Role of small RNAs and chromatin in transposable element silencing during global demethylation

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THIS DISSERTATION IS SUBMITTED FOR THE DEGREE OF
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Declaration

This dissertation is the result of my own work and includes nothing which is the outcome of work done in collaboration except as declared in the Preface and specified in the text. It is not substantially the same as any that I have submitted, or, is being concurrently submitted for a degree or diploma or other qualification at the University of Cambridge or any other University or similar institution except as declared in the Preface and specified in the text. This thesis does not exceed the word limit of 60,000 words required by the University of Cambridge School of Biological Sciences.

Rebecca V. Berrens

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Table of Acknowledgments of Assistance

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- Unbiased analysis of TE classes in total RNA-seq and ChIP-seq data, antisense analysis in *Dnmt1* KO ESCs as well as the demethylation dynamics of TEs were done by Dr. Simon Andrews

- PGCLCs were made by Dr. Ferdinand von Meyenn

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- *in vivo* PGCs were isolated by Dr. Wendy Dean

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Abstract

DNA methylation entails the addition of a methyl group to the 5-carbon of the cytosine base of the DNA. This modification is important during many biological processes such as imprinting, X-chromosome inactivation, cell differentiation as well as silencing of transposable elements (TEs). DNA methylation is dynamic during early mammalian development, despite being a more static mark in somatic cells. Global hypomethylation is a hallmark of epigenetic reprogramming in mammalian primordial germ cells (PGCs), the early embryo and in naïve embryonic stem cells (ESCs). Genome integrity is crucial during early development, as the germline DNA needs to be protected for future generations. Therefore, epigenetic reprogramming presents a critical phase for TE defence since presumably alternative silencing pathways need to be employed to limit their activity. In this thesis I investigate the role of small RNAs to control TEs during global waves of DNA demethylation in cellular reprogramming, naïve pluripotency as well as early mammalian development.

Following an introduction into the research questions, in chapter 3 I investigate the mechanism of TE regulation in an in vitro model of Dnmt1 deletion in mouse ES cells to recapitulate in vivo epigenetic reprogramming. I find that certain classes of TEs become transcriptionally upregulated and subsequently resiledenced by a mechanism independent of DNA methylation. I identify ARGONAUTE 2 (AGO2) bound siRNAs as the prominent mechanism to control certain classes of TEs, while others appear to be regulated by redistribution of repressive histone modifications.

In chapter 4, I construct Dicer constitutive and conditional KO ESCs in the background of the Dnmt1 flo/flo ESCs using CRISPR-Cas9. I dissect the role of DNA methylation and
of DICER dependent small RNAs on transcriptional changes of ESCs. Additionally, I find that DICER dependent small interfering RNAs (siRNAs) re-silence transcriptionally active TE classes.

Finally, in chapter 5, I examine the role of small RNAs in TE silencing in different models of global hypomethylation in vivo and in vitro PGCs, during iPSC reprogramming and in a transition from serum to 2i culturing of mouse ESCs.
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Chapter 1

Introduction

The relationship between genotype and phenotype during development has been a question in biology since over 60 years (Waddington, 1942). The whole organism arises from one cell and shares the same genotype, however a plethora of cell types, with specialised cellular functions, are needed to generate a functional multicellular organism (reviewed in Allis and Jenuwein, 2016). Cellular identities are generated through tight regulation of transcriptional programmes. Transcription factor (TF) networks direct cell differentiation and lineage commitment, however “epigenetic modifications” are necessary to reinforce cell-fate decision to prevent reversion into preceding cellular states (reviewed in Allis and Jenuwein, 2016). Epigenetic modifications can be inherited from one cell to the next without any variation in the DNA sequence (reviewed in Allis and Jenuwein, 2016). This propagation of “epigenetic landscapes” during cell divisions rather than genetic inheritance creates cellular identities and drives development (Waddington, 1957). Epigenetic modifications such as DNA methylation, histone modifications, higher order chromatin structure, chromatin interacting factors and non-coding RNAs build the epigenome of a cell. Those epigenetic modifications play together as well as autonomously from each other to form the epigenome. Disruption of proper modifications can lead to a disease phenotype, loss of cellular identity and incomplete early development (reviewed in Allis and Jenuwein, 2016).
1.1 Chromatin

The higher order chromatin structure is governed by an interplay of all epigenetic modifications. DNA is tightly packed in macromolecules, termed chromatin (Flemming, 1882). This organisation allows the compaction of the human two-metre long DNA molecule into the nucleus. The identification of a phosphorus-rich acid, termed nucleic acid as well as the proteins which bind to the DNA are the main building blocks of DNA (Miescher, 1871, Kossel, 1884).

In 1944 it was found that not the proteins but the nucleic acid contains the genetic information (Avery et al., 1944). The molecular basis of genetic inheritance was resolved when the structure of the DNA as a double-helix was identified (Watson and Crick, 1953).

The genome is largely organised in two distinct chromatin states: (1) heterochromatin is densely packed in mitosis (Heitz, 1928) is a silenced chromatin environment, while (2) euchromatin is the diffusely organised chromatin during interphase. Heterochromatin has been referred to as the inactive state of the DNA (Heitz, 1929, Heitz, 1932). This hypothesis was confirmed by the observation that a gene close to heterochromatin was silenced and could be activated by decondensation of the DNA region in Drosophila melanogaster (Schultz, 1936). Heterochromatin was further subdivided into (1) constitutive heterochromatin, of constantly condensed regions in all cell types, typically associated with centromeres and telomeres, and (2) facultative heterochromatin that contains regions that switch between active and inactive states during development, like the X chromosome in females as well as gene imprinting (Brown, 1966). To the present day this genome organisation is valid as heterochromatin is found to be typically a gene-poor and transcriptionally silenced part of the genome, while euchromatin is gene-rich and allows transcriptional activity. The DNA is organised in these two distinct chromatin states by post-translational modification of the histone proteins and by modification of the DNA itself (reviewed in Allis and Jenuwein, 2016). Additionally, non-coding RNAs are involved in organisation of the chromatin; the long non-coding RNA Xist has been found to be invaluable for the silencing of the X chromosome (Galupa and Heard, 2015). Furthermore, spatial organisation of the chromatin in the
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nucleus is not random. Local and long-range interaction of sequences in the DNA is
highly controlled and the DNA can be divided into discrete topological domains of
100kb to 1Mb in length (Dixon et al., 2012, Nora et al., 2012) and genes that sit in do-
 mains which are bound by the nuclear lamina have been found to be transcriptionally
silent (Guelen et al., 2008, Pickersgill et al., 2006).

Even though hundreds of chromatin modifications have been identified to date, the
modifications associate in preferential combinations with each other and hence give
rise to a small number of chromatin states (Rando, 2012, van Steensel, 2011). Indeed,
four chromatin states were identified in Arabidopsis thaliana, five states were found in
Drosophila melanogaster and the human chromatin organises in six chromatin com-
partments (Rao et al., 2014).

1.2 DNA methylation

DNA methylation entails the only covalent modification of the DNA itself and presents
the most studied epigenetic mark. DNA methylation is the addition of a methyl group
to the 5-carbon of the cytosine base in the DNA (5mC) (Holliday and Pugh, 1975,
Riggs, 1975). In mammals 4-6% of cytosines are methylated in somatic cells. DNA
methylation (5mC) plays a key role in developmental processes in mammals. This epi-
genetic modification controls genome regulation, genome stability, X-inactivation, ge-
nomic imprinting as well as transposon silencing (reviewed in Reik and Walter, 2001).
In the mammalian genome, 5mC is mostly found in cytosine residues of CpG dinu-
cleotides, while methylation of other DNA bases has been found at low levels (Guo
et al., 2014). CpG dinucleotides are mostly found in CpG islands (CGIs), while the
majority of the genome is depleted of CpG sites (Bird et al., 1985, reviewed in Goll and
Bestor, 2005). CGIs are mostly associated with promoters and remain unmethylated
while CpGs randomly distributed in the genome are methylated (Meissner et al., 2008,
reviewed in Suzuki and Bird, 2008).
1.2.1 Propagation of DNA methylation

DNA methylation is established and maintained by DNA methyltransferases (DNMTs). DNMTs are the enzymes that catalyse the addition of a methyl group to the DNA using S-adenosylmethionine as a donor for an activated methyl group (reviewed in Bestor, 2000).

Three major DNMTs exist to maintain DNA methylation in the mammalian genome. The establishment of de novo DNA methylation is accomplished by the two active DNA methyltransferases DNMT3A and DNMT3B as well as their catalytically inactive partner DNMT3L. However, during every cell division the newly synthesised DNA strand has to be remethylated to maintain symmetrical DNA methylation patterns - both strands of a CpG dinucleotide are methylated - by the DNA maintenance methyltransferase DNMT1 (Figure 1.1). The DNA methylation can be lost passively, through dilution during replication in the absence of the maintenance machinery, or actively, through enzymatic demethylation.

Another DNMT enzyme, DNMT2 has been found to methylate RNAs, specifically tRNAs, instead of DNA (Goll and Bestor, 2005).

Furthermore, DNMT3C has been discovered to be involved in transposable element silencing of specifically young TE classes in the mouse genome (Barau et al., 2016).

Figure 1.1: DNA methylation. DNMT3A and DNMT3B work together with DNMT3L to establish de novo DNA methylation. DNMT1 together with E3 ubiquitin-protein ligase (UHRF1) is responsible for the maintenance of DNA methylation after DNA replication (blue). If the maintenance methylation is impaired DNA methylation gets lost through passive dilution (yellow). Figure from Wu and Zhang, 2014.
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**De novo** methylation

*De novo* methylation is known as the establishment of methylation patterns in mammals. DNA *de novo* methylation is established by the methyltransferases DNMT3A and DNMT3B ([Okano et al., 1999; Li et al., 1992](#)). Both enzymes can bind hemimethylated as well as non-methylated DNA ([Okano et al., 1999](#)), but unlike DNMT1, they do not preferentially bind hemimethylated DNA. DNMT3A and B both contain a PWWP motif, a PHD-related domain called ADD (Atrx-Dnmt3-Dnmt3l) and a catalytic domain ([Okano et al., 1998](#)).

Deletion of *Dnmt3a* and *Dnmt3b* during early mammalian development showed that these two enzymes overlap in their functions. The knock out of *Dnmt3b* leads to demethylation of pericentric satellite DNA and severe developmental phenotypes which result in embryonic lethality from mid-gestation onwards ([Okano et al., 1999](#)). Knock out of *Dnmt3a* leads to offspring, with prominent developmental defects which result in their death a few weeks after birth ([Okano et al., 1999](#)). However, the double knock-out of *Dnmt3a* and *Dnmt3b* results in severe developmental effects and lethality from embryonic day (E)11.5 onwards ([Okano et al., 1999](#)).

Loss of *Dnmt3a* and *Dnmt3b* in mouse embryonic stem cells (ESCs) leads to gradual loss of DNA methylation ([Chen et al., 2003](#)). This hypomethylation happens over progressive cell divisions, therefore the remaining methylation is attributed to the activity of the DNA maintenance machinery ([von Meyenn et al., 2016](#)).

DNMT3L (DNMT3-like) is a truncated form of DNMT3A and DNMT3B without any enzymatic activity ([Aapola et al., 2001](#)). The enzymatic activity of those two methyltransferases is enhanced through binding of their catalytically inactive partner DNMT3L ([Chen et al., 2005; Suetake et al., 2004; Gowher et al., 2005](#)). DNMT3L plays an important role in *de novo* methylation of transposable elements during spermatogenesis. Knock out of *Dnmt3L* results in viable offspring with a severe germline defect, as well as a lack of imprinting ([Bourc’his et al., 2001](#)).
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Maintenance methylation

DNMT1 is the enzyme that maintains DNA methylation after DNA replication during S phase of the cell cycle. DNMT1 was the first DNA methyltransferase discovered in mammals (Gruenbaum et al., 1982, Bestor and Ingram, 1983). DNMT1 contains a replication foci-targeting domain (RFD) a DNA-binding CxxC domain, two Bromo-adjacent homology domains (BAH) as well as the catalytic domain, which is composed of a target recognition domain and the core catalytic motifs (Song et al., 2011, Takeshita et al., 2011). The maintenance methylase DNMT1 is targeted to the genome by the E3 ubiquitin-protein ligase (UHRF1) (previously known as NP95) which recognises hemimethylated DNA at replication foci via its SRA-domain (Sharif et al., 2007, Bo-stick et al., 2007). The methyltransferase copies the 5mC mark onto the replicated daughter strand (Bestor et al., 1988, reviewed in Bestor, 2000). The high affinity of DNMT1 to hemimethylated DNA, by binding with the RFD domain to PCNA proteins and UHRF1, has been shown in vivo (Fatemi et al., 2001). Furthermore, the CXXC domain of DNMT1 binds to unmethylated CpGs and thereby inhibits the access of the catalytic domain to those regions (Song et al., 2011).

Dnmt1 as well as Uhrf1 -null mice have reduced DNA methylation levels and die in utero (Li et al., 1992, Sharif et al., 2007). The knockout of both Dnmt1 as well as Uhrf1 in mouse embryonic stem cells (ESCs) leads to a rapid DNA demethylation of the genome. Even though self-renewal of those cells is maintained, they fail to differentiate (Lei et al., 1996, Li et al., 1992). The remaining DNA methylation level of 20%, after knockout of the maintenance methylation machinery (Meissner et al., 2005) can be attributed to the activity of the de novo methyltransferase DNMT3A and DNMT3B (Lei et al., 1996).

1.2.2 Loss of DNA methylation

The enzymes of DNA methylation are DNMT1, DNMT3a and DNMT3b, however the erasure of DNA methylation has been under debate for many years. DNA methylation
can be lost by passive dilution through the impairment of the methylation maintenance machinery - DNMT1 or UHRF1. However, rapid demethylation dynamics that cannot be explained by passive dilution have also been detected (Guo et al., 2011, Mayer et al., 2000). The discovery of Ten-Eleven-Translocation 1 (TET1) was a ground breaking finding because this enzyme enables active demethylation in mammals (Tahiliani et al., 2009). TET1, TET2 and TET3 comprise the TET dioxygenase family. These enzymes oxidise methylcytosine to 5-hydroxymethylcytosine (5hmC) and subsequently 5-formylcytosine (5fC) and 5-carboxycytosine (5caC) (Ito et al., 2011, He et al., 2011). Due to their role in DNA demethylation in plants, DNA glycosylases have been proposed to play a similar role in DNA demethylation in mammals (reviewed in Zhu, 2009). The thymine DNA glycosylase (TDG) has been found to remove 5fC and 5caC from the genome (He et al., 2011, Maiti and Drohat, 2011). Active demethylation by TET and TDG have been found during in vivo early embryonic development (Gu et al., 2011, Seisenberger et al., 2012, reviewed in Wossidlo et al., 2011). TDG can also work together with active deamination enzymes such as AID and APOBEC to lead to DNA demethylation (Kohli and Zhang, 2013). Deamination has been found to play a role during global demethylation processes in primordial germ cell (PGC) development (Popp et al., 2010), in the zygote (Santos et al., 2013) as well as during reprogramming of induced pluripotent stem cells (iPSCs) (Kumar et al., 2013).

1.2.3 Role of DNA methylation

The functional role of DNA methylation has long been affiliated to transcriptional repression (Razin and Riggs, 1980), however whole genome bisulfite sequencing has shown that the relationship is not unambiguously defined.

