Dynamics of Transcription Regulation in Human Bone Marrow Myeloid Differentiation to Mature Blood Neutrophils

Graphical Abstract

Highlights

- Dynamic acetylation enforces human neutrophil progenitor differentiation
- Neutrophils’ cytotoxic capability is established early at the (pro)myelocyte stage
- Coordinated signaling component expression prevents toxic effects to the bone marrow

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In Brief
Grassi et al. report that the establishment of transcriptional enhancers drives neutrophil differentiation. Coordinated waves of gene expression establish the cytotoxic capability of these cells at early stages of maturation. A set of super-enhancers is specifically opened at the end of the differentiation process to control neutrophil activation.

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Dynamics of Transcription Regulation in Human Bone Marrow Myeloid Differentiation to Mature Blood Neutrophils

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SUMMARY

Neutrophils are short-lived blood cells that play a critical role in host defense against infections. To better comprehend neutrophil functions and their regulation, we provide a complete epigenetic overview, assessing important functional features of their differentiation stages from bone marrow-residing progenitors to mature circulating cells. Integration of chromatin modifications, methylation, and transcriptome dynamics reveals an enforced regulation of differentiation, for cellular functions such as release of proteases, respiratory burst, cell cycle regulation, and apoptosis. We observe an early establishment of the cytotoxic capability, while the signaling components that activate these antimicrobial mechanisms are transcribed at later stages, outside the bone marrow, thus preventing toxic effects in the bone marrow niche. Altogether, these data reveal how the developmental dynamics of the chromatin landscape orchestrate the daily production of a large number of neutrophils required for innate host defense and provide a comprehensive overview of differentiating human neutrophils.

INTRODUCTION

Human neutrophils, with a daily production rate of \( \sim 10^{11} \) cells (Strydom and Rankin, 2013), are the most abundant type of leukocytes and an essential element of the innate immune
system. Patients with either acquired or inherited defects in neutrophil development, migration, or function show enhanced susceptibility to infection by opportunistic pathogens (Erlacher and Strahm, 2015; Fodil et al., 2016). Neutrophils have a variety of effector functions, such as phagocytosis of pathogens (Nauseef and Borregaard, 2014), intracellular killing in phagosomes with reactive oxygen species (ROS), and antimicrobial granule components (Darrah and Andrade, 2013; Garcia-Romo et al., 2011; Hakkim et al., 2010). They are produced in the bone marrow and reach the bloodstream across five main stages of differentiation. Myeloblasts (MBs) undergo 1 week of proliferation, followed by maturation (Bainton et al., 1971), then progressively move through promyelocytes and myelocytes (P/Ms), metamyelocytes (MMs), band neutrophils (BNs), and finally segmented neutrophils (SNs). During this time they produce different granules filled with antimicrobial compounds and develop the abilities of respiratory burst and chemotaxis. SNs represent a reserve pool from which cells are constantly released into the peripheral bloodstream, as non-dividing polymorphonuclear neutrophils (PMNs). The differentiation process exhibits a precise and tightly defined transcriptional program (Theilgaard-Mönch et al., 2005). In this study, we used total RNA sequencing (RNA-seq) to characterize and compare gene expression profiles of the various differentiation stages. Histone modifications, chromatin immunoprecipitation sequencing (ChIP-seq), and whole-genome bisulfite sequencing (WGBS) have been used to describe the epigenetic landscapes underlying these coordinated changes. For each stage we defined DNA methylation patterns and characterized four active histone marks—H3 lysine-4 trimethylation (H3K4me3), H3 lysine-7 acetylation (H3K27ac), H3 lysine-4 monomethylation (H3K4me1), H3 lysine-36 trimethylation (H3K36me3)—and two repressive marks—histone H3 lysine-9 trimethylation (H3K9me3) and histone H3 lysine-27 trimethylation (H3K27me3). The integration of these analyses reveals a high degree of coordination between the dynamics of histone modifications and transcriptional changes. The activation of PMNs triggers the host defense. This process is accomplished through some functional key responses as respiratory burst, cell migration, and degranulation. Integrating our functional assays with the epigenomic analysis gave us the opportunity to further characterize the timing of these important processes.

