

The background features a light gray gradient. On the left side, there are several thick, overlapping, swirling lines in shades of maroon, brown, and blue. Scattered across the entire page are numerous small, semi-transparent dots in various colors, including red, orange, yellow, light blue, and pink, creating a dynamic, particle-like effect.

# Genome Binding and Gene Regulation by Stem Cell Transcription Factors

Johannes Hendrik Brandsma

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Johannes Hendrik Brandsma

**Colofon**

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# **Genome Binding and Gene Regulation by Stem Cell Transcription Factors**

Binding aan het genoom en genregulatie door  
stamceltranscriptiefactoren

## **Proefschrift**

Ter verkrijging van de graad van doctor aan de  
Erasmus Universiteit Rotterdam  
op gezag van de rector magnificus  
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en volgens besluit van het College voor Promoties

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Overige leden: Prof.dr. J.N.J. Philipsen  
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Prof.dr. W.L. de Laat

Copromotor: Dr. R.A. Poot

“At every step, he felt as a man might feel who, after admiring the smooth, cheerful motion of a boat on the water, actually gets into the boat himself. He saw that apart from having to sit steadily in the boat without rocking, he also had to keep in mind, without forgetting for a moment where he was going, that there was water beneath his feet, that he had to row, that his unaccustomed hands hurt, and that it was easy only when you looked at it, but that doing it, though it made you very happy, was very hard.”

*Leo Tolstoj, Anna Karanina , Bantam classic reissue (English translation), 2006*



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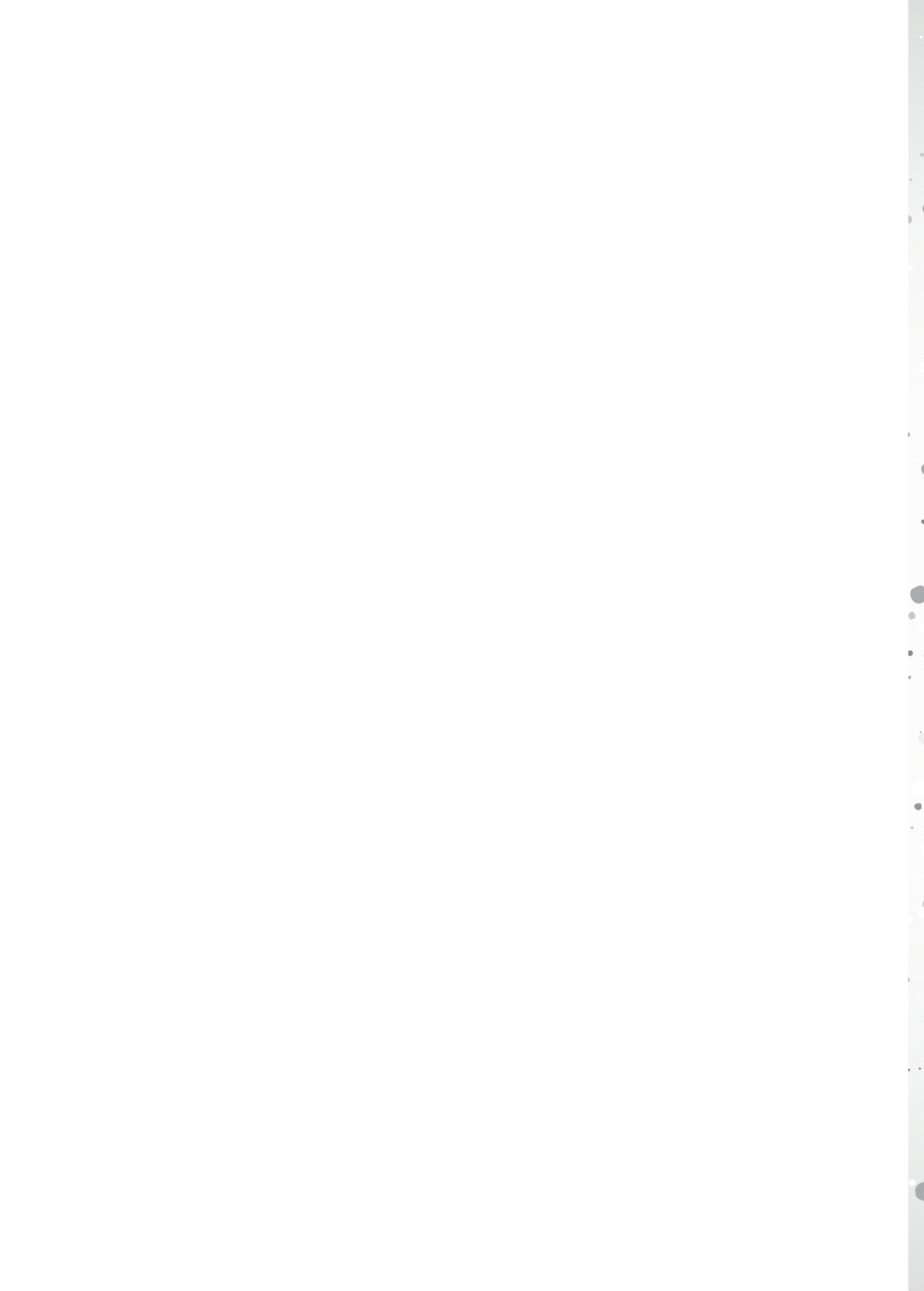
## LIST OF ABBREVIATIONS

ASD	autism spectrum disorders
bp	basepair
CCHS	Congenital Central Hypoventilation Syndrome
ChIP	Chromatin Immunoprecipitation
ChIP-MS	ChIP followed by Mass spectrometry
ChIP-qPCR	ChIP followed by quantitative PCR
ChIP-seq	ChIP followed massively parallel DNA sequencing
DNA	Deoxyribonucleic acid
E	Embryonic day
emPAI	Exponentially modified protein abundance index
ESC	Embryonic Stem Cell
GO	Gene ontology
GWAS	Genome-wide association study
ID	Intellectual disability
IP	Immunoprecipitation
IP-MS	Immunoprecipitation followed by mass-spectrometry
iPSC	Induced pluripotent stem cells
kDa / kD	kilo Dalton
KD	Knock Down
KO	Knock Out
LOF/LoF	loss of function
MD	Mental disorders
mRNA	messenger RNA
NSC	Neural Stem Cell
Oct4	Octamer-binding transcription factor 4
PCR	polymerase chain reaction
PolII	RNA Polymerase II
RNA	Ribonucleic acid
SNP	Single nucleotide polymorphism
SO-motif	Sox2-Oct4 composite DNA binding motif
Sox2	SRY (sex determining region Y)-box 2
STARR-seq	Self-transcribing active regulatory region sequencing
SZ	Schizophrenia
TAD	topological associated domain
TSS	Transcription Start Site
wt	wild type

## OUTLINE

Nearly all cells of an individual organism contain the same genome. However, each cell type transcribes a different set of genes due to the presence of different sets of cell type-specific transcription factors. Such transcription factors bind to regulatory regions such as promoters and enhancers and regulate their activity in gene transcription. Transcription factors interact with each other and form tissue-specific transcription factor networks. Identification of genome binding and gene regulation by transcription factors will enhance our understanding of how transcription factors specify cell types.

**Chapter 1** serves as a general introduction into eukaryotic transcription regulation and describes the role of enhancers and transcription factors. Chapters 2 – 4 describe the experimental work of this thesis. **Chapter 2** describes Chromatin Immunoprecipitation followed by Mass Spectrometry (ChIP-MS), a technique used to identify transcription factors and other genome binding proteins that bind to promoters, enhancers or heterochromatin. **Chapter 2** specifically describes the genome-binding pattern of transcription- and pluripotency factor Dppa2, which we identify by ChIP-MS and ChIP-seq to bind to promoters outside the classical pluripotency network. **Chapter 3** describes a transcription factor interaction network of over 200 proteins in neural stem cells, which was assembled by the identification of interaction partners of four mental disorder-associated transcription factors. **Chapter 4** describes work on the genome-wide localization of Oct4, Sox2 and Nanog in embryonic stem cells and investigates the genome localization of these transcription factors upon depletion of Oct4 or Sox2 from embryonic stem cells. The last chapter, **Chapter 5**, summarizes the results from Chapter 2-4 and provides additional discussion to these chapters, concerning its implications and future directions.





# Chapter1

Introduction

## INTRODUCTION

The most abundant class of cis-regulatory elements are enhancers<sup>1</sup>. Around half a million putative enhancers have been identified in the human genome<sup>2,3</sup>, vastly surpassing the number of human protein-coding genes. The number of active enhancers in a single cell type ranges in the ten thousands<sup>2</sup>. Enhancer activity is cell-type specific and controls cell identity by allowing the regulation of gene expression<sup>1,2,4-6</sup>. Enhancers are cis-acting regulatory sequences that stimulate transcription of genes by interacting with their promoters<sup>7</sup> and enhancer-promoter looping is a prerequisite for transcription<sup>8</sup>. Enhancers contain DNA motifs that allow the specific binding of certain transcription factors and are often bound by multiple transcription factors<sup>9</sup>. Transcription factors recruit factors that modify chromatin accessibility and recruit co-activators such as P300 and the Mediator complex to the enhancer. The Mediator complex interacts with transcription factors that are bound to the enhancer and with the pre-initiation complex and RNA polymerase II (PolII) that are bound to the promoter through chromatin looping and as such driving transcription (see also below)<sup>10</sup>. As enhancers play an important role in gene regulation, it can be expected that malfunction in enhancer-mediated gene regulation might be the underlying cause for several genetic diseases. Enhancers and other cis-regulatory regions harbor common trait and disease variants<sup>6,11,12</sup>. Single-nucleotide polymorphisms (SNPs) identified in Genome-wide association studies (GWAS) were found to be particularly enriched on enhancers and transcription start sites (TSSs) for several cell types<sup>2</sup>. Many more mutations in enhancers are expected to be found with the recent onset of next generation sequencing technology. This chapter aims to summarize these recent findings surrounding enhancers and to discuss the role of enhancers in disease. First the genome-wide techniques that were utilized to increase our understanding of enhancers are summarized. Then the current understanding of transcriptional enhancers and their role in the transcriptional regulatory landscape and development is discussed. Finally, the aberrant behavior of enhancers during disease is discussed and illustrated with examples from the literature.

### Identification and characterization of enhancers using NGS techniques

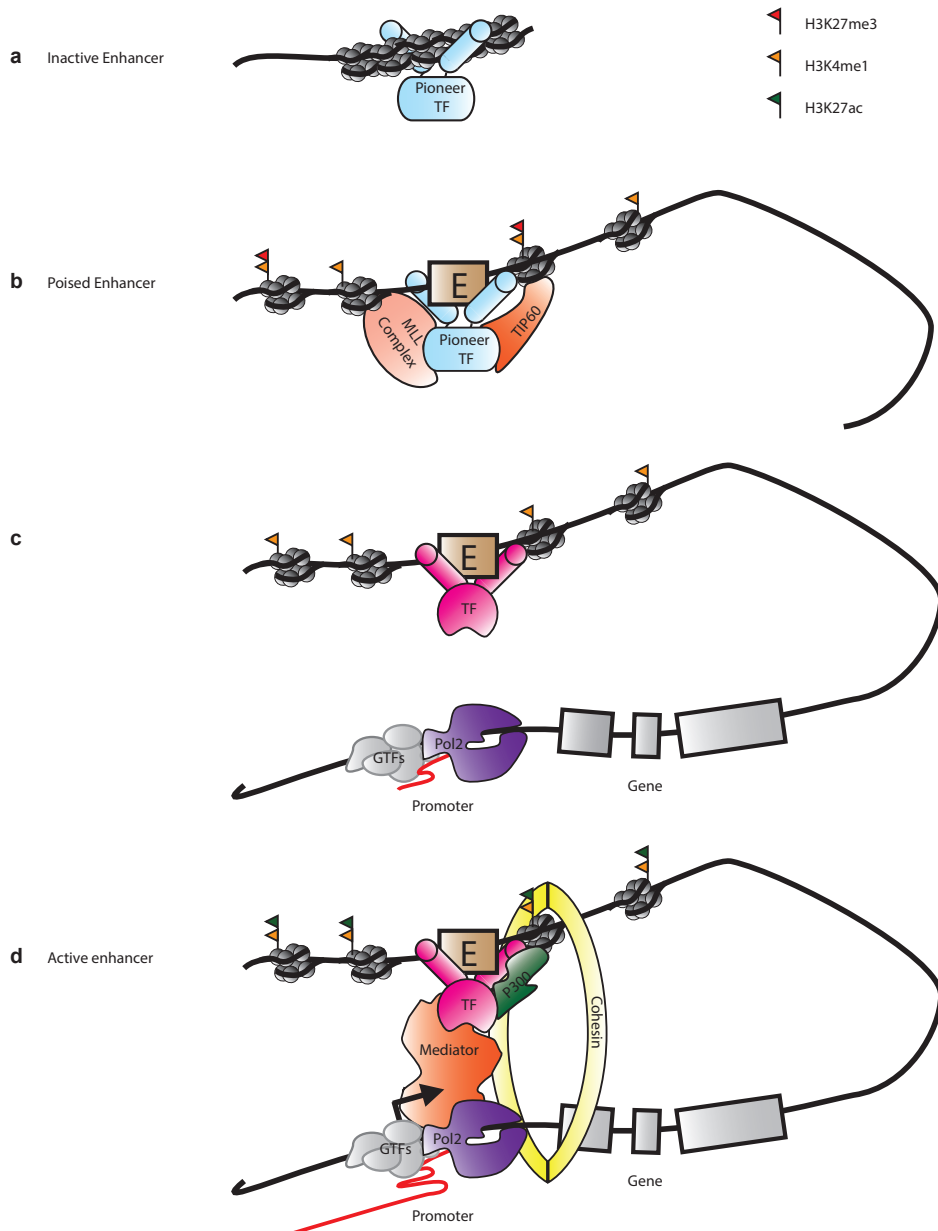
DNA is wrapped around nucleosomes. Nucleosomes are formed by an octamer of histone proteins, that is wrapped by 147 base pairs of DNA. The octamer is formed by two heterodimers consisting of H3 and H4 histones flanked by a H2A- and H2B histone. The N-terminal sequences of H3 and H4 histones, also called the histone tails, protrude from the nucleosome core and are post-translationally modified (i.e. 'histone modifications'). Histone modifications mark distinct regions of the genome and may affect chromatin structure and may attract proteins such as transcription factors and co-activators (reviewed in <sup>13</sup>). Several techniques have been developed that can be used to identify and characterize enhancers genome-wide (reviewed in <sup>14</sup>, see table 1) and the leading technique in this respect, Chromatin Immunoprecipitation followed by deep sequencing (ChIP-seq), targets histone modifications that mark enhancers. Enhancers are marked by H3 mono-methylation at Lysine 4 (H3K4me1) and for genome-wide identification of enhancers ChIP-seq can be utilized by targeting enhancer marker H3K4me1<sup>5,15</sup>. In combination with a ChIP-seq for acetylation of H3K27 (H3K27ac) active enhancers can be identified<sup>15</sup>. ChIP-seq for these histone modifications can be conveniently performed from 1 million cells. Several specialist approaches have been

developed for histone ChIP-seqs on reduced numbers of cells ranging from 100,000<sup>16</sup>, 10,000<sup>17</sup>, 5,000<sup>18</sup> to 1,000 cells<sup>18,19</sup>. ChIP-seq for co-activator P300<sup>20</sup> or subunits of the co-activator complex Mediator<sup>6</sup> have also been used to identify enhancers. Other approaches constitute of multiple ChIP-seqs against different key transcription factors of a cell type and identifying so-called multi transcription factor binding loci<sup>9,21</sup>. DNaseI hyper sensitivity assay followed by deep sequencing (DNase-seq) identifies the open regions of the genome, which includes enhancers, but also other genetic regulatory elements such as promoters, silencers and insulators<sup>22</sup>. Another technique that assesses open chromatin regions are transposase-accessible chromatin using sequencing (ATAC-seq)<sup>23</sup> and formaldehyde-assisted isolation of regulatory elements (FAIRE-seq)<sup>24</sup>.

Method	Characterization	Remarks
ChIP-seq against H3K4me1	Genomic location	Does not distinguish between active and poised enhancers
ChIP-seq against H3K27ac	Genomic location of active enhancers	H3K27ac also marks promoters
ChIP-seq against P300	Genomic location of active enhancers	P300 also binds to promoters and is sometimes present on poised enhancers
ChIP-seq against Mediator	Genomic location of active enhancers	Mediator is also present on promoters
ChIP-seq against key transcription factors	Genomic location	Does not distinguish between active and poised enhancers. Only identifies a subset of all enhancers
DnaseI Hypersensitivity Sequencing, FAIRE-seq, ATAC-seq	Genomic location of open regions	Does not distinguish between enhancers and other regulatory elements
RNA-seq and GRO-seq	Genomic location and activity	Not all enhancers produce eRNAs
HiC, CaptureC, T2C	Genomic Interaction	Often lacks the resolution to map interactions between proximal enhancers
ChIA-PET	Genomic location and interaction	
STARR-seq	Activity	Measures potential activity of genomic sequences.
ChIP-MS against H3K4me1	Enhancer binding proteins	Does not characterize individual enhancers
ChIP-MS against H3K27ac	Active enhancer binding proteins	Does not characterize individual enhancers

**Table1: Overview of OMICS methods to characterize Enhancers**

Enhancers can also be identified via RNA-seq by detection of so-called enhancer RNAs (eRNAs, see below). An interesting RNA-seq variant is Global run-on sequencing (GRO-seq)<sup>25</sup>, which instead of quantifying global transcript levels, specifically assesses ongoing transcription in the nucleus. An advantage to GRO-seq is that it can be used to identify eRNAs and simultaneously assess transcriptional activity of nearby genes<sup>26</sup>. Chromosome Conformation Capture (3C) and evolved 3C-variants, form another very important category of techniques for characterizing enhancers and other cis-regulatory regions. These techniques enable the identification of chromatin domains that are in close proximity (i.e. 'interaction of chromatin')<sup>27</sup> and can be used to confirm the interaction of an enhancer with its promoter. HiC is the only genome-wide variant within the 3C family and can be used to map all chromatin-chromatin interactions with an average resolution of 10.5 kb<sup>28</sup>. Two recently published 3C techniques known as Capture-C<sup>29</sup> and T2C<sup>30</sup> are not genome-wide, but offer an improved average resolution of 2 – 6 kb<sup>30</sup> and do not rely on ultra-deep sequencing<sup>30</sup>. Another technique that combines ChIP with 3C technology is called 'chromatin interaction analysis by paired-end tag sequencing' (ChIA-



**Figure 1: Model of enhancer activation in Eukaryotes.** (a) A DNA motif in a genomic region with closed unmarked chromatin is bound by a pioneer transcription factor. (b) The Pioneer transcription factor attracts chromatin modifying complexes like e.g. the MLL complex that monomethylates H3K4. H3K4me1 attracts the TIP60 complex that deposits H2A.Z, which leads to open and more accessible chromatin. Poised enhancers are sometimes also marked by H3K27me3. (c) The increased accessibility of the chromatin allows other transcription factors to recognize their DNA binding motifs and bind the genome. The promoter may already be bound by general transcription factors of the pre-initiation complex and Polymerase II. (d) These transcription factors recruit co-activators like p300 that acetylates H3K27 and the Mediator complex

*Legend continues on the bottom of the next page*

PET). ChIA-PET is targeted against (e.g.) a transcription factor. This allows identification of the binding regions of the transcription factor, but also identifies regions that are interacting with the binding region<sup>31</sup>. Self-transcribing active regulatory region sequencing (STARR-seq) assesses the transcriptional potential of enhancers. In this high throughput technique, potential enhancers are cloned into a construct downstream of a minimal promoter. Activation of transcription by the potential enhancers results in transcription of the enhancer itself. After a cell line has been transfected with the constructs, RNA is isolated and analyzed for over-represented potential enhancers<sup>32</sup>. To identify enhancer-bound proteins, Chromatin Immunoprecipitation followed by Mass Spectrometry (ChIP-MS) for H3K4me1 can be utilized (see Chapter 2).

### Establishment and activation of enhancers

Enhancers can be separated in three categories. Inactive enhancers, poised enhancers and active enhancers (figure 1). In contrast to poised and active enhancers, inactive enhancers cannot be distinguished from the general chromatin landscape by e.g. chromatin marks or transcription factor binding<sup>33</sup>. Poised and active enhancers are both characterized by H3K4me1<sup>5,34</sup>, whereas active enhancers also exhibit H3K27ac<sup>35,36</sup>. In addition, poised enhancers are often also enriched for H3K27me3<sup>37</sup> (figure 1). In an exemplary study using STARR-seq, a third of the potential active enhancer sequences were shown to be silenced by H3K27me3, but were able to drive transcription outside their genomic environment<sup>32</sup>. Poised and active enhancers are equally likely to interact with the TSS<sup>28</sup>.

Pioneer transcription factors play an important role in establishing poised enhancers. Pioneer transcription factors are a special class of transcription factors, which are able to engage their target sites in closed and silent chromatin lacking any evident histone marks making it accessible for other chromatin binding proteins<sup>38</sup> (reviewed in<sup>39</sup>). However, not all types of closed chromatin are permissive to pioneer factors. Heterochromatin consisting of H3K9me2 or H3K9me3 impedes pioneer transcription factor binding<sup>38,39</sup>. H3K27me3 marked silenced chromatin is probably more permissive, but this remains to be investigated. A study on a pioneer transcription factor PU.1 in macrophages and B cells demonstrated that PU.1 initiates nucleosome remodeling, which increases the accessibility to other TFs and eventually facilitates mono-methylation of H3K4<sup>40</sup>. Based on this and other data<sup>41</sup>, a general model was proposed in which pioneer transcription factors make unmarked closed chromatin accessible, allowing other transcription factors to bind to the chromatin and attract chromatin remodelers such as the MLL complex, which catalyzes the mono-methylation of H3K4<sup>39,40</sup> (figure 1). In *Drosophila* it was shown that H3K27ac signal always coincides with H3K4me1 signal on enhancers, while H3K4me1 does occur without H3K27ac<sup>36</sup>. This suggests that the poised state probably always precedes the active state in enhancers. Other studies in mouse and human do report H3K27ac apart from H3K4me1 on enhancers<sup>35,37,42</sup>, but it was suggested that this is due to under-sampling of the H3K4me1 signal<sup>36</sup>. The biological function of H3K4me1 is still

(Figure 1. Legend continues from previous page)

that bridges regulatory signals to the promoter and stimulates initiation or elongation of transcription by Polymerase II. Cohesin acts to stabilize the enhancer-promoter loop. E: Enhancer. TF: Transcription factor. GTFs: General transcription factors. PolII: RNA Polymerase II. Black line: DNA, Red line: RNA. Flags indicate presence of histone modification: H3K27me3 (red), H3K4me1 (orange), H3K27ac (green).

under debate, but has been suggested to specifically attract the TIP60/p400 complex that catalyzes deposition and acetylation of the H2A.Z, which leads to increased accessibility of enhancer chromatin<sup>43</sup>.

Poised enhancers allow for rapid activation of gene expression. Upon IFN- $\gamma$  stimulation in HeLa cells, STAT1 bound to unmarked genomic regions and to poised enhancers marked by H3K4me1. The genes associated with STAT1 bound poised enhancers showed increased expression within 30 minutes after IFN- $\gamma$  stimulation, while genes associated with unmarked STAT1 binding sites did not<sup>5</sup>. H3K4me1 may also allow for rapid reactivation of enhancers. Macrophages stimulated with LPS, activated inactive enhancers. When the macrophages were no longer stimulated with LPS, H3K27ac was rapidly lost, while H3K4me1 was sustained. Renewed LPS stimulation resulted in faster induction of gene expression than observed with the initial LPS stimulation<sup>44</sup>.

Poised enhancers can also be bound and activated by transcription factors that are expressed in later development or differentiation. For example, the enhancer for the  $\lambda 5$  gene is bound by Sox2 and Foxd3 and kept in a poised in embryonic stem cells (ESCs). Later in pro-B cells this enhancer is bound by Sox4 and the  $\lambda 5$  gene is highly expressed<sup>45</sup>. Another example is the enhancer for the *Alb1* gene, which is also kept in a permissive state by Foxd3. Upon mesodermal differentiation the *Alb1* enhancer is bound by FoxA1 and FoxA2, eventually activating transcription of the *Alb1* gene in hepatocytes<sup>46</sup>. In the case of ESC to neural differentiation, Sox2 binds enhancers in ESCs that are later activated by Sox3 in neural progenitor cells. Besides that, Sox3 also preselects for enhancers in neural progenitor cells that are later activated by Sox11 in differentiating neurons<sup>47</sup>. Transcription factors such as these are able to recognize their DNA sequence motif in open chromatin and directly bind the DNA, often by interacting with and attracting other transcription factors, co-activators and chromatin associated complexes (figure 1).