I will highlight four different ways in which DNA methylation could interfere with transcription, which are active in the genome, among others: (1) promoter methylation, (2) gene body methylation, (3) preventing the binding of transcription factors to promoters, and (4) indirect control of promoter enhancer interactions (reviewed in Klose and Bird, 2006).
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(1) DNA methylation at the Transcription start side (TSS) is implicated with gene repression. Promoter CGI methylation plays a profound role in imprinted genes and X-chromosome inactivation during development. Imprinted genes are monoallelically expressed depending on their parent of origin (reviewed in John, 2017). Imprinting is controlled by differential methylation at imprinted control regions (ICR). The methylation level of the ICR can influence single or multiple genes simultaneously (Bartolomei, 2009, Ferguson-Smith, 2011). Nevertheless, most of the CGI promoters in the genome are unmethylated through the binding of chromatin modifiers, transcription factors as well as the activity of TET enzymes (Ooi et al., 2007, Stadler et al., 2011, Williams et al., 2011). However, the plethora of unmethylated TSSs in the genome does not necessarily lead to transcriptional activation of the associated genes. Rather, they may be silenced through post transcriptional histone modifications.

(2) Although DNA methylation at the TSS is a repressive signal, gene body methylation correlates with activation of transcription (Lister et al., 2009). During transcriptional elongation histone 3 lysine 36 trimethylation (H3K36me3) recruits DNMT3B which in turn deposits DNA methylation over gene bodies (Baubec et al., 2015). Gene bodies are mostly depleted of CpGs, though intragenic CGIs exist and show tissue specific methylation patterns (Maunakea et al., 2010). Altogether, genome-wide studies showed that methylation of the promoter regions leads to transcriptional regulation and that the local CpG density plays an important role (Meissner et al., 2008, Weber et al., 2007). While methylation of high CpG-density promoters (HCP) is strongly associated with transcriptional silencing, low CpG-density promoters (LCP) do not have this property as they are mostly methylated without any effect on the transcriptional regulation (Borgel et al., 2010). Correlation between transcriptional inactivation and methylation is best described at intermediate CpG-density promoters (ICP), like germline specific promoters (Meissner et al., 2008).

(3) The binding of transcription factors (TFs) can depend on the methylation level and therefore affect transcription. The methylation of a single CpG in the binding site of the TF YY1 abolishes its binding (Kim et al., 2003). Nevertheless, the zinc finger proteins (ZFPs) with a Krueppel-Associated box (KRAB) domain solely bind to
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methylated DNA motifs (Quenneville et al., 2012). The binding of the majority of TFs seems to be unaffected by the methylation state of the binding site (Thurman et al., 2012).

(4) Another way to control gene expression is through distal regulatory regions called enhancers. Enhancers are CpG-poor, and low methylated regions (LMRs) (Stadler et al., 2011). ChIP-seq experiments of TET enzymes showed their enrichment over LMRs (Chen et al., 2013b; Williams et al., 2011). Additionally, TET enzymatic products - 5hmC and 5fC - have been found in high abundance at enhancers in ESCs (Ficz et al., 2011, Shen et al., 2013, Williams et al., 2011). Their methylation status could influence gene expression and studies of global chromatin structures will help to shed light on this part of gene regulation controlled by DNA methylation.

However, no clear picture of a role of DNA methylation in regulating gene expression can yet be drawn (Deaton et al., 2011) and further investigation has to follow to understand the regulatory function of this epigenetic mark in the genome.

1.2.4 Chromatin modifications

In addition to higher order chromatin structure as well as DNA methylation, chromatin modifications are contributing to the epigenome of a cell.

The basic unit of chromatin is the nucleosome composed of 147 bp DNA that is wound around an octamer of two histone dimers H2A, H2B and H3, H4, respectively (Kornberg, 1974, Luger et al., 1997). The nucleosomes are connected by a linker DNA of about 10-60bp which is often bound by the histone 1 (H1) linker histone (Thoma et al., 1979). A fiber of DNA under the electron microscope appears like “beads-on-a-string” first observed by Olins and Olins, 1974. The chromatin is further packaged and condensed. Chromatin binding proteins, like heterochromatin protein 1 (HP1), methyl-CpG-binding protein 2 (MeCP2) and polycomp group proteins (PcG) play a role in additional chromatin compaction (reviewed in McBryant et al., 2006).

Gene expression can be regulated through the orchestrated modification of histone tails. The posttranslational modification of histones is often found in the N-terminal histone
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tails, which represent 25-30% of the histone protein (reviewed in Zhang et al., 2015). The ability of chromatin to regulate gene expression was originally hypothesised over 60 years ago, when the first histone acetylase (HAT) was identified (Allfrey et al., 1964). Since then, a plethora of histone modifications have been found including acetylation, phosphorylation, methylation, and ubiquitination (reviewed in Arnaudo and Garcia, 2013). Histone modifications can also recruit non-histone proteins to the chromatin to control gene expression. Furthermore, histone protein variants can replace the canonical histone proteins and influence chromatin environments. H3 is replaced by CENP-A in centromeres as well as by H3.3 in actively transcribed genes (reviewed in Biterge and Schneider, 2014).

Altogether, post-translational histone modifications, and histone core organisation can change chromatin states and impact compaction/decompaction and thereby alter promoter accessibility and control gene expression (reviewed in Bannister and Kouzarides, 2011).

1.2.5 Post-translational histone modifications

Histone acetylation

Since the discovery of the first HAT, histone acetylation has been associated with transcriptional activation. The lack of histone acetylation on the inactive X chromosome was a first evidence suggesting that this post-translational modification (PTM) is marking transcriptional activity (Jeppesen and Turner, 1993). The identification of the histone acetyltransferase (HAT) as well as histone deacetylases (HDACs) confirmed the link between histone acetylation and transcription (Brownell, 1996, Kuo, 1996, Taunton et al., 1996). Histone acetylation neutralises the positive charge of the lysine residue and inhibits the binding of histone tails to the negatively charged DNA (Hong et al., 1993). This results in a chromatin decondensation and facilitates the transcription machinery to access the DNA (Norton et al., 1989, Lee et al., 1993a).

The activity of HAT and HDAC presented the first confirmation that histone modifications play a role in gene expression with HAT being the “writer” of the histone mark
and HDAC the “eraser” (Marmorstein, 2001, reviewed in Thiagalingam et al., 2003).

Histone acetylation recruits bromodomain proteins to the chromatin among these are transcriptional activators (TAFs), methyltransferases (MLLs), SWI/SNF chromatin remodellers as well as helicases (Smarca), which are commonly referred to as chromatin “readers” (reviewed in Muller et al., 2011).

Histone methylation

While histone acetylation is regarded as an activating mark, histone methylation is a more complex PTM, as it can lead to transcriptional repression as well as activation. Histone methylation is a highly specific histone modification. By mono-, di- or trimethylation of a single lysine on one of the histone is sufficient for transcriptional repression or activation (reviewed in Bannister and Kouzarides, 2011). The SET (Su(var), Enhancer of zeste, and Trtorax) domain of histone lysine methyltransferases (KMTs) contains the enzymatic activity which catalyses the transfer of the methyl group donor SAM to the amino group of a lysine residue (reviewed in Bannister and Kouzarides, 2011). KMTs are the “writers” of histone methylation (reviewed in Bannister and Kouzarides, 2011).

Histone methylation is erased by histone demethylases (KDMs) (reviewed in Bannister and Kouzarides, 2011). KDMs either have an LSD1 or an JumonjiC (JmjC)-domain to remove methyl groups from modified histones (reviewed in Bannister and Kouzarides, 2011). LSD1 demethylates the active histone mark, lysine 4 on histone 3 (H3K4), which leads to a transcriptional inhibition (Shi, 2004). LSD1 as well as several JmjC-domain containing demethylases can antagonise the repressive H3K9 methylation and thereby activate transcription (Metzger et al., 2005, Fodor, 2006, Klose, 2006). Histone methylation can attract proteins with chromo, tudor or MBT domains, which present the “readers” of this modification.
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Additional histone modifications

Next to histone methylation and acetylation, further PTMs have been studied. Histone ubiquitination is a histone modification which is, unlike the other histone modifications, mostly found on the C-terminal tail of histones. Histone ubiquitination has been found at lower levels than other modifications. Histone 2A lysine 119 ubiquitination (H2AK119ub1) is involved in gene silencing through the interaction with polycomb group proteins (Wang et al., 2004). Furthermore, histone phosphorylation is takes place during DNA damage response and phosphoruated H2A is largely enriched at sites of DNA breakage (reviewed in Allis and Jenuwein, 2016).

1.2.6 Role of histone modifications

Histone modifications are important regulators of gene transcription. In mammals, inactive genes are marked with methylation of H3K9, histone 3 lysine 27 (H3K27) and histone 4 lysine 20 (H4K20), albeit active genes are marked with high levels of acetylation and trimethylalation of histone 3 lysine 4 (H3K4me3), histone 3 lysine 36 (H3K36me3) as well as histone 3 lysine 79 (H3K79me3) (Noma et al., 2001, reviewed in Kouzarides, 2007, Barski, 2007). H3K4me3 can be found in promoter regions and H3K36me3 is found in transcribed regions (Liang et al., 2004, Pokholok et al., 2005). Genome-wide analysis of distribution of histone marks allowed for prediction of transcriptional states of promoters and enhancers (Heintzman, 2007). H3K4me1 and H3K27ac mark cell type specific enhancers while another class of enhancers has been identified by H3K122ac but lack off H3K27ac (Rada-Iglesias, 2011, Pradeepa et al., 2016). In ESCs the Nucleosome Remodelling and Deacetylation (NuRD) complex has been shown to directly regulate the expression of pluripotency genes by removal of H3K4me1 and H3K27ac from active enhancers (Reynolds et al., 2012).

In ESCs, genes important for developmental are termed “poised” and carry simultaneously activating H3K4me3 and repressing H3K27me3/H2AK119ub1 “bivalent” histone marks (Bernstein et al., 2006a, Azuara, 2006). RNA polII binding to those regions showed that they express low levels of RNA (Stock et al., 2007). This poised state
allows rapid lineage specification upon differentiation of ESCs (Bernstein et al., 2006a, Azuara, 2006). Bivalent chromatin domains are not restricted to ESCs but also exist in other cell types (Mikkelsen et al., 2007, Barski, 2007).

In the present study, I concentrated on the effect of global DNA demethylation on H3K9me2/3 and H3K27me3 genome wide, therefore I will expand on those specific modifications.

**H3K9 methylation**

H3K9 methylation is generally known as a repressive histone mark and several H3K9 methyltransferases are known to repress specific regions of the genome. H3K9 methylation silences pericentric repeats, centromeres as well as telomeres (Bannister, 2001). Pericentric repeats are a combination of large arrays of AT-rich repeats interspersed with transposable elements (TEs), also called major satellites. Centromeres represent shorter repeated DNA fragments termed minor satellites (Guenatri et al., 2004). H3K9 can be mono- di- or trimethylated and KMTs are responsible for the specific methylation level. Additionally, H3K9 modifications have been found to interact with HP1 as well as KRAB-associated protein 1 (KAP1) proteins. HP1 is a highly evolutionary conserved 25 kDa protein with a chromo- and a chromoshadow-domains. HP1 binds the chromatin with its chromodomain (Bannister, 2001, Lachner et al., 2001, Nielsen et al., 2002). With the chromoshadow-domain HP1 recruits additional chromatin binding proteins and can also attract additional HP1 molecules, resulting in a block of HP1 that inhibits transcription (Smallwood et al., 2007). HP1 is only able to bind H3K9 methylation *in vivo* if it is associated with KMTs (Chin et al., 2007, Sripathy et al., 2006, Stewart et al., 2005). The interaction of HP1 with H3K9me3 is also important for the recruitment of HDACs (Schultz et al., 2001, Stewart et al., 2005)

Even though eight KMTs for H3K9me3 have been identified in mammals only three have been attributed to specific functions in the genome.

The KMT Suv39h1/2 is important for H3K9me3 at pericentric repeats. *Suv391/2* dKO leads to a loss of HP1 binding to pericentric repeats as well as chromosomal instability (Peters et al., 2001, Peters et al., 2003, Rea, 2000, Lachner et al., 2001). Nevertheless,
Suv39h1/2 KO mice are viable and heterochromatic foci are still forming without these KTM (Pinheiro et al., 2012, Peters et al., 2001).

G9a/GLP are the main methyltransferases responsible for non-pericentric H3K9 mono- and dimethylation. G9a/GLP form heterodimers and single KO of either G9a or GLP lead to reduction of H3K9me2 and H3K9me1 (Tachibana et al., 2001, Tachibana et al., 2005). Additionally, G9a and GLP KO mice are embryonic lethal with an inability to complete meiosis in both male and female (Tachibana et al., 2002, Tachibana et al., 2005, Tachibana et al., 2007). Both G9a and GLP can read and write H3K9me1/2, which leads to a self-enforcing H3K9me2 spreading in the genome of up to 5Mb (Wen et al., 2009). H3K9me2 is highly enriched at genes bound by the nuclear lamina, and G9a KO in ESCs leads to an upregulation of those genes (Guelen et al., 2008, Yokochi et al., 2009).

ESET/SETDB1 has been studied extensively, as this KMT is responsible for most of the H3K9me3 in non-pericentric regions (Schultz et al., 2001, Wang et al., 2003). ESET is targeted to H3K9me3 by KAP1. KAP1 contains a RBCC domain, a PHD-bromo domain and an HP1 binding domain. The RBCC domain can bind to KRAB-ZFP (Friedman et al., 1996). KRAB-ZFP recruits KAP1 to specific genomic loci, KAP1 recruits ESET and NuRD and stabilises the interaction of HP1 with H3K9me3 and KAP1 (Ryan et al., 1999, Schultz et al., 2002). Kap1 and Eset KO mice die shortly after gestation (Cammas et al., 2006, Dodge et al., 2004). Mouse ESCs with Kap1 or Eset KO are not viable (Dodge et al., 2004, Rowe et al., 2010). It as been shown that KRAB-ZFPs bind to TEs specifically; and therefore, the KAP1 and ESET mediated H3K9me3 deposition seems to play a major role in TE silencing (Matsui et al., 2010, Najafabadi et al., 2015, Rowe et al., 2010).

H3K27 methylation

H3K27 methylation works in close relationship with protein complexes called polycomb repressive complexes (PRC) to induce gene silencing.

PRC1 and PRC2 are the two polycomb complexes that exist in mammals. PRC1 is
mostly involved in chromatin compaction and catalyses H2AK119ub, while PRC2 is important for methylation of H3K27 (Shao et al., 1999, Cao, 2002). The PRC2 complex is composed of the four core proteins EZH1 or 2, SUZ12, EED and RbAp46 or 48 (Cao, 2002), with the first three presenting the minimal requirement for PRC2 enzymatic activity. PRC2 is further stabilised by the interacting factor AEBP2 and JARID2 or PCL are proteins involved in recruiting PRC2 to the chromatin (reviewed in Margueron and Reinberg, 2010). EZH1 and EZH2 contain a SET domain for H3K27 di- or trimethylation. However, EZH1 depletion does not affect global H3K27me2/3 in the genome (Margueron et al., 2008), but it is impossible to generate ESCs depleted of Ezh2 (O’Carroll et al., 2001). The interaction of EED with H3K27me3 is essential for methylation of H3K27 (Margueron, 2009). PRC2 binds the chromatin through the SUZ12 and RbAp24/48 proteins (Nekrasov et al., 2005). H3K27me3 can cover large domains of about 100kb to maintain X-chromosome inactivation, as well as the highly coordinated expression of the Hox gene cluster during embryo development (Silva et al., 2003, Boyer, 2006, Bracken et al., 2006). In those cases, PRC2 is recruited to the chromatin by long non-coding RNAs (lncRNAs). *Xist* lncRNA recruits PRC2 *in cis* to the X chromosome (Plath, 2003), while lncRNA *HOTAIR*, transcribed in the *HoxC* locus, tethers PRC2 *in trans* to the *HoxD* locus (Rinn et al., 2007).