**RESULTS**

**Transcriptional Changes in Neutrophil Development**

To better understand neutrophil functions and their regulation, we analyzed the transcriptomes of circulating neutrophils and their progenitor populations (Figure S1A). Our analysis revealed quantitative and qualitative differences in expression among PMNs, SNs, and the other differentiation stages. We used cumulative distribution of expression to explore the complexity of the transcriptome (Melé et al., 2015) and found that in PMNs and SNs, the genes with higher expression give a smaller contribution to the global transcriptional output than in other stages (Figure 1A). To deepen the transcriptome comparisons, we performed differential gene expression analysis between subsequent stages and between the initial and the final stage (Figure 1B; Table S1). Downregulated genes in the P/M-PMNs comparison enriched Gene Ontology (GO) categories related to cell functions, such as metabolism, translation, cell cycle, DNA replication, mitochondria, and cell divisions (Figures S1B and S1C). On the other hand, the upregulated genes enriched fewer categories, related mainly to immune system functions and vesicle transport processes (Figures S1B and S1C).

The analysis of differentially expressed transcripts across all stages found 108 genes with significant changes in transcription starting site (TSS) use (Table S1). Only for 12 genes did this lead to changes in the open reading frame (Table S1). Cluster analysis and gap statistic (Tibshirani et al., 2001) summarized the differentially expressed genes in at least one of the five comparisons (P/M-MM, MM-BN, BN-SN, SN-PMN, or P/M-PMN) in seven main clusters (r1–r7; Figure 1C; Table S1). Genes in clusters r1 and r2 showed an increase in expression toward the latter...
stages of differentiation and enriched GO terms related to immune system and signal transduction (Table S1). Genes in clusters r5, r6, and r7 displayed a decreasing expression trend with differentiation progression and enriched GO terms related to translation, mitochondria, and cell cycle (Table S1). The remaining two clusters (r3 and r4) were made by genes whose expression increased during the intermediate stages of differentiation and then decreased at the PMN stage. These enriched GO terms related to vesicle transport and signal transduction (Table S1).

Epigenetic Dynamics in Neutrophil Development
In order to define the chromatin dynamics underlying the changes in gene expression during neutrophil differentiation, we determined the epigenetic landscape in the five differentiation stages and integrated them with the results of the transcriptome characterization. Principal-component analysis (PCA) of the four active histone modifications (H3K4me3, H3K27ac, H3K4me1, and H3K36me3) revealed a good separation among the five neutrophil development stages (Figure S2A). In contrast, PCA for the two repressive marks (H3K9me3 and H3K27me3) revealed donor-specific, rather than stage-specific, clusters (Figure S2A). Moreover, PCA analysis of DNA methylation profiles did not reveal differences among the five differentiation stages (Figure S2B), and only a few differentially methylated regions (DMRs) were identified between subsequent stages of differentiation (Figure S2C; Table S2). Altogether, histone mark and DNA methylation results indicate a modest contribution of repressive epigenetic mechanisms in the described phases of neutrophil differentiation.