One such co-activator that is attracted by transcription factors to active enhancers is the Mediator complex. The Mediator complex bridges regulatory signals of DNA binding transcription factors on enhancers to PolIII bound to the promoters as part of the pre initiation complex and as such affects the initiation, pausing and elongation of PolIII (reviewed in<sup>10</sup>) (figure 1). The binding of transcription factors to Mediator evokes conformational changes in the Mediator complex<sup>48</sup>, which facilitates interaction with PolIII<sup>49</sup> and triggers activation of stalled PolIII<sup>50</sup>. Mediator subunit CDK8 has been speculated to be part of Mediator after pausing of PolIII, attracting CDK9 as part of the super elongation complex, which in turn phosphorylates PolIII, promoting release of PolIII and transcription elongation<sup>10</sup>. Mediator also stabilizes promoter-enhancer loops by stimulating deposition of cohesin, which may form Cohesin ring-like structures between different DNA elements<sup>51</sup> like observed between sister chromatids<sup>52</sup>.

CTCF together with Cohesin is also required for promoter-enhancer looping<sup>53,54</sup>. CTCF binds the genome via its specific DNA binding motif<sup>55</sup>. The orientation of CTCF binding motifs on the genome determine the direction of chromatin looping<sup>56</sup>. Two opposite oriented CTCF binding motifs with CTCF bound in opposite topology are believed to establish chromatin loops that facilitate interaction between promoters and enhancers

via cohesin deposition<sup>56</sup>. Transcription itself is not required for the establishment of enhancer-promoter loops<sup>57</sup>.

P300 and CBP are also co-activators that are attracted by transcription factors to enhancers (figure 1), and over 400 known transcription factors interact physically and functionally with P300<sup>58,59</sup>. P300 and CBP catalyze acetylation of H3K27<sup>60,61</sup>. Like H3K27ac<sup>35,36</sup>, P300 binding is also used as a specific marker for active enhancers<sup>62</sup>. However, not all P300 binding sites are active enhancers and binding of P300 to enhancers does not always lead to acetylation of H3K27 as a subset of P300-bound poised enhancers lacks H3K27ac and/or are marked by H3K27me3<sup>37</sup>. This suggests that recruitment and the enzymatic activity of P300 are separately regulated<sup>43</sup>.

H3K27ac is recognized by bromodomain containing proteins. P300 and other histone acetyl transferases contain bromodomains and therefore amplify their own binding<sup>43</sup>. Also some subunits of the TFIID complex contain bromodomains and although these and other proteins are initially probably recruited through bromodomain-independent mechanisms, recognition of acetylated lysines may stabilize and amplify activating signals<sup>43</sup>. Also Brd4 and other members of the BRD protein family recognize H3K27ac. Brd4 has been shown to assist in transcription elongation with PolIII, but this is not dependent on the bromodomain<sup>63</sup>. A suggested role of histone acetylation is its potential to affect chromatin structure and interactions, because of its charge-nullifying ability and thus might regulate enhancer-promoter communication<sup>43</sup>.

### **Transcription factors cooperate in genome binding**

Transcription factors play an important role in establishing and activating enhancers (see above). It has been estimated that there are around 2600 DNA binding transcription factors encoded by the human genome<sup>64</sup>, of which 200 – 300 are expressed in each cell type<sup>65</sup>. Many of these transcription factors interact<sup>66,67</sup> and form cell- and tissue-specific transcription factor networks, in which tissue-specific transcription factors interact with ambiguously expressed transcription factors<sup>66</sup>.

DNA binding transcription factors often specifically bind to a specific sequence on the genome, also called the DNA binding motif. Depending on the expression of an interaction partner, a transcription factor can be targeted to different genomic targets. For example, in ESCs Oct4 (Pou5f1) and Sox2 interact and cooperatively bind to a DNA motif that consists of joint motif between a known POU binding and Sox binding motif<sup>68</sup> (see chapter 4), maintaining pluripotency of ESCs<sup>69</sup>. When Oct4 partners up with other transcription factors it leads to differentiation. Increased expression of Sox17, as seen in the inner cell mass during mesodermal differentiation, invokes Oct4 to switch interaction partners from Sox2 to Sox17 and thereby targets genes that trigger endodermal differentiation<sup>70</sup>. Expression of Otx2, as is observed when ESCs exit pluripotency, leads Oct4 to bind Otx2 and results in recruitment of Oct4 by Otx2 leading to the activation of enhancers associated with neural differentiation<sup>71</sup>. Also Sox2 is directed to different targets depending on its interaction partner. Besides that Sox2 interacts with Oct4 in ESCs (see above), it also interacts with Nanog in ESCs and this regulates ESC self-renewal<sup>72</sup>. During development, Sox2 forms a DNA binding complex with Pax6 that regulates initiation of lens development for the eye<sup>73</sup>. Sox2 also induces hair cell fate in

the cochlea with Eya1 and Six1<sup>74</sup> and is involved in many more interactions<sup>75</sup>.

As mentioned above, transcription factors recruit co-regulators and chromatin modifying proteins. The effect of binding of a transcription factor on chromatin or transcription is often determined by the other proteins it attracts. For example, Oct4 has been described as a transcriptional activator, with 89% of its target genes being down regulated upon knock-down<sup>76</sup>. Correspondingly, Oct4 interacts with, amongst others, the subunits of the activating BAF complex<sup>77</sup>. The interface that Oct4 most likely uses for interaction with BAF subunit Smarca4 was found to be essential for iPS reprogramming<sup>78</sup>. However Oct4 is also associated with silenced genes<sup>79</sup> and was found to interact with subunits of the Polycomb repressive complex1 (PRC1) and Nucleosome remodeling and deacetylase (NuRD)<sup>77,80</sup> and to recruit the H3K9 methyltransferase Eset to silence trophoblast-associated genes<sup>81-83</sup>.

### **Redundancy within and between enhancers**

When addressing the role of enhancers in disease, it is important to consider the redundancy in enhancers regulating the same gene(s). It has been proposed that besides the varying levels of redundancy observed between (homologous) genes, there is also varying redundancy between regulatory elements<sup>84</sup>. While some enhancers are essential, others display varying degrees of redundancy (reviewed in<sup>85</sup>). An early publication describes closely situated enhancer elements driving expression of *Sgs4* in *Drosophila*, where each combination of two elements was sufficient to direct expression, but none of the three could act alone<sup>86</sup>. However, apparently redundant enhancers are often well conserved and should be studied in the context of development<sup>85</sup>. For example, the *Svb* gene is essential for development in *Drosophila*. The *Svb* gene is regulated by a primary essential enhancer and two secondary enhancers. The secondary enhancers (also called shadow enhancers) play only a minimal role in optimal culturing conditions, but are essential when more environmental and genetic variability is introduced<sup>87</sup>. In a related study it was proposed that a gene is often regulated by multiple enhancers, because multiple enhancers have more chance to form successful promoter-enhancer interactions, resulting in a higher robustness of target gene expression<sup>85,88</sup>. Essential and redundant enhancers both exist, but it is difficult to estimate which of the two categories is more prominent, in part due to publication bias against negative results<sup>85</sup>. However, akin to redundancy between genes, varying degrees of redundancy between enhancers will determine whether point mutations result in abnormalities and disease.

### **Enhancer RNAs**

Another interesting feature of active enhancers is PolIII mediated bi-directional transcription at the enhancer site, resulting in the non-coding RNA class enhancer RNAs (eRNAs). Transcription of eRNAs was shown to correlate with transcription of nearby active genes<sup>89</sup>. Knock-down of the eRNA associated with P53 enhancers resulted in reduced expression of surrounding genes. This demonstrated that eRNA is more than just a by-product of transcriptional activation<sup>90</sup>. Targeted RNA interference using siRNA against several eRNAs suggested a functional role for eRNAs in transcription of target genes and promoter-enhancer looping<sup>91</sup>.

### Enhancers in the context of the three dimensional structure of the genome

The genome is organized into topologically associated domains (TADs), which are self-interacting regions defined by regulatory elements called insulators<sup>92</sup>. TADs in mouse and human have a median sizes of around 800 kb<sup>92</sup>. TADs are found to be highly conserved between different cell types and also between species<sup>5,92</sup>. Interaction between genomic regions (for example enhancer to promoter) are generally restricted to genomic regions within a TAD<sup>92,93</sup> and expression patterns of genes within TADs correlate<sup>93</sup>. TADs are defined by insulator elements<sup>92</sup> and these insulators limit interactions from one TAD to another. CTCF binds these insulator elements via its specific DNA binding motif<sup>65</sup> and is required for the enhancer blocking activity of insulators<sup>94</sup>. Global depletion of CTCF increases interactions between neighboring TADs, while reducing interactions within a TAD<sup>95</sup>.

Within a TAD, genomic distance is not a restriction for interaction. Enhancers can be as distant to their gene target as 1 Mb, exemplified by an enhancer of the Sonic Hedgehog gene<sup>96</sup>. It is therefore important to understand that the real physical distance that two regions have to bridge in order to interact is not directly represented by their linear genomic distance. Instead, linear genomic distance between promoter and enhancer should be perceived in the context of genomic organization, where a TAD organizes promoters and enhancers such that the physical distance between enhancer and promoter is decreased and interaction becomes possible. Distal elements often do not interact with the closest transcription start site<sup>97-99</sup> and are instead often located hundreds of kb away from their targets. The exact range in which enhancers act varies amongst different studies and is likely to be influenced by the different methods used and the resolution these methods offer. HiC found that only 25% of enhancer promoter pairs are within a 50 kb range and that 57% span 100 kb or larger<sup>28</sup>. Another HiC study found the median distance between interacting regions to be around 120 kb genomic distance<sup>98</sup>. A ChIA-PET study of PolII, for two cell lines found a median distance between promoter and enhancers of 57 kb and 37 kb respectively (calculated from Table S5 of <sup>99</sup>), with many promoter-enhancer interactions also spanning distances of 150 kb – 200 kb, but only a few surpassing several megabases<sup>99</sup>. ChIA-PET of enhancer-binding oestrogen-receptor-alpha found 86% of all duplex interactions (2 regions interact) to be spanning distances within 100 kb, with less than 1% bridging distances beyond 1Mb, while complex interactions (2+ regions interact) were often found to span more than 100 kb<sup>31</sup>.

The majority of promoters and cis-regulatory regions interact with multiple genomic regions<sup>98,99</sup>. The enhancers of the locus control region of the  $\beta$ -globin gene have been suggested to collaborate by aggregating into a single active chromatin hub<sup>100</sup>, where the enhancers from the locus control region form loops with the nearby globin genes<sup>101</sup>. Deletion of individual enhancer elements of the locus control region resulted in only modest reduction of  $\beta$ -globin gene expression, but deletion of the entire locus control region resulted in a dramatic reduced expression of the  $\beta$ -globin gene<sup>102</sup> (reviewed in<sup>100</sup>). Several enhancers can interact with a single promoter, but also promoters interact with promoters<sup>99</sup>. Genes corresponding to interacting promoters often have correlated expression<sup>99</sup> and might be part of chromatin hubs as described for the locus control region of the  $\beta$ -globin gene or transcription factories. However, a subclass of interacting promoters display enhancer-like chromatin modifications and act as enhancers driving

the expression of other promoters<sup>32,99</sup>.

### **The evolving enhancer**

Studies across different eukaryotic species revealed that enhancers underlie evolutionary changes between species<sup>103-106</sup> and that enhancers are, in contrast to promoters, not well conserved between species<sup>103</sup>. A possible explanation is that a gene often has one promoter, while its expression can be driven by various enhancers with different levels of redundancy (see above). I.e. mutations in enhancers have less chance of being deleterious than mutations in promoters, allowing for more evolutionary flexibility in enhancers.

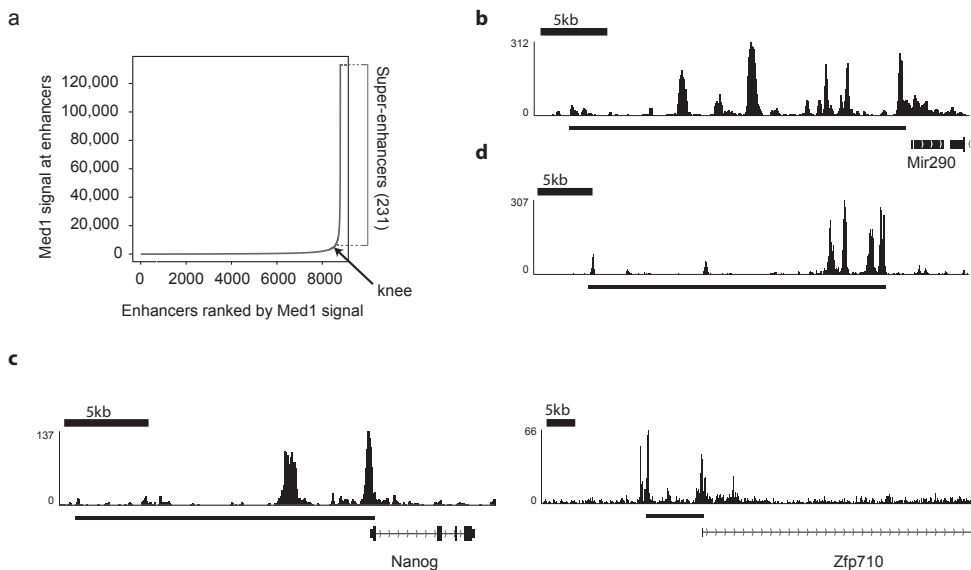
Enhancers involved in early embryogenesis and organ ontogenesis tend to be better conserved than enhancers involved in later tissue differentiation, which is reflected by the morphology of species during the different stages of embryonic development<sup>107,108</sup>. For example, conserved enhancers amongst species in liver were found near liver-specific genes<sup>103</sup>. A study across 20 mammalian species revealed that recently evolved enhancers dominate the regulatory landscape<sup>103</sup>. The majority of recently evolved enhancers derived from the adaptation of ancestral DNA<sup>103,109</sup>, frequently ancestral enhancers that are adapted to a new function<sup>109</sup>. Only a minority derived from expansion of repeat elements<sup>103</sup>.

A study on two enhancers regulating the Krüppel transcription factor in three different *Drosophila* species showed that, although total expression levels of Krüppel were conserved, the relative portion of total expression driven by either of the two enhancers varied across the species<sup>110</sup>. In other words, enhancer activity lost by one enhancer, is gained by the other enhancer. In fact, as evolutionary distance between species increases, such compensatory changes between enhancers are increasingly observed<sup>106</sup>, suggesting that this flexibility amongst enhancers leads to variation upon which evolution can act<sup>106,110</sup>.

An appealing example that illustrates recent adaptation of existing enhancers in human evolution, is an enhancer regulating Lactase-phlorizin hydrolase (*LPH*). *LPH* is for most humans specifically expressed during infancy and early childhood, allowing digestion of lactose from breast milk. Persistent expression of *LPH* (lactase persistence) during adulthood enables lifelong digestion of lactose. Although this adaptation convergently evolved for several heterogeneously distributed populations, the different 5 underlying SNPs found so far are all located in the same enhancer region 14 kb upstream of *LPH* and increase the enhancer activity of this region<sup>111-114</sup>. One of these SNPs, 13910T, completely associates with lactase persistence in the European population and creates a strong binding site for transcription factor Oct1 resulting in maintained *LPH* enhancer activity through adult life. In contrast, in the non-lactase persistence associated variant 13910C Oct1 only binds the enhancer efficiently when weaning age-specific *HNF1α* is also expressed<sup>115</sup>.

### **Super-enhancers**

A subset of enhancers was recently described as so called super-enhancers in a publication by Whyte et al.,<sup>6</sup>. Super-enhancers are domains that consist of clusters of



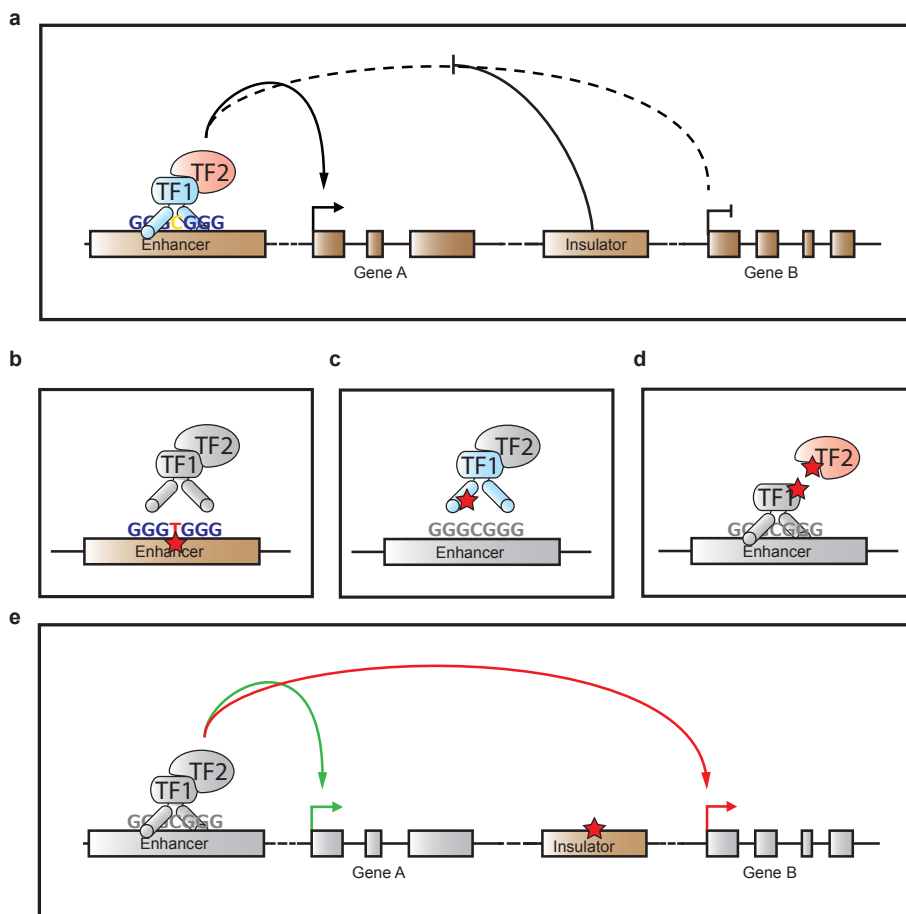
**Figure 2. Super enhancers.** (a) Distribution of Mediator ChIP-seq signal (total reads) across joint enhancers from Whyte et al.<sup>6</sup>. Figure adapted from Figure 1C in Whyte et al.<sup>6</sup>. Arrow indicates the knee in the curve, which was used as a cut-off between super-enhancers and regular enhancers. (b-d). Binding profile of the super-enhancer as determined in Whyte et al.<sup>6</sup>, y-axis displays sequence reads. Black bars indicate the region determined as super-enhancer by Whyte et al.<sup>6</sup>. (b) Binding profile of super-enhancer near *Mir290-295*. (c) Binding profile of super-enhancers near the *Nanog* and *Zfp710* genes, that display Med1 ChIP-seq signal on the promoter. (d) Binding profile of a super-enhancer downstream of *Sox2*, that contains large regions that display little ChIP-seq signal.

enhancers densely occupied by master regulator transcription factors and transcriptional co-activator Mediator and are often found near genes that define cell identity<sup>6</sup>. Super-enhancers are associated with genes that control cell identity<sup>6,116</sup> and super-enhancers at key oncogenes are acquired in cancer<sup>117</sup>.

To determine super-enhancers in a given cell type, Mediator1 (Med1) ChIP-seq signal or ChIP-seq signal of other enhancer-associated proteins and histone modifications was used to determine 'regular' enhancers. Regular enhancers within 12.5 kb were joined into a single region. These joint regions and non-joined enhancers were ranked for their cumulative ChIP-seq signal. All the enhancers beyond the knee in the curve were considered super-enhancers (figure 2a) and contain an exceptionally high density of Mediator. The 'knee of the curve' and the 12.5 kb cutoffs do not appear to be based on a biological rationale<sup>118</sup>. Citing the methods Whyte et al.<sup>6</sup>: "We first scaled the data such that the x and y axis were from 0-1. We then found the x axis point for which a line with a slope of 1 was tangent to the curve. We define enhancers above this point to be super-enhancers, and enhancers below that point to be typical enhancers". The knee of the curve as cut-off between regular and super-enhancers is sensitive to outliers. Mediator signal is also found on promoters and this is ignored by the original manuscript<sup>6</sup>, where in some cases super-enhancers also contained Mediator signal that is located on the promoter (Figure 2b). Super-enhancers often consists of several -often putative-enhancers that are separated by stretches of up to 12.5 kb that often contain very little ChIP-seq signal (Figure 2b,c).

### Diseases caused by ectopic enhancer binding

Variants that affect enhancers and are associated with disease are found on enhancers<sup>2,119</sup>, in transcription factor DNA binding domains<sup>119,120</sup> and protein-protein interaction domains<sup>120</sup>. Mutations that result in the disruption of the genomic environment of the enhancer have also been associated to genetic disease and resulted in different interactions of the enhancer with its targets (see below). Together these variants and mutations have the potential to affect normal enhancer function (figure 3). Considerations for each category are summarized in Table 2.



**Figure 3: Overview of mutations that affect gene regulation by enhancers and cause disease.** (a) Normal enhancer function. A transcription factor (TF1) binds by its DNA binding domain to a specific DNA motif on an enhancer and recruits another transcription factor (TF2). Together these transcription factors are required for enhancer-activated transcription of Gene A. The insulator element prevents contact between the enhancer and the promoter of gene B. (b-e) Mutations (red star) affecting enhancer function and causing misregulation of genes. (b) A mutation in the DNA binding motif in the enhancer element impedes binding of a transcription factor. (c) A mutation in the DNA binding domain of the transcription factor changes DNA binding specificity or abolishes DNA binding. (d) A mutation in the protein-protein interaction interface of the transcription factor impedes binding of other transcription factors. (e) Mutations that affect the position of an insulator element relative to the enhancer.

Category	effect	direct/indirect	affected genes
Enhancer element	ectopic binding of enhancer element by transcription factors	direct	locally
Genomic environment of enhancer	ectopic regulation of genes	indirect	locally
Transcription Factor: DNA binding domain	Loss of binding to enhancer elements by transcription factor	indirect	globally
Transcription Factor: Protein-protein interface	Loss of recruitment of co-activating transcription factors	indirect	globally

**Table2: Categories of mutations that affect enhancer function**

### Mutations in enhancers

Enhancer function can be disrupted by deletion of the enhancer, leading to disease. For example, a genomic deletion of the enhancer that regulates the Sclerostin gene was implicated in Van Buchem disease<sup>121</sup>. A deletion of a secondary enhancer to *ATOH7* was implicated in a local heritable form of blindness amongst Kurds living in Northern Iran<sup>122</sup>.

A wide range of DNA binding motifs facilitate binding of specific transcription factors to enhancers<sup>123</sup> (also see above) and enhancer function can be disrupted by point mutations in DNA binding motifs on enhancers. For example, a restless legs syndrome-associated SNP was demonstrated to reduce activity of the enhancers associated with the *MEIS1* locus. This SNP caused reduced binding of transcription factor CREB1 to the enhancer<sup>124</sup>.

Besides reducing binding of transcription factors to enhancers, mutations in enhancer can also lead to increased binding of transcription factors. A nucleotide duplication inside an E-Box motif was suggested to create a novel binding site for Lef1, underlying cleft lip and palate in a small Brazilian family<sup>125</sup>.

Variants associated with cancer risk have also been identified in enhancers regulating oncogenes. A study characterizing enhancers in colon cancer genome suggested that changes in the enhancers in colon cancer cells drives a specific transcriptional program that promotes colon carcinogenesis and found thousands of enhancer loci variants enriched for cancer genetic risk variants<sup>126</sup>. Individual variants in cancer have also been characterized. For example, a SNP located in the enhancer of *TOX3* is associated with breast cancer. This SNP causes increased binding of pioneer factor FOXA1 to the enhancer, resulting in the recruitment of the TLE1 repressor, diminishing enhancer activity and thereby decreasing expression of the *TOX3* gene<sup>127</sup>. In another locus a SNP associated with prostate cancer risk was also found to increase FOXA1 binding and to increase androgen responsiveness leading to increased enhancer activity<sup>128</sup>. Another cancer variant found is a colorectal cancer risk SNP in a TCF7L2 binding site on an enhancer required for tumorigenesis<sup>129</sup>. This SNP results in increased binding of the TCF7L2 transcription factor and increased luciferase activity of the enhancer interacting with the promoter of the proto-oncogene *MYC*<sup>130</sup>. Similar results with the same SNP were demonstrated in prostate cancer cells<sup>131</sup>. Also in colorectal cancer, a SNP in another enhancer of *MYC* impeded binding of the  $\beta$ -catenin transcription factor, resulting in increased expression of *MYC*<sup>132</sup>.

