CRISPR-based reagents to study the influence of the epigenome on gene expression

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Summary

The use of epigenome editing is set to expand our knowledge of how epigenetic landscapes facilitate gene expression capacity within a given cell. As epigenetic landscape profiling in health and disease becomes more commonplace, so does the requirement to assess the functional impact that particular regulatory domains and DNA methylation profiles have upon gene expression capacity. That functional assessment is particularly pertinent when analysing epigenomes in disease states where the reversible nature of histone and DNA modification might yield plausible therapeutic targets. In this review we discuss first the nature of the epigenetic landscape, secondly the types of factors that deposit and erase the various modifications, consider how modifications transduce their signals, and lastly address current tools for experimental epigenome editing with particular emphasis on the immune system.

Keywords: gene regulation, molecular biology, transcription factors

Epigenetics

The completion of the draft sequence of the human genome in 2001 [1] heralded intense interest in the management and implementation of gene expression programmes, understanding the basis of cell differentiation and cell type specification. These studies have begun to reveal how an individual cell type establishes and maintains its ability to express a distinct repertoire of genes and how this output capacity can be modulated under the influence of intrinsic and extrinsic stimuli. Consortium projects such as Roadmap [2] and Encyclopedia of DNA Elements (ENCODE) [3] have begun the job of identifying all functional elements within the human genome. This task encompasses mapping RNA transcripts in all cell types, mapping the location of DNA associating with modified histones and transcription factors and mapping modification of DNA itself by methylation. In an ENCODE update in 2012 [3], with comprehensive analysis of fewer than 20% of the known transcription factors encoded by the human genome, of half the DNA and histone modifications and potentially a similar percentage of known cell types, the data were startling. The studies have led to fundamental insights into genome organization and utility, as exemplified by findings that greater than 60% of bases in the genome are represented in long RNA molecules yet only one-tenth of these are constituents of the roughly 20,000 known genes in the genome, that there are greater than 400,000 regions of the genome that have ‘enhancer-like’ chromatin features and that there are more than 70,000 regions with ‘promoter-like’ features. Even with these incomplete projects, it is clear that there are significant gaps in our understanding of the labyrinthine influence of regulatory RNAs, transcription factors and chromatin states that contribute to gene expression pathways.

Epigenetics research is complicated by a number of factors: first, because epigenomes are cell type-specific, a challenge has been to generate purified cell types in sufficient numbers to exploit technologies that can report on the landscape in an unbiased manner. Cell types that arise during early differentiation represent a particular challenge for isolation for both numerical and ethical reasons. In adults, while cell purification is relatively straightforward in mature immune systems, emerging information from single-cell sequencing has begun to reveal an expanded repertoire of distinct cell types, as exemplified with subtypes of dendritic cells [4]. The consequence of this information is that the spectrum of cell types on which epigenetic landscapes and their associated transcriptomes might be analysed is greater than thought previously. Clear inroads are being made in this area...
through the Human Cell Atlas project [5], but even in these situations different transcriptomes may be influenced by different activation status of cells within a population. Tissue resident immune cells present particular problems; these cells, such as T resident memory cells, may lack substantial presence in the bloodstream, as demonstrated conclusively by parabiosis experiments [6]. The challenge is therefore how to collect sufficient of these cells in order to establish their epigenetic footprint, particularly in human disease scenarios.

Certain immune cells are often considered to be phenotypically plastic [4], and this property is important in the rapid adaptation to external stimuli. Functional plasticity, however, brings into question whether the gene expression patterns used to define individual cell types are sufficiently invariant to discriminate stable populations. The use of t-Distributed Stochastic Neighbour Embedding (tSNE) [7] analysis is making identification of cells having similar phenotypical characteristics more amenable and permits identification of expression signatures that may allow subsequent purification by flow cytometry.