The six different histone modifications were also used to perform a genome segmentation analysis (Ernst and Kellis, 2012), which indicated 12 representative states (Figure S3A). These were collapsed into a smaller set for ease of interpretation (Carrillo-de-Santa-Pau et al., 2017) to obtain four main categories: active, enhancer, low signal, and repressed (Figure S3A). We then focused on the changes of chromatin states, identifying between 15,685 and 27,604 at each consecutive differentiation transition (Figure S2A; Table S3). In agreement with the differential gene expression analysis results, the consecutive transitions with most changes were P/M to MM and SN to PMN. Uregulated genes were enriched in changes toward active states (i.e., from low signal/repressed to active/enhancer), while downregulated genes were enriched in transitions toward inactive states (i.e., from active/enhancer to low signal/repressed) in the majority of differentiation stages (Figure S3B). H3K27ac, strongly associated with active and enhancer states, is among the most highly informative marks about gene regulation (Karlić et al., 2010; Rada-Iglesias et al., 2011). For this reason, we decided to focus our attention on the 9,100 genomic regions characterized by changes in H3K27ac levels during differentiation (see Experimental Procedures). Using co-occupancy with H3K4me3 to identify promoter regions and co-occupancy with H3K4me1 to identify enhancers, in addition to a supervised cluster of the H3K27ac log ratio RPKM (reads per kilobase of transcript per million mapped reads), we divided these regions in two categories each made of eight subgroups (Figure 2B; Table S4). The intersection of these regions with the chromatin states revealed an overlap with active (promoter) regions in the H3K4me3-enriched clusters and an overlap with enhancers in the H3K4me1-enriched clusters (Figure S3C), corroborating our partitioning. We refer to the clusters as promoter (p1–p8) and enhancer (e1–e8). In both promoter and enhancer clusters about two-thirds of dynamic regions displayed an increase in H3K27ac (i.e., increase in activity), while about one-third displayed a reduction in H3K27ac (i.e., reduction in activity). The contingency tables made by the intersection of the genes in the RNA-seq clusters gave significant Pearson’s chi-square statistics for both acetylation clusters (enhancer chi-square = 513, p = 0.0005; promoter chi-square = 1,364, p = 0.0005). We tested the association of each gene expression cluster with each dynamic acetylation cluster (Figures 2C and 2D). Interestingly we found that expression clusters r1 and r2, containing genes with increased expression, were significantly associated with promoter clusters p1 and p2 and enhancer clusters e1 and e2. Expression clusters r6 and r7, formed by genes with decreased expression, were significantly associated with promoter clusters p5 and p6 and enhancer clusters e5 and e6, all characterized by reduced histone acetylation levels. The expression cluster r4 included genes with an increased expression in the PM-to-MM transition, and the same trend was observed in the associated acetylation clusters e4 and p4.

Dynamically acetylated enhancers of the same cluster had a genomic distance closer than expected by chance (permutation test Z = 21). This observation prompted us to investigate the presence of super-enhancers (SEs), regulatory elements made by clusters of enhancer that specify cell identity (Whyte et al., 2013; Witte et al., 2019). We made use of the established ROSE algorithm to identify 629 SEs, which could be clustered in three main groups (Figure 2A; Table S4). SE cluster 1 (SE1) was made by elements more acetylated in the P/M state, showing a decreasing trend of acetylation in the subsequent stages of differentiation; SE cluster 2 (SE2) had an opposite trend, with acetylation increasing in the first stages and then decreasing at the SN-PMN transition. Finally, SE cluster 3 (SE3) contained SEs activated at the end of differentiation process. Genes controlled by SEs belonging to cluster 3 were the only ones showing an enrichment for functional categories, all of them related to PMN’s activation (Figures 2B and 2C).

Transcription Factors and Release from Bone Marrow
Transcription factors (TFs) bind regulatory elements (promoters and enhancers) and enforce transcriptional programs (Lambert et al., 2018; White et al., 2013). To identify the TFs that have an important role in granulopoiesis, we searched for TF family consensus binding sites, enriched in regions with increased acetylation (clusters p1–p4 and e1–e4) compared with regions with decreased acetylation (clusters p5–p8 and e5–e8) and vice versa. We found enrichments for basic leucine zippers (bZIP), SOX, zinc fingers (ZF), nuclear receptors (NR), basic helix-loop-helices (bHLH), NF-κB family members (REL), homeodomains, and ETS families, with stronger signal in regions with increased acetylation (Figure 4A). Regions with decreased H3K27ac signal showed enrichment for MYB- and GATA-binding motifs. We integrated these results with gene expression analysis results and identified for each enriched motif TF genes

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differentially expressed in at least one of the comparisons between differentiation stages (Figure 4B). The majority of these TFs (81 of 95) were differentially expressed in only one of the four comparisons. Twelve of the remaining 14 genes, differentially expressed at multiple transitions, were always either up- or downregulated. Only 2 genes, differentially expressed at multiple transitions, had a trend inversion: TGIF2 was downregulated in the P/M-MM transition and upregulated in the BN-SN transition, and ETS1 was upregulated in the P/M-MM transition and downregulated in the SN-PMN transition.