In summary, variation and mutations in enhancers have been reported in relation to

disease. Point mutations in DNA binding motifs can result in increase or decrease of transcription factor binding or enhancers can be deleted entirely, impeding enhancer function and causing misregulation of genes leading to disease. Enrichment of disease-associated SNPs in enhancers<sup>2</sup> suggests that enhancers are often the underlying cause of human disease.

#### Mutations in DNA binding domains of transcription factors

Mutations in DNA binding domains may influence binding of transcription factors to enhancers, impeding enhancer function. For example, two mutations in the DNA binding homeo-domain of LHX4 were found in patients with pituitary hormone deficiency<sup>133</sup>. Both mutations prevent LHX4 from binding enhancers and lead to loss of gene activation<sup>119,133</sup>. Several deletions and missense mutations in the DNA binding domain of the CBFA1 transcription factor abolished DNA binding and are associated with cleidocranial dysplasia which is characterized by skeletal abnormalities<sup>134,135</sup>. Mutations in the DNA binding homeo-domain of repressing transcription factor ARX are associated with various syndromes all characterized by intellectual disability. These mutations lead to loss of enhancer binding by ARX and loss of gene repression<sup>136</sup>.

Mutations in the DNA binding domain of transcription factors may not necessarily lead to complete loss of binding. For example, Klf1 is a hematopoietic C2H2 zinc finger transcription factor that plays a role in erythroid gene expression. The Nan-mutant mouse contains an amino acid substitution E339D in the second zinc finger of Klf1. While the wild type KLF1 binds to GC-box motif containing element GGGG[C/T]GGGG, the Nan-KLF1 mutant only binds GGGGCGGGG. This results in misregulation of Klf1 bound genes with GGGGTGGGG elements and severe hemolytic anemia in *Nan* heterozygous mutant mice<sup>137</sup>.

Gain of genomic binding was also suggested to be a disease causing mechanism associated with mutations in DNA binding domains<sup>119,120</sup>, but is supposedly rare<sup>120</sup>. In summary, mutations in the DNA binding domains of transcription factors can cause disease by abrogating genomic binding entirely or by causing binding to a different (sub) set of genomic targets (see also<sup>119</sup>). This may in turn lead to ectopic enhancer function resulting in misregulation of genes.

#### Mutations that result in disruption of the genomic environment of the enhancer

Mutations affecting the genomic environment surrounding enhancers can disrupt the activation of enhancer targets. Enhancers can be targeted to the wrong promoters when their TADs are disrupted. For example, families with rare limb malformations were shown to contain genomic rearrangements. These genomic rearrangements misplaced a cluster of enhancers relative to an insulator/boundary element of a TAD. This resulted in inappropriate interactions with promoters otherwise insulated from interaction with these enhancers, resulting in expression of genes involved in limb formation causing limb malformation<sup>138</sup>. Misplacement of enhancers has also been described to cause cancer. A frequently observed chromosomal rearrangement associated with acute myeloid leukemia causes a distal *GATA2* enhancer to ectopically activate the proto-oncogene *EVI1*<sup>139</sup>.

Mutations that introduce a novel regulatory element can disrupt enhancer regulation. A SNP between  $\alpha$ -globin genes and their upstream regulatory elements created a promoter-like element, which interfered with the normal activation of all downstream  $\alpha$ -globin genes, causing the genetic disease  $\alpha$ -thalassemia<sup>140</sup>.

Mutations affecting CTCF and Cohesin might also influence the enhancer landscape and its interaction. Mutations in several Cohesin subunits cause Cornelia de Lange syndrome and this disease has been suggested to be the direct result of enhancer-mediated processes<sup>141</sup>. Although disruption of individual CTCF binding sites in relation to disease remains to be reported, it is tempting to speculate that displacement of CTCF by a mutation in its binding site might locally lead to inappropriate gene activation by enhancers and disease. Inversion of CTCF binding sites was shown to influence the specificity of promoter-enhancer interaction<sup>56</sup> (see also above) and might one day also be discovered to underlie disease.

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# Chapter 2

## Proteins that bind regulatory regions identified by histone modification chromatin immunoprecipitations and mass spectrometry

Erik Engelen<sup>1\*</sup>, **Johannes H. Brandsma<sup>1\*</sup>**, Maaïke J. Moen<sup>1</sup>, Luca Signorile<sup>1</sup>, Dick H. W. Dekkers<sup>2</sup>, Jeroen Demmers<sup>2</sup>, Christel E.M. Kockx<sup>3</sup>, Zehila Ozgür<sup>3</sup>, Wilfred F. J. van IJcken<sup>3</sup>, Debbie L.C. van den Berg<sup>1,4</sup>, Raymond A. Poot<sup>1</sup>

<sup>1</sup> Department of Cell Biology, Erasmus MC, Wytemaweg 80, 3015 CN Rotterdam, The Netherlands

<sup>2</sup> Proteomics Center, Erasmus MC, Wytemaweg 80, 3015 CN, Rotterdam, The Netherlands

<sup>3</sup> Center for Biomics, Erasmus MC, Wytemaweg 80, 3015 CN, Rotterdam, The Netherlands

<sup>4</sup> The Francis Crick Institute, Mill Hill Laboratory, The Ridgeway, London NW7 1AA, United Kingdom

\* These authors contributed equally to this work

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## ABSTRACT

The locations of transcriptional enhancers and promoters were recently mapped in many mammalian cell types. Proteins that bind those regulatory regions can determine cell identity but have not been systematically identified. Here we purify native enhancers, promoters or heterochromatin from embryonic stem cells by chromatin immunoprecipitations (ChIP) for characteristic histone modifications and identify associated proteins using mass spectrometry (MS). 239 factors are identified and predicted to bind enhancers or promoters with different levels of activity, or heterochromatin. Published genome-wide data indicate a high accuracy of location prediction by ChIP-MS. A quarter of the identified factors are important for pluripotency and includes Oct4, Esrrb, Klf5, Mycn and Dppa2, factors that drive reprogramming to pluripotent stem cells. We determined the genome-wide binding sites of Dppa2 and find that Dppa2 operates outside the classical pluripotency network. Our ChIP-MS method provides a detailed read-out of the transcriptional landscape representative of the investigated cell type.

## INTRODUCTION

A mammalian genome supports the generation of the hundreds of different cell types in an organism. These cell types display distinct gene expression profiles as a direct consequence of differences in the activation state of their gene promoters and distal cis-regulatory elements called transcriptional enhancers. The ENCODE project has generated a wealth of data on the genome-wide chromatin landscape of many different mouse and human cell types<sup>1,2</sup>. In particular, the genome-wide identification of regulatory regions such as transcriptional enhancers and promoters and their state of activity has the potential to increase our understanding of how cell type identity is acquired and maintained. From reprogramming experiments it has become increasingly clear that the identity of cells is to a large extent determined by transcription factors, which bind enhancers and promoters<sup>3-6</sup>. It is therefore of interest to purify native transcriptional enhancers and promoters of a given cell type and identify the proteins that bind to these regulatory regions. Here we performed chromatin immunoprecipitations for histone modifications associated with promoters, enhancers or heterochromatin in mouse embryonic stem cells (ESCs) and identified the proteins present in the different precipitated fractions by mass spectrometry, a method that we named ChIP-MS.

Our ChIP-MS experiments identified 239 factors that we could predict to bind to promoters, enhancers or heterochromatin. Among these factors are subunits of several chromatin modifying complexes and proteins that play a role in different aspects of transcriptional regulation. We also find key ESC transcription factors such as Oct4, Esrrb, Dppa2 and Klf5 that are not only important for maintaining ESC self-renewal but also facilitate the reprogramming of somatic cells to induced pluripotent stem cells (iPSCs)<sup>4,7-9</sup>. Genome-wide data sets were available for 28 ChIP-MS-identified factors and correlated well with the ChIP-MS-based predictions for these factors, suggesting a high level of accuracy of location prediction by ChIP-MS.

For many of the detected factors, the genome-wide localization has not yet been

determined and our ChIP-MS results provide the first evidence of their binding preference for a particular type of regulatory DNA. To illustrate that ChIP-MS can identify factors with an interesting genome-wide location, we determined the genome-wide binding sites of pluripotency marker and reprogramming factor Dppa2. We show that Dppa2 is not part of the classical pluripotency transcriptional network and that Dppa2 target genes reach full activation only later in development.

## RESULTS

### ChIP-MS rationale and procedure.

Transcriptional enhancers and promoters can be recognized by the chemical modifications of their associated histones, especially histone H3. Promoters of transcribed genes were found to contain histone H3 tri-methylated at lysine 4 (H3K4me3)<sup>10,11</sup> and the level of their activity correlates with the level of H3K27 acetylation (H3K27ac) present<sup>12,13</sup>. Enhancers contain histone H3 mono-methylated at lysine 4 (H3K4me1)<sup>14</sup> and active enhancers can be recognized by the presence of the H3K27ac mark<sup>15,16</sup>. Inactive (hetero)chromatin is marked by H3K9me3<sup>17</sup>. The presence or absence of these and other chromatin marks was used to postulate fifteen different chromatin regions in the mammalian genome, including promoters and enhancers with different levels of activity<sup>13</sup>.

We anticipated that ChIP for H3K4me3 would precipitate active promoters and ChIP for H3K4me1 would precipitate enhancers. ChIP for H3K27ac would preferentially precipitate the most active promoters and enhancers, whereas ChIP for H3K9me3 would precipitate heterochromatin. Accordingly, we performed large scale ChIPs in biological duplicate for H3K4me3, H3K4me1, H3K27ac or H3K9me3, and for GFP as a control, in mouse embryonic stem cells (ESCs, Fig. 1a,b). Crosslinking of the chromatin was performed with Disuccinimidyl glutarate (DSG), a protein-protein crosslinker, followed by standard formaldehyde crosslinking, to increase the crosslinking efficiency of genome-bound factors to the chromatin<sup>18-20</sup>. ChIP wash steps were performed in low adherence tubes to increase protein yield and reduce background<sup>20,21</sup>. Bound protein factors were de-crosslinked and eluted by prolonged heating in protein denaturing conditions, separated on an SDS-polyacrylamide gel, tryptic peptides isolated and analyzed by mass spectrometry. A representative protein gel showed the (unresolved) histones precipitated with each histone modification antibody but not with the GFP control (Fig. 1c). Analysis by Western blot revealed that comparable amounts of chromatin were precipitated in the different histone-modification ChIPs, as indicated by the total content of histone H3 (Fig. 1d). ChIP against H3K4me1, H3K4me3 or H3K27ac precipitated chromatin with these respective histone modifications (Fig. 1d). Minor amounts of H3K4me1 were observed in the H3K4me3 ChIP and vice-versa. This is to be expected as H3K4me1 is present at low levels around active promoters<sup>15</sup>. H3K9me3 ChIP precipitated H3K9me3-marked chromatin but no significant amounts of the other histone modifications (Fig. 1d). We conclude that the histone modification ChIPs efficiently precipitated the intended chromatin fractions.

Subsequently, we tested whether our modified ChIP protocol, that we use for ChIP-MS, still precipitated the intended genomic regions, as compared to conventional ChIP. DNA

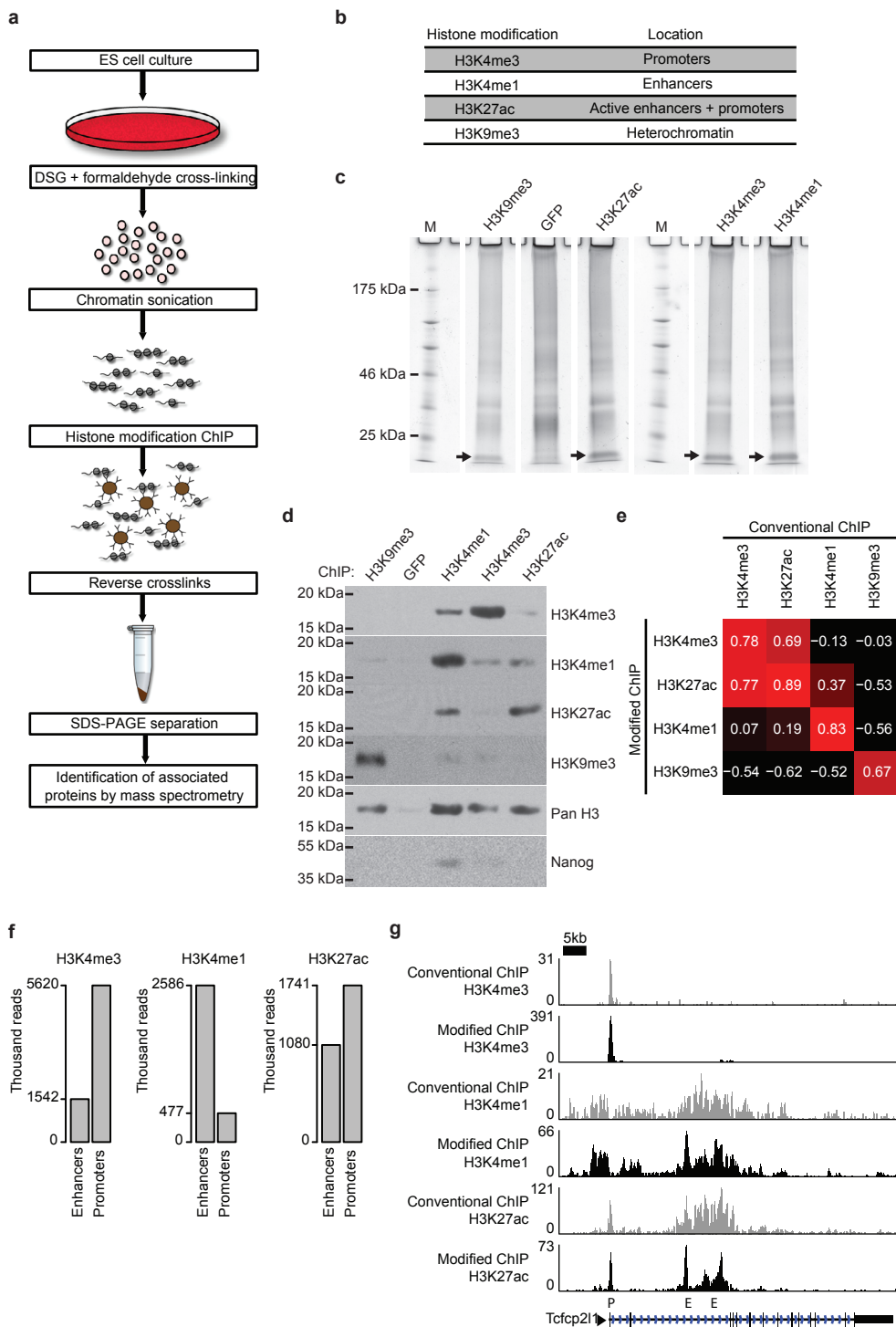


Figure 1. (Legend at the bottom of the next page)

precipitated by modified ChIP for the different histone modifications was sequenced (ChIP-seq) and mapped to the genome. DNA precipitated by modified ChIP correlated well with the corresponding published conventional ChIP-seq for all four histone modifications (Fig. 1e). Modified H3K4me3 ChIP predominantly precipitated promoters and modified H3K4me1 ChIP precipitated predominantly enhancers, as intended (Fig. 1f). Examples of histone modification tracks around pluripotency genes *Tcfcp2l1* (Fig. 1g) and *Nanog* (Supplementary Fig. 1) show high similarity between our modified ChIP and conventional ChIP. We conclude that the inclusion of additional crosslinker DSG has not significantly altered the genomic regions precipitated by our ChIP protocol, as compared to conventional ChIP.

### Prediction of genome localization of identified factors.

We analyzed the different precipitated chromatin fractions and GFP-control fractions by mass spectrometry for an unbiased identification of the protein factors present in each fraction. We identified 249 factors that have at least a 3-fold difference in Exponentially Modified Protein Abundance Index (emPAI) score, a measure for the amount of protein present<sup>22</sup>, in the ChIPs for one histone modification compared to the ChIPs for one or more of the other histone modifications. Included factors should have no or very low presence (more than five-fold lower emPAI score) in any of the GFP control ChIPs (Supplementary Tables 1 and 2, and Methods). These two selection steps were included to exclude proteins that bind to chromatin indiscriminately of the tested histone modifications, or are background of the ChIP-MS procedure, respectively. Of the 249 factors, 10 factors were only present in the H3K27ac fraction, which does not discriminate between promoters and enhancers, leaving 239 factors for which we could predict their binding to promoters, enhancers or heterochromatin. We assigned to identified factors the locations “promoter”, “enhancer” and “heterochromatin” according to the fraction (H3K4me3, H3K4me1 and H3K9me3, respectively) in which they have the highest emPAI value (Supplementary Tables 1, 2 and 3, and Methods). This annotation is not absolute, as factors can be present in more than one location, but it does provide clarity and facilitates a more systematic validation with published genome-wide localization data (see below).

We also indicated the presence of a factor in the H3K27Ac fractions by calculating the ratio of its average emPAI value in the H3K27ac fractions over its H3K4me3 emPAI score or its H3K4me1 emPAI score, whichever one is the highest, a ratio that we call the H3K27ac ratio. Presenting the ChIP-MS association of a factor with the H3K27ac modification in this way compensates for the considerable differences in ChIP-MS detection levels for

**Figure 1. Outline and initial validation of the ChIP-MS protocol.** (a) Flowchart of the ChIP-MS protocol. (b) Histone modifications used in ChIP-MS and their predominant location on the genome. (c) Representative 10% polyacrylamide gel with proteins from ChIPs for the indicated histone modifications and the GFP control ChIP. Arrows indicate unresolved histones in the histone modification ChIPs, which are absent in the GFP control ChIP. Molecular weight markers are depicted by M. (d) Western blot analyses of the histone modification content, histone content and the presence of Nanog in the immunoprecipitated chromatin fractions. Different ChIPs are indicated at the top, antibodies used for the different Western blot analyses are on the right. (e) Correlation between DNA regions precipitated by modified ChIP and conventional ChIP for H3K4me3, H3K27ac, H3K4me1 or H3K9me3. (f) Overlap of DNA precipitated with modified ChIP for H3K4me3, H3K4me1 or H3K27ac with promoters and enhancers. Number of ChIP-seq reads overlapping with promoters or enhancers is indicated. (g) ChIP-seq tracks for modified ChIP or conventional ChIP for H3K4me3, H3K4me1 or H3K27ac around pluripotency gene *Tcfcp2l1*. Sequence reads were plotted relative to chromosomal position. Genome location of *Tcfcp2l1* is shown, scale bar indicates 5 kb of genome. P indicates promoter, E indicates putative enhancer.

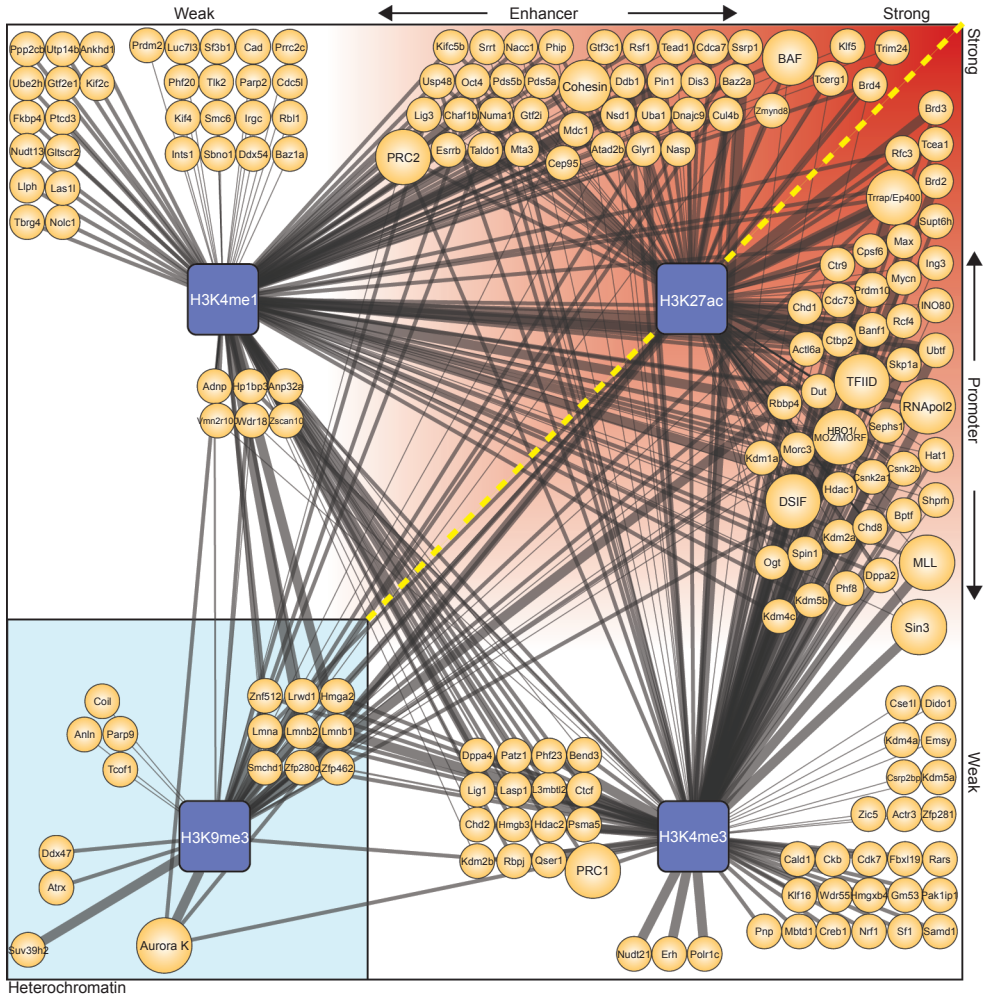
different proteins (Supplementary Table 1) and is therefore more informative than its H3K27ac emPAI value per se. Small emPAI values for H3K4me3 or H3K4me1 increase the uncertainty of the H3K27ac ratio value. We used the H3K27ac ratio as a predictor for the level of H3K27ac, and thereby the activity, of the promoters or enhancers bound by the factor (Supplementary Table 1).

For a visual representation (Fig. 2), identified factors and complexes were allocated according to their predicted binding to promoters, enhancers or heterochromatin. The calculated H3K27ac ratios were used to position predicted enhancer-binding factors on the upper horizontal axis reflecting the level of activity of bound enhancers or position predicted promoter binders on the right vertical axis reflecting the level of activity of bound promoters (Fig. 2).

An early indication that the predictions from our ChIP-MS experiments were valid came from the “promoter” prediction of all 5 identified RNApol2 subunits and 7 identified TFIID subunits (Supplementary Table 3). The H3K9 methyltransferase Suv3-9 binds pericentric heterochromatin<sup>23</sup> and was indeed observed solely in the H3K9me3 fraction (“heterochromatin” prediction, Fig. 2 and Supplementary Table 1). We identified a large number of subunits of established chromatin modifying complexes including the BAF complex, Sin3 complex and MLL complex (Fig. 2 and Supplementary Table 3). Strikingly, the localization prediction for different subunits within the same complex was nearly 100% identical (Supplementary Table 3), indicating a high level of consistency in the predictions.

Among the ChIP-MS identified factors with the highest H3K27ac ratio, predicting binding to highly active regulatory regions, were chromatin factors of the BET family; Brd2, 3 and 4 (Fig. 2 and Supplementary Table 1). BET family members were shown to bind to hyperacetylated chromatin<sup>24</sup>. Brd4 was recently identified as a key factor in the marking and functional maintenance of exceptionally large and active “super enhancers”, which regulate the expression of cell fate determining genes<sup>25,26</sup>. ChIP-MS classified Brd4 to bind predominantly to enhancers and Brd2 and 3 to bind promoters. Using ChIP-seq, Brd2 and Brd3 were indeed shown to bind promoters, whereas Brd4 was also present at enhancers<sup>27</sup>. The finding that from all ChIP-MS detected proteins, several BET family members are among the proteins with the highest H3K27ac ratio, validates the use of the H3K27ac ratio as an indicator of the level of activity of bound promoters or enhancers.

