Polycomb eviction as a new distant enhancer function

Citation for published version:

Digital Object Identifier (DOI):
10.1101/gad.16985411

Link:
Link to publication record in Edinburgh Research Explorer

Document Version:
Publisher's PDF, also known as Version of record

Published In:
Genes & Development

Publisher Rights Statement:
Open Access Article

General rights
Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy
The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact openaccess@ed.ac.uk providing details, and we will remove access to the work immediately and investigate your claim.
RESEARCH COMMUNICATION

Polycomb eviction as a new distant enhancer function

Douglas Vernimmen,1 Magnus D. Lynch,1 Marco De Gobbi,1 David Garrick,1 Jacqueline A. Sharpe,1 Jacqueline A. Sloane-Stanley,1 Andrew J.H. Smith,1,2 and Douglas R. Higgs1,3

1MRC Molecular Haematology Unit, Weatherall Institute of Molecular Medicine, University of Oxford, John Radcliffe Hospital, Oxford OX3 9DS, United Kingdom; 2Institute for Stem Cell Research, University of Edinburgh, Edinburgh EH9 3JQ, United Kingdom

Remote distal enhancers may be located tens or thousands of kilobases away from their promoters. How they control gene expression is still poorly understood. Here, we analyze the influence of a remote enhancer on the balance between repression (Polycomb—PcG) and activation (Trithorax—TrxG) of a developmentally regulated gene associated with a CpG island. We reveal its essential, nonredundant role in clearing the PcG complex and H3K27me3 from the CpG island. In the absence of the enhancer, the H3K27me3 demethylase (JMJD3) is not recruited to the CpG island. We propose a new role of long-range regulatory elements in removing repressive PcG complexes.

Supplemental material is available for this article.

Received May 9, 2011; revised version accepted July 5, 2011.

It is still not fully understood how genes are switched on or off during differentiation and development. However, the role of remote regulatory elements in this process is thought to be critical [Bulger and Groudine 2011]. One important epigenetic mechanism regulating many genes involves the opposing effects of the Polycomb [PcG, repressive] and Trithorax group [TrxG, activating] proteins, which play critical roles in stem cell biology, development, and cancer (Simon and Kingston 2009). The promoters of most PcG/TrxG target genes are associated with CpG islands that remain unmethylated in all cell types [Mendenhall et al. 2010; MD Lynch, AJ Smith, M De Gobbi, M Flenley, JR Hughes, D Vernimmen, H Ayyub, JA Sharpe, JA Sloane-Stanley, L Sutherland, et al., in prep.]. During lineage commitment and differentiation, PcG complexes are cleared from silenced genes and replaced by TrxG complexes as these genes are switched on. Specific histone methyltransferases in these complexes create characteristic chromatin signatures when PcG [H3K27me3] or TrxG [H3K4me3] complexes are bound to chromatin. Bivalent chromatin patterns [H3K27me3 and H3K4me3] may be seen prior to commitment, and these two chromatin marks are then resolved during differentiation [Sawarkar and Paro 2010; Surface et al. 2010]. Bivalent domains may also arise de novo during differentiation [Roh et al. 2006; Mohn et al. 2008]. The recruitment and removal of PcG and TrxG complexes are a dynamic process. However, the potential of remote cis-acting elements to influence the balance between repression and activation throughout development and differentiation has not been explored. To address these issues, we characterized changes in the recruitment of transcription factors (TFs), cofactors, and chromatin-associated proteins [including PcG and TrxG], and associated chromatin modifications to the human α-globin CpG island in the presence or absence of a distant enhancer.

A major problem in analyzing these aspects of gene regulation arises when perturbation of the gene in question [e.g., a key TF] leads to a perturbation of cell fate. In this case, the cellular phenotype alters as regulation of the gene under investigation is altered. To circumvent this, such experiments are often done in cell lines whose environment, the normal and mutant cells are directly comparable (Vernimmen et al. 2008). The recruitment and removal of PcG and TrxG nuclease during differentiation (Roh et al. 2006; Mohn et al. 2008). The recruitment and removal of PcG and TrxG complexes are a dynamic process. However, the potential of remote cis-acting elements to influence the balance between repression and activation throughout development and differentiation has not been explored. To address these issues, we characterized changes in the recruitment of transcription factors (TFs), cofactors, and chromatin-associated proteins [including PcG and TrxG], and associated chromatin modifications to the human α-globin CpG island in the presence or absence of a distant enhancer.