Small domains of H3K27me3 are enriched in promoter regions of ESCs. H3K27me3 promoter methylation is found in 10% of genes in ESCs. Developmental regulatory genes are mostly enriched for H3K27me3 (Boyer, 2006, Marks et al., 2012, Zhao et al., 2007). PRC2 and H3K27me3 generate the chromatin environment to allow gene expression by recruiting PRC1 to genes that need to be silenced.

PRC1 is a multiprotein complex and a key component of the complex are the E3 ubiquitin ligases RING1a and RING1b. The E3 ubiquitin ligases catalyse the H2AK119 ubiquitination. While *Ring1b* KO is embryonic lethal in mice, *Ring1a* KO mice survive (del Mar Lorente et al., 2000, Voncken et al., 2003). Another member of the PRC1 complex is one of the six PCGF protein, giving rise to six PRC1 complexes. (Gao et al., 2012). Additionally, either CBX or RYBP proteins are members of the PRC1 complex (Wang et al., 2010). CBX proteins contain a chromodomain and can therefore
bind H3K27me3, which has been confirmed in vitro (Bernstein et al., 2006b, Fischle et al., 2003). The interaction of CBX with H3K27me3 recruits PRC1 to PRC2 targets to drive chromatin compaction and thereby gene repression. In ESCs depleted of Cbx proteins PRC2 targets are largely expressed (Leeb et al., 2010). The exact mechanism by which PRC1 leads to transcriptional repression is still under investigation. Ubiquitination is one form of silencing developmental genes and while Hox genes are repressed independently of this histone modification, Cbx KO leads to an activation of the Hox genes (Endoh et al., 2012, Morey et al., 2012).

RYBP-PRC1 complexes can bind to chromatin independent of PRC2 and H3K27me3, however in this thesis I am concentrating on the H3K27me3 modification, therefore I will not describe this role of PRC1.

1.3 Histone modifications and DNA methylation

Post-translational histone modifications and DNA methylation can be attracted to the same chromatin loci and work together to control gene expression.

Negative feedback loops enforce repressive chromatin, for example ESET catalyses H3K9me3 at pericentric repeats, which in turn recruits HP1 to the chromatin. HP1 recruits the de novo methyltransferases to the chromatin. The DNA methylation is then recognised by the DNA binding protein MECP2. MECP2 has a methyl binding domain (MBD), which is also able to recruit SUV39 histone methylase (Meehan et al., 1992). This closes the circle through further H3K9me3 deposition and heterochromatic persistence at pericentric repeats (reviewed in Zhang et al., 2015). While the knock out of the H3K9me3 methyltransferase Swi39h1/2 does not lead to a complete global loss of DNA methylation, hypomethylation was found in heterochromatic repeats (Lehnertz et al., 2003). Cooperative effects of DNA methylation and histone modifications can be induced by proteins which multiple binding domains.

UHRF1 can bind to hemimethylated DNA, via a SET and RING-associated (SRA) domain but this protein can also bind to methylated lysine 9 of histone H3 (H3K9me2/3),
via a tudor domain (Arita et al., 2008, Avvakumov et al., 2008). Conditional double knock out studies of \textit{Dnmt1} as well as \textit{Uhrf1} suggested an interaction between UHRF1 and the histone 3 lysine 9 (H3K9) methyltransferase ESET. In a hemimethylated genome, due to deletion of \textit{Dnmt1} the binding affinity of UHRF1 to hemimethylated DNA confounds DNA binding of ESET and thereby H3K9me3 deposition (Sharif et al., 2016) (Figure 1.2).

**Figure 1.2: Maintenance DNA methylation and H3K9me3.** The DNMT1 protein has an RFD domain (brown), an CXXC domain (yellow), a BAH1 domain (pink), a BAH2 domain (blue) and the enzymatic domain (green). UHRF1 has a UBL domain (brown), a tandem tudor domain (light blue) a PHD domain (pink) an SRA domain (dark blue) and a RING domain (orange). UHRF1 binds with its SRA domain to hemimethylated CG sites (hmCG) and directs DNMT1 to the sites by binding it with the RFD domain. UHRF1 can also bind H3K9me3 with its RING domain. Figure from Du et al., 2015.

Additionally, KMTs can directly interact with DNMTs to repress transcription. G9a/GLP interacts directly with the maintenance methylase DNMT1 (Estève et al., 2006) and is needed for \textit{de novo} DNA methylation of retrotransposons in ESCs (Dong et al., 2008).

Histone modifications can recruit DNMTs to repress transcription. Both DNMT3A and DNMT3B can bind to H3K36me3 via their PWWP domain (Baubec et al., 2015, Dhayalan et al., 2010). This interaction has been shown in yeast, where DNMT3B is recruited to sites of transcriptional elongation by H3K36me3 (Morselli et al., 2015).
However, histone modifications can also antagonise DNA methylation and the same region. The enzymatically inactive isoform DNMT3L can bind to the tails of histone H3 through the ADD domain. This interaction is disrupted if the lysine 4 (K4) at the N terminus of histone H3 is methylated (Ooi et al., 2007). Therefore, DNA methylation is found at regions depleted of H3K4 methylation. This occupation of different regions in the genome is visible at CGIs, where low levels of DNA methylation but high levels of H3K4 methylation is found (Weber et al., 2007) (Figure 1.3).

Figure 1.3: De novo DNA methylation and H3K4 methylation. The DNMT3A protein has a PWWP domain (blue), an ADD domain (brown) and the enzymatic domain (green). DNMT3L has a ADD domain (brown), a an enzymatically inactive DNA methyltransferase domain (pink). DNMT3A can bind to unmethylated DNA and lead to de novo methylation which is enhanced by binding to DNMT3L. DNMT3L can also bind H3K4 in its unmethylated state. Figure from Du et al., 2015.

1.4 Non-coding RNAs

Non-coding RNAs are also involved in shaping the epigenome of a cell.

The genome comprises only 1.5% protein coding genes, about three-quarters of the genome produces non-coding RNAs (ncRNAs) (Djebali et al., 2012). The size of ncR-
NAs ranges from 21-24nt of small RNAs to 2.3 to 17.2 kilobases of long non-coding RNAs (lncRNAs) (reviewed in Costa, 2007, Fu, 2014).

1.4.1 lncRNAs

lncRNAs can have cis or trans regulatory functions on gene activity (Wang et al., 2014, Lu et al., 2014, Fort et al., 2014, reviewed in Wang and Chang, 2011, Melé and Rinn, 2016). The first lncRNA of the H19 locus was discovered in 1991 (Brannan et al., 1990). lncRNAs can work as scaffolds for higher-order chromatin organisation (reviewed in Nagano and Fraser, 2011). The lncRNA X-inactive specific transcript (Xist) plays an important role in X-chromosome inactivation. The inactivation of one X chromosome in female embryos is done to obtain dosage compensation. The silencing is accomplished by the binding of the noncoding RNA Xist with a preceding histone remodelling by loss of the active histone marks H3K4 di- and trimethylation concurrent with the gain of the repressive histone marks H3K9me2 and H3K27me3, and ubiquitination of H2A (Brown et al., 1991), reviewed in Galupa and Heard, 2015).

1.4.2 small RNAs

Since the discovery in 1998, RNA interference (RNAi) has been found to be important in gene regulation, heterochromatin formation as well as regulation of transposable elements (TEs) (reviewed in Malone and Hannon, 2009). RNAi mediated mechanisms are defined as mechanisms in which small RNAs (20-30 nucleotides long) serve for recognition and regulation of its target RNA. Small RNAs are furthermore defined by association with ARGONAUTE (AGO) family proteins. At least three different classes of small RNAs exist in mammals: endogenous short-interfering RNAs (endosiRNAs), microRNAs (miRNAs) and PIWI-interacting RNAs (piRNAs). Small RNAs play a role in post-transcriptional gene silencing by direct base-pairing with its targets. However, small RNAs can also work as transcriptional silencers (TGS) by directing chromatin modifications to their targets (Figure 1.4) (reviewed in Slotkin and Martienssen, 19...
1.4.3 miRNAs

miRNAs have been found to play a role in almost all cellular processes ranging from development to oncogenesis through post-transcriptional gene regulation (Wightman et al., 1993, reviewed in Ameres and Zamore, 2013). miRNAs are the most abundant class of small RNA in somatic tissues in mammals (miRNAs) (reviewed in Ha and Kim, 2014). The first miRNA *lin-4* was discovered in *Caenorhabditis elegans*, (Lee et al., 1993b). Over 1500 miRNAs are expressed in the mouse genome (Griffiths-Jones, 2004) and it has been shown that most mRNAs contain evolutionary conserved miRNA target sites (Friedman et al., 2009).

![Small RNA pathway in mice](image)

**Figure 1.4: Small RNA pathway in mice.** Different small RNA pathways are active in the mouse genome. The miRNAs (pink) and endosiRNAs (esiRNAs) (pink) are dependent on DICER, while piRNAs (blue) are dependent on MIWI, MILI and MIWI2. Figure from Iwasaki et al., 2015.
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1.4. Non-coding RNAs

Post-transcriptional regulation by miRNAs

miRNAs control gene expression by post-transcriptional gene silencing (PTGS). PTGS of mRNAs is accomplished by partial or full complementary binding of the miRNA and AGOs are acting as the effector proteins. By recruitment of GW182 proteins AGOs induce translational repression, mRNA deadenylation, mRNA decapping and mRNA decay by 5’-to-3’ degradation (reviewed in Ameres and Zamore, 2013). The 5’ pairing of nucleotide 2-7 of the miRNA to the target mRNA is conserved and called the miRNA “seed” (Bartel, 2009; Grimson et al., 2007). Additionally to the seed pairing a 3’ pairing at nucleotide 13-16 enhances target repression further repression is accomplished by a 5’ adenosine (Grimson et al., 2007; Wee et al., 2012; Lewis et al., 2005; Nielsen et al., 2007). miRNAs mostly bind to the target mRNAs in their 3’ UTR (Grimson et al., 2007; Forman and Coller, 2010). Additionally, the general accessibility of the miRNA binding site in the mRNA sequence is of importance. However, Watson-Crick complementarity of the seed region to the target mRNA suffices for post-transcriptional gene repression, therefore half of all protein-coding genes in mammals can be regulated by this class of small RNAs (reviewed in Ameres and Zamore, 2013). In plants miRNAs which bind with perfect complementarity to their targets trigger mRNA cleavage (Tang et al., 2003; Llave et al., 2002), albeit mammalian miRNAs mostly block translation by lack of complementarity to the target mRNA (Shin et al., 2010; Yekta et al., 2004; Davis et al., 2005). Furthermore, out of the four mammalian AGO proteins, three are catalytically inactive and only AGO2 has endonuclease functions. Ribosome profiling showed, that miRNAs are mostly translationally repressed through mRNA decay by miRNAs in mammals (Baek et al., 2008). Nevertheless about 15% of miRNAs block translational initiation (Guo et al., 2010).

Although miRNAs are described as translational repressors, binding of certain miRNAs to the target mRNA leads to activation of transcription. miR128 in the mammalian brain controls the nonsense-mediated decay machinery and hence increases mRNA
1.4. Non-coding RNAs

In general, miRNAs seem to not completely disrupt gene expression but rather “tunes” their targets (Bartel, 2009). Furthermore, miRNAs can serve as diagnostic tools and potentially interfere with disease outcomes as was shown for hepatitis C infection (reviewed in Shrivastava et al., 2015).

miRNA biogenesis

Several key proteins are involved in the biogenesis of miRNAs. The miRNAs are encoded as miRNA genes in the genome. RNA polymerase II (Pol II) is responsible for the transcription of miRNA genes into primary miRNA (pri-miRNA). The pri-miRNA is capped with a 7-methylguanosine cap (m7Gppp) and a 3′ poly(A) tail. The miRNA transcripts folds into an integral hairpin structure which incorporates the miRNA sequence. The RNase III endonuclease DROSHA cleaves the pri-miRNA into the 60nt long stemloop precursor miRNA (pre-miRNA) (Lee et al., 2003, Denli et al., 2004, Gregory et al., 2004). DROSHA can cleave more than one pre-miRNA out of the pri-miRNA stem loop.

However, DROSHA is part of the multisubunit complex called the microprocessor complex. The microprocessor complex can be made of a large complex or the smaller heterodimer complex comprised of DROSHA bound to the dsRNA-binding protein DiGeorge syndrome critical region gene 8 (DGCR8). DGCR8 can bind dsRNA and therefore directs DROSHA to the pri-miRNA (Landthaler et al., 2004, Han et al., 2004).
Gregory et al., 2004, Denli et al., 2004). Animal pri-miRNAs are processed in the nucleus (Lee et al., 2003, Billy et al., 2001, Provost et al., 2002, Lee et al., 2002). The pre-miRNA is then exported from the nucleus by the RAN-GTP protein EXPORTIN-5 (EXP-5) (Bohsack et al., 2004, Lund et al., 2004, Okada et al., 2009). Dicer is a cytoplasmic ribonuclease which generates 22 nucleotide miRNA/miRNA* duplex from the pre-miRNA (Bernstein et al., 2001, Hutvágner et al., 2001, Grishok et al., 2001). Even though both miRNAs are produced as duplexes only one of the strands is finally loaded into the RNA-induced silencing complex (RISC) (Hammond et al., 1999, Tuschl et al., 1999). ARGONAUTE (AGO) proteins are making up the core of the RISC. AGOs have two domains, the PIWI and the PAZ domain. The PIWI domain encodes an RNase H protein, and harbours the nuclease activity, while the PAZ domain binds to the 3′ overhang of two nucleotides of an siRNA (Lingel et al., 2003, Song et al., 2003, Song et al., 2004) (Figure 1.5).

1.4.4 Endogenous siRNAs

Endogenous siRNAs (endosiRNAs) present another class of small RNAs involved in PTGS. EndosiRNAs were first discovered in Drosophila melanogaster (Czech et al., 2008, Kawamura et al., 2008, Okamura et al., 2008). EndosiRNAs are around 21nt long and originate from convergent sense/antisense transcription of genomic loci and can be formed through inter- or intramolecular interactions. TEs can give rise to the double-stranded RNA (dsRNA) that feeds into the endosiRNA pathway.

Post-transcriptional gene silencing by endosiRNAs

EndosiRNAs can control transcription of both protein-coding genes and TEs by PTGS. Mutations in proteins important for the endosiRNA pathway in flies does not impair fertility and leads to viable offspring (Förstemann et al., 2005, Okamura et al., 2008, Lee et al., 2004, Liu et al., 2003), even though TEs are silenced by endosiRNAs in vitro.
The idea arose that piRNAs and endosiRNAs might cooperate to compensate for each other’s loss of function (reviewed in Malone and Hannon, 2009). In mammals, the endosiRNA pathway is able to take over the role of the piRNA pathway in the female germline (Tam et al., 2008, Watanabe et al., 2008). The MT transposon family is highly upregulated in Dicer KO in oocytes (Murchison et al., 2007), although Piwi KO does not lead to transcriptional upregulation of this TE family. dsRNA feeding into the siRNA pathway can be generated through sense transcription of protein-coding transcripts while the antisense species can arise from pseudogene transcripts in the mouse oocyte (Tam et al., 2008, Watanabe et al., 2006).

**endosiRNA biogenesis**

The generation of endosiRNAs is similar to miRNA biogenesis. However the two small RNAs originate from different sources. While the precursor of miRNAs is encoded in the DNA, endosiRNAs arise from double stranded transcription. DICER is an essential player in the biogenesis of siRNAs (Grishok et al., 2001, Hutvágner et al., 2001). Dicer processes double stranded RNA (dsRNA) into 21-23nt long duplexes of siRNAs (Bernstein et al., 2001). siRNAs are produced as duplexes and only one strand is loaded into the RNA-induced silencing complex (RISC). The so-called guide strand recognises its target mRNAs by Watson-Crick base pairing. Perfect pairing leads to endonucleolytic cleavage of the target mRNA which is also referred to as slicing (Figure 1.6).