Many studies have identified factors that are important for maintaining ESC pluripotency<sup>28-30</sup> or factors that reprogram somatic cells towards ESC-like induced pluripotent cells (iPSCs)<sup>4,6,7</sup>. We found that more than a quarter of our ChIP-MS identified factors (63 out of 239) have a role in pluripotency acquisition or maintenance (Supplementary Table 1). ChIP-MS-identified factors included established reprogramming factors Oct4, Esrrb, Klf5 and Mycn (Fig. 2 and Supplementary Table 1), which as part of a 3-4 factor mix, reprogram somatic cells to iPSCs<sup>4,7,8</sup>. Our ChIP-MS data predicted that Oct4, Esrrb and Klf5 bind predominantly to enhancers and Mycn predominantly binds to promoters, in agreement with published genome localization data<sup>3,31</sup>. Nanog, another well-known pluripotency factor, is difficult to detect by mass spectrometry<sup>21,32</sup> and was indeed not identified by ChIP-MS. Western blot analysis of our ChIP-MS samples showed that



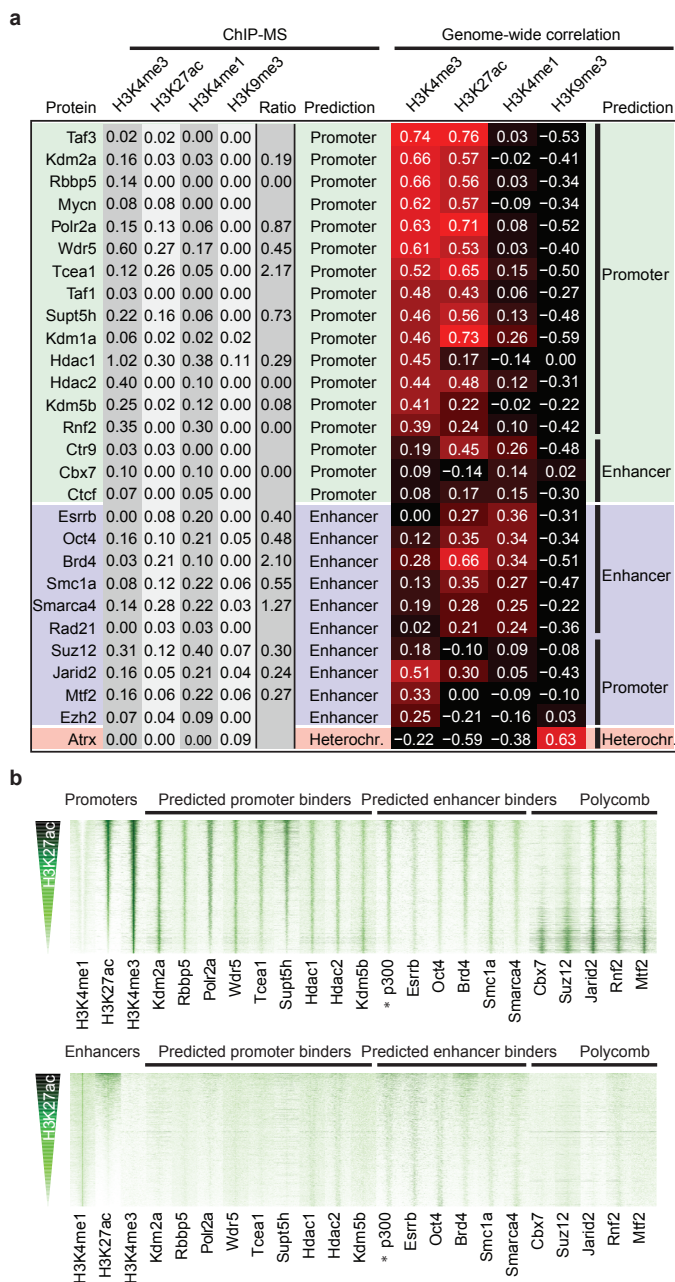
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**Figure 2. ChIP-MS predicted locations of identified factors and complexes.** Visual representation of factors (small orange circles) and complexes (large orange circles) identified by ChIP-MS for four different histone modifications (blue squares). Thickness of the edges indicates average emPAI score of a factor or complex in histone modification ChIP. Factors and complexes are positioned according to their ChIP-MS location prediction. To the left of the yellow dashed line are predicted enhancer binders, positioned horizontally from weak activity enhancers (left) to strong activity enhancers (right) according to their H3K27ac ratio. To the right of the yellow dashed line are predicted promoter binders positioned vertically from weak activity promoters (bottom) to strong activity promoters (top) according to their H3K27ac ratio. In the left bottom square are factors and complexes predicted to bind heterochromatin.

Nanog was present in the H3K4me1 fraction (Fig. 1d), suggesting it binds to enhancers, in agreement with published data<sup>3</sup>.

**Estimation of ChIP-MS prediction accuracy.**

Our list of 239 ChIP-MS assigned factors includes 28 factors for which the genome-wide binding sites have been determined in mouse ESCs by ChIP-seq (Fig. 3a), which provides an opportunity to probe the accuracy of our localization prediction. In a first



**Figure 3. Validation of ChIP-MS predictions with published genome-wide location information.** (a) Comparison of location prediction by ChIP-MS with location prediction by correlation of genome-wide binding sites with the indicated histone modifications on the genome. Protein factors for which genome-wide locations in mouse ESCs are determined by ChIP-seq are listed on the left, according to their ChIP-MS prediction as promoter binder (top, green panel), enhancer binder (middle, blue panel), or heterochromatin binder (bottom, red panel). Indicated in columns from left to right are: protein factor, its average emPAI values in the different histone modification ChIPs, its H3K27ac ratio (if highest

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analysis, we compared our ChIP-MS-based predictions (Fig. 3a) with location predictions derived from the correlation of factor binding sites with the different histone marks on the genome (Fig. 3a). Of the 17 factors predicted by ChIP-MS to be promoter-associated, 14 factors (82%) were indeed most associated on the genome with the active promoter mark H3K4me3 (Fig. 3a). In the case of Ctr9, Ctf and Cbx7, the ChIP-MS prediction was not conform the location prediction by genome-wide correlation. However, in these cases both the ChIP-MS values and genome-wide correlations for H3K4me3 and H3K4me1 (which differentiates between “promoter” and “enhancer” prediction) were very similar (Fig. 3a). Moreover, the correlation with any of the four tested histone marks was low for Ctf and Cbx7.

Of the 10 factors predicted by ChIP-MS to be predominantly associated with enhancers, 6 had indeed the highest association on the genome with enhancer mark H3K4me1 (Fig. 3a). These include Oct4 and Esrrb, two key pluripotency and reprogramming transcription factors, and Smarca4 (Brg1), the catalytic subunit of the SWI-SNF chromatin modifying complex (Fig. 3a). All the wrongly annotated factors (Suz12, Jarid2, Mtf2, Ezh2) are members of the Polycomb family of repressor proteins, which had “Enhancer” ChIP-MS predictions but were assigned “Promoter” by the genome-wide correlation of their binding sites (Fig. 3a). Polycomb factors are abundant in ESCs and often bind broadly at relatively inactive promoters with similarly low levels of H3K4me3 and H3K4me1 (see below)<sup>33,34</sup>. Both histone marks are indeed faithfully detected by ChIP-MS but in the above cases this leads to wrong predictions, albeit by small margins in the H3K4me3 and H3K4me1 ChIP-MS values (Fig. 3a). Atrx was correctly assigned by ChIP-MS to bind H3K9me3-containing heterochromatin (Fig. 3a). We conclude from this analysis that ChIP-MS predicted factor location to promoters, enhancers or heterochromatin with high accuracy, with the few false identifications in the expected grey areas.

Subsequently, we assessed whether the H3K27ac ratio of a factor correlates with the association of its binding sites with H3K27ac on the genome (Fig. 3a). From the factors with genome-wide location information (Fig. 3a), we took factors with a highest emPAI value of 0.1 or higher, to be well above the detection limit of our ChIP-MS experiments. We calculated for each of these factors their H3K27ac ratio (Fig. 3a). These ratios were compared to the correlation of genome-wide binding of these factors with H3K27ac-marked regions (Fig. 3a). Indeed, promoter-predicted factors Tcea1, Polr2a and Supt5h have the highest H3K27ac ratios and have relatively high correlations with H3K27ac on the genome (Fig. 3a). Promoter-predicted factors Hdac1 and 2, Kdm5b, Rnf2 and Cbx7 have low H3K27ac ratios and are factors with relatively low genome-wide associations

*(Figure 3. Legend continues from previous page)*

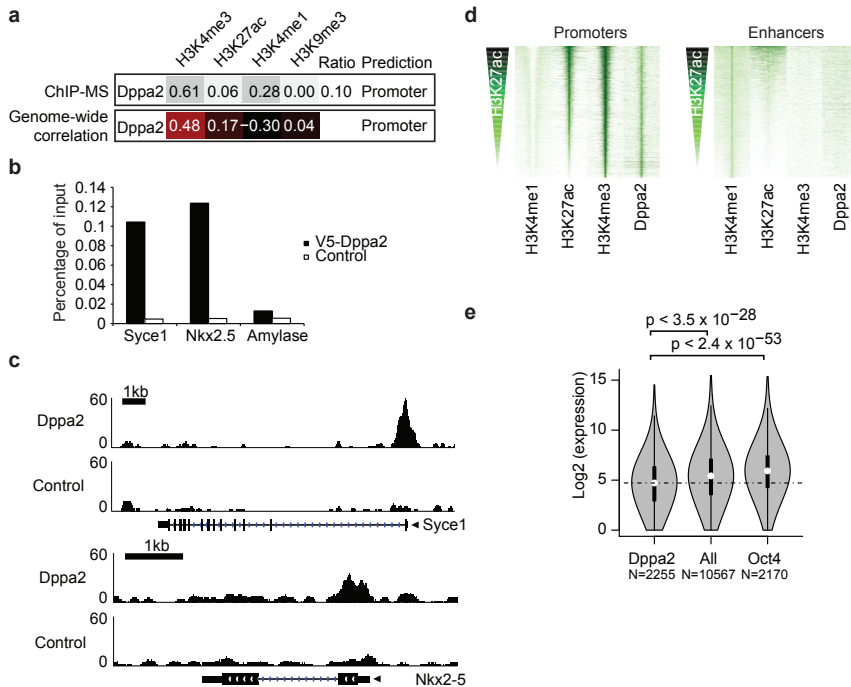
emPAI value  $\geq 0.1$ ), ChIP-MS location prediction, correlation of genome-wide binding sites with the indicated histone modifications and location prediction by highest correlation with a histone modification, according to Figure 1b. **(b)** Binding of selected protein factors to promoters and enhancers in mouse ESCs. Heatmaps of 12913 promoters (upper panel) or 30564 enhancers (lower panel), centered on H3K4me3 signal (Promoters) or H3K4me1 signal (Enhancers), ranked on H3K27ac content from top to bottom. Displayed is 8 kb around the center of the promoter or enhancer. Normalized ChIP-seq reads representing the level of H3K4me1, H3K27ac and H3K4me3 histone modifications are indicated in the first three lanes. Normalized ChIP-seq reads representing relative binding intensity to promoters (upper panel) and enhancers (lower panel) of protein factors from Figure 3a (highest emPAI value  $\geq 0.1$ ) are displayed in lanes 4-12 and 14-20. Factors are arranged according to binding prediction or Polycomb factor identity. \*p300 was not predicted by ChIP-MS but its genome-wide location was included in lane 13 for comparison.

with H3K27ac (Fig. 3a). Kdm2a and Rbbp5 have low H3K27ac ratios but still have high genome-associations with H3K27ac. Hence, in these two cases the H3K27ac ratios do not correlate well with H3K27ac association on the genome. Brd4 is the enhancer-predicted factor with the highest H3K27ac ratio and indeed has the highest correlation with H3K27ac on the genome (Fig. 3a). Oct4, Esrrb and Smc1a have intermediate H3K27ac ratios and intermediate genome-wide association with H3K27ac. Smarca4 has a high H3K27ac ratio, which in this case was not a good predictor, as Smarca4 has an intermediate level of genome-wide association with H3K27ac. We conclude from our above analyses that H3K27ac ratios provide a good, albeit not flawless, indication of the level of localization to H3K27ac-marked regions on the genome.

In a second approach to assess the accuracy of ChIP-MS, we investigated the presence of the above factors at promoters and enhancers in mouse ESCs. We assigned promoters as being present at the start of a gene and containing H3K4me3. Enhancers were assigned by their H3K4me1 content in the absence of H3K4me3. Promoters and enhancers were ranked top to bottom by their H3K27ac content (Fig. 3b). Promoter-predicted factors Kdm2a, Rbbp5, Polr2a, Wdr5, Tcea1, Supt5h and Kdm5b were indeed observed to only bind promoters and not enhancers (Fig. 3b). Hdac1 and Hdac2 predominantly bound to promoters but also showed binding to enhancers. Enhancer-predicted factors Esrrb, Oct4, Brd4 and Smc1a all showed strong binding to enhancers (Fig. 3b). These factors also showed binding to promoters to varying degrees. For comparison, we also included published genome-wide localization data of archetypal enhancer binder p300 in Figure 3b. Remarkably, p300 showed binding to enhancers and promoters (Fig. 3b), as previously observed in human ESCs<sup>33</sup>. Polycomb factors Rnf2, Jarid2, Suz12, Cbx7 and Mtf2 bind promoters with no correlation (Rnf2, Jarid2) or an anti-correlation (Cbx7, Suz12 and Mtf2) for H3K4me3 and H3K27ac content (Fig. 3b). The broad binding of Polycomb factors to promoters with relatively low H3K4me3 levels would explain their similar ChIP-MS values for H3K4me3 and H3K4me1 (Fig. 3a) and the associated ChIP-MS prediction uncertainties. The above analysis suggests that ChIP-MS-mediated prediction of binding to promoters or enhancers has high accuracy, with Polycomb factors again being an exception.

### **Dppa2 is not part of the classical ESC pluripotency network.**

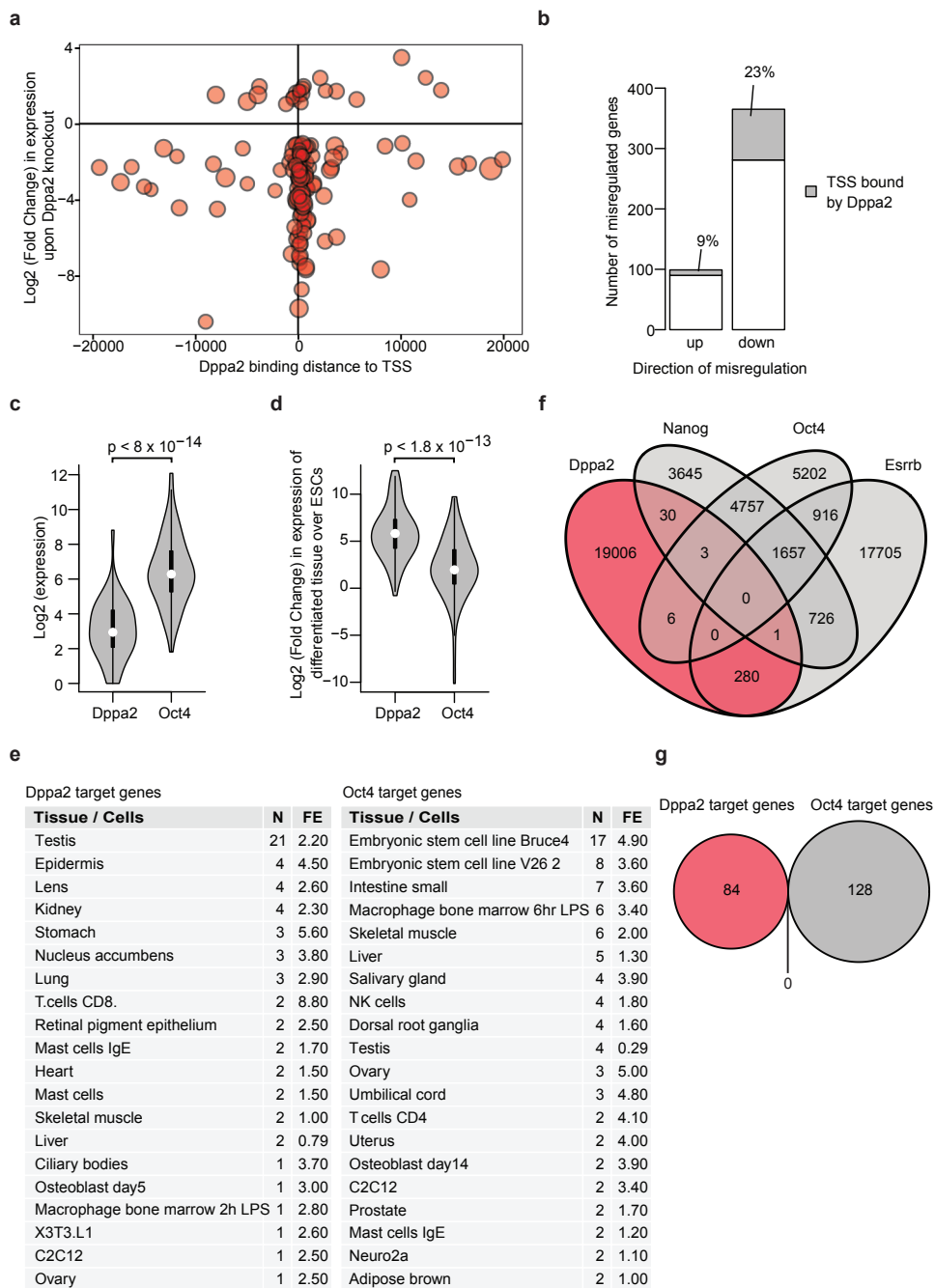
For many of the ChIP-MS detected factors the genome localization has not yet been determined by genome-wide ChIP and our experiments provide the first information on their genome binding preferences. As an example that ChIP-MS can identify factors with an unusual and therefore interesting genomic distribution, we focused on Dppa2 (Developmental PluriPotency Associated 2). Dppa2 is a member of family that also contains Dppa3 (Stella) and Dppa4, which all harbor a SAP DNA binding domain. The genome-wide binding sites of members of this family have not been determined so far. Dppa2 is exclusively expressed in the inner cell mass of the early embryo and later in the developing germ line and in cells derived from these tissues, such as embryonic stem cells and primordial germ cells<sup>35</sup>. Furthermore, Dppa2 expression was identified as an early marker for successful reprogramming towards iPSCs<sup>9</sup>. Dppa2 knockout ESCs have a slower proliferation rate and Dppa2 knockout mice die after birth from respiratory defects<sup>36</sup>. Recently it was shown that Dppa2, in combination with Lin28, Sall4 and Esrrb, drives the reprogramming of fibroblasts into iPSCs<sup>9</sup>. We selected Dppa2 because its



**Figure 4. Analyses of genome-wide binding sites of Dppa2.** (a) Comparison of location prediction for Dppa2 by ChIP-MS with location prediction by the correlation of identified Dppa2 genome-wide binding sites with the indicated histone modifications on the genome. Indicated in the upper panel from left to right are: Dppa2 average empAI values in the different histone modification ChIPs, its H3K27ac ratio and ChIP-MS location prediction. Indicated in the lower panel from left to right are: the correlation of Dppa2 genome-wide binding sites with the indicated histone modifications and location prediction by highest correlation with a histone modification, according to Figure 1b. (b) Binding of Dppa2 to the promoters of the indicated genes, detected by anti-V5 ChIP on V5-Dppa2 expressing ESCs or control ESCs. Precipitated DNA for the indicated genes is shown as percentage of input, the Amylase gene is used as a negative control region. (c) Localization of Dppa2 on the promoter of Syce1 (upper panel) or Nkx2-5 (lower panel). Sequence reads from anti-V5 ChIP-seq on V5-Dppa2 expressing ESCs (Dppa2) or control ESCs (Control) were plotted relative to chromosomal position. Genome locations of Syce1 gene (upper panel) and Nkx2-5 gene (lower panel) are shown, scale bars indicate 1 kb of genome. (d) Binding of Dppa2 to promoters and enhancers in mouse ESCs. Heatmaps of 12913 promoters (left panel) or 30564 enhancers (right panel), centered on H3K4me3 signal (Promoters) or H3K4me1 signal (Enhancers), ranked on H3K27ac content from top to bottom. Displayed is 8 kb around the center of the promoter or enhancer. Normalized ChIP-seq reads representing the level of H3K4me1, H3K27ac and H3K4me3 histone modifications are indicated in the first three lanes. Normalized V5-Dppa2 ChIP-seq reads representing the relative binding intensity of Dppa2 to promoters (left panel) and enhancers (right panel) are displayed in the fourth lane of each panel. (e) Distribution of absolute expression levels of H3K4me3 marked genes in mouse ESCs that (from left to right) are bound at the promoter by Dppa2, all genes, and bound within 20kb around the promoter by Oct4. Shown is a violin plot where the white dot indicates the median and the thick black bar indicates 50% of the genes. Log2 value of the absolute expression, derived from published RNAseq data, the number of genes in each category and p-values by Mann-Whitney test are indicated.

ChIP-MS profile was unusual for a pluripotency-inducing factor. Dppa2 had the highest empAI score for H3K4me3 but a low H3K27ac ratio, suggesting it binds predominantly to promoters with low activity (Fig. 4a). This is different for other pluripotency-inducing factors, such as Oct4 and Esrrb, which predominantly bind moderately active enhancers (Fig. 3b). To identify the genome-wide binding sites of Dppa2, we established an ESC line that expressed V5-tagged Dppa2 and we performed anti-V5 Dppa2 ChIP and





**Figure 5. Dppa2 target genes and their overlap with the pluripotency network.** (a) Bubble plot indicating the positions of Dppa2 binding sites relative to the transcription start site (TSS) on genes that are either up-regulated  $\geq 2$  fold (upper part) or down-regulated  $\geq 2$  fold (lower part) upon Dppa2 gene knockout in mouse ESCs. Log<sub>2</sub> of the fold change in expression upon Dppa2 KO is indicated on the Y-axis. Distance of Dppa2 binding sites from the transcription start site is indicated on the X-axis. Size of

*Legend continues on the bottom of the next page*

sequenced the precipitated genomic DNA (ChIP-seq). We verified that V5-Dppa2 binds to the promoters of *Syce1* and *Nkx2-5* (Fig. 4b,c), the only known Dppa2 genomic binding sites<sup>36</sup>, which suggested that our V5-Dppa2 ChIP identified bona-fide Dppa2 binding sites. Dppa2 binding sites had the highest genome-wide association with H3K4me3, whereas the association with H3K27ac (Fig. 4a) is lower than that of Oct4 and Esrrb (Fig. 3a). Subsequently, we investigated the presence of Dppa2 at promoters and enhancers in mouse ESCs. We found that Dppa2 binds promoters but is absent from enhancer regions (Fig. 4d). Interestingly, Dppa2 promoter binding displayed no correlation with H3K27ac content (Fig. 4d), a binding pattern that otherwise was only observed with the repressor Kdm5b and the Polycomb repressor proteins (Fig. 3b). These analyses suggest that our Dppa2 ChIP-MS location prediction was correct and that the binding pattern of Dppa2 is different compared to other reprogramming factors.

As expected, Dppa2 binds the promoters of genes with a lower median expression than other H3K4me3-marked promoters and Oct4-bound genes in ESCs (Fig. 4e and Supplementary Table 4). The gene expression profile of Dppa2 knockout ESCs was recently determined and it was observed that far more genes were down-regulated than up-regulated, compared to wild-type ESCs<sup>36</sup>. Dppa2 binds the promoters of nearly a quarter of the down-regulated genes but much less to promoters of the up-regulated genes (Fig. 5a,b). This suggests that Dppa2 maintains the expression of its putative target genes (Supplementary Table 5) by binding at their promoter. In contrast, Oct4 maintains the expression of its target genes by binding mostly outside promoters (Supplementary Fig. 2). We find that the median expression of Dppa2 target genes is 10-fold lower than the median expression of Oct4 target genes in ESCs (Fig. 5c). Nearly all Dppa2 target genes are higher expressed in tissues other than ESCs (Fig. 5d and Supplementary Table 5). The largest minority of Dppa2 target genes is highest expressed in testes, but many Dppa2 target genes have their highest expression in other tissues (Fig. 5e and Supplementary Table 5). For Oct4, the pattern is very different, as many Oct4 target genes have their highest expression in ESCs (Fig. 5d,e). Considering that Dppa2 and Oct4 appear to regulate different sets of genes, we determined the overlap in genome-wide binding sites between classical pluripotency factors such as Oct4, Nanog

(Figure 5. Legend continues from previous page)

the bubbles correlates with fold difference of Dppa2 ChIP peak over control. **(b)** Bar diagram showing the total number of up-regulated and down-regulated genes upon Dppa2 knockout and the number of these genes bound by Dppa2 within 1 kb from the TSS (grey areas). The number of Dppa2 bound genes is also indicated as a percentage of the total number of up-regulated or down-regulated genes. **(c)** Distribution of absolute expression levels in mouse ESCs of Dppa2 target genes and Oct4 target genes. Dppa2 target genes are bound by Dppa2 within 1 kb of the TSS and  $\geq 2$ -fold down-regulated upon Dppa2 knockout, Oct4 target genes are bound by Oct4 within 20 kb of the TSS and  $\geq 2$  fold down-regulated after 24 hrs of Oct4 depletion. Shown is a violin plot where the white dot indicates the median and the thick black bar indicates 50% of the genes. Log2 of the absolute expression, derived from published RNAseq data and p-value by Mann-Whitney test are indicated. **(d)** Distribution of the fold change in expression of Dppa2 target genes and Oct4 target genes in the differentiated tissue with the highest expression versus expression in mouse ESCs. Shown is a violin plot where the white dot indicates the median and the thick black bar indicates 50% of the genes. Log2 of the fold change in expression, derived from published RNAseq data and p-value by Mann-Whitney test are indicated. **(e)** Lists of tissues or cells where Dppa2 target genes (left panel) or Oct4 target genes (right panel) are highest expressed. Tissue or cells, the number of genes (N) and fold enrichment (FE) of a tissue/ cell type within Dppa2 target genes or Oct4 target genes are indicated. The 20 tissues/ cells with the highest number of gene overlap and fold enrichment are shown. **(f)** Venn diagram showing the overlap of genomic binding sites in mouse ESCs of Dppa2, Nanog, Oct4 and Esrrb. **(g)** Venn diagram showing the lack of overlap of Dppa2 target genes and Oct4 target genes.

and Esrrb, and Dppa2. Whereas Oct4, Nanog and Esrrb showed an extensive overlap in binding sites, as previously reported<sup>3</sup>, Dppa2 showed little overlap with these factors (Fig. 5f). In particular, the overlap of Dppa2 with Oct4 and Nanog was nearly absent. Moreover, there was no overlap between Dppa2 target genes and Oct4 target genes (Fig. 5g). We conclude that Dppa2 regulates a set of genes that is separate from the set of genes regulated by the classical pluripotency transcription factors, and not ESC-specific in its expression.