Here we show that, in erythroid cells, the remote tissue-specific enhancer [MCS-R2] plays an essential, nonredundant role in clearing the PcG complex and its associated modification [H3K27me3] from the CpG island, when the α-globin genes become fully activated. Furthermore, in the absence of the enhancer, the CpG island does not recruit the H3K27 demethylase JMJD3. This demonstrates that in addition to the recruitment of TFs, cofactors, and the preinitiation complex [PIC] at the promoter, long-range regulatory elements also play a critical role in removing repressive PcG complexes.

Results and Discussion

Using the humanized mouse, we previously showed that the removal of MCS-R2 dramatically reduces the formation of chromosomal looping, which normally occurs between the upstream regulatory elements and the promoter in erythroid cells. In the absence of MCS-R2, the human α-globin genes are expressed at <2% of normal, in erythroid cells [Vernimmen et al. 2009]. Here, we first determined the effect of MCS-R2 on recruitment of key, cell type-specific, and ubiquitously expressed TFs known
Vernimmen et al.

Figure 1. Generation of humanized mice as a model to study the human α-globin locus organization and methylation at key promoters, late during erythropoiesis. Of interest, there is a progressive decrease in TF occupancy at key promoters (Supplemental Table 1). These findings are reflected by suboptimal histone acetylation [Supplemental Table 1]. Of interest, there is a progressive decrease in TF occupancy at key cis elements [MCS-R1/HS-48, MCS-R3/HS-33, and MCS-R4/HS-10] along the chromosome, with the α-globin promoter being the most affected [Supplemental Fig. 1; Supplemental Table 1]. Furthermore, in the absence of MCS-R2, ubiquitination of H2B [H2Bub, associated with activation] does not occur efficiently at the α-globin gene [Fig. 2, left panel]. Ubiquitination of H2B is thought to be required for subsequent methylation of H3K79 by Dot1 [Shilatifard 2008], and without MCS-R2, H3K79 is not efficiently methylated [Fig. 2; Supplemental Fig. 2]. As previously described [Vernimmen et al. 2007], without MCS-R2, occupancies of the PIC components [TBP, TFIIE, and TFIIB] were all low with an even more dramatic effect on PolII [Fig. 2; Supplemental Fig. 1] compared with the normal chromosome. These findings confirm that in the humanized mouse cells [as in primary human and mouse cells], a key role of the upstream element (MCS-R2) is to facilitate recruitment and/or stabilization of the PIC at the α-globin promoter, late during erythropoiesis.

We then evaluated the role of MCS-R2 on the deposition of methyl groups to the Lys 4 residue of histone H3. As in human primary erythroblasts [De Gobbi et al. 2007], the upstream elements are marked by mono- and dimethyl H3K4 in fully active humanized erythroid cells [Supplemental Fig. 3]. In contrast, the chromatin associated with the α-globin promoter and the body of the gene is modified by H3K4me3 [Fig. 2, right panel; Supplemental Fig. 3]. These findings are also consistent with the chromatin features of enhancers and promoters, as established in genome-wide analyses [Heintzman et al. 2007]. In the absence of MCS-R2, the mono- and dimethylation of H3K4 seem to occur normally [Supplemental Fig. 3, left panel]. Moreover and more unexpectedly, despite the absence of detectable H2Bub and H3K79me3 and the associated transcription (expression of only ~2% of normal), the overall level of H3K4me3 at the α-globin gene was ~50% of that seen in the fully active, highly transcribed gene [considering the amount of H3K4me3 detected on the endogenous control] [Fig. 2; Supplemental Fig. 3]. Note that, in the absence of MCS-R2, the distribution of mono-, di-, and trimethyl H3K4 becomes skewed more to the promoter than the body of the gene. This might be partially due to higher levels of H3 occupancy at the promoter in the ΔMCS-R2 compared with normal [Supplemental Fig. 4]. Of interest, in the absence of the enhancer, H3K36me3 [a marker of transcriptional elongation] is also detectable [Supplemental Fig. 5].