The P/M-to-MM transition included the majority (65 of 95) of differentially expressed TFs, followed by the bone marrow-to-blood transition (SN to PMN) with 20 differentially expressed TFs (Figure 4B). Interestingly 19 of them were downregulated, and NKX3-1, the only upregulated one, had a reported transcriptional repressor activity (Simmons and Horowitz, 2006). The TF families MYB, GATA, and E2F were exclusively overrepresented in enhancers and promoters with decreased acetylation, reflecting their role in other lineages and cell cycle progression (Bartunek et al., 2003; Ghazaryan et al., 2014). Accordingly, differentially expressed genes from these families were all downregulated (Figure 4B).

**Acquisition and Control of Cytotoxic Capabilities**

Cell proliferation is halted during neutrophil differentiation; our transcriptome and epigenome data indicated that this occurs in the bone marrow, as previously reported (Klausen et al., 2004; Theilgaard-Mønch et al., 2005). Flow cytometry quantification of Ki67, a marker for proliferation (Gerdes, 1990), showed that the majority of the signal drops in the early stages of myeloid
Figure 3. Identification of Super-enhancers Associated with Neutrophil Differentiation

(A) Heatmap of H3K27ac density at super-enhancers with dynamic acetylation during different stages of neutrophil differentiation. For each state tag density for the three replicates is included.

(B and C) GO Biological Processes (B) and GO Slim (C) categories enriched by genes associated with super-enhancers activated at the final stages of neutrophil differentiation (cluster SE3).

Figure 2. The cell-permeable PAF/fMLP (both determined by flow cytometry) increased during differentiation (Figures S4A and S4B). Additionally, we also noted that CXCR1, CXCR2, CXCR4, FPR1, FPR2, PTAFR, and C5AR1, activating G protein-coupled receptors for the major chemotactic factors in neutrophils, follow the same expression trend (Figure S4C), and are all in the expression cluster r1.

Transcriptional and Epigenetic Control of Granule Proteins

Another main function of the neutrophils is the degranulation consisting in the release of an assortment of proteins, stored in different granules. These granules are distinguishable by their contents: azurophilic granules (AG), specific granules (SG), gelatinase-containing granules (GG), ficolin-containing granules (FG), and secretory vesicles (SV) (Rørvig et al., 2013). To gain insight in the control of the biogenesis of these granules during differentiation, we examined available proteomic data (Rørvig et al., 2013) and selected the proteins whose granule assignments were supported by other existing literature (Table S6) and the granule proteins assigned to the cell membrane (CM) fraction. We found that genes encoding proteins belonging to various granules display different expression trends (Figure 6A), and in agreement with this, they showed a tendency to belong to different RNA-seq clusters (chi-square = 91.8062, p < 0.0005; Figure S5A). AG proteins were associated with clusters r6 and r7 (Fisher’s exact test corrected p < 0.05), while CM proteins were associated with cluster r1 (Fisher’s exact test corrected p < 0.005) and SG proteins with cluster r4 (Fisher’s exact test corrected p = 3.1 × 10⁻⁵).

To determine the time of appearance of AG’s proteolytic capability, we measured their combined serine protease activity, in sorted neutrophil progenitors, upon CytoB/MLP stimulation or Triton X-100 lysis. The results showed that although AG proteins have already been produced and stored at P/M stage, the protease release machinery is not in place before the BN stage. P/M cells were found to harbor the highest concentration of proteolytic activity, which was then diluted over to the MM daughter cells and remained stable in activity thereafter (Figure 6B). Genes of different granule proteins, similarly to what observed for gene expression, displayed different H3K27ac trends (Figure S5B).

In agreement with this, genes encoding soluble AG proteins, such as AZU1, PRTN3, and ELANE, had a high level of the H3K27ac in...
the P/M stage and reduced in the MM stage (Figures 6C and S5B) and were enriched in clusters p5, p6, and e5 (Fisher’s test corrected p values: for p5, \( p = 6.64 \times 10^{-3} \); for p6, \( p = 6.64 \times 10^{-3} \); and for e5, \( p = 3.65 \times 10^{-2} \)). In contrast, SG-associated genes, such as LTF, with increased acetylation in the P/M-to-MM transition, were enriched mainly in clusters p4 and e4 (Fisher’s test corrected p values: for p4, \( p = 1.7 \times 10^{-3} \); and for e4, \( p = 2.88 \times 10^{-2} \)) (Figures 6C and S5B). The SV-associated genes as MMP25 displayed increased acetylation at the final stages of differentiation (Figures 6C and S5B).