AGO proteins are making up the core of the RISC. Four AGO proteins (AGO 1-4) exist in mammals and all of them can bind small RNAs to trigger translational repression. Yet, in mammals AGO2 is the only endonuclease that possesses the enzymatic activity to cleave its target RNA (Meister et al., 2004, Liu et al., 2004, Song et al., 2004, Okamura et al., 2004, Rand et al., 2005, Matranga et al., 2005).
Figure 1.6: endosiRNA biogenesis. endosiRNAs are generated from dsRNA, which is generated either in cis by sense (black) /antisense (grey) transcription in the same locus, by generation of a hairpin structure of the transcript or in trans through sense and antisense strand derived from different genomic locations. dsRNA is then loaded into DICER in the cytoplasm (white) and processed on the siRNA guide and passenger stand. The guide strand is normally loaded into the siRISC complex with the key component AGO2 (red).

Noncanonical biogenesis of endosiRNAs and miRNAs

Next to the canonical pathways, miRNAs and endosiRNAs can also be generated through non-canonical pathways.

One of those pathways is DGCR8 and DROSHA-independent, while another one is TUTase-dependent and a third has been described to be DICER-independent (reviewed in Ha and Kim, 2014). A non-canonical biosynthetic pathway has been shown to take place for mirtrons, miRNA precursors consisting of spliced introns. As mirtrons resemble premiRNAs, they do not undergo Drosha dependent cleavage. Instead, they are directly exported to the cytoplasm where they are further processed by Dicer (reviewed in Babiarz and Blelloch, 2009). Furthermore, DICER and DGCR8-independent miRNAs and siRNAs have been found in mouse ESCs (Babiarz et al., 2008).

1.4.5 piRNAs

miRNAs and endosiRNAs can be found in differentiated as well as undifferentiated cell types, however piRNAs are confined to the germline as well as during cancer progression. piRNAs are transcribed from a long intergenic region called the piRNA cluster. They are longer than the other small RNAs (24-30nt) and have a 2′-O-methyl-group at their 3′ end. Furthermore, other than miRNAs and siRNAs, piRNAs are Dicer-independent and mainly control transposable elements (TEs). P-element-induced-
1.4. Non-coding RNAs

wimpy-testes (PIWI) proteins were originally identified in *Drosophila melanogaster* and are essential for germline development (reviewed in Thomson and Lin, 2009, Cox et al., 1998). In *Mus musculus* three PIWI proteins - MIWI, MIWI2 and MILI - exist. However, the three proteins are expressed at different stages during early development (Figure 1.7). Interestingly, mutations in PIWI proteins specifically affect the male germline, while the female germline remains unaffected (Carmell et al., 2007, Aravin et al., 2007; Kuramochi-Miyagawa et al., 2008, Aravin et al., 2008). MILI or MIWI2 knock down lead to activation of TEs of long interspersed elements (LINE) as well as the long terminal repeat class (LTR) retrotransposons (Carmell et al., 2007, Kuramochi-Miyagawa et al., 2008, Aravin et al., 2006, Girard et al., 2006, Grivna et al., 2006).

Transcriptional and post-transcriptional gene silencing of piRNAs

piRNAs can regulate transcription on the level of transcriptional gene silencing (TGS) or PTGS. While prepauchtene piRNAs mostly function through TGS by *de novo* methylation of their targets, pachytene piRNAs cleave their target and therefore work through PTGS. The main difference between the two classes is that prepauchtene piRNAs originate from TEs and are processed by MILI and MIWI2 in the so called gonocyte stages (just after cell cycle arrest). Pachytene piRNAs, on the contrary, mostly originate from piRNA clusters in unannotated regions of the genome and are bound by MILI and MIWI in round spermatids at post-natal day 14 (P14) (Aravin et al., 2007, Kuramochi-Miyagawa et al., 2008, Aravin et al., 2008, Girard et al., 2006, Lau et al., 2006). MILI and MIWI are endonucleases and therefore necessary for cleavage of the target RNA. Only in the presence of MIWI and MILI, LINE-1 (L1) TEs are cleaved in mouse testes (Reuter et al., 2011, De Fazio et al., 2011). Silencing of intracisternal A particles (IAPs) and L1 elements by *de novo* DNA methylation is abolished in *Mili* and *Miwi2* KO in the male germline (Aravin et al., 2007, Kuramochi-Miyagawa et al., 2008, Aravin et al., 2008, Bourc’his and Bestor, 2004).

The deletion *Maelstrom* - important for piRNA biogenesis - has been shown to induce spermiogenic arrest (Castañeda et al., 2014). Accordingly, PIWI protein and piRNAs

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are thought to be invaluable for genome integrity and fertility of the organism. Nevertheless, L1 transcriptional activity during spermatogenesis by deletion of a pachytene piRNA cluster does not effect germline development (Xu et al., 2008)
Thus, the regulation of TEs by PIWI-piRNAs and the effect on gametogenesis are still not unambiguously defined (reviewed in Iwasaki et al., 2015).

piRNA biogenesis

Figure 1.7: piRNA biogenesis. Mouse PIWI proteins are expressed at different stages during spermatogenesis. MILI is expressed throughout spermatogenesis (orange), MIWI2 is expressed just until P6.5 (violet) and MIWI is expressed from P14 onwards (green). piRNAs are classified into prepachytene piRNAs (red) and pachytene piRNAs (blue). MILI and MIWI2 bind to prepachytene piRNA clusters. MILI performs to ping-pong cycle to silence targets in PTGS and produce piRNAs that associates with MIWI2. MIWI2 then localises to the nucleus and leads to de novo methylation of its targets. Pachytene piRNAs regulate their targets through PTGS. Figure from Iwasaki et al., 2015.
piRNAs are generated by two distinct pathways. The primary piRNA pathway and
the secondary piRNA ping-pong cycle for amplification.
Primary piRNAs are known to have a bias for a 5′ uridine (U) overhang, and secondary
piRNAs bind with 10 nt complementarity to primary piRNAs. Additionally, secondary
piRNAs are biased towards the sense strand and have an adenosine base at the tenth
nucleotide.
Transcription of primary piRNA clusters gives rise to long single-stranded RNA tran-
scripts, which harbour several truncated TE sequences as well as transposons in an-
tisense direction to their coding strands (Brennecke et al., 2007, Prud’homme et al.,
1995, Zanni et al., 2013). Those long transcripts are exported from the nucleus and
cleaved by a cytoplasmic endonuclease MitoPLD in the mouse (Huang et al., 2011,
Watanabe et al., 2011). The resulting precursor RNAs are loaded into nuage, a struc-
ture found between aggregated mitochondria. A key component of a nuage is the RNA
helicase MOV10L1 (Frost et al., 2010, Zheng et al., 2010). A 3′-5′ endonuclease needed
for the generation of the mature piRNAs is still not identified. Deletion of Armitage
(Drosophila homologue of mouse MOV10L1) leads to reduction of germline piRNAs
(Malone et al., 2009).
A methylatransferase, not identified in mouse, then transfers the 2′-O-methyl group to
the 3′ ends of the piRNAs and the piRNAs are loaded into the PIWI-piRISC complex
(Saito et al., 2007, Horwich et al., 2007). The Piwi-piRISC complexes are transported
from the cytoplasm to the nucleus (Sienski et al., 2012).
MILI-piRNA complexes are associated with primary piRNAs and only secondary piRN-
As are associated with MIWI2. Those secondary piRNAs bound by MIWI2 are
imported to the nucleus and direct TGS by targeted DNA methylation (Aravin et al.,
2008, Kuramochi-Miyagawa et al., 2008, Reuter et al., 2011, De Fazio et al., 2011).
The details regarding ping-pong amplification of piRNAs in the mouse are still being
debated. MILI is localised in the cytoplasm and can only induce a single ping-pong
cycle, but it is thought that MILI-piRNAs can amplify piRNAs by a pathway yet to
be identified (De Fazio et al., 2011) (Figure 1.7).
1.5 Epigenetic reprogramming

Although DNA methylation is a covalent modification of the DNA and regarded as a stable epigenetic modification, during epigenetic reprogramming a global methylation erasure followed by \textit{de novo} methylation occurs in two waves in every generation during mammalian development. The first wave happens shortly after fertilisation in the early embryo and a second wave takes place in the primordial germ cells (PGCs) which are the embryonic progenitors for sperm or oocytes (reviewed in Hackett and Surani, 2013, Seisenberger et al., 2013b, Seisenberger et al., 2013a) (Figure 1.8).

![Figure 1.8: Epigenetic reprogramming in mammalian development. Schematic representation of the genome-wide waves of DNA methylation and demethylation in the mouse genome in early mammalian development. Image from (Lee et al., 2014b).](image)

1.5.1 Zygote reprogramming

In order to form the totipotent zygote, epigenetic modifications are being reprogrammed. The zygote has the potential to generate an entire organism with multiple cell types. Those cell types all have a different developmental potency and the canalisation of cells into different cell fates is defined by epigenetic features (Surani et al., 2007). Global demethylation of both female (oocyte) and male (sperm) pronuclei occurs just
after fertilisation. Nuclear exclusion of DNMT1 leads to a passive loss of DNA methylation in the female pronucleus (Howell et al., 2001). However, global demethylation in the highly methylated male pronucleus happens by active demethylation through the oxidation of 5mC to 5hmC and 5fC, shown by immunofluorescence (IF) (Gu et al., 2011, Inoue and Zhang, 2011, Iqbal et al., 2011, Santos et al., 2013, Wossidlo et al., 2011). This oxidation is most likely accomplished by TET3, which is the only TET enzyme expressed in the zygote and knockdown as well as knockout of Tet3 inhibits global demethylation of the male pronucleus (Gu et al., 2011, Peat et al., 2014).

Active base excision repair (BER) pathways acting solely on 5mC are also involved in demethylation of the paternal pronucleus, as small molecule inhibition of BER results in global hypermethylation (Hajkova et al., 2010, Santos et al., 2013).

The difference in global hypomethylation in the two pronuclei is of interest. The maternal factor Stella presents a prominent explanation. Stella protects the maternal genome and paternal imprints with the deposition of H3K9me2 (Hajkova et al., 2010, Nakamura et al., 2012). Deletion of Stella from the zygote leads to incomplete preimplantation development and demethylation of the maternal genome and paternal imprinted regions (Nakamura et al., 2006).

Another way to protect imprint demethylation is through the action of the zinc-finger protein 57 (ZFP57) and KAP1 by recruitment of the maintenance methylase DNMT (Quenneville et al., 2011, Li et al., 2008). The hypomethylated zygote has to acquire DNA methylation from blastocyst stage onwards, in order to separate trophoeoderm from the inner cell mass (ICM) (Santos et al., 2002). To reinforce lineage commitment gene promoter methylation is highly apparent at developmental genes (Borgel et al., 2010). In the transition from blastocyst to egg cylinder stage further remethylation is occurring, finally giving rise to a hypermethylated genome in the epiblast stage (Smith et al., 2012).

### 1.5.2 PGC reprogramming

In mice, PGCs are the precursors of the future gametes and in order to gain developmental potency, PGCs also need to go through a wave of global demethylation.
Furthermore, the imprinted regions have to be reset (Hajkova et al., 2002). PGCs can be found in the epiblast of the developing embryo around embryonic day (E)7.25 (Seisenberger et al., 2012, Ginsburg et al., 1990) and share their epigenetic modifications with the epiblast. At E8 PGCs are first localised at the base of the allantois before they reach the developing gonads (E11.5) by migrating through the hindgut (Anderson et al., 2000, Molyneaux et al., 2001). Methylation reprogramming in PGCs occurs in two phases during their development, an early reprogramming event during their migration phase (E7.5 - E8.5) and a late reprogramming phase in their gonadal stage (E8.5 - E10.5) (Guibert et al., 2012, Seisenberger et al., 2012, Kagiwada et al., 2013, Hackett et al., 2013, Seki et al., 2005).

The early phase of reprogramming affects all genomic features and is accomplished by passive demethylation due to impairment of the maintenance machinery by transcriptional regulation of Uhrf1 (Seisenberger et al., 2012, Hackett et al., 2013). An additional exclusion of UHRF1 from the nucleus results in a great reduction of DNMT1 activity and results in global hypomethylation. Recent in vitro studies accompanied by modelling have shown that global demethylation pathways are largely reliant on the maintenance methylation machinery, DNMT1 and UHRF1 (von Meyenn et al., 2016). However, ICR as well as CGI promoters of germ cell and meiosis specific genes further to CGIs associated with the inactive X chromosome stay methylated until the gonadal stage (Seisenberger et al., 2012, Hackett et al., 2013). The mechanism of their resistance to demethylation is not unambiguously defined.

The second phase of demethylation is accomplished through active demethylation, as the dynamics of demethylation do not allow for a passive pathway (Bagci and Fisher, 2013). During epigenetic reprogramming 5hmC levels are increasing from E10.5 onwards (Hackett et al., 2013, Yamaguchi et al., 2013), suggesting the involvement of TET enzymes. Yet, the knockout of Tet1 in PGCs resulted in minor changes of methylation levels in E13.5 PGCs (Yamaguchi et al., 2012). Furthermore, Tet1/Tet2 double KO did not abolish PGC differentiation (Vincent et al., 2013). Thus, active demethylation by TET enzymes might be important at very targeted loci, such as ICRs as well as CGIs of gametic genes (Hackett et al., 2013, Vincent et al., 2013, Yamaguchi et al., 2013). Another mechanism that might play a role in active demethylation are
activation-induced cytidine deaminase (AID) and TDG. Aid KO results in hypermethylated PGCs at stage E13.5 (Popp et al., 2010) but does not lead to any fertility problems (Kagiwada et al., 2013).

Epigenetic reprogramming does also encompass chromatin remodelling. While H3K9me3 levels stay unchanged over the course of global demethylation in PGCs, H3K9me2 levels follow global demethylation dynamics and get erased from E8 to E13.5 and global levels of H3K27me3 increase (Hajkova et al., 2008, Seki et al., 2007).

In mice, following the wave of demethylation of PGCs to about 5-8% of 5mC at E13.5, de novo methylation begins in male germ cells during embryonic days E14.5 and E16.5 and goes on until prospermatogonia phase (Davis et al., 1999, Davis et al., 2000, Kato et al., 2007). The de novo methylation rises from 7% at E13.5 to 55% at E16.5 before the beginning of meiosis and reaches a final methylation level of 90% in sperm (Kobayashi et al., 2013, Pastor et al., 2014). The remethylation is accomplished by the de novo methyltransferases DNMT3A, DNMT3B and DNMT3L (Kato et al., 2007, Bourc’his et al., 2001, Bourc’his and Bestor, 2004, Kaneda et al., 2004, Ciccone et al., 2009). TE remethylation is accomplished with the help of piRNAs (Bourc’his and Bestor, 2004). Most piRNAs in the foetus are complementary to TEs and in Dnmt3L, Mili, Miwi2 KO LINEs, ERVs are transcriptionally active resulting in meiotic catastrophe (Aravin et al., 2007, Bourc’his and Bestor, 2004). Also conditional KO of DNMT3a or DNMT3L leads to TE reactivation, as well as meiotic arrest and finally germ cell apoptosis (Bourc’his and Bestor, 2004, Kaneda et al., 2004). Male and female remethylation happens with different kinetics, as the final methylation level of the two sexes is different (Smith et al., 2012, Popp et al., 2010, Smallwood et al., 2011, Kobayashi et al., 2013) (reviewed in Seisenberger et al., 2013b, Seisenberger et al., 2013a). - The different kinetics of remethylation of the two germlines can be attributed to the activity of Dnmt3L in the male germline (Bourc’his et al., 2001, Bourc’his and Bestor, 2004, Aravin et al., 2007, Aravin et al., 2008). Additionally, Dnmt3C has been found to play a role in methylating especially young elements in the male germline in mice (Barau et al., 2016).