The H3K4me3 modification is imposed by the human TrxG proteins, which include several complexes [hSet1 and MLL1-5] [Shilatifard 2008]. It has recently been shown that H3K4 methylation at CpG islands may be mediated via the hSet1 complex, which contains a CpG island-binding protein [CBP, or Cfp1]. CBP is thought to bind unmethylated CpG-rich sequences irrespective of transcription, thereby recruiting hSet1 [Thomson et al. 2010]. This implies that chromatin at unmethylated CpG islands might be modified by H3K4me3 as a default state. Here we found that in human nonerythroid cells, CBP/Cfp1 is not detectable at the associated CpG island (as judged by two different antibodies) [Supplemental Fig. 6]. Similarly, when the α genes are severely down-regulated in the humanized mouse MCS-R2 mutant, CBP/Cfp1 is present at very low levels [Fig. 2; Supplemental Fig. 7]. Thus, CBP/Cfp1 is only readily detected in association with high levels of α-globin expression. Therefore, it seems unlikely that the relatively high levels of H3K4me3 seen in the ΔMCS-R2 mutant [when the α genes are expressed at relatively low levels] reflect a default state mediated by CBP/Cfp1. In contrast, it suggests...
that H3K4me3 might be a sensitive mark of basal transcription. Interestingly, it was previously suggested that CpG islands marked by H3K27me3 may exclude binding of CGBP (Thomson et al. 2010). We showed previously that a functionally repressive PcG complex is bound at the CpG islands associated with the human α-globin promoters in human nonerythroid cells, in which the α genes are transcriptionally silent (Garrick et al. 2008). Here, we show that CGBP is indeed excluded from CpG islands bound by PcG by comparing human nonexpressing (PcG-bound) versus expressing (PcG-unbound) cells (Supplemental Fig. 6). We therefore next determined whether the lack of CGBP recruitment in the ΔMCS-R2 mutant is associated with persistent PcG binding.

PcG complex is normally bound to both the adult (α) and embryonic (ζ) globin CpG islands in human embryonic...
stem cells and pluripotent progenitors, in which expression of the α genes is detectable at extremely low levels [De Gobbi et al. 2011]. Polycomb remains bound to the silenced embryonic α gene in mature erythroid cells. In contrast, PcG is cleared from the adjacent α-globin CpG island as this gene becomes expressed at high levels in mature erythroid cells [Garrick et al. 2008; De Gobbi et al. 2011]. This pattern is fully recapitulated in the humanized mouse system [Fig. 3, right panel; Supplemental Fig. 8]. Moreover, we show here that as the PRC2-associated chromatin modification [H3K27me3] is erased, the histone H3K27 demethylase JMJD3 is recruited [Fig. 3, right panel; Supplemental Fig. 8].

Figure 3. Removal of Polycomb repressor complexes is dependent on MCS-R2 and associated with recruitment of JMJD3. Graphs are displayed as in Figure 2. Additional amplicons flanking the embryonic α-globin gene [142, 147, and 149] have been used in these experiments to highlight the two independent PcG domains observed in the mutant human allele ΔMCS-R2 (shown in the left panel). The shaded area represents the adult α-globin CpG island, from which PcG is normally cleared in the Normal allele in erythroid cells (shown in the right panel). [Left] In the absence of MCS-R2, PcG is not cleared from the α-globin genes and thus remains present at both domains. M Gata6 represents a positive control for PcG binding. The JMJD3 antibody used recognizes the N-terminal region of the protein (Millipore).
We next studied humanized mice in which MCS-R2 had been deleted to determine whether the upstream enhancer normally plays a role in clearing the PcG complex from the \( g \)-globin CpG island during erythropoiesis. As before, in the absence of MCS-R2, the silenced \( g \)-globin gene remains bound by PcG throughout differentiation. However, in the absence of MCS-R2, PRC2 is no longer cleared from the \( g \)-globin CpG island, and JMJD3 is no longer recruited [Fig. 3, left panel, Supplemental Fig. 8]. We showed previously that PcG repression of the \( g \)-globin gene may be mediated (at least in part) by HDAC1, which is normally cleared with PcG complex during erythropoiesis [Garrick et al. 2008]. Here, in the absence of MCS-R2, HDAC1 remains bound to the \( g \)-globin genes [Fig. 3]. These contrasting observations on two adjacent CpG islands show that MCS-R2 exerts a specific effect on PcG clearance from the \( g \)-globin (but not the \( z \)-globin) gene during adult erythropoiesis.