**DISCUSSION**

Our study dissects human neutrophil differentiation through the analysis and integration of transcriptomes, epigenomes, and functional assays of five consecutive stages of maturation.

In the transcriptome of a cell, a relative small number of genes contribute to a large fraction of the expression (Melé et al., 2015). Interestingly, we found this trend to be less pronounced in the two latest stages of neutrophil maturation, SN and PMN, indicating a change in transcriptome features at the final stages of the differentiation.

The P/M-to-MN and SN-to-PMN were the consecutive transitions with the largest numbers of differentially expressed genes and changes in chromatin states. These transcriptomic and epigenomic differences reflect the important differences between these stages. Indeed, the P/M-to-MN transition denotes a passage from cells capable of dividing to cells with indented nuclei and unable to divide (Wahed and Dasgupta, 2015). Similarly, the SN-to-PMN transition denotes the passage from bone marrow to the bloodstream.

Different stages of maturation do not display relevant differences in DNA methylation. This suggests that the unique patterns of DNA methylation found in neutrophils (Schuyler et al., 2016) are established at earlier stages of differentiation, in agreement with a previous study reporting that the majority of changes in DNA methylation occur between the oligopotent common myeloid progenitor and the bipotent granulocyte monocytic progenitor, with only minor changes occurring in the P/M-to-PMN transition (Rönnblad et al., 2014). Our study extends this result, with more differentiation steps during maturation of the granulocyte progenitors, thus providing a more detailed description of the dynamics of transcriptional enhancers’ commitment. Moreover, the use of WGBS allowed a 60-fold increase of the number of CpG analyzed residues.
Figure 5. Cell Cycle and NADPH Oxidase Regulation in Neutrophil Differentiation

(A) Flow cytometry analysis of Ki67 expression in bone marrow neutrophil progenitor fractions normalized to isotope control in gray for the three biological replicates.

(B) Schematic of the assembly of the nicotinamide adenine dinucleotide phosphate (NADPH) oxidase enzyme complex on the phagosomal or plasma membrane upon neutrophil activation.

(C) Heatmap representing the expression in log2(FPKM+1) of NADPH complex subunits during neutrophil differentiation.

(D) Respiratory burst in sorted neutrophil progenitors from bone marrow and mature PMN upon stimulation with PMA and PAF/fMLP. Results are expressed as maximal rates in nmol H2O2/min $\pm 10^6$ PMNs and as mean ± SEM from the three biological replicates. t test corrected p values of the consecutive stages are indicated in the figure: *p < 0.1 and **p < 0.01. ns, nonsignificant. The t test corrected p values of all comparisons are reported in Table S5.

In line with the DNA methylation results the two repressive histone marks analyzed cannot be used to discriminate among neutrophil differentiation stages. Previous studies demonstrated that repressive H3K9me3 and H3K27me3 marks and DNA methylation have a crucial role in cell commitment (Bock et al., 2012; Hawkins et al., 2010), and our results suggest that in neutrophils, they do not play any further roles once lineage commitment is achieved.

We focused our attention on promoter and enhancer regions subject to dynamic acetylation across differentiation. These regions, subdivided in clusters, showed significant overlaps of target genes with the clusters of differentially expressed genes, highlighting a strong correlation between epigenomic and transcriptomic profiles at the various differentiation stages. The study of dynamically acetylated regions led us to identify SEs differently modulated across differentiation; a consistent group of these is activated at the end of differentiation and have as potential targets the genes involved in immunological response and neutrophil activation.

To further characterize the dynamically acetylated regions, we looked for enriched TF binding sites and integrated them with the differential expression results. This approach aimed to highlight TFs playing a potential role in the differentiation process. CEBPA is a TF of the bZIP family with a major role in the neutrophil differentiation program, and it is known to be crucial for the development of acute myeloid leukemia (Avellino and Delwel, 2017). Our results show that CEBPA is highly expressed in the bone marrow and then significantly downregulated in the SN-to-PMN transition. CEBPE, another member of the bZIP family, is also highly expressed in the bone marrow and is downregulated in the SN-to-PMN transition but also in the preceding transition, BN to SN. Interestingly CEBPE modulates some of the lactoferrin-containing SG genes (Khanna-Gupta et al., 2007) and regulates the expression of GG and FG granule genes (Gombart et al., 2001), both characterized by a decreasing trend of expression at the PMN stage.