Although the whole genome becomes hypomethylated during epigenetic reprogram-
Chapter 1. Introduction

1.6 Transposable elements (TEs)

Transposable elements (TEs) are mobile genomic regions that are able to change their location within the genome by transposition. TEs were discovered in plants in the 1950s \((\text{McCLINTOCK}, 1951)\). The presence of mobile elements was also found in bacteria and in \textit{Drosophila melanogaster}, where the cross of a wild strain of fruit flies to a lab strain led to an increase of chromosomal rearrangements, recombination effects and sterility \((\text{Shapiro}, 1969, \text{Kidwell et al.}, 1977)\). The incompatibility of the two fruit fly genomes, termed “hybrid dysgenesis”, could be explained by the activity of the DNA transposon called P element TEs \((\text{Rubin et al.}, 1982\text{ reviewed in Majumdar and Rio, 2015})\). The P element had expanded in genomes of wild fruit flies, while the genome of the lab strain had been protected \((\text{Anxolabéhère et al.}, 1988)\).

The non-coding genome was termed “junk” DNA in the 1970s \((\text{Ohno}, 1972)\). Due to the accumulation of large amounts of non-coding DNA the term “selfish DNA” arose and TEs were called “parasites of the genome” \((\text{Doolittle and Sapienza}, 1980, \text{Orgel and Crick}, 1980)\).

Whole genome sequencing allowed a closer examination of the TE contribution to the genome, and it was found that TEs comprise \(\sim70\%\) of the mammalian genome \((\text{Lander et al.}, 2001, \text{Chinwalla et al.}, 2002, \text{de Koning et al.}, 2011)\).
While TEs are still regarded as the main danger for genome integrity, their regulatory function and role in evolution are being progressively discovered (Fedoroff, 2012b, Chuong et al., 2016, Elbarbary et al., 2016, Ecco et al., 2016).

The mouse genome is comprised of ancestral TEs as well as newly acquired TEs. The mouse genome (2.7 Gb) is about 0.5 Gb smaller than the human genome (3.2 Gb). The lineage separation of rodent and primates happened 65 to 100 Myrs ago (Chinwalla et al., 2002). As the mutation rate in the mouse is very high only 5% of ancestral TEs persisted in the mouse genome. 44% of the mouse genome are TEs and 85% of those are mouse specific. In general, mouse repeats are younger than human TEs, most of them have integrated in the last 25 Myrs and still contain around 1000 active LINEs and ERVs, while most of the human TEs are inert (Chinwalla et al., 2002).

1.6.1 TE families

TEs can be divided in to two different classes, according to their mechanism of transposition: Class I transposons, which are also called retrotransposons, work in a “copy and paste” mechanism as an RNA intermediate is first produced and the cDNA is then reinserted in another place of the genome (reviewed in Smit, 1996) and class II transposons are called DNA transposons and can directly move by a “cut and paste” mechanism. While the major class are class I transposons with about 47% of the genome, only about 3% of the genome is comprised of DNA transposons (Lander et al., 2001). Class I transposons, or retrotransposons are further classified into endogenous retroviruses (ERVs) as well as long interspersed elements (LINEs) and short interspersed elements (SINEs) (Okamura and Nakai, 2008).

DNA transposons

The different coding sequences of DNA transposons define their mechanism of transposition. DNA transposons are mostly 1.5-5kb in long. They encode a transposasase gene which is flanked by tandem inverted repeats (TIRs). The transposasase allows
DNA transposons to move by excising and reintegrating in a new genomic location (Muñoz-López and García-Pérez, 2010). The integration leads to a target site duplication (TSDs) which is the addition of a short sequence of DNA, about 4-8bp, at either site of the DNA transposon. All DNA transposons transpose by this mechanism with the exception of Helitrons and Mavericks that replicate through a single stranded DNA intermediate.

Activity of DNA transposons is linked to transcription, and retrotransposition leads to a gain of a transposon copy while excision leads to DNA double-stranded breaks that need repair. The break can be repaired by homologous recombination and results in the regeneration of the transposon copy, or non homologous end joining (NHEJ), with only the TSDs remaining. These TSD sequences in the genome are called transposon “footprints” (reviewed in Feschotte and Pritham, 2007). DNA transposons can be classified by their different transposases into eleven superfamilies (Wicker et al., 2007) - with the main ones being Tc1/mariner, hAT, piggyback, Maverick and Helitrons. Tc1/mariner existing in most eukaryotic species (Capy et al., 1996, Plasterk et al., 1999). Miniature inverted repeat transposable elements (MITEs) have lost their transposase but have accumulated in many genomes as they use transposases encoded elsewhere in the genome for mobilisation (reviewed in Feschotte et al., 2002).

**Endogenous retroviruses**

LTR-retrotransposons make up approximately 8-10% of mouse genomes (Chinwalla et al., 2002). While they were thought to be relics of primary infectious retroviruses which lost their Env gene and became bona fide endogenous genomic sequence (reviewed in Smit, 1996), LTR transposons have been found to acquire Env genes and become infection competent during evolutionary time (Malik et al., 2000). Due to this, another name for LTR retrotransposons is endogenous retroviruses or short ERVs. Interestingly, not one single colonisation event, but rather several separate events, can be attributed to the accumulation of the highly polymorphic class of 400 ERV families in the genome (Maksakova et al., 2013, Zhang et al., 2008, reviewed in Goodier and Kazazian, 2008).
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The ERV life-cycle is characterised by an RNA intermediate. In mammals, ERV transposons are a few hundred bp to 7kb long. All full-length ERVs encode at least two genes, *Gag* and *Pol*. While the *Gag* gene encodes a polymorphic capsid protein, the *Pol* gene encodes three proteins: a protease (PR), a reverse transcriptase (RT) with an upstream RNAseH domain and an integrase (IN) (Wicker et al., 2007). Transcription of the ERV mRNA is accomplished by RNA pol II which binds to the promoter located in the 5’LTR. Both *Gag* and *Pol* genes are transcribed into one mRNA. Depending on the host the mRNA is either (1) translated by fusion into one open reading frame (ORF) that is subsequently cleaved to generate two open reading frames or (2) ribosomal frame shifting enables the translation of both proteins (Craig, 2015).

Following the translation of the ERV proteins, the reverse-transcription initiates downstream of the 5’LTR at the primer binding site (PBS). Host tRNAs are used as primers resulting in double stranded cDNA (Lund et al., 2000). The integration of ERVs also creates TSDs (Mager and Stoye, 2015). A lot of ERVs lack ORFs and therefore are unable to replicate. However, ERV retrotransposition is solely dependent on the LTR sequences as well as the PBS sequence for reverse transcription, therefore ERVs can mobilise *in trans*. In mammals, the VL30 and MaLR families are so called non-autonomous ERV elements (reviewed in Stocking and Kozak, 2008). Furthermore, ERV copies can be fixed through homologous recombination of the two LTRs as a solitary LTRs in the genome (Mager and Goodchild, 1989, reviewed in Sverdlov, 1998). Those LTRs can be identified in the genome by their TSDs, and a polyadenylation signal (reviewed in Smit, 1996).

Several families of ERVs have been classified. Phylogenetic analysis of the RT, as well as the organisation of RT and IN domain in the *Pol* gene, allowed the classification of ERVs into *gypsy*-like or *copia*-like transposons (Xiong and Eickbush, 1990, reviewed in Havecker et al., 2004). It has long been thought that in vertebrates, ERVs have likely originated from the endogenisation of a retrovirus (reviewed in Boeke and Stoye, 1997), however phylogenetic analyses of LTR transposons suggests that these elements can acquire *Env* genes which allow them to gain infectious potential (Malik et al., 2000). Depending on their original retrotransposon endogenisation, ERV transposons in mammals can be further subdivided into three subclasses. The Class I ERVs are
most closely related to gamma- and epsilonretroviruses. Class II ERVs are related to lentiviruses as well as alpha- beta- and deltaretroviruses, while spumaviruses share highest sequence homology with class III ERVs (Gifford et al., 2005; Jern et al., 2005; Bénit et al., 1997; Cordonnier et al., 1995; Ono et al., 1986; Peters and Glover, 1980). ERVs make up around 12% of the mouse genome.

Class I ERVs comprise only 1.2% of the mouse genome and only about 10 to 300 full-length copies are still active (reviewed in Mager and Stoye, 2015). The mouse leukemia virus (MLV) family is the best described ERV1 element. MLVs integrated into the mouse genome about 1.5Myrs ago and are still able to infect other cells. However, MLVs are undergoing endogenisation and therefore differing amounts of MLVs (10 to 100 copies) can be found in different mouse strains (Stocking and Kozak, 2008). RLTR6 and VL30 are also ERV1 elements and share the same LTR elements. While RLTR6 encodes gag, pol and env genes, VL30 does not encode any protein coding genes. Therefore, RLTR6 is autonomous and VL30 coopts the RLTR6 coding genes to transpose in trans (Bromham et al., 2001; French and Norton, 1997).

Class II ERVs make up 4.9% of the mouse genome and comprise 8 members which account for around 2000 active ERV2 elements (reviewed in Mager and Stoye, 2015). ERV2 elements are sometimes also referred to as ERVK elements, because the first identified ERV2 class uses lysine-tRNAs as primers. MMTV was initially identified as a bona fide retrovirus, however only 2-3 full length MMTV exist in the mouse genome (Nusse and Varmus, 1982). Early transposons (ETn) and MusD transposons are highly related and make up around 300 copies of the mouse genome. ETns are highly active, however they lack coding sequences. Therefore, their transposition depends on the expression of the MusD proteins (Baust et al., 2003; Mager and Freeman, 2000, reviewed in Mager and Stoye, 2015).

Intracisternal A-type particles (IAPs) are the most studied mouse ERVs. IAPs were first discovered as viral particles in the endoplasmatic reticulum of several cancer cells (Dalton et al., 1961; De Harven and Friend, 1958). MMERVK10C as well as RLTR10 are also members of the ERV2 class, with MMERVK10C representing an internally
deleted RLTR10 element. IAP and ETn families are two high copy number ERVs responsible for most of the insertional germ-line mutations described in mice (Maksakova et al., 2006, Zhang et al., 2008).

**Class III ERVs** are the most abundant as well as most ancient ERV elements in the mouse genome (5.9%). ERV3 elements are sometimes referred to as ERVL elements as human retrotransposition starts with a leucine-tRNAs. As they have resided in the mouse genome for a long time mostly solo-LTRs of ERV3 elements play an important role in embryonic development as well as gametogenesis as early activators of developmental genes important for two-cell stage embryos. Mouse ERV with a leucine tRNA primer-binding site elements (MERVL) and mammalian apparent LTR-retrotransposons (MaLRs) are the two members of the ERV3 class. Both elements share a similar LTR sequence, hence it is thought that they arose from a common ancestor (McCarthy and McDonald, 2004). MERVLs have the ability to retrotranspose while MaLR elements are non-autonomous (Bénit et al., 1997, Smit, 1993). The two main MaLR elements - ORR1 and MT - are about 2-2.4kb in length and are ten times more abundant in the genome than MERVL elements.

**LINE elements**

LINE elements are autonomous TEs that lack LTRs. Based on the RT domain eleven superfamilies of LINE elements have been identified in eukaryotes: CRE, R2, R4, L1, RTE, Tad, R1, LOA, I, Jockey, CR1 (Malik et al., 1999, Kapitonov et al., 2009, Wicker et al., 2007). LINE elements are dominant in mammals and comprise 20% of the genome. L1 elements in human and mouse are the most studied LINE elements. The full-length human L1 is about 6kb long and composed of an 900nt long 5′ untranslated region (UTR), two ORFs (ORF1 and ORF2) as well as a 3′ UTR with a polyadenylation signal and a polyA tail (Babushok and Kazazian, 2007). The ORF2 encodes 150kDa protein with an reverse transcriptase (RT) domain and an endonuclease domain (EN), while ORF1 encodes for a 40kDa protein with unknown function (Feng et al., 1996, Mathias et al., 1991, Martin et al., 2003, Holmes et al., 1992). R2
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and RTE LINE elements only encode ORF2, while most other LINE elements encode both ORFs. To allow retrotransposition both ORF1 and ORF2 are required (Moran et al., 1996). In mice, the promoter of L1 sits in the 5’UTR and the amount of GC-rich tandem repeat monomers in the UTR is directly linked to the promoter activity (De-Berardinis and Kazazian, 1999). Furthermore, the 5’UTR also harbours an antisense promoter with yet unknown function (Li et al., 2014, Speck, 2001). In humans deletion of this promoter and knock-down of Dicer led to increased L1 retrotransposition (Yang and Kazazian, 2006) and in mouse 20-24nt siRNAs complementary to L1 sequences were found in the mouse germline (Watanabe et al., 2006). The two ORFs encode a bicistronic mRNA, even though internal ribosome entry sites (IRES) in front of each ORF in mice have been found (Li et al., 2006). After translation ORF1 and ORF2 form ribonucleoprotein (RNP) complexes and associate in cis with their encoding mRNA (Babushok and Kazazian, 2007, Kulpa and Moran, 2006). The RNP complex is transported into the nucleus and the L1 endonuclease nicks the host genome to generate a primer for the L1 reverse transcriptase (Jurka, 1997). The reaction is known as target primed reverse transcription (TRPT) and the cDNA is generated at the site of integration (Cost et al., 2002, Luan et al., 1993). Integration creates TSDs and leaves a often non-functional L1 without a 5’UTR as 5’ truncation or inversion of the 5’ sequence happen during integration (Szak et al., 2002). Pseudogenes are sometimes created through an improper association of the LINE proteins with their mRNA but instead transcribe genic mRNAs in trans (Esnault et al., 2000, Garcia-Perez et al., 2007).

L1 insertions account for 20% of the mouse genome, with about 600,000 copies. Of those, about 20,000 L1s are full-length and around 3,000 are competent to retrotranspose (Goodier et al., 2001). The active L1s are divided into L1A and L1F, while the L1V class is extinct. This classification is based on the L1 promoter. About 6.4 Myrs ago the L1F class arose by a replacement of the V promoter with an F promoter, this got further replaced by an A-promoter and then replaced back to an F-promoter, giving rise to the active classes of L1A, L1Gf and L1Tf-types (Sookdeo et al., 2013). In the mouse genome 900 L1A, 400 L1Gf and 1800 L1Tf elements are presumably still active (Goodier et al., 2001).
SINE elements

SINE elements are non-autonomous TEs that rely on the enzymatic machinery of LINE elements. Most of the SINE elements’ 5’ ends are derived from small cellular RNAs which are transcribed by polymerase III. Mouse SINE elements can be divided into four classes: B1, B2, ID and B4, depending on their RNA pol III promoter. The pol III promoter originates either from tRNAs, ribosomal 5S RNA or signal recognition particle 7SL RNA (Nishihara et al., 2006, reviewed in Kramerov and Vassetzky, 2005). The 7SL RNA gave rise to most abundant SINEs: B1s (650000 copies) and Alu elements (1.1 million copies) in mouse and human, respectively. The ID and B2 SINEs in mouse (350000 copies) harbour a tRNA-related RNA pol III element (Vassetzky and Kramerov, 2013). The B4 SINE elements arose through a fusion of the B1 and ID elements (Kramerov and Vassetzky, 2001).