A further challenge is technical, and derives from the fact that sequencing-driven techniques such as ChiPseq and ATACseq enable population-level analysis which need careful interpretation in order to determine for how many cells in that population the landscapes derived are accurately representative. Epigenome-wide analysis from single cells and small numbers of cells is now the subject of numerous applications [8–10], and data derived from single-cell epigenome analysis will no doubt be of great importance in understanding the mechanism of cell differentiation.

Chromatin organization

The ability to control transcriptional output of any cell is impacted at a number of levels. Experiments on higher-order chromatin have started to reveal the complex communications that exist between chromosome domains and the establishment of chromosome territories within the nucleus [11]. Chromosome conformation analysis has determined the presence of topologically associated domains (TADs) and lamina-associated domains (LADs) [12]. Genes within TADs are more likely to be regulated by domains within the same TAD, and at the borders of these lie boundary elements that separate individual TADs from one another. LADs are regions of chromatin that associate with the nuclear lamina and tend to be enriched with repressed genes (Fig. 1). Within these distinct chromatin regions, gene expression is impacted by the immediate epigenetic landscape. ENCODE [3] and Roadmap [2] have worked to discover what combination of epigenetic marks are found at individual genes, and what transcription factors and co-factors assemble to deposit, maintain and modify these marks. While ENCODE sought to catalogue the regulatory elements of human cells grown in culture, Roadmap epigenomics expanded this repertoire by studying cells derived directly from human tissues in health and disease.

The two primary determinants of epigenetic landscapes remain the histone code and DNA methylation. Consequently, the substrates are the histone proteins which constitute the protein component of the nucleosome and, predominantly, the cytosine : guanosine dinucleotide (CpG) of DNA. In this review, we refer only to the canonical targets, histones H2A, H2B, H3 and H4 and modifications thereof, but there is a large body of work studying replacement histones such as H2AZ and H3.3, let alone the linker histone H1, along with non-canonical targets for DNA methylation.

Understanding what effect specific post-translational modification of histone proteins has upon gene expression capacity remains an active area of research [13]. For the illustrative purposes of this review a compendium of the modifications that assemble around an individual gene and the functions they bestow is not possible. However, consideration of the combinatorial impact of histone methylation, acetylation, phosphorylation, ubiquitylation, crotonylation and a number of other modifications lies at the heart of our ability to understand the impact upon gene expression.

Despite these challenges, expressed genes or genes for which expression is permissive upon receipt of appropriate stimuli within the cell tend to be marked epigenetically in a broadly similar way. Epigenetic landscapes can reveal capacity for both current and predictive future gene transcription compared to RNA signals, which are a record of either current or historical expression which remains in the cell until those RNAs are degraded. An average expressed gene would tend to display a complex pattern of histone modifications which includes but is far from limited to including trimethylation of lysine 4 on histone H3 at the nucleosomes that pack approximately 3 kb of DNA around the transcriptional start site (TSS) [14]. Histones around TSSs tend to be acetylated at H3K27 [15], while those that demarcate transcriptional regulatory domains such as enhancers may be decorated by both H3K27acetylation and H3K4 monomethylation [16,17], and these distal regulatory domains may be in anything from within kilobases to megabases away from the genes whose expression they regulate. H3K36 trimethylation is enrichment at transcribed exons [18] and histone acetylation at H3K9, 12 and 14 is also found around expressed genes [19]. Expressed genes also display H4K20, H3K27
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and H3K9 monomethylation [14] and H3K79 dimethylation [3]. A summary of the potential roles of some of these modifications is shown in Table 1.

In terms of genes that are epigenetically silenced, histones associated with the transcribed region and 3′ and 5′ intergenic regions of DNA are often decorated by H3K27 and H3K9 trimethylation [14] and H2AK119 ubiquitylation [20]. Thus, it becomes apparent that depending upon the nature of the modification, specific amino acids such as K9 and K27 on histone H3 can convey signals of both permissibility and prevention of transcription. How this antagonism of regulatory complexes is regulated underpins gene expression potential. How these modifications are assembled in a temporal sense is not clear, nor is it clear how precise recruitment of the dizzying numbers of catalytic activities that shape the landscape is enabled. Abundant evidence demonstrates the fact that numerous epigenome regulators exist within multi-protein complexes such as polycomb, trithorax and Spt-Ada-Gcn5 acetyltransferase (SAGA) which regulate silencing, expression capacity and nucleosome remodelling, to name but three [21–23].