The transcriptome and epigenome characterizations depicted a precise differentiation program that, in agreement with previous findings (Theilgaard-Mønch et al., 2005), progressively activates immune functionalities, turning off essential cell processes. The study of five differentiation stages gave us the opportunity to detect with better resolution where the cell cycle inactivation takes place. We have shown that the P/M stage is the most proliferative, with Ki67 signal dimming in the following MM and BN stages and completely disappearing in the SN stage. These data, alongside with the expression of key cell cycle regulators, indicate that the cell cycle is arrested after the P/M stage.

The antimicrobial activity of neutrophils is achieved through ROS production by the NADPH oxidase enzyme complex. We showed that this enzyme complex activity reaches its peak only at the end of differentiation, although its expression is already detectable at the MM stage. The observed difference in ROS production during differentiation between PMA and PAF/fMLP kinetics highlights that G protein-coupled receptor and its signaling machinery are required to trigger the process. Indeed, we showed that when the surface expression of fMLP receptor FPR1 is low, at early stages of development, the response to PAF/fMLP is impaired.

Along the same line, we noticed that genes encoding for the proteins contained in the various granules displayed different
transcriptional and epigenetic dynamics. Indeed, the proteases were released by cellular activation only at the mature stage, because the signaling cascade and surface expression of activating GPCR receptors were not present at earlier stages.

The results presented in this study shed light on the neutrophil differentiation processes and represent a useful resource for future studies on as yet uncharacterized primary immune disorders or neutrophil-related disorders such as congenital neutropenia syndromes, Chédiak-Higashi syndrome, and lazy leukocyte syndrome (Dinauer, 2014; Nunoi et al., 1999).

**EXPERIMENTAL PROCEDURES**

**Study Design**

Bone marrow neutrophils were collected from three different healthy donors, and mature neutrophils were collected from other three other different healthy donors. ChIP-seq, RNA-seq, and WGBS experiments were done on samples from the same donors.

**Cell Isolation and Library Preparation**

The study was approved by the local ethics committee of Sanquin Blood Supply Organization and the Academic Medical Center (Amsterdam, the Netherlands).

PMNs were enriched by gradient centrifugation and purified by negative selection with EasySep Human Neutrophil Enrichment Kit. Bone marrow neutrophil progenitors were purified by fluorescence-activated cell sorting (FACS) with CD11b and CD16 (see Supplemental Experimental Procedures for further details). ChIP-seq and total RNA-seq libraries were prepared according to the BLUEPRINT protocols (http://www.blueprint-epigenome.eu).

**ChIP-Seq Analysis and Genome Segmentation**

Sequenced reads were mapped against the GRCh38 genome assembly with BWA (Li and Durbin, 2009). ChromHMM software (version 1.10) (Ernst and Kellis, 2012) was then used to segment the genome in 200 bp intervals and assign epigenetic states. Chromatin states consistently identified in all three replicates were represented with a Sankey diagram at each stage of development using the “makeRiver” and “riverplot” functions included in the riverplot R package version 0.5.