There is increasing evidence that CpG islands, such as those associated with the \( g \)-globin promoter, constitute at least one element that can mediate recruitment of PcG and TrxG complexes to mammalian promoters [Mendenhall et al. 2010; MD Lynch, AJ Smith, M De Golbi, M Flenley, JR Hughes, D Vernimmen, H Ayyub, JA Sharpe, JA Sloane-Stanley, L Sutherland, et al., in prep.]. As previously noted, PcG-binding sites are dynamic, are nucluesome-depleted, and have a rapid histone turnover [the residency time of PcG is in the order of a few minutes] [Ficz et al. 2005]. PcG binding is therefore thought to be dynamic and sensitive to the antagonistic action of TrxG proteins together with positive and negative input from other TFs and cofactors. However, it is not known whether the eviction of PcG silencing from its targets, seen during development and differentiation, depends on the presence of distal regulatory elements or only on (co)factors acting at proximal \( cis \) elements. In this study, we used a mouse experimental model to analyze the CpG island associated with the human \( g \)-globin promoter in two states: without and with its interacting distant enhancer, both in terminally differentiated erythroid cells. We also compared the recruitment of CGBP in nonexpressing versus expressing human cells. In nonerythroid cells, the unmethylated, nucluese-insensitive CpG island associated with the \( g \)-globin gene is bound by PcG and is transcriptionally silenced [referred to as the “silent state”] [Fig. 4A]. In erythroid cells without MCS-R2 [referred to as “basal state”], and in contrast to nonerythroid cells [Garrick et al. 2008], the promoter becomes accessible to some TFs and is associated with some active chromatin modifications [e.g., H3K4me3 with relatively low levels of transcription (~2% of normal)] [Fig. 4B]. Nevertheless, the PcG complex with its associated modification (H3K27me3) is still prominent at the \( g \)-globin CpG island. We also demonstrate here that PcG and CGBP binding are mutually exclusive [cf. Figs. 2 and 3]. In erythroid cells with MCS-R2 (referred to as “active state”), PcG complexes are completely removed from the CpG island [Fig. 4C]. Furthermore, the histone H3K27 demethylase JMJD3, which may remove H3K27me3 and thereby facilitate transcription, is also recruited at high levels. Recruitment of the SAGA complex [e.g., PCAF and GCN5] becomes prominent and the downstream effects [e.g., deposition of H2Bub and H3K79 methylation] are established. At this stage, high levels of transcription are associated with binding of CGBP. We thus show that the recruitment of the demethylase JMJD3 and full clearance of the PcG-repressive complex [including PRC2 and HDAC1] at the \( g \)-globin CpG island depend on one or more activities mediated by the remote regulatory element and are associated

---

**Figure 4.** Long-range control of epigenetic regulation. (A) In nonerythroid cells, the CpG island is entirely silenced by PcG and HDAC1, associated with the repressive histone mark H3K27me3. The promoter “P” is not sensitive to DNaseI, and transcription does not occur. (B) In erythroid cells lacking the enhancer, the gene remains repressed by PcG and marked by H3K27me3. At this basal level of expression, the promoter becomes accessible to some TFs and chromatin-modifying enzymes and is marked by moderate levels of H3K4me3, which reflect very low levels of transcription. (C) In the presence of the enhancer, PcG is evicted and the H3K27me3 histone mark is erased by recruitment of demethylase JMJD3. Acetylation [H3ac and H4ac], H3K79me3, and H2Bub increases with spreading of HAT and Bre (SAGA) along the coding sequence “C.” At this activated stage, the remaining TFs, including Pol II, are now fully recruited, and a high rate of transcription occurs. The CpG island at this stage is also bound by CGBP.
with the transition between basal and fully activated transcription.