SINEs are about 500-800bp long and are non-autonomous TEs. While LINE and ERV elements have the ability to retrotranspose, the retrotransposition of SINE elements depends on the activity of LINE elements. This hypothesis was first formed because of the 3’ end sequence homology between LINE and SINE elements in the polyA tail. Cell culture studies confirmed a dependency of SINE retrotransposition on LINE activity (Kajikawa and Okada, 2002) and studies have since then found L1 activity responsible for the activity of over a million SINE elements (Zhang et al., 2003, Vinckenbosch et al., 2006). However, not all SINE 3’ UTRs are similar to LINE polyA tails, and the short tandem repeat or polyT tail of Alu elements in humans are essential for their mobility (Dewannieux et al., 2003). SINE RNAs are transcribed by their own polymerase, the RNAs are transported to the cytoplasm where they are incorporated into LINE RNPs, and integrated by coopting the LINE EN and RT machinery. Specifically the 3’ poly A tail of the Alu elements bind to the L1ORF2p (Doucet et al., 2015, Ahl et al., 2015). Only B1 and B2 SINE elements are still active in the mouse genome and about 1.4 million SINEs are integrated in the mouse genome and occupy 8% of the mouse genome (Gibbs et al., 2004, Gilbert et al., 2004).
1.6.2 Lifecycle of TEs

The accumulation of TEs in the genome is divided into a burst of newly acquired TE activity, followed by a period of decay. Vertical or horizontal transfer leads to acquisition of new TEs in the genome. Vertical transfer is the derivation of a new TE through modification of an existing one while horizontal transfer happens through endogenisation of TEs from other species or through viral integration. TEs are masters of vertical transfer. It has be proposed that the 5’UTR of LINE elements is not conserved in LINE elements and LINEs acquire new regulatory units by incorrect template switching in the retrotransposition event (Hayward et al., 1997). In the mouse genome L1 families have been found to exchange coding sequences to give rise to mosaic elements which in turn can regain activity (Saxton and Martin, 1998; Sookdeo et al., 2013). While in most species, TEs horizontal transfer is leading to an increasing variety of TE species, mammals are subjected to relatively fewer instances of horizontal transfer. This might explain why mammals have fewer TE families which in itself are made up of many different members. 50% of LINE elements in both human and mouse originated from different variants of the LINE family through vertical gene transfer (Khan et al., 2006). An example of horizontal gene transfer is the endogenisation of retrotransposons to give rise to ERVs in vertebrates, like the Bov-B LINE element which was transferred from snakes to cows (Kordis and Gubensek, 1999).

The accumulation of TEs in the host genome is controlled by the insertion rate versus the loss of their functional activity (Charlesworth and Charlesworth, 1983). TEs transpose through insertion of an identical genetic copy into a new genomic region while the original sequence becomes inactivated through genetic drift. This efficient process leads to the distribution of identical copies of TEs, which decay over time by accumulating mutations and blending into the background (reviewed in Smit, 1996). As TEs are not positively selected for in the genome, they acquire mutations over evolutionary time. Therefore, only small amounts of TEs are still being active and account for most of the retrotransposition events with only 80 - 100 L1s being active and 4 - 5 very active (HOT) L1s are still active in the human genome (Brouha et al., 2003). TEs are responsible for 10-15% mutations in mice (Kazazian and Moran, 1998; Maksakova et al., 2013).
TEs can initially proliferate in the genome, as the genome does not recognise them. However, the genome defence pathways are subsequently set in place and finally TEs acquire mutations which inactivate them. The activity of TEs can lead to mutations in both host genome and TE, therefore TE accumulation is controlled by natural selection (Charlesworth and Charlesworth, 1983). A lot of TEs in the genome are inactive. The genome of mouse and human share about 165Mb of repeat sequences from 100Myrs ago (Chinwalla et al., 2002).

The accumulation of TEs is not random in the genome. Gene-rich regions are mostly spared from TE integration. While L1s were thought to be at higher frequency at pericentric repeats in humans, this could not be confirmed by whole genome sequencing (Lander et al., 2001, Laurent et al., 1997, Schuder et al., 2001). L1s and ERVs are highly enriched at AT-rich sites, while SINEs are enriched at GC-rich sites in the human and mouse genome (Smit, 1999, Soriano et al., 1983). L1 transgene insertion in mouse and human genomes showed that they inserted randomly in the genome (Babushok et al., 2006, Gilbert et al., 2002, Symer et al., 2002). Furthermore, new insertions of LINEs are mostly found throughout the genome, with no preference (Ovchinnikov et al., 2001), and while SINEs are integrate preferentially in AT-rich regions, they are selected for in GC-rich regions (Lander et al., 2001, Chinwalla et al., 2002). ERVs transgenes as well as bona fide retroviruses insert preferentially in genic regions, although they are selected for preferentially intergenically (Brady et al., 2009, Mitchell et al., 2004, Medstrand et al., 2002 reviewed in Lander et al., 2001).

1.6.3 TEs and the genome: Friends or foe?

The genome is comprised of genes as well as regulatory sequences. Studying the mosaic patterning of maize Barbara McClintock already defined TEs as mutable loci which have the ability to regulate the expression of nearby genes (McCLINTOCK.
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1951).

Genome evolution by TEs

TEs are creating genetic variation due to their transposition and have therefore been described as drivers of genomic evolution (Fedoroff, 2012a). In the mouse genome, L1 insertions lead to 2-3% of mutations while ERV insertions make up 10-12% of mutations (Druker and Whitelaw, 2004, Maksakova et al., 2006). Human TEs are largely silent with only 0.3% of mutations in human occurring through transposition of LINEs and SINEs (Callinan and Batzer, 2006). TEs can directly insert into protein-coding regions or regulatory regions that disrupt transcription (Kazazian et al., 1988). TEs can interfere with splicing, lead to early termination of a gene transcript, work as an alternative enhancers or work as an alternative promoters (Nekrutenko and Li, 2001, Kreahling and Graveley, 2004, Chen et al., 2009, Cohen et al., 2009, Gifford et al., 2013). An example of this presents the insertion of an LTR element into the Dicer mRNA which leads to an alternative Dicer isoform in the mouse oocyte (Flemr et al., 2013). LINE elements can mobilise SINE as well as genic mRNAs in trans and lead to the distribution of cellular RNA in the host genome (Esnault et al., 2000, Wei et al., 2001). TSDs are also described as the footprints of TEs, however L1s and Alus in humans can also transpose independent of the L1 endonuclease and therefore lack this TSD. The retrotransposition is rather linked to DNA double-stranded break repair, as the TEs insert into regions and repair the break (Morrish et al., 2002). Furthermore, insertions of L1s can lead to additional deletions of 1bp to 20kb in the host genome (Symer et al., 2002), while TE flanking genomic sequences can also transpose together with the TE into new locations (Moran et al., 1999, Han et al., 2005). It is referred to as 5’ or 3’ transduction and happens when TEs make use of alternative 5’ promoters or 3’ polyadenylation signals. 3’ transduction happens with high frequency and 15-20% carry an additional 200bp of genomic DNA sequence into new locations (Goodier et al., 2000, Pickeral et al., 2000, Tubio et al., 2014).

TEs can have a profound effect on the genome when homologous recombination of
non-allelic TEs leads to chromosomal rearrangements. If the two TEs are located on different chromosomes it leads to chromosomal translocations and if it takes place on the same chromosome it results in deletions, duplication or inversions (Sasaki et al., 2010). Throughout human evolution 27% of duplication, 44% of inversions and and over 500 deletion events were linked to improper homologous recombination of L1 and Alus (Bailey et al., 2003, Lee et al., 2008, Han et al., 2008, Sen et al., 2006).

TEs as disease-forming mutations

Insertions of TEs are mostly found in intergenic regions and therefore do not affect the host genome. In humans, the activity of LINEs and SINEs was directly linked to 50 genetic diseases ranging from muscle and blood disorders to immunodeficiencies, visual impairment and cancer (Kaer and Speek, 2013, Lee et al., 2012). While Alu insertions into the Breast cancer genes 1 and 2 (BRCA1-2) can directly cause breast cancer, TE mobilisations can also increase by disruption of cellular defence mechanisms that control TEs activity (Ross et al., 2010).

TEs as regulatory elements

TE insertions can lead to the derivaition of protein-coding genes resulting in beneficial functions in evolution. This happens mostly through gene duplications (Zhang et al., 2003). Furthermore, L1 in trans mobilisation of genic mRNA can lead to intron-less genes without a promoter. In the mouse genome about 3000 so called “retnogenes” have been found; 1000 of those have been found to be transcribed and 120 are now bona fide protein coding genes (Vinckenbosch et al., 2006, Zhang et al., 2003).

Furthermore, TEs have been endogenised by the host genome and evolved new functions. The recombination activating Rag1/Rag2 protein was created by the domestication of a DNA transposon 500Myrs ago (Kapitonov and Jurka, 2005, Thompson, 1995). VDJ recombination driven by Rag1 and Rag2 is a key component of the adaptive immune system in vertebrates. Furthermore, the imprinted genes Peg10 and Rtl1
arose through endogenisation of ERV transposons during placenta formation (Youngson et al., 2005). Additionally the sycytins are envelope genes of retroviral origins and are expressed specifically in the mammalian placenta (Cornelis et al., 2014).

1.6.4 Major Satellites

Transposable elements are able to retrotranspose and can present a danger to genome integrity, however the activity of additional repetitive regions in the genome can result into genome instability. Pericentric repeats present AT-rich long repetitive domains also called major satellites (Wong and Rattner, 1988, Joseph et al., 1989). Major satellites are 6Mb long and composed of 234bp repetitive units (Choo, 1997). These regions play a very important role throughout organisation of chromosomes during interphase of mitosis (reviewed in Comings, 1980, Manuelidis, 1990). In ESCs, major satellites are highly enriched for H3K9me3 which works together with HP1 to maintain heterochromatin of the centromeric region (Peters et al., 2001, Taddei et al., 2001). While H3K9me3 and DNA methylation are lost on pericentric repeats upon Suv39h1/2 KO, H3K27me3 gets enriched at those repeats, which suggests a compensatory mechanism to silence these repetitive elements (Lehnertz et al., 2003, Peters et al., 2003). Additionally, in hypomethylated ESCs, H3K9me3 is unaffected, yet H3K27me3 becomes enriched at pericentric heterochromatin (Marks et al., 2012, Saksouk et al., 2014, Cooper et al., 2014). In Dnmt triple KO ESCs, major satellites are not transcriptionally active but are enriched for PRC2. It was shown that the DNA binding protein BEND3 recruits PRC2 to major satellites upon DNA demethylation (Dai et al., 2013, Saksouk et al., 2014). BEND3 binding to major satellites is impaired upon DNA methylation (Dai et al., 2013).
1.7 Epigenetic modifications of TEs

Transposable elements (TEs) comprise around 70% of the mammalian genome (de Koning et al., 2011). TEs can randomly transpose in the genome, therefore they are sometimes described as the parasites of the host genome. Mobilised TEs are able to lead to chromosomal breakage, large-scale genomic rearrangements as well as interfere with gene expression by disrupting protein coding genes, as well as by altering transcriptional regulatory networks (McCLINTOCK, 1951, Chuong et al., 2016, Elbarbary et al., 2016).

In order to preserve genome integrity, the cell must guard the host genome by discriminating TEs from protein-coding genes. However, the plethora of TEs classes and very little sequence similarity between them makes this a difficult task for the genome (reviewed in Malone and Hannon, 2009).

1.7.1 Methylation of TEs

DNA methylation is a stable epigenetic mark that ensures the silencing of transposable elements. Studies suggest that DNA methylation evolved to protect the genome from TEs (Yoder et al., 1997). DNA methylation is thought to play a main role in TE silencing in differentiated tissues. However, differentiated cells are not viable without DNA methylation. Dnmt1 KO mice show 1000 fold upregulation of IAPs (Walsh et al., 1998, Sharif et al., 2016). Furthermore, KO of Dnmt3L lead to activation of LINEs and ERVs in the male germline (Bourc’his and Bestor, 2004). Dnmt3C has been found to be involved in methylating especially young TEs in the male germline (Barau et al., 2016).

Additionally, Tet1 KO has been found to lead to transcriptional activation of LINE-1 elements in mouse ESCs (de la Rica et al., 2016). As reported before, TEs are active in severely hypomethylated cancer types (Schulz, 2006). Although Dnmt1 KO leads to demethylation of IAP elements (Arand et al., 2012), Dnmt1 constitutive KO in ESCs does not result in IAP activity (Hutnick et al., 2010). The long term deletion of DNA methylation at TEs in ESCs seem to be compensated for by deposition of repressive chromatin modifications (Karimi et al., 2011, Matsui et al., 2010). However, in acute
global hypomethylation of ESCs IAPEZ, LINEs, MERVL as well as MMERGLN are upregulated (Walter et al., 2016).

1.7.2 Histone modifications of TEs

In addition to DNA methylation, histone modifications have been implicated in TE regulation. The KO of Eset in mouse ESCs led to upregulation of a large class or ERV1 and ERV2, as well as an upregulation of L1s (Karimi et al., 2011, Matsui et al., 2010). Also Kap1 KO transcriptionally activated all ERV classes, as well as L1 elements (reviewed in Rowe and Trono, 2011). Furthermore, chimeric transcripts derived by TEs used as alternative promoters for genes arise upon Eset KO in mouse ESCs (Karimi et al., 2011). Therefore, H3K9me3 plays a major role in TE silencing through ESET and KAP1 pathways, as KRAB-ZFP targets the histone methylase to TE regions. However, HP1 plays only a minor role in keeping TE elements transcriptionally inactive (Maksakova et al., 2013). KRAB-ZFP bind in the 5′ region of TEs; and therefore, H3K9me3 is highly enriched at the promoter region of TEs. MLV TEs are silenced by binding of ZFP809 to the proline tRNA binding site (PBS) and recruitment of KAP1 and ESET (Wolf and Goff, 2007, Wolf and Goff, 2009). KRAB-ZFPs evolve very quickly in the genome and are thought of as a genome defense against TEs. They are regarded as an example of the arms race of the genome against TEs; as new ERVs evolved, KRAB-ZFPs appeared (Thomas and Schneider, 2011), therefore young L1s might have still escaped ZFP targeting now.

Two L1 classes, L1PA3 and L1PA4, in the human genome escaped KRAB-ZFP control as they deleted the binding site of the zinc finger protein (Jacobs et al., 2014). On the other hand, ancient L1s, that were highly active before KRAB-ZFPs evolved, are not recognised by ZFP elements, but guarded by KAP1 and H3K9me3 (Bulut-Karslioglu et al., 2014).

SUV39H plays a major role in controlling LINE elements. Suv39h dKO in ESCs leads to depletion of H3K9me3 at ERVs as well as L1 elements, but while ERVs do not get transcriptionally upregulated, L1 type A are strongly expressed (Bulut-Karslioglu et al., 2014). TE silencing by H3K9me3 has not been found in differentiated cells yet,
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as KO of Suv39h, Eset and Kap1 in mouse embryonic fibroblasts or upon differentiation of ESCs did not lead to transcriptional upregulation of TEs (Bulut-Karslioglu et al., 2014, Matsui et al., 2010, Rowe et al., 2013). ERV3 seem to be silenced by H3K9me2, as MERVL elements are enriched for H3K9me2 in ESCs and G9a KO leads to transcriptional upregulation of MERVL elements (Maksakova et al., 2013).

H3K9me3 is not enriched at ERV3 transposons, neither in differentiated or undifferentiated cells, nevertheless KO of Kap1, Suv39 as well as Eset led to minor MERVL transcription. Therefore, it is thought that these elements are silenced by all three H3K9 methylation pathways but through an indirect yet to be identified mechanism (Bulut-Karslioglu et al., 2014, Maksakova et al., 2013). Additionally, demethylation of histone methylation has been found to activate ERV elements; the deletion of the KDM Lsd1/Kdm1a of H3K4me1/2 led to upregulation of MERVL elements (Macfarlan et al., 2012). Finally, upon global hypomethylation in ESCs MERVL elements have been found to be silenced by redistribution of H3K27me3 (Walter et al., 2016).