**RNA-Seq Analysis**

Trim Galore (version 0.3.7) (http://www.bioinformatics.babraham.ac.uk/projects/trim_galore/) with parameters “--q 15 -s 3 --length 30 -e 0.05” was used to trim PCR and sequencing adapters. Trimmed reads were aligned to the Ensembl version 80 (Cunningham et al., 2015) human transcriptome with Bowtie 1.0.1 (Langmead et al., 2009) using the parameters “-a –best –strata -S -m 100 -X 500 –chunkmbs 256 –novf –fr.” MMSEQ (version 1.0.8a) (Turro et al., 2014; Turro et al., 2011) was used with default parameters to quantify gene expression. Genes and transcripts with posterior probability > 0.5 (calculated by MMDiff), absolute fold change > 2, and FPKM (fragments per kilobase of transcript per million mapped reads) > 1 in at least one of the two compared cell types were considered differentially expressed. Differentially expressed genes in at least one of the comparisons P/M-MM, MM-BN, BN-SN, SN-PMN, and P/M-PMN (Figure S1) were clustered using the “kmeans” (Hartigan and Wong, 1979) clustering R function. Gene expression FPKMs were log 2-transformed, and for each gene the Z score was calculated. We set the number of centers to seven according to the results of the gap statistic (Tibshirani et al., 2001) by using the “clusGap” function of the cluster R package.
SE Identification
SEs in each sample were predicted by the ROSE algorithm (Whyte et al., 2013) using H3K27ac as the surrogate mark. Briefly, all H3K27ac peaks within ±2.0 kb around TSSs were first excluded. The remaining peaks closer than default distance of 12.5 kb were stitched together and subsequently ranked by normalized H3K27ac level corrected by input background. Finally, SEs were separated from typical enhancers on the basis of the inflection point of H3K27ac signal curve.

Functional Enrichment Analysis
Functional enrichment analysis was done by using FIDEA (D’Andrea et al., 2013) with default settings.

Promoter/Enhancer/SE Gene Assignment
For each transcript annotated in Ensembl version 80 (Cunningham et al., 2016), we defined a promoter as the 2,000 bp (±1,000 bp) around its TSS. All the regions defined as promoter by the genome segmentation analysis and the H3K27ac dynamic analysis were assigned to the genes of the overlapping promoters. All the regions defined as enhancer by segmentation analysis and the H3K27ac analysis were assigned to the gene with the closest promoter (distant less than 10,000 bp) and to genes with promoters overlapping their interacting regions (CHiCAGO score > 5 in neutrophils) derived from the study of Javierre et al. (2016). The “findOverlaps” and “distanceToNearest” functions of the GenomicRanges (Lawrence et al., 2013) R package were used to make region comparisons.

Motif analysis was performed with the HOMER (Heinz et al., 2010) program. WGBS was carried out to minimum genome coverage of 30x with established protocols (Kulis et al., 2012).

Differential methylation was estimated for (1) individual CpG sites in pairwise comparisons by using methyl_dif (Flainer et al., 2014) and across regions in group-wise comparison by using replicates with metilene (Juhling et al., 2016).

Association between Acetylation and Expression Clusters
We built a contingency table with the number of genes for each acetylation cluster (rows) and gene expression cluster (columns). The Pearson’s chi-square test was used to verify the non-independence of the two categorical variables. The significance of the association of each gene expression cluster with each dynamic acetylation cluster was tested by a hyper-geometric test, and the derived p values were corrected using the Benjamini-Hochberg method (Hochberg and Benjamini, 1990).

NADPH-oxidase activity was assessed with an Amplex Red kit (Drewniak et al., 2013; Kuipers et al., 2005). Protease release after degranulation was measured with DQ-green BSA (see Supplemental Experimental Procedures for further details).

DATA AND SOFTWARE AVAILABILITY
The web site https://blueprint.haem.cam.ac.uk/neutrophil collects the analyzed data presented in this article. All the genome-wide data generated in this study, as well as the sequencing details, can be accessed via http://dcc.blueprint-epigenome.eu/.

SUPPLEMENTAL INFORMATION
Supplemental Information includes Supplemental Experimental Procedures, five figures, and six tables and can be found with this article online at https://doi.org/10.1016/j.celrep.2018.08.016.

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AUTHOR CONTRIBUTIONS
F.P. and T.W.K. designed the study, performed and supervised research, and wrote the manuscript. L.G. analyzed the data and wrote the manuscript. S.U., A.M., F.W., E.C.d.S.P., G.Y., and J.P. analyzed the data. I.H.H., A.T.J.T., E.M., E.J.-M., K.B., H.K., E.H., M.G., M.L.Y., and M.L. performed experiments. E.L., A.D., L.C., and P.F. performed and supervised data management. M.V., D. Roos, T.K.v.d.B., S.H., D. Rico, M.K., I.G., A.V., W.H.O., and H.G.S. provided expert supervision. M.F. provided expert supervision and wrote the manuscript. J.H.A.M. designed the study, analyzed the data, supervised research, and wrote the manuscript. All authors read and approved the final version of the manuscript.

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