These findings demonstrate for the first time that the pattern of PcG binding at a CpG island may be affected by cis-acting elements located far away from the associated promoter. In contrast, the chromatin modification associated with TrxG activity [H3K4me3] appears to be more dependent on local changes at the CpG island that occur in the context of basal transcription. Future studies will address how long-range enhancers exert these effects. It is possible that transcriptional activation per se competes with the competitive binding of PcG complexes and is responsible for the clearance of these complexes [MD Lynch, AJ Smith, M De Gobbi, M Flenery, JR Hughes, D Vernimmen, H Ayyub, JA Sharpe, JA Sloane-Stanley, L Sutherland, et al., in prep.]. The second is that upstream elements also deliver new proteins [e.g., JMJD3] or modify proteins [e.g., histones] that facilitate the removal of PcG. In the past, detailed analysis of the globin genes has established many of the general principles underlying mammalian gene regulation, and it therefore seems probable that this new role of distal regulatory elements in removing PcG from their target promoters will be of considerable general importance.

Materials and methods

Primary cells

Ter119-positive mature primary mouse erythroid cells [humanized] were obtained by automagnetic-activated cell sorting, as previously described (Vernimmen et al. 2007, 2009; Wallace et al. 2007). Primary human erythroid blasts were obtained from peripheral blood mononuclear cells (PBMCs) collected from blood donors and expanded in a two-phase system as previously described [De Gobbi et al. 2007, Epstein-Barr virus (EBV)-transformed lymphoblastoid cell lines were derived from healthy subjects.

Chromatin immunoprecipitation assay (ChIP)

ChIP was performed as previously described [Vernimmen et al. 2007]. For histone-modifying enzymes, chromatin was first cross-linked with EGS [Pearce, Thermo Scientific, product no. 21565] in PBS at a final concentration of 2 mM for 60 min at room temperature. Formaldehyde was then added at a final concentration of 1% for 15 min at room temperature, and samples were sonicated for 20 min at 4°C to fragment genomic DNA [Bioruptor, Diagenode]. The antibodies used were Pol II (H-224), JMJD3 (N-24), GCN5 (H-424), PCAF (H-369), CBP (A-22) (purchased from Santa Cruz Biotechnology); H3K79me3 (ab2621), H3K27ac (ab1791), CGBP (ab56035), H3K4me3 (ab8085), and Suz12 (ab12073) [purchased from abcam]; DOT1L (A300-954A) [purchased from Rethyl Laboratory, Inc.]; HDAC1 (06-720), H3ac (06-599), H4ac (06-866), H3K9ac (07-352), H3K14ac (07-353), H3K27me3 (07-449), H3K4me1 (07-436), H3K4me2 (07-030), H3K4me3 (05-745R), H4 (07-108), H2A (06-13923), H2B (06-1790), JMJD3 (07-1533), and HDAC1 (07-1534) [purchased from Millipore]; H2Bub [MM-0029] [purchased from Medimabs]; and Ezh2 (PAB0649) [purchased from Abnova]. Real-time PCR using primers and probes (S FAM–3′TAMRA) for murine and human α-globin locus were described previously [Anguita et al. 2004, De Gobbi et al. 2007].

Acknowledgments

We are very grateful to Dave Skalnik for CGBP antibody. Many thanks to Jane Mellor, Tom Milne, Nick Proudfoot, Roger Patien, Maggie Walmsey, and Richard Gibbons for critically reading the manuscript. Furthermore, we thank Sue Butler and Christina Rode for technical assistance. This work was funded by the Medical Research Council and the National Institute for Health Research [NIHR] Biomedical Research Centre Program.

References