## DISCUSSION

We describe here CHIP-MS, a method to predict the binding of factors to enhancers or promoters. Our experimental set-up is straightforward and does not rely on metabolic labeling for quantification by mass spectrometry. Nevertheless, using emPAI<sup>22</sup> and the simple rule that the CHIP fraction in which a factor has the highest emPAI score decides its binding prediction, we achieved a remarkably high percentage of correct predictions when comparing to published CHIP-seq data. We employed crosslinking with protein-protein crosslinker DSG, followed by standard crosslinking with formaldehyde. We previously used this “double crosslinking” procedure to improve CHIP efficiency<sup>19-21</sup>. Aside from our extended crosslinking procedure, our CHIP-MS procedure is based on standard CHIP protocols and mass spectrometry procedures, which should facilitate its application in other cell types.

A number of studies have been performed to screen for protein factors that bind to individual histone modifications by using modified histone peptides<sup>37-39</sup> or *in vitro* assembled modified nucleosomes<sup>40</sup>, or identify protein factors that bind native chromatin harboring specific histone modifications by conventional ChIP combined with mass spectroscopy<sup>41</sup>. Binding to different tri-methylated lysines was assessed in these studies, but binding to enhancer marks or activity marks, such as the H3K4me1- or H3K27ac-marked chromatin that we interrogated with CHIP-MS, has not been addressed yet. Previous studies identified a number of ubiquitously expressed factors that we also observed in our ChIPs. However, our CHIP-MS procedure is sufficiently sensitive to also detect the sequence-specific transcription factors, such as Oct4, Esrrb, and Klf5, that determine ESC identity. These factors will be different in other cell types, which makes our procedure highly suitable to study the changing spectrum of regulatory region-associated factors during cell differentiation.

The list of proteins for which we predict genome localization by CHIP-MS (Fig. 2 and Supplementary Table 1) contains factors that play a role in a number of cellular processes, including all levels of transcriptional regulation and chromatin organization. CHIP-MS detected chromatin modifying complexes with clear location predictions, such as the BAF complex (enhancer), Aurora kinase complex (heterochromatin), Trrap complex (promoter), MLL complex (promoter) and Sin3 complex (promoter). The annotation of subunits of the Polycomb complexes PRC1 and PRC2 was more ambiguous, as PRC1 and PRC2 bind broad areas around inactive promoters marked to a similar extent by H3K4me3 and H3K4me1, leading to false “enhancer” predictions for several PRC subunits. Fortunately, Polycomb factors are well characterized<sup>34,42</sup> and would therefore be easy to recognize and treated with caution in any CHIP-MS prediction list.

Figure 2 ranks detected factors by the activity of their bound promoters or enhancers. To our knowledge, such an analysis has not been performed yet and provides a read-out on an important criterion for a large set of factors in ESCs. The ranking is based on the H3K27ac ratio; the ratio of the H3K27ac empAI score over the highest of the H3K4me3 empAI score or H3K4me1 empAI score, which we anticipated would be the more informative value than the H3K27 empAI score per se, as it compensates for the considerable differences in ChIP-MS detection levels for different factors. Perhaps the clearest indication that the H3K27ac ratio performs well in differentiating factors by the activity of their bound regions is that out of the nearly 240 factors detected by ChIP-MS, 3 members of the family of BET proteins, including Brd4, are among the proteins with the highest H3K27ac ratios. Brd4 was recently identified as a functional component and marker of “super enhancers”<sup>25,26</sup>, arguably the most active enhancers in the genome of a cell. The ranking of the chromatin modifying complexes follows common sense. The activating BAF chromatin remodeling complex and Trrap histone acetylase complex have higher H3K27ac ratios than the Sin3 repression complex and the PRC1 and PRC2 complexes. The good performance of ChIP-MS on factors with a known genome-wide location suggests that also the localization predicted for the many factors without genome-wide ChIP data will in most cases be accurate. Our data set, graphically represented in Figure 2, therefore provides valuable new information on the transcriptional network in ESCs. Importantly, ChIP-MS can detect factors with an unusual, and therefore interesting, genome localization that can then be further investigated, Dppa2 being an example.

We performed our ChIP-MS experiments in ESCs. The establishment and maintenance of pluripotency, as well as the exit from pluripotency, is intensely studied in ESCs and several large data sets of relevant factors for the above processes are available. We find that a quarter (63 factors) of the ChIP-MS detected factors contributes to maintaining pluripotency. ChIP-MS detected Oct4, Esrrb, Klf5, Dppa2 and Mycn, factors which, as part of a 3-4 factor mix, reprogram somatic cells to iPSCs<sup>4,7,8</sup>. Intriguingly, ChIP-MS predicts that these factors do not all bind to the same type of locations on the genome. Oct4 and Esrrb were predicted to bind moderately active enhancers, whereas Klf5 binds to highly active enhancers. Mycn and Dppa2 were predicted to bind to high and low activity promoters, respectively. This suggests a division of labor between the different factors in the reprogramming process.

We found that Dppa2 had a different ChIP-MS profile compared to other pluripotency factors and accordingly we determined its genome-wide binding sites by ChIP-seq. Indeed, Dppa2 turned out to be an unusual pluripotency factor. Dppa2 binding sites and target genes do not overlap with Oct4, suggesting that Dppa2 is not part of the classical pluripotency circuit. Furthermore, Dppa2 target genes were found to be much lower expressed than Oct4 target genes in ESCs and higher expressed later in development. It is an intriguing question how Dppa2 can be an early marker and factor for reprogramming to iPSCs, as was recently shown<sup>9</sup>, without actually regulating ESC-specific genes. Dppa2 was proposed as a factor that binds target genes to maintain an active chromatin structure and facilitate their later expression<sup>36</sup>. This epigenetic marking hypothesis is consistent with the expression pattern of the identified Dppa2 target genes. However, we did not find that genes bound by Dppa2 in ESCs were preferentially down-regulated in

Dppa2 knockout lungs, using a published gene expression set<sup>36</sup>.

In conclusion, we established here a method to annotate factors to enhancers and promoters with different activities. ChIP-MS is straightforward in its set-up, which should facilitate its application to other cell types and growth conditions, provided sufficient cell quantities can be obtained. We showed that ChIP-MS data add to our knowledge and understanding of the transcriptional circuitry that determines cell identity.

## METHODS

### Cell Lines and constructs.

Mouse embryonic stem cell line CGR8 was grown on gelatin-coated dishes without feeders in GMEM (Glasgow Minimum Essential Medium) supplemented with LIF (leukemia inhibitory factor), 15% FBS, 0.25% sodium bicarbonate, 1mM glutamine, 1mM sodium pyruvate, non-essential aminoacids, 50µM β-mercaptoethanol and penicillin/streptomycin, as previously described<sup>19</sup>. The coding sequence for Dppa2 was amplified from mouse ES cell cDNA and cloned with an N-terminal V5-tag into a pPyCAG driven expression vector. CGR8 cells were transfected with the V5-Dppa2 expression vector using Lipofectamine 2000 (Invitrogen), clones were selected with 1µg/ml puromycin (Sigma) and stable expression of V5-tagged Dppa2 tested by Western blot analysis with anti-V5 antibody (1:2000) (Invitrogen).

### ChIP-MS procedure.

For each histone modification ChIP, 300x10<sup>6</sup> ESCs were used. For chromatin preparation, cells were washed on plate three times with PBS and incubated with 2 mM Disuccinimidyl glutarate (DSG, Thermo Scientific) in PBS for 45 min at room temperature. Subsequently, ESCs were washed in PBS three times, 0.1 volume of 11% formaldehyde (Merck) in 50 mM HEPES-KOH pH 7.5, 100 mM NaCl, 1 mM EDTA, 0.5 mM EGTA was added, mixed and incubated for 12 min at room temperature, washed two times in 4 °C PBS and collected by centrifugation. All subsequent steps were performed on ice with pre-cooled buffers. Cell lysis was performed as described<sup>43</sup>. In brief, cells were collected and resuspended in LB1 (50 mM HEPES-KOH, pH 7.5, 140 mM NaCl, 1 mM EDTA, 10% glycerol, 0.5% NP-40, 0.25% Triton X-100). After 10 minutes incubation, cells were pelleted by centrifugation and resuspended in LB2 (10 mM Tris-HCl, pH 8.0, 200mM NaCl, 1mM EDTA, 0.5mM EGTA). After 10 min incubation, cells were pelleted and resuspended in 3 ml freshly prepared LB3 (10 mM Tris-HCl, pH 8.0, 100 mM NaCl, 1 mM EDTA, 0.5 mM EGTA, 0.1% Na-deoxycholate, 0.5% N-lauroylsarcosine) and sonicated on a Soniprep 150 (MSE), 27 cycles 15 s on, 45 s off on amplitude 7. After sonication, enriched DNA fragment size was confirmed to be between 200 and 1000 bp. 300 x 10<sup>6</sup> ESCs yielded approximately 10 mg chromatin (as measured by DNA content).

Antibodies used in the different histone modification ChIPs or GFP control ChIP are against H3K4me1 (ab8895, Abcam), H3K4me3 (ab8580, Abcam), H3K27Ac (ab4729, Abcam), H3K9me3 (ab8898, Abcam) and GFP (sc8334, Santa Cruz Biotechnology). To prevent immunoglobulin elution and subsequent interference with the mass spectrometry analysis, 50 µg antibodies were crosslinked to 500 µl Protein A magnetic bead solution (15 mg beads, Life Technologies) with Dimethyl Pimelimidate (Sigma). Cross-linked

antibody-bead complexes were equilibrated in LB3 buffer and subsequently blocked with 0.5 mg/ml BSA (New England Biolabs) and 0.2 mg/ml sonicated salmon sperm DNA (Stratagene) for one hour. The antibody-bead mixture was rotated overnight with approximately 10 mg chromatin at 4°C. Beads were transferred to 1.5 ml no stick tubes (Alpha laboratories) and washed five times 5 minutes in RIPA buffer (50 mM Hepes-KOH pH 7.6, 500 mM LiCl, 1mM EDTA, 1% NP-40, 0.7% Na-deoxycholate). After washing, the beads were boiled for 35 min at 95 °C in 2x SDS sample buffer (100 mM Tris-HCl pH 6.8, 200 mM DTT, 4% SDS, 20% Glycerol, 0.2% Bromophenol blue) and supernatant was transferred to a fresh tube. ChIP-MS samples were run on 10% precast SDS-PAGE gels (NuPage Invitrogen) and stained with colloidal coomassie stain (Invitrogen). Gel lanes were sliced, in-gel digested with trypsin to yield peptides and proteins identified by analyses on an LQT-Orbitrap mass spectrometer (Thermo), as described<sup>21</sup>. For Western blot analyses, ChIP samples were separated on a 4-12% polyacrylamide gel (Novex) and nitrocellulose blots probed with antibodies against the used histone modifications (see above, 1:500 dilution, pan histone H3 antibody (Abcam 1791, 1:1000 dilution) and Nanog (Cosmo Bio Ltd., 1:2000 dilution). Non-cropped versions of the Western blot panels in Figure 1d can be found in Supplementary Fig. 3.

#### ChIP-MS inclusion and prediction criteria.

Two independent ChIPs were performed for each tested histone modification and for GFP, as control ChIPs, and analyzed by mass spectrometry. For inclusion into the ChIP-MS list of identified proteins (Supplementary Tables 1 and 2), factors needed to be identified by mass spectrometry with a Mascot score of 50 or higher in at least one histone modification ChIP. A Mascot score of at least 45 in any of the other ChIPs was annotated in the ChIP-MS list. In case of Mascot scores between 45 and 60, individual peptide MS/MS spectra were checked manually and interpreted as valid identifications or discarded. In addition, the Mascot program was used to determine the Mascot peptide significance threshold ( $P < 0.05$ ) in the ChIP-MS samples. Significance thresholds were H3K4me3 (Experiment 1); Mascot score 28, H3K4me3 (Experiment 2); Mascot score 28, H3K4me1 (Experiment 1); Mascot score 28, H3K4me1 (Experiment 2); Mascot score 28, H3K27ac (Experiment 1); Mascot score 28, H3K27ac (Experiment 2); Mascot score 28, H3K9me3 (Experiment 1); Mascot score 28, H3K9me3 (Experiment 2); Mascot score 29, GFP (Experiment 1); Mascot score 28, GFP (Experiment 2); Mascot score 28. For inclusion into the ChIP-MS list (Supplementary Tables 1 and 2), protein identifications had to be based on peptides with a Mascot score at or above the Mascot peptide significance threshold of the sample in which the peptides were observed. For assessment of the quantity of the identified proteins in the ChIP-MS samples we used emPAI a calculation method based on the number of peptide spectra identified by MS, normalized for the number of peptides that theoretically should be identifiable for that protein<sup>22</sup> (to compensate for large proteins, which likely have more MS peptides). Inclusion into the ChIP-MS list required an at least 5-fold higher average emPAI score in the ChIPs for at least one histone modification compared to any of the anti-GFP control ChIPs. Inclusion into the ChIP-MS list further required a factor to have an at least 3-fold higher average emPAI score in the two ChIPs for one histone modification compared to the two ChIPs for one or more of the other histone modifications, to exclude factors that bind chromatin indiscriminately of the tested histone modifications. Cytoskeletal and cytoplasmic proteins were excluded. Average emPAI scores were calculated from the two independent ChIP-MS experiments.

Localisation prediction was according to the following criteria; highest average emPAI score in H3K4me3 ChIP samples gives “promoter” prediction, highest average emPAI score in the H3K4me1 ChIP samples gives “enhancer” prediction, highest average emPAI score in the H3K9me3 ChIP samples gives “heterochromatin” prediction. In case average emPAI scores for the H3K4me3 and H3K4me1 ChIP samples were equal, the prediction was “promoter”. The H3K27ac ratio of a factor is defined as the ratio of its average H3K27ac emPAI score over the average H3K4me3 emPAI score or average H3K4me1 emPAI score, whichever one is the highest.

### **Chromatin immunoprecipitation and sequencing.**

Anti-V5 ChIPs were performed as described<sup>21</sup>. For V5-Dppa2 ChIP, CGR8 ESCs stably expressing V5-Dppa2 (see above) were used, for the control ChIP, the parental CGR8 parental ESC line was used. For each ChIP,  $100 \times 10^6$  ESCs were used. Precipitated DNA was analyzed by quantitative PCR or used for library generation followed by next generation sequencing on an Illumina Genome analyzer, as described<sup>20</sup>.

### **Data Analysis.**

Sequences with low complexity that are unlikely to map uniquely to the genome were removed from the Dppa2 ChIP-seq, modified ChIP-seq experiments for the used histone modifications and published ChIP-seq datasets (Supplementary Table 6), using prinseq-lite with the dust method with 7 as threshold<sup>44</sup>. The remaining sequences with a Phred score  $< 70$  were mapped to the mm9 reference genome using Bowtie<sup>45</sup> v0.12.7, where we used a seed length of 36 in which we allowed a maximum of 2 mismatches. If a read had multiple alignments only the best matching read was reported. Duplicated reads were removed. MACS<sup>46</sup> v1.4.2 was used for peak calling of Esrrb, Nanog, Oct4, Polr2a, P300, H3K4me1 and H3K4me3 using default settings. ChIP-seq datasets with multiple replicates were merged. For peak calling the Polr2a, P300, H3K4me1, H3K4me3 ChIP-seq datasets, the sequenced input was used as control. For peak calling Esrrb, Nanog and Oct4, the GFP ChIP was used as a control (Supplementary Table 6). For Dppa2, peak calling, we provided MACS1.4.2 with a shift size of 75 base pairs. Peaks with a p-value  $\leq 1 \times 10^{-10}$  were retained for Dppa2, Oct4, Nanog, Polr2a, H3K4me3, H3K4me1 and sites with a p-value  $\leq 2 \times 10^{-10}$  for Esrrb and P300. For Dppa2, only peaks with at least 100 aligned reads were retained. Dppa2, Oct4, Nanog and Esrrb peaks were considered to overlap if their peak summit was within 125 base pairs of each other. Venn diagrams, violin plots and bubble plots were created in R using the VennDiagram, vioplot and ggplots2 packages, respectively. To calculate the overlap between the mapped reads from the modified ChIP-seq experiments for H3K4me3, H3K4me1, H3K27ac and promoters or enhancers. Promoters were defined as the regions from -1 kb to +1 kb of the summits of a significant RNAPol2 binding sites (Polr2a) within 1 kb of a transcription start site (TSS). Enhancers were defined as the regions from -1 kb to +1 kb of the summits of a significant P300 binding sites that were not within 1kb of a TSS. The sequencing profiles of the conventional ChIP and modified ChIP for the used histone modifications and the Dppa2 ChIP-seq experiments (Fig. 1g, Fig. 4c and Supplementary Fig. 1) were created in the IGV browser<sup>47</sup>.

### **Genome-wide correlation.**

To calculate the genome wide correlation between the published histone modification

ChIP-seq datasets and the protein factors, we divided the entire genome in bins of 1000 base pairs and calculated Reads Per Million (RPM) for all bins in all datasets. The input was subtracted. For each histone modification and protein factor we selected the 4000 bins with the highest RPM. A unified list was created for each individual protein factor, containing the selected bins of the four used histone modifications and that of the protein factor itself. The Spearman correlation coefficients of the protein factor with the different histone modifications were calculated from this list. To calculate the correlations between conventional and modified ChIP-seq experiments for the used histone modifications, we used a unified list that included the 4000 bins with the highest RPM for each conventional and modified ChIP-seq experiment.

2

### Heatmaps.

To assign promoter regions, H3K4me3 peak summits, as determined by MACS (see above) were required to be within 1 kb range of a TSS, resulting in 12913 promoter regions. To assign enhancer regions, H3K4me1 peak summits were filtered against the presence of H3K4me3 signal in a region from -4.1 kb and + 4.1 kb around the H3K4me1 peak summit, resulting in 30564 enhancer regions. Promoters were sorted for number of H3K27ac reads present in the central 2 kb of the promoter region. Enhancers were sorted for the number of H3K27ac reads in the central 8.2 kb of the enhancer region, to also include broad enhancers. For both promoters and enhancers we displayed a region from -4.1 kb to +4.1 kb around the peak summits, divided into 51 bins of 160 bp each. Promoter and enhancer heatmaps for each protein factor or histone modification were normalized by calculating the RPM based on the sum of all reads found in the displayed promoter and enhancer region for that factor.

### Expression of Dppa2 and Oct4 target genes.

Dppa2-bound genes contained a MACS-called Dppa2 peak (see above) within 1 kb from the TSS. Dppa2 target genes were defined as Dppa2-bound genes with at least 2-fold difference in expression in Dppa2 knockout ESCs, compared to wild-type ESCs and an adjusted p-value of  $\leq 0.10$ , in a Dppa2 knockout microarray dataset<sup>31</sup>. The GEO2R script, as provided by the authors on the Gene Expression Omnibus (GEO), was used to calculate fold change in expression and adjusted p-value for each probe. Multiple probes to the same gene were aggregated by taking the average fold change. Oct4-bound genes contained a MACS-called Oct4 peak (see above) within 20 kb from the TSS. Oct4 target genes were defined as Oct4-bound genes with at least 2-fold difference in expression between 24 hrs and 0 hrs after Oct4 knockdown<sup>48</sup> in ZHBTc4 ESCs that have their only intact Oct4 gene under doxycycline control. The used microarray dataset<sup>48</sup> was already normalized by the authors. A published RNA sequencing dataset consisting of two replicates<sup>49</sup> was used to calculate the mean expression of Dppa2- or Oct4-bound genes and Dppa2- or Oct4-target genes. Both replicates were mapped against mouse reference NCBIM37.67 using Tophat<sup>50</sup> v2.0.11 with default settings and a segment length of 20. The aligned exon reads were counted and normalized using Bioconductor DESeq2 package in R. Replicates were normalized by dividing the counts by their sizefactors. The expression level per gene was calculated by taking the average of both replicates and calculating the reads per kb for each gene. To calculate the fold change of Dppa2- and Oct4-target genes in differentiated tissues over ESCs, we used the BioGPS mouse MOE430 Gene Atlas<sup>51</sup>. The same database was used to determine the tissue or cell line

in which the Dppa2 and Oct4 target genes were highest expressed.

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### **Author contributions**

E.E. performed the ChIP-MS and Dppa2 experiments and analyzed the data. J.H.B. performed the modified ChIP-seq experiments and all bioinformatics analyses. M.J.M. performed the Western analyses and cloned Dppa2 cDNA into an expression vector. L.S. cloned Dppa2 cDNA into an expression vector. D.H.W.D and J.D. performed the mass spectrometry analyses. C.E.M.K., Z.O. and W.F.J.v.IJ. performed Illumina sequencing of ChIP material. D.L.C.v.d.B. conceived the study and performed pilot experiments, R.A.P. conceived the study and designed experiments. R.A.P., E.E., J.H.B and D.L.C.v.d.B. wrote the manuscript.

### **Author information**

The authors declare no competing interests. ChIP sequencing data are available through the Gene Expression Omnibus (NCBI), accession code GSE58113.

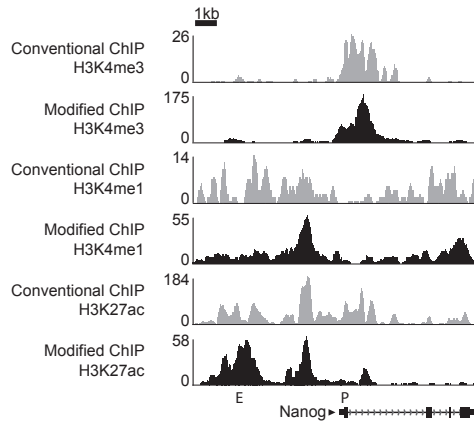
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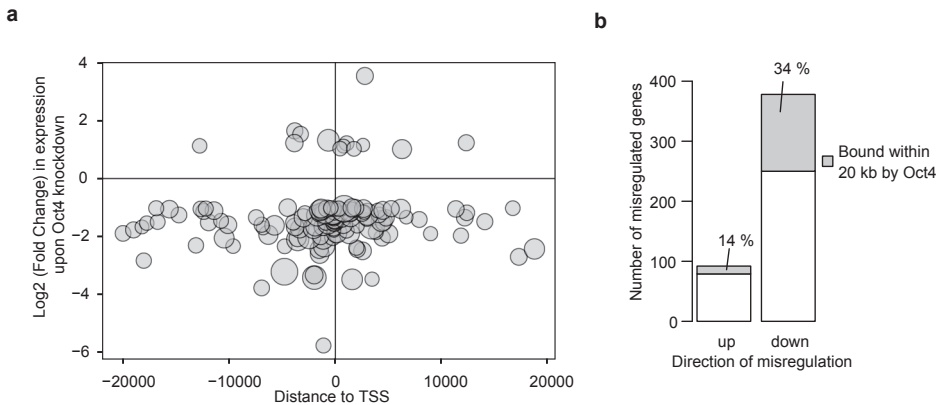
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SUPPLEMENTARY INFORMATION

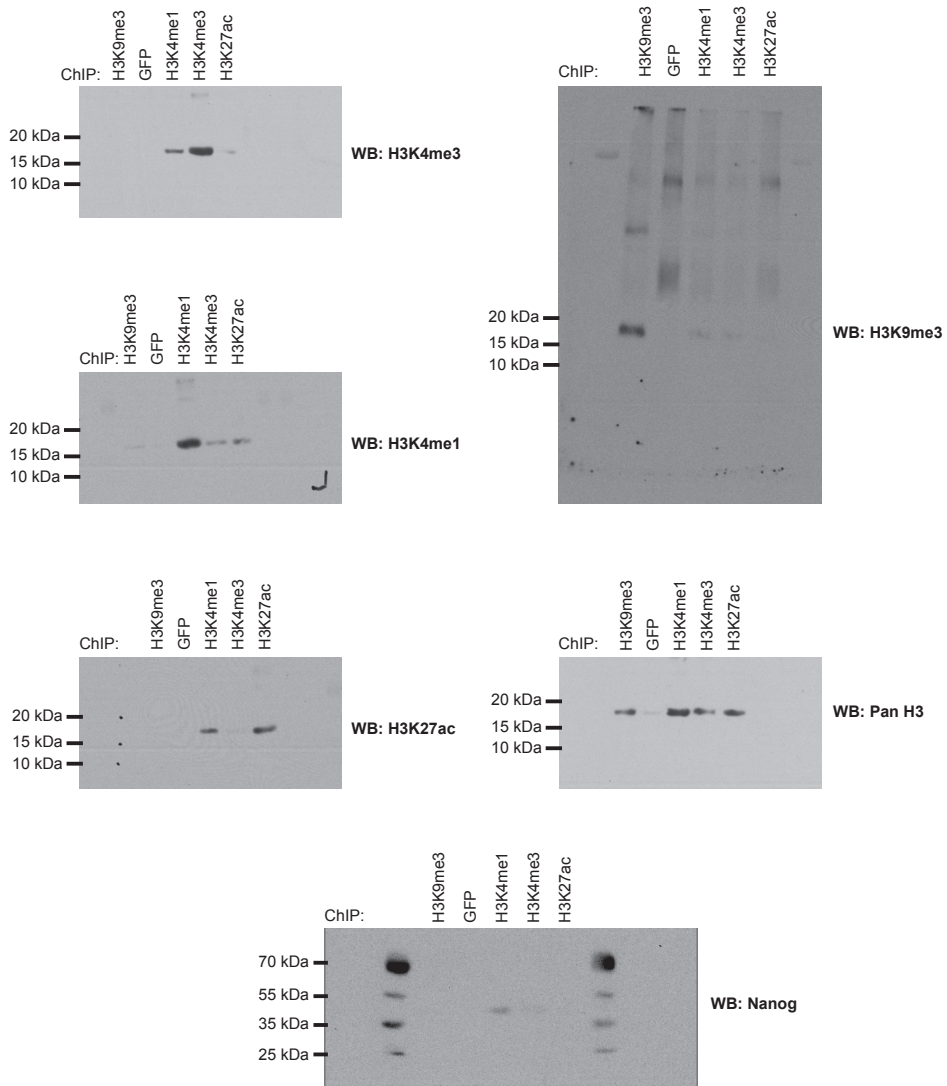


**Supplementary Figure 1. Modified ChIP-seq tracks around pluripotency gene *Nanog*.** ChIP-seq tracks for modified or conventional ChIP for H3K4me3, H3K4me1 or H3K27ac around pluripotency gene *Nanog*. Sequence reads were plotted relative to chromosomal position. Genome location of *Nanog* is shown, scale bar indicates 1 kb of genome. P indicates Promoter, E indicates putative enhancer.