1.7.3 Regulation of TEs by small RNAs

Further to DNA methylation and histone modifications, small RNAs have been described as playing a role in TE control. Two recent studies in the mouse oocyte have indicated a role of miRNAs as well as endogenous siRNAs (endosiRNAs) to control TE expression (Flemr et al., 2013, and Stein et al., 2015). In the mouse oocyte and in preimplantation embryos, endosiRNAs have been found to control IAPs and MERVL elements (Tam et al., 2008, Watanabe et al., 2006, Svoboda et al., 2004). In ESCs SINE derived endosiRNAs were discovered; two tandem inverted SINE elements give rise to a hairpin siRNA that feeds into the endosiRNA pathway (Babiarz et al., 2008). Additionally, studies have shown that upon Dicer KO in ESCs, L1 and IAP mRNA were upregulated (Kanellopoulou et al., 2003) and in human cancer cell lines LINEs have been found to be regulated by endosiRNAs (Yang and Kazazian, 2006). Artificially generated L1 siRNAs have been found to reduce retrotransposition of L1 in cell culture (Soifer et al., 2005). Furthermore, miRNAs have been found to potentially play a role in TE silencing as 12% of the miRNAs so far reported share homology
with TEs (Smalheiser and Torvik, 2005). In the male germline IAPs are repressed by so-called piwi-interacting small RNAs (piRNAs) which lead to transcriptional gene silencing (TGS) through de novo DNA methylation (Aravin et al., 2008, Kuramochi-Miyagawa et al., 2008). The PIWI proteins MILI, MIWI and MIWI2 are important during spermatogenesis and Dnmt3L, Mili, or Miwi2 KO leads to LINEs, ERVs and meiotic catastrophe, with impairment of the de novo methylation of the promoters of IAPs as well as L1 (Aravin et al., 2007, Bourc’his and Bestor, 2004). Also KO of Maelstrom, which is important for perinuclear chromatoid body formation important for piRNA biogenesis, leads to impaired spermatogenesis. Furthermore, the KO of the MAELSTROM leads to retrotransposition of IAP and L1 elements in cell culture (Soper et al., 2008).

1.7.4 TEs during epigenetic reprogramming

During early mammalian development, TEs become transcriptionally activated and contribute to genome regulation. 15-20% of total capped RNA in the oocyte and at blastocyst stage stems from TEs, with LINEs as well as ERVs being expressed (Fadloun et al., 2013). IAP, MusD and MERVL virus particles have been detected in the mouse embryo (Kuff and Lueders, 1988, Ribet et al., 2008). ERV3 elements are highly active during the first wave of global demethylation in early mammalian development. 13% of RNA in the unfertilised oocyte is comprised of MaLR RNAs (Peaston et al., 2004). MERVL elements present the best studied ERV during embryonic development. Already eight hours after fertilisation, at the onset of zygotic genome activation, MERVL are the earliest transcripts in the oocyte. MERVL are upregulated 300-fold in the oocyte and two-cell embryo, while transcription decreases rapidly in the following cell divisions (Kigami et al., 2003, Macfarlan et al., 2012). Interestingly, this ERV3 upregulation also controls about 300 endogenous genes, which get expressed as chimeric transcripts between MERVL LTR and their exons. MERVL elements are potentially important early activators in development as most of the genes they regulate are confined to the two-cell stage. Furthermore, in serum-grown ESCs a small proportion of cells express high levels of MERVL as well as genes controlled by
1.8 Embryonic stem cells

Mouse embryonic stem cells (ESCs) can be isolated from the inner cell mass of the early blastocyst and cultured in vitro and serve as an invaluable tool for in vitro studies in developmental biology (Evans and Kaufman, 1981, Martin, 1981).

Even though ESCs can self-renew infinitely without immortalisation, which is normally attributed to a tumorigenic potential (Suda et al., 1987), ESCs are still able to contribute to an embryo after prolonged in vitro culturing.

It is important to culture ESCs in optimal conditions to preserve the self renewal and pluripotency to prevent differentiation. Mouse ESCs are cultured on a layer of inactivated mouse embryonic fibroblasts (MEFs) that provide nutrients. The addition of serum and Leukaemia Inhibitory Factor (LIF) preserves pluripotency of ESCs (Smith et al., 1988, Williams et al., 1988) and allows culturing ESCs without MEFs. Nevertheless, ESCs need to be grown on plates coated with gelatine to provide elasticity of the surface, which is invaluable to maintain pluripotency (Skardal et al., 2013). Although ESCs are derived from the inner cell mass (ICM) of the blastocyst, which is depleted of DNA methylation, cultured ESCs have a methylation landscape largely similar to somatic cells, with about 75% of CpGs methylated (Stadler et al., 2011, Popp et al., 2010).

The addition of two small molecule inhibitors of FGF and GSK signalling (2i inhibitors,
PD0325901 and CHIR99021) allows ESC culturing in serum-free conditions (Ying et al., 2008, Kunath et al., 2007, Wray et al., 2010). Not only do 2i grown ESCs show a more homogenous morphology than serum grown ESC (Marks et al., 2012), but also the transcriptional profiling led to the conclusion that those 2i grown ESCs resembled the “ground state” of pluripotency (Ying et al., 2008). Additionally, 2i grown ESCs have a much lower methylation level than serum grown ESCs, more closely resembling the methylation of the ICM (Ficz et al., 2013, Angermueller et al., 2016, von Meyenn et al., 2016).

1.8.1 Transcription factor networks regulate ESC pluripotency

Self-renewal is a key property of ESCs which is tightly regulated by a network of transcription factors. The pluripotency network in ESCs is governed by the master regulator OCT4, encoded by the Pou5f1 gene (reviewed in Pan et al., 2002). Knockout of Pou5f1 leads to embryonic lethality at the blastocyst stage (Nichols et al., 1998), while deletion in ESCs leads to differentiation. NANOG is the second member of the pluripotency network (reviewed in Saunders et al., 2013). ESC depleted of NANOG start to differentiate but can be maintained (Chambers et al., 2007). However, NANOG knock out mouse embryos die at blastocyst stage (Mitsui et al., 2003, Silva et al., 2009). This is not surprising as OCT4 and NANOG work jointly and regulate the same genes (Loh et al., 2006). The transcription factor SOX2 can regulate Nanog expression (Rodda et al., 2005). And OCT4, NANOG as well as SOX2 have been found to control the same genes by co-occupying the same promoters (Boyer et al., 2005, Chambers et al., 2009). Therefore, we can think of the regulation of pluripotency genes as a highly organised network (Boyer et al., 2005).

In mouse ESCs 5 to 20% of the binding sites of NANOG and OCT4 have been found in ERV2 TEs (Kumarso et al., 2010). Therefore, ERV2 might also be important drivers of the pluripotent state. In human, naïve ESCs primate-specific ERVs provide functional binding sites for pluripotency transcription factors and can thereby drive expression of alternative long non-coding (lncRNAs) transcripts (Wang et al., 2014, Lu et al., 2014, Fort et al., 2014).
1.8.2 H3K9 and H3K27 methylation in ESCs

In mouse ESCs, DNA methylation and the repressive histone marks H3K9me2, me3 and H3K27me3 mark distinct genomic loci. Even though, H3K9 methylation largely overlaps with DNA methylation, H3K9me2/3 is highly enriched in TEs, telomeres, centromeres and pericentric regions and depleted in CGIs. In ESCs lacking DNA methylation, H3K9me3 is unaffected, however H3K27me3 becomes enriched at pericentric heterochromatin (Marks et al., 2012, Saksouk et al., 2014, Karimi et al., 2011, Tsumura et al., 2006). Additionally, an even broader enrichment of H3K27me3 can be found at chromocentres upon depletion of SUV39h in Dnmt triple KO ESCs (Schmitges et al., 2011). This is very intriguing, as the repressive marks H3K27me3 and H3K9me3 are normally mutually exclusive in pluripotent and differentiated cells (Hawkins et al., 2010, Mikkelsen et al., 2007). The mutually exclusive enrichment of those two repressive histone marks might be explained by the ability of HP1 proteins to prevent CBX-PRC1 and PRC2 binding in the zygote (Tardat et al., 2015).

Additionally, DNA methylation and H3K27me3 are mutually exclusive at CGIs in ESCs (Brinkman et al., 2012, Statham et al., 2012). PRC2 is attracted to GC-rich unmethylated regions and therefore H3K27me3 is enriched at GC-sites (Jermann et al., 2014, Lynch et al., 2012, Mendenhall et al., 2010). One way to prevent DNA methylation at promoters in ESCs is the ability of PRC2 to recruit TET enzymes to these genomic regions in ESCs (Neri et al., 2013). Although H3K27me3 is confined to CGIs in ESCs, in mouse differentiated cells, H3K27me3 expands away from CGIs into methylated regions and can encompass 10% of the genome with 15-40kb domains (Tanay et al., 2007, Hawkins et al., 2010, Pauler et al., 2009). Furthermore, upon ESC differentiation DNA methylation gets enriched at promoter regions (Mohn et al., 2008, Ohm et al., 2007). PRC could be important to silence the genes during development with DNA methylation being a long-term silencer in differentiated cells.
Chapter 2

Material and Methods

2.1 Materials

<table>
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<td>NanoDrop® Technologies</td>
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<tr>
<td>780 AxioObserver point scanning confocal microscope</td>
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<tr>
<td>Sonicator 3000</td>
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<td>CD4 pull down</td>
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<td>anti-IAP</td>
<td>Dr. Cullens lab</td>
<td>IF</td>
</tr>
<tr>
<td>anti-rabbit Alexa-Fluor 568</td>
<td>Life-Molecular Probes</td>
<td>IF</td>
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<tr>
<td>anti-AGO2</td>
<td>Dr. O’Carrolls lab</td>
<td>AGO2 IP</td>
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<td>anti-H3K9me3</td>
<td>Active Motif, Clone</td>
<td>ChIP-seq</td>
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<td></td>
<td>MABI0319</td>
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<td>anti-H3K27me3</td>
<td>Active Motif, 39155</td>
<td>ChIP-seq</td>
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### Table 2.4: Laboratory materials

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<td>Pechiney Plastic Packaging</td>
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<td>Starlab</td>
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<td>Tissues</td>
<td>Kimwipes</td>
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<td>Phase Lock tubes</td>
<td>5-Prime</td>
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<td>1.8 ml Cryotube</td>
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<td>Cell strainer 40 µm</td>
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<td>96 well PCR plates</td>
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<td>384 well qRTPCR plates</td>
<td>Biorad</td>
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<td>96 well qRTPCR plates</td>
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<td>Scalpel</td>
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<td>MACS columns</td>
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2.1. Materials

Chapter 2. Material and Methods

Table 2.5: Kits

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<td>Qiamp kit</td>
<td>Qiagen</td>
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<td>TURBO DNA-free kit</td>
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<td>PicoGreen® dsRNA kit</td>
<td>Thermo Fisher Scientific</td>
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<td>Platinum SYBR Green qRT PCR</td>
<td>Life Technologies Ltd</td>
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<td>small RNA library kit</td>
<td>Illumina</td>
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<td>NEBNext</td>
<td>New England Biolabs</td>
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<tr>
<td>Imprint kit</td>
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<td>QuickExtract</td>
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### Table 2.6: Chemicals and Reagents I

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## 2.2 Oligonucleotides

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### Table 2.10: qRTPCR Primer and sequences

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<th>Primer</th>
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Table 2.11: PCR Primer and sequences

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<td>Dicer_20_21_screen_RV1</td>
<td>TGACCAGAATAAGAAGGAGCGGA</td>
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</table>

2.3 Methods

2.3.1 Cell culture

ESCs were cultured in standard serum-containing media (DMEM (Gibco, 11995-040), 4 mM L-glutamine, 110 µg/ml sodium pyruvate, 15% fetal bovine serum, 1 µg/ml Penicillin- Streptomycin (Gibco, 15140-122), 0.1 mM nonessential amino acids (Gibco, 11140-050), 50 µM beta-mercaptoethanol (Sigma, M6250) and 103 U/ml mouse LIF (Stem Cell Institute, Cambridge).

For serum-free culturing of ESCs under 2i culturing conditions (Ying et al., 2008), the medium was replaced by serum-free N2B27, composed of 1:1 DMEM/F12 (Gibco, cat. 21331-020) and Neurobasal (Gibco, 21103-049), 1X N2 (Stem Cell Sciences, cat. SF-NS-01-005), 1X B27 (Gibco cat. 17504-044), 1% L-glutamine (Gibco, 25030-081), 1 µg/ml Penicillin- Streptomycin (Gibco, cat 15140-122), and 50 µM beta-mercaptoethanol (Sigma, M6250), 103 U/ml LIF and the two inhibitors (MEK inhibitor PD0325901 (1 µM, Stem Cell Institute Cambridge) and GSK3 inhibitor CHIR99021 (3 µM, Stem Cell Institute Cambridge).
ESCs were maintained in an incubator at 37 °C with 5% CO2 humidified atmosphere. ESCs were grown on immortalised mouse embryonic fibroblasts (MEFs) or on 0.1% gelatine. A tamoxifen-inducible Cre recombinase was used to induce recombination of LoxP sites in the floxed cell lines using tamoxifen (800 nM) (Sigma).

**Passaging cells**

The medium of ESCs was changed every day and ESCs were split (1:4) every 2 days. For splitting ESCs, the cells were washed with 1 x PBS (Thermo Fisher Scientific). TrypLE Express (Thermo Fisher Scientific, 12604013; for serum-free ESCs) or Trypsin (Gibco; for serum-grown ESCs) was added for 3 min at 37 °C and was inactivated by adding 10 x ESC medium. The cells were spun down at 300 x g at room temperature for 3 min and resuspended in ESC medium and passaged to a new gelatine-coated or MEF coated plate.

**Thawing cells**

To thaw ESCs from liquid nitrogen, 15 ml falcons with 10 ml ESC medium were prepared. Cells were thawed in a 37 °C water bath, resuspended in ESC medium and spun down at 300 x g at room temperature for 3 min. The ESCs were resuspended in ESC medium and plated into 10 cm dishes.

**Freezing cells**

Cells from a 10 cm dish were split (1:4) and resuspended in 0.5 ml ESC medium. Two times ESC freezing medium (20% DMSO, 60% FBS and 20% ESC medium) was added and the mixture cells were transferred into cryotubes. Cryotubes were transferred into an isopropanol freezing container and frozen at -80 °C overnight (ON). The following day the frozen vials were transferred to the liquid nitrogen storage tank.
2.3.2 Reprogramming of MEF to iPSCs

Wildtype or AidKO MEF cultures were established from female 13.5 to 14.5 dpc embryos. For each transfection, 0.8x10^6 MEF were nucleofected using Amoxa Nucleofection Technology (Lonza AG; program A-023), according to the manufacturer’s instructions, with 1 µg of each plasmid. Plasmids for reprogramming pB-TRE-OCKS, pBASE, pB-CAG-rtTA were obtained from Sanger Institute’s plasmid repository. Reprogramming was performed in ES cell medium in the presence or absence of Doxycycline, in a 5 % O2 incubator. The medium was refreshed every other day. Colonies were picked on day 6 of reprogramming and expanded for at least 29 passages.

2.3.3 In vitro PGC like cells

PGC like cells (PGCLC) were generated following the protocol in (Meyenn et al., 2016).

2.3.4 In vivo PGC collection

All embryonic samples for library preparation were collected from timed matings of C57Bl/6J female mice PGCs carrying the Oct4-GFP transgene expressed in the developing gonad (Yoshimizu et al., 1999) described in Seisenberger et al., 2012. Animal work carried out as part of this study is covered by a project license (to W.R.) under the 1986 animal (scientific procedures) act, and is further regulated by the Babraham Institute Animal Welfare, Experimentation, and Ethics Committee.