Cytosine methylation is the most prevalent modification of DNA. In canonical DNA methylation, methyl groups are added to position 5 of the pyrimidine ring of deoxy-cytosine within the context of a CpG dinucleotide. The methylation status of CpG dinucleotides is not uniform among individual genes; rather, CpGs show position-specific variation in methylation. While the DNA regions proximal to TSSs of genes that are permissive for transcription tend to be depleted of DNA methylation, the areas flanking the TSS may display a greater degree of variation in methylation [24]. At genes displaying a greater concentration of CpG dinucleotides around their TSSs (CpG islands) the tendency is for CpG dinucleotides at the flanks (or shores and shelves) of the islands to show most variation in methylation status in distinct cell populations. Genes that are epigenetically silenced may have promoters that are either enriched for methylated cytosines at TSSs or at CpG island shores.

It is likely is that these two gross transcriptional states, being epigenetically silenced or actively transcribed, represent the extremities of the possible gene expression potentials, and an individual gene in any given cell is unlikely to be able to switch easily and rapidly between the two. Variation of transcriptional activity may reflect different regulatory domains being revealed or not revealed in distinct cell states. Alternatively, or additionally, transient expression of key transcription factors such as Jun, Fos and nuclear factor kappa B (NF-κB) might act upon and modify further a chromatin landscape that is already broadly permissive for transcription.

Writers and erasers: protein motifs that catalyse deposition and removal of post-translation histone modifications

One of the key advances to understanding the causal role of epigenetic landscapes on gene expression and cellular phenotype has been the identification of proteins that
deposit, remove or signal transducers of histone and DNA modifications, the so-called writers, erasers and readers of the epigenetic codes.

**Histone methylation**

The identification of the SET (Su(var)3-9, Enhancer-of-zeste and Trithorax) domain allowed the enzymes that methylate distinct lysine residues in histones to be characterized [25]. Histone methyltransferases now include proteins that are able to mono-, di- or trimethylate both lysine and arginine residues and non-SET-containing methyl transferases have also been reported such as the arginine-targeting protein arginine methyltransferases (PRMT) family [26]. These proteins use s-Adenosyl methionine as a co-enzyme and methyl donor.

In contrast, Lsd1 (KDM1A), a flavin-containing amino oxidase, was the first enzyme identified that had the ability to demethylate mono- and dimethylated histone H3 lysine 4 histone substrates [27]. Subsequently, a second, larger family of Fe (II) and 2-oxoglutarate oxygenases, which contain a catalytic domain which has been termed the 'Jumonji domain', was identified [28]. Like the SET domain, the Jumonji domain was found to be present in a large number of histone demethylase proteins, with individual proteins having the ability to attack specific residues of the histone proteins.

**Histone acetylation and crotonylation (HAT)**

Both histone acetyl transferase and deacetylase families are characterized by their sequence homologies. There are upwards of 30 distinct proteins in humans with histone acetyl transferase activity, a catalytic activity that transfers an acetyl group from acetyl coenzyme A (CoA) to the ε-amino group of a histone lysine residue. Some of these proteins can also use crotonyl CoA as their substrate. These HATs are categorized into families, first by their cellular distribution, which comprises either nuclear or cytoplasmic HATs. The nuclear HATs are categorized further by structural homology into three main groups, the GNAT (GCN5-related N-acetyltransferases) [29], MYST (MOZ: monocytic leukaemia zinc finger protein), Ybf2/Sas3, Sas2 and Tip60 (Tat interacting protein) [30] and CBP/p300 families [31].

Four classes of histone deacetylase (HDAC) proteins have been identified, and these classes are again demarcated based upon sequence homologue. Class I HDACs tend to be expressed ubiquitously and compartmentalized to the nucleus [32], whereas class IIA/IB and 4 shows some cell type restriction; these proteins have a zinc-dependent catalytic domain [33] (class III HDACs are the sirtuins, which are NAD+-dependent [34]).