**Supplementary Figure 2. Position of Oct4 binding sites on Oct4 target genes.** (a) Bubble plot indicating the positions of Oct4 binding sites relative to the transcription start site (TSS) on genes that are either up-regulated  $\geq 2$  fold (upper part) or down-regulated  $\geq 2$  fold (lower part) after 24 h of Oct4 depletion in mouse ESCs. Log<sub>2</sub> of the fold change in expression is indicated on the Y-axis, Oct4 binding distance from TSS (in base pairs) is indicated on the X-axis. Size of the bubbles correlates with fold difference of Oct4 ChIP peak over control. (b) Bar diagram showing the total number of up-regulated and down-regulated genes upon Oct4 depletion and the number of these genes bound by Oct4 within 20 kb from the TSS (grey areas). The number of Oct4-bound genes is also indicated as a percentage of the total number of up-regulated or down-regulated genes.

2



**Supplementary Figure 3. Western Blots (full scans) for ChIP-MS samples (as in Figure 1d).** Western blots for histone modifications and Nanog for ChIP-MS samples. ChIP-MS samples (top), molecular weight markers (left) and antibody used for Western blot (right) are indicated.

**Supplementary Table 1.** ChIP-MS identified proteins, predictions and phenotypes

Protein	Complex <sup>a</sup>	H3K4me3 avrg emPAI	H3K27ac avrg emPAI	H3K4me1 avrg emPAI	H3K9me3 avrg emPAI	GFP avrg emPAI	H3K27ac ratio	Prediction <sup>b</sup>	Pluripotency phenotype <sup>c</sup>
Act16a	BAF + INO80 + Trrap/Ep400	0,29	0,16	0,12	0,08		0,55	Promoter	-
Actr3	Arp2/3	0,04					0	Promoter	-
Actr5			0,03					Unclear	-
Adnp				0,03	0,02		0	Enhancer	-
Ankhd1				0,10			0	Enhancer	-
Anln					0,03			Heterochromatin	-
Anp32a	SET	0,17		0,18			0	Enhancer	-
Arid1a	BAF		0,07	0,06			1,17	Enhancer	yes
Arid4a	Sin3	0,12					0	Promoter	-
Ash2l	MLL	0,25					0	Promoter	yes
Atad2b			0,03	0,04	0,04		0,75	Enhancer	-
Atrx					0,09			Heterochromatin	-
Aurkb	Aurora K	0,22	0,10	0,10	0,65			Heterochromatin	yes
Banf1		0,66	0,47	0,63			0,71	Promoter	yes
Bap18	MLL	0,63					0	Promoter	-
Baz1a	ACF			0,01			0	Enhancer	-
Baz2a	NoRC	0,06	0,22	0,20	0,07		1,10	Enhancer	-
Bend3		0,24		0,04			0	Promoter	-
Bptf	NURF	0,08	0,02	0,02			0,25	Promoter	yes
Brd1	MOZ/MORF	0,03	0,05	0,10			0,50	Enhancer	-
Brd2		0,37	0,75	0,18			2,03	Promoter	-
Brd3		0,12	0,32				2,67	Promoter	-
Brd4		0,03	0,21	0,10			2,10	Enhancer	-
Brms1	Sin3	0,14					0	Promoter	-
Brms1l	Sin3	0,47					0	Promoter	-
Brpf1	MOZ/MORF	0,16					0	Promoter	-
Brpf3	MOZ/MORF	0,03					0	Promoter	-
Cad				0,04			0	Enhancer	-
Cald1		0,06					0	Promoter	-
Cbx7	PRC1	0,10		0,10			0	Promoter	yes
Cdc5l	Prp19			0,04			0	Enhancer	-
Cdc73	PAF1	0,10	0,10				1,00	Promoter	yes
Cdca7			0,09	0,09			1,00	Enhancer	-
Cdca8	Aurora K	0,28	0,13	0,24	1,17			Heterochromatin	-
Cdk7	TFIIH	0,05					0	Promoter	-
Cdk9	P-TEFb		0,09					Unclear	yes
Cep95			0,02	0,04			0,50	Enhancer	-
Chaf1b	CAF1		0,23	0,52			0,44	Enhancer	yes
Chd1		0,26	0,24	0,14	0,02		0,92	Promoter	yes
Chd2		0,05		0,03			0	Promoter	-
Chd8		0,15	0,03				0,20	Promoter	-
Ckb		0,14					0	Promoter	-
Coil					0,03			Heterochromatin	-
Cpsf6	CFIm	0,07	0,07				1,00	Promoter	-
Creb1		0,11					0	Promoter	-
Cse1l		0,04					0	Promoter	-
Csnk2a1		0,89	0,27	0,21			0,30	Promoter	-
Csnk2b		0,82	0,26	0,08			0,32	Promoter	-
Csrp2bp	ATAC	0,04					0	Promoter	-
Ctbp2		0,46	0,30	0,43	0,04		0,65	Promoter	yes
Ctcf		0,07		0,05			0	Promoter	-
Ctr9	PAF1	0,03	0,03				1,00	Promoter	yes
Cul4b	DDB1/Cul4 ubiquitin ligase	0,17	0,18	0,18			1,00	Enhancer	-
Ddb1	DDB1/Cul4 ubiquitin ligase	0,19	0,24	0,29			0,83	Enhancer	-
Ddx47					0,15			Heterochromatin	-
Ddx54				0,04			0	Enhancer	-
Dido1		0,02					0	Promoter	-
Dis3			0,05	0,05			1,00	Enhancer	-
Dmap1	Trrap/Ep400	0,07		0,04			0	Promoter	yes
Dnajc9		0,07	0,14	0,14			1,00	Enhancer	-
Dpf2	BAF		0,22	0,12			1,83	Enhancer	-
Dppa2		0,61	0,06	0,29			0,10	Promoter	yes
Dppa4		0,58		0,39			0	Promoter	yes
Dpy30	MLL	0,18					0	Promoter	yes
Dut		0,41	0,18	0,18			0,44	Promoter	-
Emsy		0,02					0	Promoter	-
Ep300			0,05					Unclear	yes
Ep400	Trrap/Ep400		0,02					Unclear	yes



Protein	Complex <sup>a</sup>	H3K4me3 avrg emPAI	H3K27ac avrg emPAI	H3K4me1 avrg emPAI	H3K9me3 avrg emPAI	GFP avrg emPAI	H3K27ac ratio	Prediction <sup>b</sup>	Pluripotency phenotype <sup>c</sup>
Erh		0,37					0	Promoter	-
Esrb			0,08	0,20			0,40	Enhancer	yes
Ezh2	PRC2	0,07	0,04	0,09			0,44	Enhancer	yes
Fam60a	Sin3	0,24					0	Promoter	-
Fbxl19	SCF ubiquitin ligase	0,05					0	Promoter	-
Fkbp4				0,07			0	Enhancer	-
Gltscr2				0,07			0	Enhancer	-
Glyr1		0,19	0,24	0,26	0,07		0,92	Enhancer	-
Gm53		0,19					0	Promoter	-
Gtf2e1	TFIIE			0,08			0	Enhancer	-
Gtf2i			0,04	0,07			0,57	Enhancer	-
Gtf3c1	TFIIIC	0,01	0,01	0,02			0,50	Enhancer	-
Hat1		0,11	0,04	0,04			0,36	Promoter	-
Hcfc1	MLL	0,05					0	Promoter	yes
Hdac1	NuRD + Sin3 + REST	1,02	0,30	0,38	0,11		0,29	Promoter	yes
Hdac2	NuRD + Sin3 + REST	0,40		0,10			0	Promoter	-
Hdgfrp2			0,05					Unclear	-
Hmga2		0,14	0,14		0,28			Heterochromatin	-
Hmgb3		0,11		0,1			0	Promoter	-
Hmgxb4		0,20					0	Promoter	-
Hp1bp3		0,26		0,38	0,23		0	Enhancer	-
Incenp	Aurora K	0,06			0,26			Heterochromatin	-
Ing1	Sin3	0,06					0	Promoter	-
Ing2	Sin3	0,11					0	Promoter	-
Ing3	Trrap/Ep400	0,04	0,04				1,00	Promoter	-
Ing4	Trrap/Ep400	0,14		0,07			0	Promoter	-
Ing5	MOZ/MORF	0,58	0,13	0,20			0,22	Promoter	yes
Ino80	INO80	0,05	0,05				1,00	Promoter	yes
Ints1	Integrator			0,02			0	Enhancer	-
Irgc				0,04			0	Enhancer	-
Jarid2	PRC2	0,16	0,05	0,21	0,04		0,24	Enhancer	yes
Kdm1a	BHC	0,06	0,02	0,02	0,02		0,33	Promoter	yes
Kdm2a	SCF ubiquitin ligase	0,16	0,03	0,03			0,19	Promoter	-
Kdm2b	PRC1	0,09		0,01			0	Promoter	yes
Kdm4a		0,03					0	Promoter	-
Kdm4c		0,65	0,03	0,13			0,05	Promoter	yes
Kdm5a		0,02					0	Promoter	-
Kdm5b		0,25	0,02	0,12			0,08	Promoter	yes
Kif2c				0,05			0	Enhancer	-
Kif4				0,03			0	Enhancer	-
Kifc5b			0,07	0,15			0,47	Enhancer	-
Kif16		0,16					0	Promoter	-
Kif5			0,07	0,04			1,75	Enhancer	yes
L3mbtl2		0,08		0,07			0	Promoter	yes
Las1l	5FMC			0,06			0	Enhancer	-
Lasp1		0,29		0,13			0	Promoter	-
Lig1		0,08		0,02			0	Promoter	-
Lig3			0,02	0,05			0,40	Enhancer	-
Liph				0,13			0	Enhancer	-
Lmna		0,13	0,08	0,08	0,48			Heterochromatin	-
Lmnb1		0,99	1,60	1,83	3,15	0,37		Heterochromatin	-
Lmnb2		0,12	0,24	0,30	0,98			Heterochromatin	-
Lrwd1	ORC	0,14		0,11	0,35			Heterochromatin	-
Lsm2	U6 SnRNP		0,18					Unclear	-
Luc7l3				0,04			0	Enhancer	-
Max		0,11	0,11	0,10			1,00	Promoter	yes
Mbtd1		0,06					0	Promoter	-
Mdc1			0,03	0,05			0,60	Enhancer	-
Meaf6	Trrap/Ep400	0,51	0,19	0,19			0,37	Promoter	-
Men1	MLL	0,32	0,03	0,15			0,09	Promoter	-
Mil2	MLL	0,21		0,02		0,01	0,00	Promoter	-
Morc2a			0,05					Unclear	-
Morc3		0,70	0,24	0,24	0,02		0,34	Promoter	-
Mta3	NuRD	0,24	0,14	0,27			0,52	Enhancer	-
Mtf2	PRC2	0,16	0,06	0,22	0,06		0,27	Enhancer	yes
Mycn		0,08	0,08				1,00	Promoter	yes
Myst2	HBO1	0,56	0,36	0,39	0,06		0,64	Promoter	yes
Myst3	MOZ	0,08					0	Promoter	yes
Myst4	MORF	0,04					0	Promoter	-
Nacc1		0,04	0,07	0,13			0,54	Enhancer	yes
Nasp			0,02	0,02			1,00	Enhancer	-
Nfrkb			0,03					Unclear	yes

Proteins that bind regulatory regions identified by histone modification chromatin immunoprecipitations and mass spectrometry

Protein	Complex <sup>a</sup>	H3K4me3 avrg emPAI	H3K27ac avrg emPAI	H3K4me1 avrg emPAI	H3K9me3 avrg emPAI	GFP avrg emPAI	H3K27ac ratio	Prediction <sup>b</sup>	Pluripotency phenotype <sup>c</sup>
Nolc1				0,05			0	Enhancer	-
Nrf1		0,08					0	Promoter	-
Nsd1		0,01	0,04	0,05			0,80	Enhancer	-
Nudt13				0,10			0	Enhancer	-
Nudt21	CFIm	0,48					0	Promoter	-
Numa1		0,22	0,15	0,29	0,10		0,52	Enhancer	-
Oct4		0,16	0,10	0,21	0,05		0,48	Enhancer	yes
Ogt		0,15	0,02	0,05			0,13	Promoter	yes
Pak1ip1		0,19					0	Promoter	-
Parp2				0,03			0	Enhancer	-
Parp9					0,04			Heterochromatin	-
Patz1		0,05		0,05			0	Promoter	-
Pbrm1	BAF	0,02	0,11	0,11			1,00	Enhancer	-
Pds5a	Wapl		0,03	0,05			0,60	Enhancer	-
Pds5b	Wapl	0,02	0,01	0,02			0,50	Enhancer	-
Phc1	PRC1			0,04			0	Enhancer	-
Phf16	HBO1	0,04					0	Promoter	-
Phf20				0,02			0	Enhancer	yes
Phf23		0,53		0,09			0	Promoter	yes
Phf8		0,23	0,02				0,09	Promoter	-
Phip		0,04	0,07	0,13			0,54	Enhancer	-
Pin1			0,11	0,11			1,00	Enhancer	yes
Pnp		0,20					0	Promoter	-
Polr1c	Pol I	0,3					0	Promoter	-
Polr2a	Pol II	0,15	0,13	0,06			0,87	Promoter	-
Polr2b	Pol II	0,43	0,33	0,07			0,77	Promoter	-
Polr2c	Pol II	0,57	0,16	0,11			0,28	Promoter	-
Polr2e	Pol II	0,85	0,24	0,08			0,28	Promoter	-
Polr2g	Pol II	0,22					0	Promoter	-
Ppp2cb				0,11			0	Enhancer	-
Prdm10		0,03	0,03				1,00	Promoter	-
Prdm2				0,02			0	Enhancer	-
Prrc2c				0,01			0	Enhancer	-
Psma5	Proteasome	0,07		0,07			0	Promoter	-
Ptcd3				0,12			0	Enhancer	-
Qser1		0,03		0,01			0	Promoter	-
Rad21	Cohesin		0,03	0,03			1,00	Enhancer	yes
Rars		0,05					0	Promoter	-
Rbbp4	NuRD/CAF1	1,51	0,62	1,09	0,37	0,12	0,41	Promoter	-
Rbbp5	MLL	0,14					0	Promoter	yes
Rbl1				0,03			0	Enhancer	-
Rbpj		0,30		0,20	0,14		0	Promoter	-
Rfc3		0,05	0,10				2,00	Promoter	-
Rfc4		0,20	0,15	0,19			0,75	Promoter	-
Rnf2	PRC1	0,35		0,30			0	Promoter	yes
Rsf1	RSF	0,05	0,11	0,12			0,92	Enhancer	-
Ruvbl2	Ttrap/Ep400	1,36	1,91	0,41	0,16		1,40	Promoter	yes
Samd1		0,26					0	Promoter	-
Sap130	Sin3	0,18					0	Promoter	-
Sap30	Sin3	0,29					0	Promoter	-
Sap30l	Sin3	0,09					0	Promoter	-
Sbno1				0,01			0	Enhancer	-
Sephs1		0,14	0,05				0,36	Promoter	-
Sf1		0,14					0	Promoter	-
Sf3b1				0,04			0	Enhancer	-
Shprh		0,04	0,01	0,03			0,25	Promoter	-
Sin3a	Sin3	1,34	0,15	0,30			0,11	Promoter	yes
Sin3b	Sin3	0,09					0	Promoter	-
Skp1a	SCF ubiquitin ligase	0,45	0,23	0,10			0,51	Promoter	-
Smarca4	BAF	0,14	0,28	0,22	0,03		1,27	Enhancer	yes
Smarcb1	BAF		0,09	0,05			1,80	Enhancer	yes
Smarcc1	BAF	0,10	0,46	0,31	0,07		1,48	Enhancer	yes
Smarcd1	BAF		0,50	0,35			1,43	Enhancer	-
Smc1a	Cohesin	0,08	0,12	0,22	0,06		0,55	Enhancer	yes
Smc6	Smc5/Smc6			0,03			0	Enhancer	yes
Smchd1			0,02	0,03	0,04			Heterochromatin	-
Spin1		0,72	0,13	0,13			0,18	Promoter	-
Srrt		0,04	0,04	0,08			0,50	Enhancer	-
Ssrp1	FACT	0,68	0,93	0,73	0,29	0,13	1,27	Enhancer	-
Suds3	Sin3	0,41					0	Promoter	-
Supt4h2	DSIF	0,34					0	Promoter	yes
Supt5h	DSIF	0,22	0,16	0,06			0,73	Promoter	-



Chapter2

Protein	Complex <sup>a</sup>	H3K4me3 avrg emPAI	H3K27ac avrg emPAI	H3K4me1 avrg emPAI	H3K9me3 avrg emPAI	GFP avrg emPAI	H3K27ac ratio	Prediction <sup>b</sup>	Pluripotency phenotype <sup>c</sup>
Supt6h		0,07	0,09				1,29	Promoter	-
Suv39h2					0,31			Heterochromatin	yes
Suz12	PRC2	0,31	0,12	0,40	0,07		0,30	Enhancer	yes
Taf1	TFIID	0,03					0	Promoter	yes
Taf2	TFIID	0,06	0,02				0,33	Promoter	yes
Taf3	TFIID	0,02	0,02				1,00	Promoter	yes
Taf4a	TFIID	0,11					0	Promoter	yes
Taf5	TFIID	0,05					0	Promoter	yes
Taf6	TFIID	0,20	0,14				0,70	Promoter	yes
Taf7	TFIID	0,16	0,11				0,69	Promoter	yes
Taldo1			0,05	0,11			0,45	Enhancer	-
Tbrg4				0,09			0	Enhancer	-
Tcea1	TFIIS	0,12	0,26	0,05			2,17	Promoter	-
Tcerg1		0,02	0,03	0,02			1,50	Enhancer	-
Tcof1					0,02			Heterochromatin	-
Tead1			0,04	0,04			1,00	Enhancer	-
Tlk2				0,06			0	Enhancer	-
Trim24			0,25	0,04			6,25	Enhancer	-
Trim33			0,08					Unclear	-
Trp53			0,06					Unclear	-
Trrap	Trrap/Ep400	0,05	0,05	0,01			1,00	Promoter	yes
Uba1			0,03	0,03			1,00	Enhancer	-
Ube2h				0,24			0	Enhancer	-
Ubtf		0,79	0,44	0,43			0,56	Promoter	-
Usp48		0,15	0,08	0,21	0,05		0,38	Enhancer	-
Utp14b				0,07			0	Enhancer	-
Vmn2r100		0,02		0,06			0	Enhancer	-
Wdr18	5FMC	0,04		0,12	0,08		0	Enhancer	-
Wdr5	MLL	0,60	0,27	0,17			0,45	Promoter	yes
Wdr55		0,05					0	Promoter	-
Zfp280c		0,05			0,22			Heterochromatin	-
Zfp281		0,04					0	Promoter	yes
Zfp462		0,01			0,02			Heterochromatin	-
Zic5		0,03					0	Promoter	-
Zmynd8	Integrator		0,05	0,04			1,25	Enhancer	-
Znf512		0,13		0,24	0,25			Heterochromatin	-
Zscan10		0,04		0,09			0	Enhancer	yes

<sup>a</sup> Subunit of indicated protein complex

<sup>b</sup> Prediction of genome localization based on our ChIP-MS criteria

<sup>c</sup> ESC pluripotency phenotype (references in supplementary material)

Supplementary Table 2. ChIP-MS identified proteins

Protein	H3K4me3						H3K27ac						H3K4me1						H3K9me3						GFP											
	Experiment 1			Experiment 2			Experiment 1			Experiment 2			Experiment 1			Experiment 2			Experiment 1			Experiment 2			Experiment 1			Experiment 2								
	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b				
Actf6a	305	0.57	6	25																																
Actr3	56	0.08	1	2																																
Actr5								79	0.05	1	1																									
Adnp																																				
Ankhd1																																				
Anln																																				
Anp32a	75	0.17	2	3	47	0.17	1	2																												
Arid1a																																				
Arid4a	79	0.24	2	4																																
Ash2l	195	0.35	5	15	77	0.14	2	3																												
Atad2b																																				
Atrx																																				
Aurkb					238	0.44	4	18																												
Banf1	107	0.93	2	6	59	0.39	1	2																												
Bap18	166	0.79	3	30	119	0.47	2	8																												
Baz1a																																				
Baz2a	57	0.04	2	2	239	0.07	5	7	363	0.21	10	19	579	0.23	11	23																				
Bend3	196	0.26	5	7	206	0.21	5	5																												
Bpif	422	0.11	10	17	173	0.04	4	10	102	0.03	3	3																								
Brd1	70	0.06	2	5					57	0.03	1	2	88	0.06	2	2																				
Brd2	471	0.51	10	45	273	0.23	5	11	527	0.57	11	30	750	0.93	14	53																				
Brd3	119	0.15	3	5	81	0.09	2	4	278	0.44	7	14	174	0.2	4	11																				
Brd4	72	0.05	2	5					313	0.23	9	15	323	0.18	6	33																				
Brms1	81	0.28	2	2																																
Brms1l	179	0.6	5	5	177	0.33	3	5																												



Protein	H3K4me3						H3K27ac						H3K4me1						H3K9me3						GFP										
	Experiment 1			Experiment 2			Experiment 1			Experiment 2			Experiment 1			Experiment 2			Experiment 1			Experiment 2			Experiment 1			Experiment 2							
	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b			
Brip1	233	0.23	7	11	91	0.08	3	3																											
Brip3					86	0.05	2	5																											
Cad												204	0.08	6	9																				
Cald1	61	0.12	2	2																															
Cbx7	50	0.19	1	4								56	0.19	1	1																				
Cdcs1												58	0.08	2	2																				
Cdc73	82	0.2	3	3	84	0.13	2	2	62	0.06	1	2																							
Cdca7									64	0.18	1	2																							
Cdca8									62	0.25	2	3	97	0.23	2	5	56	0.25	2	2	344	1.17	7	24	392	1.17	7	40							
Cdk7																																			
Cdk9					61	0.18	2	2																											
Cep95									45	0.04	1	14	68	0.07	2	8																			
Chaf1b									45	0.45	1	1	109	1.04	2	2																			
Chd1	672	0.34	15	48	382	0.18	8	29	239	0.16	8	14	641	0.32	14	47	212	0.11	6	11	323	0.16	7	28											
Chd2	177	0.09	4	22													96	0.05	3	7															
Chd8	516	0.23	14	33	199	0.07	4	12																											
Ckb	66	0.18	2	4	60	0.09	1	1																											
Coll																																			
Cpsf6					88	0.13	2	13					131	0.13	2	14																			
Creb1	60	0.22	2	2																															
Cse1l					72	0.07	2	64																											
Csnk2a1	383	1.4	10	19	115	0.37	3	6	106	0.27	3	4	117	0.25	3	5	58	0.17	2	3															
Csnk2b	238	1.32	6	14	80	0.32	2	3					135	0.52	3	5					51	0.15	1	2											
Csrp2bp	66	0.08	2	9																															
Ctbp2	452	0.67	7	68	154	0.24	4	8	229	0.44	5	6	72	0.16	2	4	309	0.51	6	9	188	0.34	4	6											
Ctcf	87	0.14	3	5																															
Ctcf9	108	0.06	2	2									91	0.06	2	2																			

