modulate GATA1 occupancy (Fig. 2c); indeed, the presence of additional factors is supported in the DNase I footprint (FP5) itself in addition to the clear conservation of its underlying sequence elements (Fig. 1c). As noted above, the vast majority of genome-editing events found in cells with low γ-globin expression created alleles that did not ablate a functional GATA1 binding site.

Our results show that the functional impact of individual TF binding sites within cis-regulatory DNA can be efficiently interrogated by coupling the spectrum of alleles triggered by an engineered nuclease with expression of a downstream transcript or protein target. In the context of the distal BCL11A enhancer, our results pinpoint a single TF recognition site that can be targeted via genome editing to effect a potentially therapeutically meaningful outcome; editing the enhancer has functionally similar consequences on γ-globin mRNA expression to those of ablating the coding region of BCL11A itself (Supplementary Fig. 7c), both of which do not measurably affect erythroid differentiation (Supplementary Fig. 2).

Functional footprinting thus encompasses a simple and generalizable strategy to dissect the function of individual cis-regulatory elements such that any molecular sensor (i.e., protein, RNA, etc.) can be linked to the function of individual base pairs within non-coding DNA. Although methods exist that can assess the function of large libraries of synthetic alleles in both coding and feasibly noncoding DNA in vivo, the low efficiencies of allele integration achieved via homology-directed repair necessitate the use of large amounts of starting material and selectable markers to enrich for edited cells 14. As such, application of these methods is limited to conventional cell lines. In contrast, functional footprinting does not rely on homology-directed repair for the introduction of a library of synthetic alleles, thus enabling application in primary cells and developmental systems for which material and time are important considerations. Furthermore, this paradigm is agnostic to particular genome-editing and downstream product–detection approaches; although the ZFN and TALEN platforms were used herein because of efficiency and downstream considerations such as the potential for therapeutic translation, such efforts could, in
principle, rely on any genome-editing platform to attain a similar outcome\(^2\), subject to inherent limitations in design and efficiency. Similarly, functional footprinting is, in principle, compatible with any molecular readout (i.e., RNA, protein, post-translational modification levels, etc.) that can be efficiently sorted, and, critically, the detection of the sensor can occur on fixed (nonviable) cells, permitting use of a wide array of current and emerging technologies such as FACS on protein or RNA levels\(^{15}\).

METHODS

Methods and any associated references are available in the online version of the paper.

Note: Any Supplementary Information and Source Data files are available in the online version of the paper.

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AUTHOR CONTRIBUTIONS

K.-H.C., S.S.-S., Y.Z., S.J.H., D.E.P., L.Z., C.M.O., A.H.S., A.K.M. and N.P. performed the experiments and data collection. J.V., A.R., K.-H.C., Y.R.B., P.-Q.L., G.L., M.C.H., E.J.R., T.P., G.S. and F.D.U. analyzed the data. D.E.B. and S.H.O. provided critical insights. J.V., J.A.S. and F.D.U. wrote the manuscript with input from A.R., K.-H.C., E.J.R., F.D.U., T.P. and G.S.

COMPETING FINANCIAL INTERESTS

The authors declare competing financial interests: details are available in the online version of the paper.

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ONLINE METHODS

Nuclease design and validation. ZFNs were designed and assembled using an archive of prevalidated two-finger modules into expression vectors bearing obligate heterodimer forms of the FokI endonuclease. TALENs were designed as described and cloned into expression vectors bearing truncated forms of the TALE domain. In brief, the nuclease consists of a triple flag domain, a nuclear localization signal, the engineered DNA-binding domain and the FokI nuclease domain containing the obligate heterodimer mutations. The designed sequence recognition domains are provided in Supplementary Tables 1 and 2. Nuclease were first assessed for editing efficiency by transient transfection of expression constructs into K562 cells followed by genotyping of the target locus using the Surveyor/Cel1 endonuclease. ORFs for maximally active nucleases against each genomic position were recloned into an expression vector optimized for mRNA production bearing a 5′ and 3′ UTRs and a synthetic poly(A) signal. The mRNAs were generated using the mMessage mMachine T7 Ultra kit (Ambion) following the manufacturer’s instructions. In vitro synthesis of nuclease mRNAs used either (i) a pVAX-based vector containing a T7 promoter and the nuclease proper, which requires enzymatic addition of a poly(A) tail following the transcription reaction or (ii) a pGEM-based vector containing a T7 promoter, the nuclease proper, a 3′ UTR and a 64-bp poly(A) stretch.

Purification and genome editing of human CD34+ cells. Human mobilized CD34+ cells (adult) were purchased from AllCells. Small-scale mRNA transfections were performed with a BTX device (Harvard Apparatus), using the CD34+ cell program per manufacturer’s instructions. Either 2 µg of mRNA for each ZFN or 4 µg of mRNA for each TALEN were transfected into 200,000 cells in a 100-µl volume. Large-scale transfections were performed using the MaxCyte device according to manufacturer’s instructions, using 3 million cells in a total volume of 100 µl and 6 µg of each nuclease mRNA. After transfection, the cells were exposed to transient hyperthermia for 16 h and then cultured at 37 °C at 5% CO2. The cell lines were not tested for mycoplasma contamination or authenticated for these experiments.