**DNA methylation and demethylation**

The three mammalian DNA methyl transferases (DNMTs), DNMT1, 3a and 3b, catalyse *de novo* and maintenance DNA methylation. DNMT1 is the key maintenance methyl transferase utilizing hemimethylated DNA as its substrate, but can also undertake *de novo* methylation. DNMT3A and B are both *de novo* methyltransferases but have a lower activity than DNMT1 [35].

The ten-eleven translocation methylcytosine dioxygenase (TET) family initiate a chain of reactions that can ultimately demethylate 5 methycytosine [36,37]. This process

| Histone modification | Signal characteristic | Proposed function |
|----------------------|-----------------------|-------------------|
| H3K4me1              | Peak/region           | Marks regulatory domains, also enriched downstream of transcriptional starts |
| H3K4me2              | Peak                  | Marks regulatory elements at transcription starts |
| H3K4me3              | Peak                  | Marks regulatory domains preference for active promoters |
| H3K9Ac               | Peak                  | Marks regulatory domains |
| H3K9me1              | Region                | Marks actively transcribed genes, preference for 5’ ends |
| H3K9me3              | Peak/region           | Repressive mark |
| H3K27Ac              | Peak                  | Mark of regulatory domains, distinguishes active enhancers and promoters |
| H3K27me1             | Region                | Marks active promoters |
| H3K27me3             | Region                | Repressive marks found at silenced loci deposited by PRC2 |
| H3K36me3             | Region                | Elongation mark associated with transcribed exons |
| H3K79me2             | Region                | Transcription-associated mark |
| H3K20me1             | Region                | Marks 5’ end of active genes |
| H2AK119Ub            | Region                | Repressive mark found at silenced loci deposited by PRC1 |

PRC = polycomb repressive complex.
generates a number of intermediates (5-hydroxy methylcytosine, 5-formylcytosine and carboxylcytosine) that may also have roles as epigenetic marks [38].

Many of the writers and erasers of histone and DNA modifications have additional motifs outside of their catalytic domains which are responsible for the targeting of the protein to chromatin. These domains include chromodomains which recognize methylated histones, bromodomains which recognize acetylated histone, plant homeodomain (PHD) fingers, TUDOR and WD40 domains (reviewed in 39), demonstrating that both reader and writer functionality can reside within the same protein. In addition, histone and DNA-modifying proteins assemble into multi-protein complexes such as Complex of Proteins Associated with Set1 (COMPASS), SAGA and polycomb.

In terms of DNA methylation, both methylated and unmethylated CpG dinucleotides are recognized by specific binding proteins. MBD (methyl-binding domain-containing) and methyl CpG-binding proteins (MECP) recognize methylated DNA [40], while unmethylated CpG at TSSs are recognized by the H3K36 demethylase KDM2A [41], providing further evidence for the interplay between histone post-translational modifications and DNA methylation. The bromodomain-containing protein BRD4, which is found as a translocation product with the nuclear protein in testis (NUT) protein in human squamous carcinoma, has been targeted using bromodomain inhibitors [42]. These compounds block the binding of BRD4 to acetylated histone 4, and a therapeutic strategy using bromodomain inhibitors has been used for mixed-lineage leukaemia (MLL)-fusion leukaemia [43].

Cataloguing the epigenetic landscape in health and disease highlights regions of chromatin at which differences occur, but there needs to be a mechanistic correlate or analysis of cause or consequence to determine how these modifications influence gene expression.

Collectively, the identification of the writers and erasers of the epigenetic landscape has presented an opportunity to begin to interrogate the precise influence of epigenetic landscapes upon cellular phenotype and gene expression potential. However, with such a large number of enzymes catalysing post-transcriptional modification of histones in a tightly controlled temporal manner histone, a key challenge to being able to alter the epigenetic landscape at will has been how to deliver precisely the right catalytic domains of histone or DNA-modifying enzymes at the right time.