Protein	H3K4me3						H3K27ac						H3K4me1						H3K9me3						GFP												
	Experiment 1			Experiment 2			Experiment 1			Experiment 2			Experiment 1			Experiment 2			Experiment 1			Experiment 2			Experiment 1			Experiment 2									
	Mascot	emPAL	Unique pept. a	Unique pept. b	Mascot	emPAL	Unique pept. a	Unique pept. b	Mascot	emPAL	Unique pept. a	Unique pept. b	Mascot	emPAL	Unique pept. a	Unique pept. b	Mascot	emPAL	Unique pept. a	Unique pept. b	Mascot	emPAL	Unique pept. a	Unique pept. b	Mascot	emPAL	Unique pept. a	Unique pept. b	Mascot	emPAL	Unique pept. a	Unique pept. b					
Cul4b	262	0.26	7	9	79	0.07	2	239	0.22	6	8	195	0.14	4	4	229	0.13	4	8	275	0.22	7	11														
Ddb1	349	0.22	8	10	198	0.15	5	339	0.29	9	9	317	0.19	6	7	414	0.31	10	15	339	0.26	8	14														
Ddx47																																					
Ddx54																																					
Dldd1	120	0.04	3	12																																	
Dis3																																					
Dmap1	100	0.14	2	3																																	
Dnajc9					48	0.13	1	3																													
Dpf2																																					
Dppa2	255	0.86	6	33	157	0.36	3	13																													
Dppa4	231	0.92	7	17	75	0.24	2	22																													
Dpy30	50	0.36	1	4																																	
Dut	117	0.64	3	8	58	0.18	1	4	49	0.18	1	2	48	0.18	1	3	53	0.17	2	6	69	0.18	1	5													
Ermsy	56	0.03	1	1																																	
Epp300																																					
Epp400																																					
Erh	71	0.74	2	2																																	
Esrb																																					
Ezh2	99	0.09	2	5	73	0.04	1	2	47	0.04	1	1	98	0.04	1	2	79	0.04	1	2	141	0.14	3	4													
Fam60a	75	0.48	2	2																																	
Fbxl19					75	0.1	2	4																													
Fkbp4																																					
Glsr2																																					
Glyr1	136	0.19	3	6	147	0.19	2	9	181	0.34	4	6	111	0.13	2	4	292	0.32	5	16	172	0.19	3	11													
Gms3																																					
Gtf2e1					59	0.37	2	3																													
Gtf2i																																					



Protein	H3K4me3						H3K27ac						H3K4me1						H3K9me3						GFP							
	Experiment 1			Experiment 2			Experiment 1			Experiment 2			Experiment 1			Experiment 2			Experiment 1			Experiment 2			Experiment 1			Experiment 2				
	Mascot	empAI	Unique pep <sub>a</sub>	Pept. count <sub>b</sub>	Mascot	empAI	Unique pep <sub>a</sub>	Pept. count <sub>b</sub>	Mascot	empAI	Unique pep <sub>a</sub>	Pept. count <sub>b</sub>	Mascot	empAI	Unique pep <sub>a</sub>	Pept. count <sub>b</sub>	Mascot	empAI	Unique pep <sub>a</sub>	Pept. count <sub>b</sub>	Mascot	empAI	Unique pep <sub>a</sub>	Pept. count <sub>b</sub>	Mascot	empAI	Unique pep <sub>a</sub>	Pept. count <sub>b</sub>	Mascot	empAI	Unique pep <sub>a</sub>	Pept. count <sub>b</sub>
Glf3c1	55	0.02	1	1				53	0.02	1	1					97	0.03	2	4													
Hat1	88	0.15	2	4	45	0.07	1	2								63	0.07	1	6													
Hcf1	137	0.09	4	7																												
Hdac1	453	1.65	9	75	239	0.38	5	24	118	0.3	3	7	170	0.3	4	9	176	0.28	4	7	244	0.48	6	23	87	0.14	2	7	60	0.07	1	3
Hdac2	222	0.79	5	52												114	0.2	3	6													
Hdgfrp2								67	0.1	2	2																					
Hnga2					82	0.28	1	7																								
Hngb3	52	0.21	1	6																												
Hngxb4	60	0.39	2	2												63	0.2	1	6													
Hpl1bp3	45	0.12	1	3	104	0.39	3	6																								
Incep					152	0.11	3	9																								
Ing1	52	0.12	1	1																												
Ing2	57	0.11	1	15	57	0.11	1	6																								
Ing3					70	0.08	1	1					74	0.08	1	1																
Ing4	60	0.28	2	2																												
Ing5	168	0.87	4	22	71	0.29	1	2	46	0.13	1	3	63	0.13	1	1	72	0.27	2	3	73	0.13	1	2								
Inc80	147	0.06	3	5	79	0.04	3	3	140	0.09	4	5																				
Ints1																																
Irgc																																
Jarid2	263	0.17	8	13	309	0.14	5	11	102	0.05	2	2	128	0.05	2	4	281	0.19	7	11	424	0.23	8	19	70	0.03	2	2	104	0.05	2	3
Kdm1a	94	0.08	2	2	91	0.04	1	1					57	0.04	1	1																
Kdm2a	435	0.24	9	43	158	0.08	3	15					66	0.06	2	2	51	0.03	1	2	62	0.03	1	3								
Kdm2b	267	0.16	6	13	71	0.02	2	5																								
Kdm4a	67	0.06	2	3																												
Kdm4c	861	0.87	18	86	647	0.43	11	48					119	0.06	2	4	73	0.06	2	2	270	0.2	6	16								
Kdm5a	61	0.04	2	2																												
Kdm5b	519	0.3	13	38	424	0.2	9	16					107	0.04	2	3	68	0.05	1	5	347	0.18	8	18								





Protein	H3K4me3						H3K27ac						H3K4me1						H3K9me3						GFP										
	Experiment 1			Experiment 2			Experiment 1			Experiment 2			Experiment 1			Experiment 2			Experiment 1			Experiment 2			Experiment 1			Experiment 2							
	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b			
Phlp	152	0.05	4	7	63	0.02	1	1	134	0.07	4	4	187	0.07	4	6	248	0.1	7	9	295	0.15	7	17											
Pin1													57	0.21	1	3					48	0.21	1	2											
Php	129	0.39	3	5																															
Poir1c	174	0.6	5	8																															
Polr2a	460	0.18	11	21	321	0.11	6	9	333	0.12	8	9	498	0.14	9	15	129	0.05	3	3	159	0.07	4	5											
Polr2b	709	0.67	18	54	258	0.18	6	14	407	0.34	11	17	421	0.31	10	17	63	0.05	2	2	148	0.08	3	4											
Polr2c	308	0.93	7	16	147	0.21	2	2				158	0.32	3	5					100	0.21	2	2												
Polr2e	252	1.37	6	12	100	0.33	2	2	57	0.33	2	2	71	0.15	1	2					55	0.15	2	2											
Polr2g	114	0.44	2	2																															
Ppp2cb																																			
Prdm10	63	0.06	2	2								87	0.06	2	5																				
Prdm2																																			
Prrc2c																																			
Psma5	79	0.14	1	1																															
Ptcd3																																			
Qser1	135	0.06	4	4	46	0.02	1	1																											
Rad21																																			
Rars																																			
Rbbp4	697	2.02	14	108	476	1	9	71	288	0.62	7	13	304	0.62	6	28	507	1.18	12	25	488	1	9	48											
Rbbp5	144	0.27	4	9																															
Rbl1																																			
Rbpj	169	0.3	4	8	142	0.3	4	9																											
Rtc3																																			
Rtc4	94	0.3	3	4	52	0.09	1	1	103	0.3	3	3	74	0.19	2	3																			
Rnl2	212	0.6	5	12	52	0.1	1	1																											
Rsf1	81	0.07	2	3	76	0.02	1	4	192	0.09	4	7	279	0.12	5	16	127	0.09	3	9	325	0.14	6	19											
Ruvb12	729	2.3	14	92	240	0.42	5	7	497	1.02	9	28	716	2.8	15	24	291	0.49	6	10	170	0.32	4	4											





Protein	H3K4me3						H3K27ac						H3K4me1						H3K9me3						GFP								
	Experiment 1			Experiment 2			Experiment 1			Experiment 2			Experiment 1			Experiment 2			Experiment 1			Experiment 2			Experiment 1			Experiment 2					
	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	Mascot	empAI	Unique pept. a	Pept. count b	
Suz12	229	0.37	6	11	251	0.25	5	73	0.09	2	193	0.14	4	6	294	0.29	6	10	455	0.5	8	19	144	0.14	3	6							
Taf1	100	0.05	3																														
Taf2	149	0.12	4	5			1	47	0.03	1																							
Taf3	79	0.04	1	7			1	68	0.04	1																							
Taf4a	144	0.22	4	4																													
Taf5	65	0.09	2	2																													
Taf6	230	0.29	5	6	136	0.11	2	97	0.11	3	168	0.17	3	3																			
Taf7	84	0.2	2	5	70	0.11	1	48	0.11	1	59	0.11	1	2																			
Talor1																																	
Tbig4																																	
Tee1	78	0.23	2	6				133	0.52	4	4																						
Tee1g1					57	0.03	1				125	0.06	2	3																			
Tcof1																																	
Tead1																																	
Tlk2																																	
Trim24																																	
Trim33																																	
Trp53																																	
Trrap	204	0.06	5	148	0.04	4	4	98	0.04	3	314	0.05	6	9																			
Uba1																																	
Ube2h																																	
Ubf1	732	1.07	19	73	457	0.5	9	374	0.5	10	14	359	0.38	8	18	342	0.35	8	14	457	0.5	10	28										
Usp48	195	0.13	4	11	162	0.16	3																										
Utp14b																																	
Vmm2r100																																	
Wdr18	52	0.08	1	1																													
Wdr5	202	0.57	4	22	170	0.62	5	86	0.21	2	3	133	0.33	3	10	119	0.24	3	3														





**Supplementary Table 3.** CHIP-MS identified complex subunits and predictions

Complex / subunits	H3K4me3 avrg emPAI	H3K27ac avrg emPAI	H3K4me1 avrg emPAI	H3K9me3 avrg emPAI	GFP avrg emPAI	H3K27ac ratio	Prediction <sup>a</sup>
BAF complex	0,04	0,25	0,17	0,01		1,43	Enhancer
Arid1a (Baf250a)		0,07	0,06				Enhancer
Dpf2 (Baf45d)		0,22	0,12				Enhancer
Pbrm1 (Baf180)	0,02	0,11	0,11				Enhancer
Smarca4 (Brg1)	0,14	0,28	0,22	0,03			Enhancer
Smarca1 (Baf47)		0,09	0,05				Enhancer
Smarcc1 (Baf155)	0,10	0,46	0,31	0,07			Enhancer
Smarcd1 (Baf60a)		0,50	0,35				Enhancer
Sin3 complex	0,32	0,01	0,03			0,04	Promoter
Sin3a	1,34	0,15	0,30				Promoter
Sin3b	0,09						Promoter
Arid4a (Rbbp1)	0,12						Promoter
Brms1	0,14						Promoter
Brms1l	0,47						Promoter
Fam60a	0,24						Promoter
Ing1	0,06						Promoter
Ing2	0,11						Promoter
Sap130	0,18						Promoter
Sap30	0,29						Promoter
Sap30l	0,09						Promoter
Suds3	0,41						Promoter
PRC1 complex	0,15		0,14			0	Promoter
Cbx7	0,10		0,10				Promoter
Phc1			0,04				Enhancer
Rnf2	0,35		0,30				Promoter
PRC2 complex	0,17	0,07	0,23	0,04		0,29	Enhancer
Ezh2	0,07	0,04	0,09				Enhancer
Jarid2	0,16	0,05	0,21	0,04			Enhancer
Mtf2	0,16	0,06	0,22	0,06			Enhancer
Suz12	0,31	0,12	0,40	0,07			Enhancer
TFIID complex	0,09	0,04				0,47	Promoter
Taf1	0,03						Promoter
Taf2	0,06	0,02					Promoter
Taf3	0,02	0,02					Promoter
Taf4a	0,11						Promoter
Taf5	0,05						Promoter
Taf6	0,20	0,14					Promoter
Taf7	0,16	0,11					Promoter
Aurora Kinase complex	0,19	0,08	0,11	0,69			Heterochromatin
Aurkb	0,22	0,10	0,10	0,65			Heterochromatin
Cdca8	0,28	0,13	0,24	1,17			Heterochromatin
Incenp	0,06			0,26			Heterochromatin



Complex / subunits	H3K4me3 avrg emPAI	H3K27ac avrg emPAI	H3K4me1 avrg emPAI	H3K9me3 avrg emPAI	GFP avrg emPAI	H3K27ac ratio	Prediction <sup>a</sup>
Pol II complex	0,44	0,17	0,06			0,39	Promoter
Polr2a	0,15	0,13	0,06				Promoter
Polr2b	0,43	0,33	0,07				Promoter
Polr2c	0,57	0,16	0,11				Promoter
Polr2e	0,85	0,24	0,08				Promoter
Polr2g	0,22						Promoter
MLL complex	0,34	0,04	0,05			0,13	Promoter
MLL2	0,21		0,02		0,01		Promoter
Ash2l	0,25						Promoter
Bap18	0,63						Promoter
Dpy30	0,18						Promoter
Hcfc1	0,05						Promoter
Men1	0,32	0,03	0,15				Promoter
Rbbp5	0,14						Promoter
Wdr5	0,60	0,27	0,17				Promoter
Trrap/Ep400 complex	0,30	0,39	0,09	0,03	0,00	1,33	Promoter
Trrap	0,05	0,05	0,01				Promoter
Ep400		0,02					Unclear
Dmap1	0,07		0,04				Promoter
Ruvbl2	1,36	1,91	0,41	0,16			Promoter
HBO1/MOZ/MORF complex	0,20	0,07	0,09	0,01		0,34	Promoter
Myst2	0,56	0,36	0,39	0,06			Promoter
Myst3	0,08						Promoter
Myst4	0,04						Promoter
Brd1	0,03	0,05	0,10				Enhancer
Brpf1	0,16						Promoter
Brpf3	0,03						Promoter
Meaf6	0,51	0,19	0,19				Promoter
Ing4	0,14		0,07				Promoter
Ing5	0,58	0,13	0,20				Promoter
Phf16	0,04						Promoter
Cohesin complex	0,04	0,07	0,12	0,03		0,60	Enhancer
Rad21		0,03	0,03				Enhancer
Smc1a	0,08	0,12	0,22	0,06			Enhancer
DSIF complex	0,28	0,08	0,03			0,29	Promoter
Supt4h2	0,34						Promoter
Supt5h	0,22	0,16	0,06				Promoter

<sup>a</sup> Prediction of genome localization based on our CHIP-MS criteria

**Supplementary Table 4.** Dppa2 target genes

Gene symbol	Ensembl gene ID	Tissue/cells with highest expression	FC over Embryonic stem cells
Pdgfb	ENSMUSG00000000489	Macrophage bone marrow 2hr LPS	83.7
Gnmt	ENSMUSG00000002769	Liver	3404.7
Gstm5	ENSMUSG00000004032	Testis	247.1
Taf7l	ENSMUSG00000009596	Placenta	633.1
Slc47a1	ENSMUSG00000010122	Kidney	64.5
Dazl	ENSMUSG00000010592	Testis	27.1
Mov10l1	ENSMUSG00000015365	Testis	26.4
Cd83	ENSMUSG00000015396	Macrophage peri LPS thio 1hrs	2432.4
Nkx2-5	ENSMUSG00000015579	Heart	26.8
Nuak1	ENSMUSG00000020032	MEF	46.2
Sycp3	ENSMUSG00000020059	Testis	19.2
Fgf22	ENSMUSG00000020327	Epidermis	1.8
Nipal4	ENSMUSG00000020411	Epidermis	56.4
Myocd	ENSMUSG00000020542	Umbilical cord	121.9
Cmpk2	ENSMUSG00000020638	Macrophage peri LPS thio 7hrs	2480.9
Rasgrf2	ENSMUSG00000021708	Hypothalamus	160.9
Ddx4	ENSMUSG00000021758	Testis	13.1
Galnt14	ENSMUSG00000024064	Kidney	11.1
Lipo1	ENSMUSG00000024766	Lacrimal gland	33.4
Rin1	ENSMUSG00000024883	Nucleus accumbens	19.1
Tdrd1	ENSMUSG00000025081	Testis	82.6
Syce1	ENSMUSG00000025480	Testis	24.4
1500015O10Rik	ENSMUSG00000026051	Osteoblast day21	2756
Tnfrsf11a	ENSMUSG00000026321	RAW 264 7	79.7
Cybrd1	ENSMUSG00000027015	Stomach	58.2
Bfsp1	ENSMUSG00000027420	Lens	5785.3
Adad1	ENSMUSG00000027719	Testis	1618.9
Sycp1	ENSMUSG00000027855	Testis	108.1
Hormad1	ENSMUSG00000028109	Not Available	Not available
Spaca1	ENSMUSG00000028264	Testis	1382.4
Slc10a4	ENSMUSG00000029219	Not Available	Not available
Figla	ENSMUSG00000030001	Lens	1.7
Mesp2	ENSMUSG00000030543	Testis	3.3
Chst15	ENSMUSG00000030930	Mast cells IgE	73.9
Cryab	ENSMUSG00000032060	Lens	157.1
Itga11	ENSMUSG00000032243	Not Available	Not available
Kank1	ENSMUSG00000032702	Epidermis	61.2
Ptprm	ENSMUSG00000033278	Lung	118.1
Cdcp1	ENSMUSG00000035498	Cornea	36.9
C530008M17Rik	ENSMUSG00000036377	Testis	50.3
Kcna6	ENSMUSG00000038077	Cerebral cortex prefrontal	26.2
Spon1	ENSMUSG00000038156	Lens	391.8
Mpp6	ENSMUSG00000038388	Mast cells IgE	3.6



Gene symbol	Ensembl gene ID	Tissue/cells with highest expression	FC over Embryonic stem cells
Tspyl5	ENSMUSG00000038984	Nucleus accumbens	98
D1Pas1	ENSMUSG00000039224	Testis	6.5
Dusp26	ENSMUSG00000039661	Dorsal root ganglia	95.3
Fkbp6	ENSMUSG00000040013	Testis	12.6
Sec1	ENSMUSG00000040364	Testis	21.3
Mael	ENSMUSG00000040629	Testis	89.3
Rsph6a	ENSMUSG00000040866	Testis	262.4
Prima1	ENSMUSG00000041669	Neuro2a	9.9
Uggt2	ENSMUSG00000042104	Testis	9.2
Abcb4	ENSMUSG00000042476	Liver	190.9
Disc1	ENSMUSG00000043051	Not Available	Not available
Npy5r	ENSMUSG00000044014	Nucleus accumbens	11
Pcsk9	ENSMUSG00000044254	mIMCD.3	20.9
Mettl24	ENSMUSG00000045555	Lung	11.5
Olig3	ENSMUSG00000045591	Thymocyte DP CD4.CD8.	2
She	ENSMUSG00000046280	Lung	19.1
Gm11554	ENSMUSG00000048294	Embryonic stem cells Bruce4+ V26, average	1
Zfp697	ENSMUSG00000050064	Kidney	40.5
Tacstd2	ENSMUSG00000051397	Epidermis	640.2
Kcnf1	ENSMUSG00000051726	Olfactory bulb	263.7
Fes	ENSMUSG00000053158	Mast cells	99.2
Sh2d4a	ENSMUSG00000053886	Stomach	106.8
Kbtbd13	ENSMUSG00000054978	Skeletal muscle	29.6
H2-Q5	ENSMUSG00000055413	T.cells CD8.	783.5
Gulp1	ENSMUSG00000056870	Retinal pigment epithelium	87.5
Unc13d	ENSMUSG00000057948	Mast cells	73.8
Kl	ENSMUSG00000058488	Kidney	1051.7
Kcng2	ENSMUSG00000059852	Heart	80.1
Nkpd1	ENSMUSG00000060621	Stomach	8.2
Rnf217	ENSMUSG00000063760	C2C12	25.6
Zar1	ENSMUSG00000063935	Ovary	4.1
Dmrtc1c2	ENSMUSG00000067561	Not Available	Not available
4930444P10Rik	ENSMUSG00000067795	Testis	249.5
Slc25a31	ENSMUSG00000069041	Testis	27.1
Atxn1l	ENSMUSG00000069895	Ciliary bodies	3.9
Hsf5	ENSMUSG00000070345	Testis	303.5
Zfp783	ENSMUSG00000072653	Retinal pigment epithelium	22.4
Klhl40	ENSMUSG00000074001	Skeletal muscle	253
Tspyl3	ENSMUSG00000074671	T.cells CD8.	38.2
Chst14	ENSMUSG00000074916	X3T3.L1	9.7
VglI3	ENSMUSG00000091243	Osteoblast day5	16.7

**Supplementary Table 5:** Accession numbers of studies used for genome-wide correlations

Histone modifications	GEO dataset accession number
H3K27ac	GSE24165
H3K4me1	GSE24165
H3K4me3	GSE24165
H3K9me3	GSE12241
Input	GSE24165

Protein	GEO dataset accession number
Atrx	GSE22162
Brd4	GSE36561
Cbx7	GSE42466
Ctcf	GSE49847
Ctr9	GSE20530
Esrrb	GSE11431
Ezh2	GSE49178
GFP	GSE11431
Hdac1	GSE27844
Hdac2	GSE27844
Jarid2	GSE19708
Kdm1a	GSE27844
Kdm2a	GSE21202
Kdm5b	GSE31968
Mtf2	GSE16526
Mycn	GSE11431
Nanog	GSE44286
Oct4	GSE44286
P300	GSE49847
Polr2a	GSE49847
Rad21	GSE33346
Rbbp5	GSE22934
Rnf2	GSE26680
Smarca4	GSE14344
Smc1a	GSE22557
Supt5h	GSE20485
Suz12	GSE48122
Taf1	GSE31270
Taf3	GSE30959
Wdr5	GSE22934

Protein	Bioproject accession number
Tcea1	PRJEB2674

**Supplementary Dataset**

Excel table Available at:  
<http://www.nature.com/ncomms/2015/150520/ncomms8155/extref/ncomms8155-s2.xlsx>