Target loci genotyping following genome editing. Forty-eight hours following electroporation, genomic DNA was extracted from 50,000 CD34+ cells using a MasterPure kit (Epicentre). Deletions were genotyped by PCR and nondenaturing PAGE. Targeted locus disruption was measured by Surveyor/Cel1 (ref. 17) or deep amplicon sequencing on the Illumina platform. For the latter, the target locus was amplified in a two-step PCR from ~100 ng genomic DNA. The initial PCR used primers bearing (3′ to 5′) a locus-specific region, a randomized region and an adapter sequence compatible with the second PCR step; the second PCR used primers (3′ to 5′) bearing a stretch that anneals to the first-round amplicon, an amplicon-specific barcode and the Illumina flow cell–specific sequences. All generic primer sequences followed manufacturer’s instructions for the MiSeq sequencer. After sequencing, FASTQ sequence reads were filtered via fastq_quality_filter (http://hannonlab.cshl.edu/fastx_toolkit/) for sequences where all bases had Q ≥25 (Phred score). Sequences were further filtered for those matching 23 bp on the 5′ and 3′ end of the amplicon to exclude oligonucleotide synthesis–based deletions and primer dimers before alignment to the intended amplicon. High-quality sequences were then grouped, first according to their indel score (the deviation from wild-type length) and then according to the location of deletions or insertions that account for the difference. Single-base-pair substitutions are not analyzed as they represent an artifact of the sequencing platform (a bona fide signature of the targeted gene disruption process is a small insertion or deletion).

In vitro erythropoiesis. The protocol is based on work from the Donay laboratory. After electroporation the cells were treated as follows. Day 0 to day 7: 4 × 104 CD34+ cells/ml were cultured in EDM (IMDM, human holo-transferrin (330 µg/ml); insulin (10 µg/ml); heparin (2 IU/ml), 5% plasma) in the presence of 10−6 M hydrocortisone, 100 ng/ml SCF, 5 ng/ml IL-3 and 3 IU/ml Epo. On day 4, 1 volume of cell culture was diluted in 4 volumes of fresh medium containing SCF, IL-3, Epo and hydrocortisone. Day 7 to day 11: the cells were resuspended at 4 × 106 cells/ml in EDM supplemented with SCF and Epo. Day 11 to day 15 and out to day 20: the cells were cultured in EDM supplemented with Epo alone. Cell counts were adjusted to between 7.5 × 104 and 1 × 106 on day 11 and cells were harvested on day 13 or 14 for mRNA analysis and day 20 for immunofluorescence staining of γ-globin and FACS.

Immunofluorescence staining of γ-globin was modified from a previously described method. Briefly, cultured erythroid cells were fixed with 4% formaldehyde (Sigma) in PBS and then permeabilized with acetone (Sigma). Cells were washed once with PBS supplemented with 0.5% bovine serum albumin (Sigma) and stained with R-phycocerythrin–conjugated anti-γ-globin antibody (Santa Cruz Biotechnology; cat no. SC-21756PE). Cells were washed and resuspended in NucRed (Life Technologies)–containing PBS–0.5% BSA. Cell sorting was performed using BD Aria III with FACSDiva v6 (BD Biosciences). Erythroblasts were first gated on the basis of positive staining for NucRed. γ-globin–high (top ~10%) and γ-globin–low (bottom ~10%) erythroblasts were sorted and sequenced via amplicon sequencing as described above.

Determining functional effects of nuclease edits. To determine significant effects of nuclease cleavage on γ-globin mRNA expression, we modeled the relationship between editing efficiency determined by amplicon sequencing and relative γ-globin mRNA levels using a robust linear model using the function lnrmob from the “robustbase” package in R. We then computed the model residuals for each editing experiment and selected a threshold of 1σ for significance.

Fetal erythroblasts generation. Fetal livers (50- to 100-d gestation) were obtained from the fetal tissue repository (University of Washington Birth Defects Research Laboratory) with permission of the University of Washington Institutional Review Board. The erythroid culture of dissociated fetal livers and characterization of fetal liver–derived erythroblasts have been described previously. These cells were subjected to DNase I treatment for profiling for DHSs and transcription factor binding footprints as described.

Real-time RT-qPCR measurement of globin mRNA levels. Whole-cell RNA was isolated from in vitro–generated erythrocytes using a High Pure RNA Isolation Kit (Roche). The levels

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of mRNA for the individual globin genes (α, β and γ) were then measured by real-time RT-qPCR on an ABI 7300 RT-PCR machine mode using the following manufacturer-provided probe sets: α-globin (HBA), Hs00361191_g1; β-globin (HBB), Hs00758889_s1; γ-globin (HBG), Hs00361131_g1; BCL11A, (Hs01093197_m1); 18S rRNA (18S) Hs99999901_s1.

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