This has led to a number of approaches. In the first, the precise nature of the catalytic enzymes is disregarded and the goal is merely to deliver epigenetic marks that are either permissive or refractory for gene expression. Platforms such as zinc finger nucleases [44] and transcription activator-like effector nucleases (TALENs) [45] were the first reagents to employ these technologies enabling delivery to precise genomic locations. Their use has been eclipsed by the clustered, regularly interspaced, short palindromic repeats (CRISPR)-Cas9 system [46]. In a modification of this system, a nuclease dead version of Cas9 is cloned in frame with transcriptional activators such as VP16, the herpes simplex virus protein vmw65, to deliver transcriptional activation to a particular genomic location in a guideRNA-dependent manner[47]. In terms of gene repression, CRISPRinterference (CRISPRi) was described nearly half a decade ago, and refined by use of the Kruppel-associated box (KRAB) domain, a repressor of transcription which can mediate efficient silencing in mammalian cells [48]. KRAB domains are present in almost 400 human proteins, and exert their repressive capacity via recruitment of KRAB-associated protein-1 (Kap1) and heterochromatin protein-1 (HP1) to mediate H3K9 trimethylation (Fig. 2a). These reagents can bypass the epigenetic constraints of gene expression within a cell and have been used for a variety of genomewide screens to efficiently silence either single or multiple genes or silence the influence of regulatory domains in the native genomic context [49–58].

Guide RNA-directed dCas9 fusion proteins have certain features that are advantageous over short interfering and short hairpin RNAs. A particular benefit is that, as regulators of the epigenome, the duration of efficacy is potentially longer than methodologies that target RNA degradation. When combined with the use of inducible promoters such as tetracycline-induced transcriptional activation, precise control of modulation can be achieved.

More refined analysis, where the objective is to understand the functional consequence of delivery of specific methylation or acetylation modifying activities, are also beginning to be deployed. This approach has been conducted in fewer studies than have been undertaken using a more unbiased approach. Proteins that function as epigenetic regulators often have multiple modules mediating, among others, protein interaction and catalytic activity. Ideally, these modules might be isolated from one another to achieve specificity. Researchers have used the catalytic domain of p300 or an inactive mutant thereof to reconfigure the enhancer landscape and prove the involvement of histone acetylation at precise locations to alter gene expression potential [59,60] (Fig. 2b). Similarly, fusion of histone or DNA methyl transferases to Cas9 has been used to prove the influence of specific DNA methylation events [61–66].

Within the immune system only a few gene loci have been subjected to epigenome editing. Forkhead box protein 3 (FoxP3) regulatory domains were targeted with either dCas9.TET1 or dCas9.p300 catalytic domains in order to
demethylate or deposit H3K27acetylation. While altered DNA methylation did not display a potent effect, histone acetylation promoted FoxP3 expression and induction of FoxP3 target genes [67,68].

While fusion proteins of catalytic domains coupled to dCas9 can be used to deliver a particular histone or DNA modification artificially onto a chromatin template, they do not address how, in diseased cells, the altered chromatin landscape initiates, what transcription factors are differentially recruited and precisely which enzymes are the propagators of the altered landscape. These altered landscapes can, however, be interrogated to determine transcription factor binding-site enrichment. To further

**Outlook and summary**

CRISPR.Cas9-mediated epigenome editing is beginning to be employed to understand the epigenetic constraints upon gene expression, and in particular the impact of altered epigenomes in diseased cells. For such studies, however, numerous considerations arise. What is the cell type of interest? Is it possible to isolate those cells in health and disease? How many subjects should be studied? Are there distinct endotypes of disease that must be considered? Is there likely to by an impact of SNP variation on disease? Which histone modification should be studied? With the emergence of technologies permitting analysis of either single or few cells, this field of research is likely to be fertile, as the focus of analysis shifts from descriptive to analysis of functional consequence.

**Disclosures**

The authors declare no competing financial interests.

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