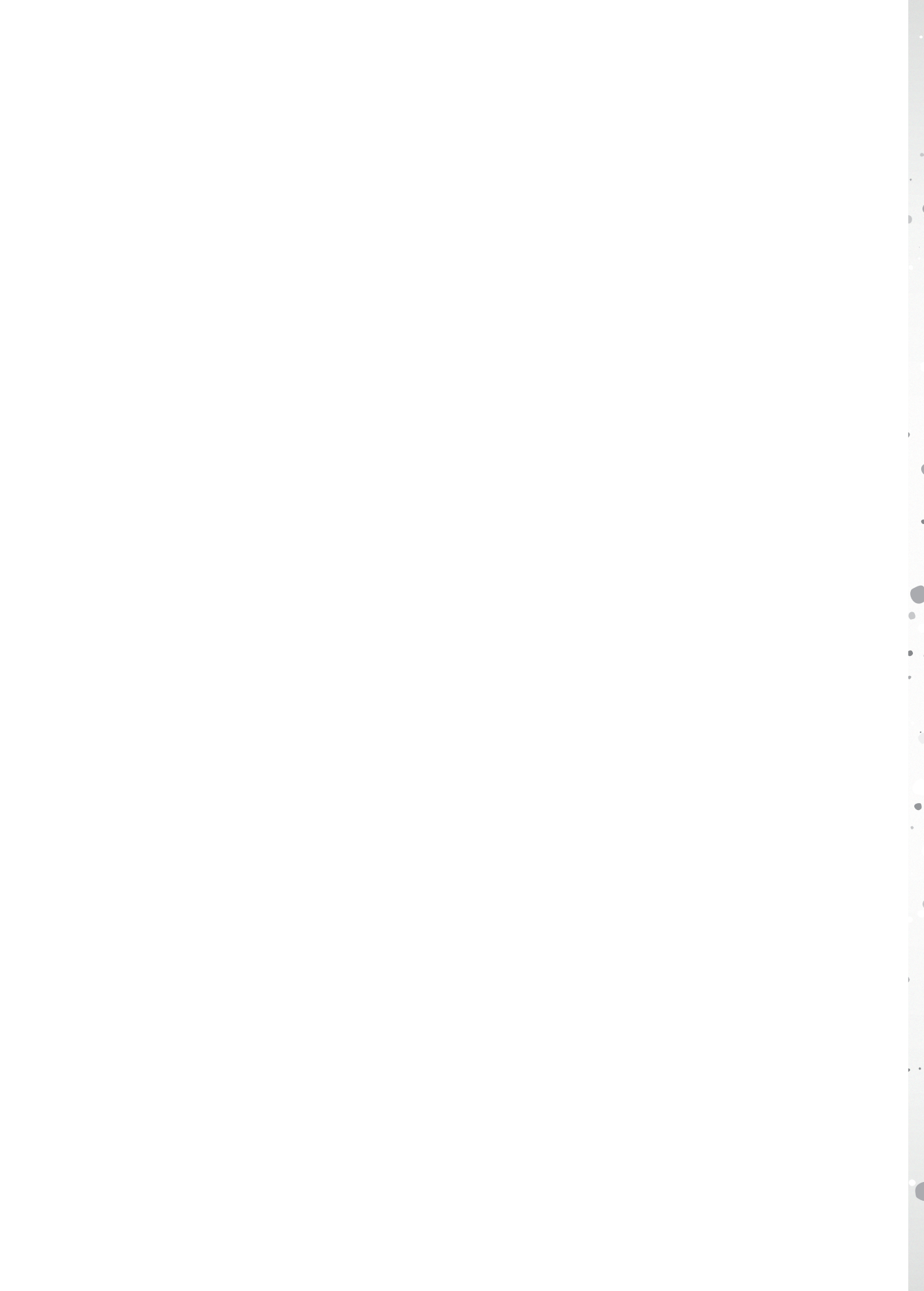


**References for pluripotency phenotype of factors identified by ChIP-MS**

Arid1a<sup>1</sup>, Ash2l<sup>2</sup>, Aurkb<sup>3</sup>, Banf1<sup>4</sup>, Bptf1<sup>5</sup>, Cbx7<sup>6</sup>, Cdc73<sup>7</sup>, Cdk9<sup>8</sup>, Chaf1b<sup>9</sup>, Chd1<sup>10</sup>, Ctbp2<sup>11</sup>, Ctr9<sup>8,12</sup>, Dmap1<sup>13</sup>, Dppa2<sup>14</sup>, Dppa4<sup>14</sup>, Dpy30<sup>15</sup>, Ep300<sup>16</sup>, Ep400<sup>13</sup>, Esrrb<sup>17</sup>, Ezh2<sup>18</sup>, Hcfc1<sup>19</sup>, Hdac1<sup>20</sup>, Ing5<sup>8</sup>, Ino80<sup>19</sup>, Jarid2<sup>21</sup>, Kdm1a<sup>22</sup>, Kdm2b<sup>23</sup>, Kdm4c<sup>24</sup>, Kdm5b<sup>25</sup>, Klf5<sup>26</sup>, L3mbtl2<sup>27</sup>, Max<sup>28</sup>, Mtf2<sup>29</sup>, Mycn<sup>30</sup>, Myst2<sup>31</sup>, Myst3<sup>31</sup>, Nacc1<sup>32</sup>, Nfrkb<sup>19</sup>, Oct4<sup>33</sup>, Ogt<sup>34</sup>, Phf20<sup>35</sup>, Phf23<sup>12</sup>, Pin1<sup>36</sup>, , Rad21<sup>37</sup>, Rbbp5<sup>15</sup>, Rnf2<sup>8,13,37</sup>, Ruvbl2<sup>13</sup>, Sin3a<sup>38</sup>, Smarca4<sup>39</sup>, Smarcb1<sup>40</sup>, Smarcc1<sup>13</sup>, Smc1a<sup>8,13</sup>, Smc6<sup>12,13</sup>, Supt4h2<sup>13</sup>, Suv39h2<sup>19</sup>, Suz12<sup>41</sup>, Taf1<sup>42</sup>, Taf2<sup>42</sup>, Taf4a<sup>42</sup>, Taf5<sup>42</sup>, Taf6<sup>42</sup>, Taf3<sup>42,43</sup>, Taf7<sup>19</sup>, Tpr<sup>13,19</sup>, Trrap<sup>13</sup>, Wdr5<sup>44</sup>, Zfp281<sup>32,45</sup>, Zscan10<sup>46</sup>.

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# Chapter 3

## An interaction network of mental disorder proteins in neural stem cells

Maaïke J. Moen<sup>1</sup>, Hieab H.H. Adams<sup>1\*</sup>,  
**Johannes H. Brandsma<sup>1\*</sup>**, Dick H.W. Dekkers<sup>2</sup>, Umut  
Akinci<sup>1</sup>, Sofia Karkampouna<sup>1†</sup>, Christel E.M. Kockx<sup>3</sup>,  
Zeliha Ozgür<sup>3</sup>, Wilfred F.J. van IJcken<sup>3</sup>, Jeroen Demmers<sup>2</sup>,  
Raymond A. Poot<sup>1</sup>

<sup>1</sup> Department of Cell Biology, Erasmus MC, Wytemaweg  
80, 3015 CN Rotterdam, The Netherlands

<sup>2</sup> Center for Proteomics, Erasmus MC, The Netherlands

<sup>3</sup> Center for Biomics, Erasmus MC, The Netherlands

**\* These authors contributed equally to this work**

<sup>†</sup> Present address: Department of Molecular Cell  
Biology, Leiden University Medical Center, Leiden, The  
Netherlands

Manuscript submitted



# Chapter 4

## Interdependency of Oct4, Sox2 and Nanog localization on the Embryonic Stem Cell Genome

**Johannes H. Brandsma**<sup>\*1</sup>, Debbie L.C. van den Berg<sup>\*1,2</sup>, Florian Halbritter<sup>3</sup>, Mike Dekker<sup>1</sup>, Zeliha Ozgür<sup>4</sup>, Christel E.M. Kockx<sup>4</sup>, Wilfred F.J. van IJcken<sup>4</sup>, Simon R. Tomlinson<sup>3</sup>, Raymond A. Poot<sup>1</sup>

<sup>1</sup> Department of Cell Biology, Erasmus MC, Wytemaweg 80, 3015 CN Rotterdam, The Netherlands

<sup>2</sup> The Francis Crick Institute, Mill Hill Laboratory, The Ridgeway, London NW7 1AA, United Kingdom

<sup>3</sup> MRC Centre for Regenerative Medicine, Institute for Stem Cell Research, School of Biological Sciences, University of Edinburgh, Edinburgh, EH9 3FF, United Kingdom

<sup>4</sup> Center for Biomics, Erasmus MC, Wytemaweg 80, 3015 CN, Rotterdam, The Netherlands

**\* These authors contributed equally to this work**

**Work in progress**



# Chapter 5

## General Discussion

# Addendum



## SUMMARY

Cell type-specific sets of transcription factors control cell identity by implementing cell type specific gene expression. Transcription factors bind to regulatory regions such as promoters and enhancers on the genome. Transcription factors regulate gene expression, often together with other transcription factors. In this thesis the genome localization and interaction partners of transcription factors are studied in Embryonic Stem Cells (ESCs) and Neural Stem Cells (NSCs), leading to an improved understanding of their molecular environment and mode of action.

Proteins that bind to regulatory regions such as promoters and enhancers have not been identified systemically. In Chapter 2 active enhancers, promoters or heterochromatin are purified from ESCs by Chromatin Immunoprecipitations for specific histone modifications and co-purified proteins are identified by mass spectrometry, a method we name ChIP-MS. ChIP-MS identifies 239 proteins that are predicted to localize to promoters or enhancers with different levels of activity, or heterochromatin. These localization predictions are verified using published genome-wide datasets that were available for 28 ChIP-MS identified proteins and indicate a high accuracy of genome localization prediction by ChIP-MS. 63 out of 239 ChIP-MS identified proteins are important for pluripotency and include Oct4, Esrrb, Klf5, Mycn and Dppa2, which are reprogramming factors for induced pluripotent stem cells. We select Dppa2 for further investigation, because ChIP-MS data suggests that Dppa2 binds to promoters with low activity, which is unusual for a pluripotency-inducing factor. By a ChIP-seq of V5 tagged Dppa2, we confirm that the ChIP-MS data correctly identifies Dppa2 as a transcription factor that binds to promoters with low activity. Dppa2 is specifically expressed in ESCs or related pluripotent stem cells and Chapter 2 shows that Dppa2 is most likely a transcriptional activator, but that nearly all Dppa2 target genes have their highest expression in other tissues than ESCs. In addition, Dppa2 binding sites show no overlap with the binding sites of pluripotency factors Oct4, Nanog and Esrrb and we therefore conclude that Dppa2 is not part of the classical pluripotency network. Chapter 5 shows that in addition to histone modification H3K4me3 and H3K27ac, Dppa2 bound promoters are also enriched for H3K27me3.

In Chapter 3 the interacting proteins of four transcription factors in mouse NSCs are identified and assembled into a network of over 200 proteins, which is the first transcription factor interaction network in a neural system and includes 68 proteins that are associated with mental disorders. The network is highly enriched for proteins encoded by evolutionarily constrained genes in human. Evolutionarily constrained genes are more frequently mutated in disease, including Autism Spectrum Disorders and this suggests that the network probably contains undiscovered Mental Disorder proteins and can be used as a discovery tool for novel mental disorder genes and to establish the molecular connections for known mental disorder proteins. Chapter 3 also shows that proteins that interact or share a network are also more likely to overlap on the genome. Tcf4 and Ascl1 have a strong overlap in human phenotypes and we show that they interact, have strong overlaps in genomic binding sites and share target genes, amongst which *Phox2b*. Activation of *PHOX2B* during development is likely affected by mutations in TCF4 and ASCL1 and this would explain the overlapping abnormal breathing phenotypes observed in Pit Hopkins syndrome and Congenital Central Hypoventilation Syndrome, that can be

caused by *tcf4* and *ascl1*, respectively.

Transcription factors can cooperatively bind to the genome and regulate gene expression. Transcription factors Oct4, Sox2 and Nanog maintain pluripotency and self-renewal of ESCs and often co-localize on the ESC genome, but cooperative binding of Sox2 and Oct4 was never investigated genome-wide. Chapter 4 shows that genome-wide binding of Oct4 and Sox2 to the ESC genome is interdependent on binding sites that contain the composite Sox2-Oct4-motif, but that binding of Oct4 or Sox2 is not reciprocally reduced upon depletion of Sox2 or Oct4, respectively. It further more shows that Oct4 directly recruits Sox2 and that Nanog is recruited via Sox2 to binding sites that contain the composite Sox2-Oct4 motif and that Sox2 and Nanog also bind the ESC genome independent Oct4 on binding sites that contain the Sox2-motif only.

Finally, Chapter 5 provides additional discussion to the previous chapters and puts the findings of these chapters in broader context and discusses new directions for further research.



## SAMENVATTING

Celtype-specifieke transcriptiefactoren beheersen de identiteit van cellen door het bewerkstellen van celtype-specifieke genexpressie. Transcriptiefactoren binden aan regulerende regio's op het genoom, zoals promotoren en enhancers. Transcriptiefactoren reguleren genexpressie, vaak in combinatie met andere transcriptiefactoren. In dit proefschrift worden de lokalisatie op het genoom en de interacterende eiwitpartners van transcriptiefactoren in embryonale stamcellen (ES cellen) en neurale stamcellen (NS cellen) nader onderzocht en dit leidt tot een verbeterd inzicht van de moleculaire omgeving en werking van transcriptiefactoren.

Eiwitten die binden aan regulerende regio's op het genoom zijn nog niet systematisch geïdentificeerd. In hoofdstuk 2 worden enhancers, promotoren en heterochromatine opgezuiverd uit ES cellen doormiddel van Chromatine-immunoprecipitaties van bepaalde histonmodificaties. Vervolgens wordt de identiteit van co-gepurificeerde eiwitten vastgesteld doormiddel van massaspectrometrie. Deze methode noemen wij ChIP-MS. Doormiddel van ChIP-MS voorspellen we voor 239 eiwitten dat deze binden aan promotoren of enhancers van verschillende activiteit of heterochromatine. Deze lokalisatievoorspellingen zijn getoetst met behulp van gepubliceerde genoombrede datasets die voor 28 eiwitten beschikbaar waren en hieruit blijkt een hoge nauwkeurigheid van de uit ChIP-MS verkregen lokalisatievoorspellingen. 63 van de 239 doormiddel van ChIP-MS geïdentificeerde eiwitten zijn belangrijk voor pluripotentie van ES cellen, inclusief Oct4, Esrrb, Klf5 en Dppa2 die behoren tot de transcriptiefactoren die het herprogrammeren van somatische cellen naar pluripotente stamcellen mogelijk maken. Dppa2 is nader onderzocht, omdat ChIP-MS erop duidt dat Dppa2 aan minder actieve promotoren bindt en dit is ongebruikelijk voor transcriptiefactoren die kunnen herprogrammeren. Doormiddel van een ChIP-seq van Dppa2 bevestigen we dat Dppa2 inderdaad aan minder actieve promotoren bindt. Dppa2 wordt specifiek in ES cellen of gerelateerde pluripotente stamcellen tot expressie gebracht. Hoofdstuk 2 laat zien dat Dppa2 waarschijnlijk een transcriptionele activator is, maar ook dat bijna alle Dppa2 gereguleerde genen het hoogst tot expressie komen in andere weefsels dan ES cellen. Ook overlappen Dppa2 bindingsplekken op het genoom niet met die van pluripotiefactoren zoals Oct4, Nanog en Esrrb, waardoor we de conclusie trekken dat Dppa2 niet deel uit maakt van het klassieke pluripotentienetwerk in ES cellen. Hoofdstuk 5 laat zien dat door Dppa2 gebonden promotoren naast verrijkt te zijn voor histonmodificaties H3K4me3 en H3K27ac, ook verrijkt zijn voor H3K27me3.

In hoofdstuk 3 worden de interacterende eiwitten van vier transcriptiefactoren in muizen NS cellen geïdentificeerd en samengevoegd tot een netwerk van meer dan 200 eiwitten. Dit is het eerste interactienetwerk van transcriptiefactoren in een neuraal celtype en bevat 68 eiwitten die in verband worden gebracht met psychische aandoeningen. Het netwerk is verrijkt voor eiwitten die gecodeerd worden door in de mens evolutionair invariante genen. Mutaties in evolutionair invariante genen leiden vaak tot ziektes, waaronder Autisme spectrum stoornissen. Dit doet vermoeden dat het beschreven netwerk waarschijnlijk nieuwe genen bevat die in verband kunnen worden gebracht met psychische aandoeningen en dat het netwerk dus als hulpmiddel kan worden gebruikt voor het ontdekken van zulke genen. Daarnaast kan het ook worden gebruikt

voor het vaststellen van moleculaire verbanden tussen bestaande genen die in verband worden gebracht met psychiatrische aandoeningen. Hoofdstuk 3 laat ook zien dat het voor eiwitten die interacteren of onderdeel uitmaken van het netwerk waarschijnlijker is dat ze ook overlap vertonen in hun bindingsplaatsen op het genoom. Tcf4 en Ascl1 hebben een grote overlap van fenotypen in de mens en hoofdstuk 3 laat zien dat ze met elkaar interacteren en hun bindingsplaatsen op het genoom een grote overlap vertonen waaronder bij het ziekte-relevante *Phox2b* gen. Mutaties in TCF4 en ASCL1 hebben waarschijnlijk invloed op de activatie van *PHOX2B* tijdens de ontwikkeling en dit zou de abnormale overlappende ademhalingsfenotypen verklaren die worden waargenomen in Pit Hopkins syndroom en aangeboren centraal hypoventilatiesyndroom en die respectievelijk veroorzaakt kunnen worden door tcf4 en ascl1.

Transcriptie factoren kunnen coöperatief binden aan het genoom en zo genexpressie gezamenlijk reguleren. Transcriptiefactoren Oct4, Sox2 en Nanog behouden de pluripotentie en zelf-vernieuwing van ES cellen en binden vaak op dezelfde bindingsplaatsen op het genoom, maar het coöperatief binden van Oct4, Sox2 en Nanog is nog nooit genoombreed onderzocht. Hoofdstuk 4 laat zien dat genoombrede binding van Oct4 and Sox2 wederzijds afhankelijk is op bindingsplaatsen die het samengestelde Sox2-Oct4 bindingsmotief bevatten, maar dat binding van Oct4 en Sox2 niet in dezelfde mate worden beïnvloed door depletie van respectievelijk Sox2 of Oct4. Het laat daarnaast zien dat Oct4 direct Sox2 rekruteert naar bindingsplaatsen met het samengestelde Sox2-Oct4 motief en dat Nanog door Sox2 mede wordt gerekruteerd. Daarnaast zijn Sox2 en Nanog ook onafhankelijk van Oct4 in staat om te binden aan bindingsplaatsen die een Sox2 motief bevatten.

Tenslotte vindt er hoofdstuk 5 aanvullende discussie plaats met betrekking tot de voorgaande hoofdstukken, plaatst het de bevindingen van deze hoofdstukken in een bredere context en bediscussieert het de mogelijkheden voor vervolgonderzoek.



## **CURRICULUM VITAE**

### **PERSONAL DETAILS**

Name                   Johannes Hendrik Brandsma  
Date of birth        29 April 1986  
Place of birth:     Heerenveen, The Netherlands

### **EDUCATION**

2011-2016         PhD program at Erasmus Medical Centre  
                      Department of Cell Biology, Erasmus Medical Centre, Rotterdam, The  
                      Netherlands

2009-2001         Master of Science in Molecular Biology & Biotechnology  
                      University of Groningen, Groningen, The Netherlands

2005-2008         Bachelor of Science in Molecular Biology  
                      University of Groningen, Groningen, The Netherlands

1998-2005         VWO Natuur & Gezondheid  
                      RSG Tromp Meesters, Steenwijk, The Netherlands

### **RESEARCH**

2011 – 2016        PhD research  
                      Department of Cell Biology, Erasmus MC, Rotterdam, The Netherlands  
                      (Prof.dr. F.G. Grosveld and Dr. R.A. Poot)

2011                MSc research project  
                      Department of Medical Biology, University Medical Centre Groningen,  
                      The Netherlands  
                      (Prof.dr. M.G. Rots)

2010                MSc research project  
                      Department of Microbiology, University of Groningen,  
                      Groningen, The Netherlands  
                      (Prof.dr. A.J.M Driessen)

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## PUBLICATIONS

Engelen E\*, **Brandsma JH\***, Moen MJ, Signorile L, Dekkers DHW, Demmers J, Kockx CEM, Ozgür Z, Van IJcken WFJ, Van den Berg DL, Poot RA. Proteins that bind regulatory regions identified by histone modification chromatin immunoprecipitations and mass spectrometry. *Nat Commun* 6, 7155 (2015)

Benjamin D et **Social Science Genetic Association Consortium**. Education-associated SNPs are enriched for brain function and disorders. (under revision)

Moen MJ, Adams HH\*, **Brandsma JH\***, Dekkers DHW, Akinci U, Karkampouna S, Kockx CEM, Ozgür Z, Van IJcken WFJ, Demmers J, Poot RA. A protein interaction network of mental disorder factors in neural stem cells. (submitted)

Wang W, Xu L, **Brandsma JH**, Wang Y, Hakim MS, Zhou X, Yin Y, Fuhler GM, Van der Laan LJW, C. Van der Woude J, Sprengers D, Metselaar HJ, Smits R, Poot RA, Peppelenbosch MP, Pan Q. Convergent Transcription of Interferon-stimulated Genes by TNF- $\alpha$  and Interferon- $\alpha$  Augments Their Antiviral Activity against HCV and HEV (under revision).

Wang W, Wang Y, Zhou X, Yin Y, Xu L, Debing Y, Carrillo EH, **Brandsma JH**, Sprengers D, Poot RA, Metselaar HJ, Smits R, Berkhout B, Neyts J, Peppelenbosch MP, Pan Q PKC $\alpha$ /AP-1 cascade directly drives transcription of interferon-stimulated genes and exerts broad antiviral activity (submitted)

(\* equal author contribution)



## PHD PORTFOLIO

Name student	Johannes H. Brandsma
Erasmus MC Department	Cell biology
Research school	Graduate School MGC
PhD Period	September 2011 – February 2016
Promoter	Frank G. Grosveld
Co-promoter	Raymond A. Poot

### PhD training

#### Courses

2011	Biochemistry and Biophysics (Rotterdam)
2011	Safely working in the laboratory (Leiden)
2012	Cell and Developmental Biology (Rotterdam)
2012	Genetics (Rotterdam)
2012	Next Generation Sequencing data analysis (Rotterdam)
2012	Analysis of microarray gene expression data using R/BioC and web tools (Rotterdam)
2013	Scripting for life science researchers (Leiden)
2013	Literature course (Rotterdam)
2014	RNA-seq workshop (Oslo, Norway)

#### Workshops, Symposia and Conferences

2011	1st Chromatin Symposium: Chromatin Changes in Differentiation and Malignancies, Giessen, Germany
2012	19th MGC PhD workshop, Dusseldorf, Germany
2012	NIRM/ISD conference Stem Cells, Development and Regulation, Amsterdam, The Netherlands
2013	3rd Winter School of the Collaborative Research Centre TRR81, Kleinwalsertal, Austria (oral presentation)
2013	SBBCD-BVCOB: Experimental models of human diseases, Luik, Belgium
2013	20th MGC PhD workshop, Luxemburg, Luxemburg (poster presentation)
2013	23rd MGC Symposium, Rotterdam, The Netherlands
2014	4th Winter School of the Collaborative Research Centre TRR81, Kleinwalsertal, Austria (oral presentation)
2014	4th IUAP DevRepair meeting, Brussel, Belgium (oral presentation)
2014	21st MGC PhD workshop, Munster, Germany (oral presentation)
2014	24th MGC Symposium, Rotterdam, The Netherlands
2014	5th IUAP DevRepair meeting, Rotterdam, The Netherlands
2015	5th Winter School of the Collaborative Research Centre TRR81, Kleinwalsertal, Austria (oral presentation)
2015	8th annual meeting of the DSSCR (oral presentation), Utrecht, The Netherlands
2015	FASEB: Transcription, Chromatin, and Epigenetics, Palm Beach, Florida, USA (poster presentation)
2015	3rd Chromatin Symposium: Chromatin Changes in Differentiation and Malignancies, Marburg, Germany (poster presentation)

#### Additional Activities

2013	Junior science Program, Erasmus MC, 3 high school students
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## DANKWOORD

Although only my name is on the cover of this thesis, it also includes the hard work of many others and I would like to use the next few pages to especially thank them and others for their hard work, support, advice, collegiality and friendship. I attribute this thesis to them.

First, I thank my co-promoter and daily supervisor Raymond. I value the many conversations we had, in which you always kept your calm, even if I lost mine. Your feedback and advice resulted in me making better decisions, whether this concerned the latest version of a figure I squeezed out of our data, experiments, a presentation, writing this thesis or career advice. I thank you for giving me the opportunity to work in your group, I learned a lot and had a good time. I wish you all the best!

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The members of the reading committee, Sjaak Philipsen, Wouter de Laat and Joost Gribnau. Thank you for being in my reading committee and for promptly reading my thesis and for the provided feedback. Sjaak, I also want to thank you for organizing the winterschool in Kleinwalsertal, which were definitely (the) highlights of my PhD! Joost, thank you (and also Raymond) for organizing the seminar series 'Frontiers in Science in the Low Countries'.

I also thank the other members of my committee, Robert Hofstra, Danny Huylebroeck and Harmen van de Werken. Danny, I always appreciated your feedback. I really enjoyed the biannual Belgium beer tastings and thank you for inviting me to speak at the 4th IUAP DevRepair meeting in Brussel. Harmen, I thank you for the many interesting discussions we had during lunch. To desperation of the others in our lunch group, this often concerned our endeavors in bioinformatics.

My Paranimfen Maaïke and Agnese, it is a honor to have you both standing at my side. Agnese, although our work together has not made it into my thesis, I really enjoyed working with you. I hope the fact that my paranimfen only have X sex chromosomes, partly compensates the whole male defense committee. Sorry Agnese, one more that adds to the shameful bias of your list. Maaïke, we went through all the trials and tribulations of the PhD together. I especially thank you for the work that is part of Chapter 3 of this thesis and for often taking the initiative in organizing social activities, such as Sinterklaas, for our lab. Maaïke, you are next!

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Absolutely essential for the realization of this thesis was learning to do data analysis of Next Generation Sequencing data. This would not have been possible without online communities such as, for example, biostars.org and seqanswers.com. Thank you to all members of these communities that ever asked or answered questions about programming or data analysis.

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Grtz,

Johan