2.3.5 DNA extraction

The Qiamp kit from Qiagen was used according to supplier’s protocol. Elution was done with 100 µl EB buffer. If DNA was used for CRISPR-Cas9 screening or polymerase chain reaction (PCR) purposes, the QuickExtract (Epicentre, QE09050) solution was used following the supplier’s protocol.
2.3.6 RNA extraction

RNA was extracted using Trizol isolation reagent (Tri-Reagent, Sigma) and Phase Lock tubes following manufacturer’s instructions (5-Prime), and was subjected to DNase treatment using the Ambion DNA-free kit according to the manufacturer’s instructions (Invitrogen).

2.3.7 Measurement of RNA and DNA concentration

The concentration of DNA and RNA was estimated using the NanoDrop 2000 UV-Vis Spectrophotometer (ThermoScientific). The genomic DNA concentration was measured using the Picogreen kit according to manufacturer’s protocol. For library preparation, DNA and RNA were analysed on a Bioanalyzer (Agilent) following the manufacturer’s protocols.

2.3.8 qRTPCR

100 ng -1 μg of DNase treated RNA was reverse transcribed (Thermo RevertAid K1622) using random hexamer primers. Platinum SYBR Green qRTPCR (Life Technologies Ltd) was used for qRTPCR reaction. The cDNA was diluted 200 times and 2 μl was used for each qRTPCR reaction. The qRTPCR was run in the Biorad 384 well machine following the manufacturer’s condition and measuring SYBR signal. Endogenous controls (Atp5b, Hspcb, U1) were used to normalise expression. Primers are listed in Materials (Table 2.10).

2.3.9 CRISPR KOs

For CRISPR KO the protocol published by Ran et al., 2013 was followed with the subsequent changes. Guide RNAs were designed using the online tools http://crispr.mit.edu/ and http://crispr.dbcls.jp/ as well as
https://chopchop.rc.fas.harvard.edu/. The gRNAs which were predicted best by all three prediction tools, taking into consideration off target effects, were used.

**Constitutive Dicer KOs**

The gRNAs for the DICER KO (Table 2.9) were then cloned into the pSpCas9(BB)-2A-GFP plasmid (Addgene plasmid ID: 48138). ESCs were cultured on gelatinised plates and transfected with 1 μg of gRNA construct. Transfection was done using Lipofectamine 2000 transfection reagent (Invitrogen).

Two days after transfection, ESCs were split and resuspended in 1 ml of 1 x PBS with 1% FBS. The cells were then pipetted through a cell strainer (40 μm, Corning, Product #352340) into a round bottom polypropylene tube of flow cytometry. The cells were sorted on the LSR Fortessa Cell Analyser (BD Biosciences). In order to distinguish live from dead cells, DAPI incorporation was measured. Single cells were sorted for GFP expression into 96-well plates which had been gelatine-coated and filled with 100 μl ESC medium. Medium was changed every day and colonies were grown until confluent. The cells were then split 1:2, with half of the colonies being expanded and DNA extractions carried out from the other half using QuickExtract. Clones were screened by PCR using screening primer pairs for Dicer_23_24_screen (Table 2.11) and MyTaq Redmix following manufacturer’s protocols. Positive colonies identified by PCR were send for Sanger sequencing.

**Conditional Dicer KOs**

The conditional KOs were generated the same way as the constitutive KOs with the following changes. The gRNAs for Intron 14_15 and Intron 20_21 of Dicer (Table 2.8) were cloned into the pSpCas9(BB)-2A-CD4 plasmid. Cells were cultured on feeder plates and transfected with 1 μg of gRNA construct as well as 100 ng donor single stranded DNA for either Intron 14_15 (Dicer_14_15_donor_loxP) or Intron 20_21 (Dicer_20_21_donor_loxP). Cells were trypsinised and resuspended in 70 μl PBS and stained with human CD4 microbead antibody (Miltenyl Biotec, Cat. No. 130-045-101)
according to manufacturer’s instructions. The CD4 positive cells were enriched using MACS columns. Negative cells were collected from flow through. The cells were eluted in 500 µl PBS, spun down and resuspended in ESC medium. 4000 to 5000 cells were plated in 10 cm dishes to have single cells. The medium was changed every day and after 1 week the colonies were picked. The colonies were picked into Trypsin in 96 well round bottom plates. The colonies were then divided into two. Half of the colonies were plated in 96 well plates coated with gelatine and containing 200 µl ESC medium, and the other half were transferred into a 96 well PCR plate for DNA extraction and screening. For screening, 96 well plates were centrifuged for 10 min at 3000 x g at room temperature. The ESC medium was discarded and the cells were resuspended in 100 µl H₂O. 10 µl of Proteinase K was added and cells were incubated for 30 min at 55°C. Afterwards 5 µl of the DNA was used for PCR using MyTaq Redmix and screening primers for intron 14_15 and intron 20_21 of Dicer (Table 2.11). As a diagnostic test, 10 µl of PCR product were digested with PvuI in the case of intron 14_15 or EcoRI for intron 20_21. 20 µl of the product was run on a 2 % agarose gel. Positive colonies identified by PCR were send for Sanger sequencing.

**Cloning of gRNA constructs**

gRNAs primers were ordered with sticky end overhangs and ligated according to \cite{Ran2013} into the Cas9 plasmid using T4 Ligase (New England Biolabs) following to manufacturer’s instructions. The ligation mix was incubated for 10 min at room temperature. The ligation batch was mixed with 100 µl of Top10-bacteria (Invitrogen) and incubated on ice for 10 min. The heat shock was performed for 45 sec at 42°C and the samples were subsequently put on ice for 2 min. After having added 200 µl of LB-medium (1 % (w/v) Bacto tryptone, 0.5 % (w/v) Bacto yeast extract, 170 mM NaCl, adjusted to pH 7.6 with NaOH) without ampicillin the bacteria were shaken for 30 min at 300 rpm and 37°C for recovery on the thermocycler. They were plated on LB-agar plates with ampicillin (100 µg/mL) which had been prewarmed at room temperature (1.5 % (w/v) Bacto agar in LB medium, PBS (130 mM NaCl, 100 mM Na₂HPO₄, pH 7.0). The plates were incubated over night at 37 °C. The next day, 5 ml
LB-medium with 100 µg/mL ampicillin was inoculated in a round bottom tube (BD Biosciences) with single bacterial colonies, which grew ON on a shaker at 37 °C. The plasmid DNA was isolated using QiAprep Spin Miniprep kit from Qiagen and following the manufacturer’s protocol. The U1 reverse primer (Table 2.11) was used to test for positive cloning of gRNAs into the plasmid. The PCR product was send for Sanger sequencing at Sigma.

**Testing gRNA efficiency**

Firstly, the T7 Endonuclease assay was performed to test for the cutting efficiency of the gRNAs. Thereafter, gRNAs cloned into the Cas9 plasmid were transfected into ESCs. After two days the cells were either sorted for GFP or CD4 expression and the DNA was isolated out of the bulk population of the positive culture using 300 µl of QuickExtract (Epicentre, QE09050) following supplier’s instructions. Afterwards the DNA was incubated with the T7 endonuclease for 5 min and 10 µl of the sample were run on a 10% agarose gel. The agarose gel was stained with SYBR Gold and analysed using a gel imager to calculate the gRNA efficiency.

**Screening for CRISPR clones**

For each PCR reaction of 30 µl sample volume: forward Primer (100 µM) 0.5 µl, reverse Primer (100 µM) 0.5 µl, template 50 ng, 2 x MyTaq Polymerase mix 15 µl, H₂O (to 15 µl). The PCR was run on the Biorad thermocycler with the following conditions: 2 min denaturation at 95°C, 35 cycles with 30 sec denaturation at 95°C, 1 min annealing at 55°C and 30 sec elongation at 72°C were completed. Before cooling to 4°C, an additional elongation step for 5 min at 72°C was accomplished.

**Agarose gel electrophoresis**

Agarose flat-bed gels in various concentrations (0.6 - 2 % agarose in 1 x TE buffer) and sizes were run to separate DNA fragments in an electrical field (10-20 V/cm) for analytical or preparative use. The desired amount of agarose was boiled in 1X TE buffer
until it was completely dissolved. After it cooled to approximately 55 °C, SYBR green (0.5 g/ml final concentrations) was added to the liquid agar, which was then poured in a flat-bed tray with combs. As soon as the agarose solidified, the running buffer (1 x TE buffer) was added. DNA in the 6 x loading buffer (1.5g Ficoll 400, Orange G dye, H₂O to 10mL) was loaded into the wells and separated electrophoretically. 10 x TE: 400 mM Tris, pH 8.0, 10 mM EDTA, acetic acid for pH titration.

2.3.10 Small RNA qRTPCR

For small RNA qRTPCR Taqman miRNA kits were used according to the manufacturer’s instructions for mmu_miR93 (Taqman, Cat. No. TM001090), mmu_miR7081_mat (Taqman, Cat. No. TM467052_mat) and snoRNA202 (Taqman, Cat. No. 001232) was used as a positive control.

2.3.11 AGO2 immunoprecipitation

Cells were cultured on 15 cm dishes and harvested in 1x PBS. Pellets were frozen at -80°C until further processing. ESCs were resuspended in 300 µl lysis buffer (50 mM Tris, pH 8, 150 mM NaCl, 5 mM MgCl₂, 15 % Glycerol, 1 mM DTT, 0.5 % Na deoxycholate, 0.5 % Triton X-100, Protease inhibitor cocktail (Roche), 50 µg/ml yeast tRNA, 2 mM Vanadyl ribonucleoside complex) and cells were pelleted at 10,000 rpm, 10 min, 4°C. The supernatant was used as whole ESC extract. 25 µl Protein G-coupled Dynabeads (Thermo Fisher Scientific, Cat. No. 10003D) were washed 3 x with 1 ml of Wash Buffer (10 mM Tris pH 8, 150 mM NaCl, 1 mM MgCl₂, 0.01% NP-40). 50 µl of purified AGO2 antibody (kind gift of the Dr. O’Carroll lab) was added, filled up to 1 ml with Wash Buffer and incubated ON at 4°C in a rotating wheel. On the next day the beads were washed three times with wash buffer and the negative control (beads with extract but without serum) was prepared. The ESC extract was prespun to remove precipitated proteins and 200 µL extract was added to the beads and filled up to 600 µL with Lysis buffer. The mix was incubated for 2-4 h at 4°C in a rotating wheel and subsequently washed five times with wash buffer. The IP was
eluted with 300 µL Proteinase K buffer (10 mM Tris pH 7.5, 0.5% SDS, 5 mM EDTA, 1 µl Proteinase K/reaction) after 30 min of 50°C incubation on the thermomixer, at 850 rpm. RNA was isolated by phenol extraction and eluted in 8 µl H2O.

2.3.12 siRNA knock down

For the siRNA knockdown, the Dharmacon SMARTpool siRNAs were used according to the manufacturer’s description. For the first round of transfection, cells were split and plated at a density of 1 x 10^5 onto gelatine coated 12 well plates. 24 h after plating, cells were transfected with siRNAs using Lipofectamine 2000 (Invitrogen, Cat. No. 11668019) and following manufacturer’s instructions: 2.5 µl of 10 µM siRNAs in 100 µl Optimem were transfected using 2 µl of Lipofectamine 2000 per well. Dharmacon siGENOME SMARTpool siRNA against mouse Dicer (Dharmacon, Cat. No. MU-040892-01-0005), Dgcr8 (Dharmacon, Cat. No. M-051365-00) or Ago2 (Dharmacon, Cat. No. MU-058989-01-0005) and siGENOME non-targeting siRNA2 (Dharmacon, Cat. No. D-001210-02-05) were used for transfection in triplicates. The second transfection was done 48h after the first transfection. Cells were split onto 6 well plates. The second transfection was done with the following conditions: 5 µl of 10 µM siRNAs in 200 µl Optimem using 5 µl Lipofectamine 2000 per well. Cells were harvested 48 h after the second transfection and RNA was extracted using Tri-Reagent.

2.3.13 Immunofluorescence

Cells were grown directly on sterile cover slips coated with 1 % gelatine and were fixed in 2 % PFA (Polysciences, Inc. Cat 18814) for 30 minutes at room temperature, washed in PBS and permeabilised in PBS 0.5 % Triton X-100 for 30 minutes at room temperature. After washing, they were blocked in PBS 0.05 % Tween-20, 1 % BSA (blocking solution, BS) ON at 4°C and subsequently incubated 60 minutes at room temperature with a rabbit polyclonal anti-IAP antibody (kind gift of the Dr. Cullen lab) diluted 1:200 with BS. After 60 minutes of washing in BS, the cells were incubated with anti-rabbit Alexa-Fluor 568 (Life-Molecular Probes) diluted 1:500 with BS for 60
minutes at room temperature in the dark. After washing ON at 4°C in PBS 0.05 % Tween® 20, the cells were stained with DAPI (5 µg/mL) and mounted with SlowFade Gold (Life-Molecular Probes). Slides were imaged on a Zeiss 780 AxioObserver point scanning confocal microscope with a Plan Apochromat 63x/1.40 oil objective. Single optical slices were captured. Images were pseudocoloured using Adobe Photoshop CS4 and levels were adjusted according to internationally accepted guidelines for image manipulation.

2.3.14 Next generation sequencing library preparation

Whole Genome Bisulfite sequencing libraries

For whole genome bisulfite sequencing (WGBS-seq) library preparation, 100 ng of DNA (measured by PicoGreen) were fragmented by sonication (Covaris E220 sonicator), with the following conditions: duty factor, 10; peak incident power, 140; cycles/burst, 200; time, 55 sec; temperature, at 12°C. The fragmented DNA was end-repaired, A-tailed and the adapters were ligated following the NEBNext Ultra kit protocol (NEB, cat: E7370L). The adapter-ligated DNA was treated with bisulfite reagent following the two step protocol from the Sigma Imprint kit (Sigma, cat: MOD50). The libraries were amplified using the HiFi Uracil+ ReadyMix (KapaBiosystems, cat. KK2801) for 16 cycles and iPCRtagged indexing primers. Subsequently the libraries were purified using AmPure beads. The library concentration was quantified using the Kapa Library quantification kit (KapaBiosystems, KK4844) and equimolar amounts of each indexed library were pooled to run two WGBS-seq libraries on one lane of HiSeq 2000 Illumina system sequencing 100 bp paired-end.

Stranded Total RNA sequencing libraries

Total RNA was extracted using Tri-Reagent (Sigma). 100 ng - 1 µg of RNA with a RNA integrity number (RIN) of 8-10 were used for total RNA-seq libraries. After Ribozero depletion, stranded total RNAseq libraries were prepared using the TruSeq
RNA library preparation kit v2 (Illumina) following manufacturer’s instructions. The RNA-seq libraries were indexed and 50 bp single end sequencing was performed using TruSeq reagents (Illumina) according to manufacturer’s instructions. Libraries were sequenced on Illumina HiSeq 2000.

Small RNA-seq libraries

Small RNA-seq libraries were produced according to the Illumina protocol (RS-200-0012), with the following changes: 10 ng or 1 µg RNA with a RIN of 8-10 were used as input material. The instructions were followed until the cDNA purification. In order to purify the cDNA, the samples were run on 10 % Novex PAGE gel. DNA sample buffer (5x Qiagen sample buffer, miniprep kit) was used, and the Illumina custom and high resolution markers were prepared following supplier’s instructions. A 10 % agarose gel was run for 1h at 145 volts. The gel was stained for 5 min in SYBR gold and visualised under UV light. Four holes were punched into 200 µl PCR tubes using the tip of a 21 G needle. The entire area between the 145 and 160 bp markers was excised and placed in the 200 µl PCR tubes. The 200 µl PCR tube was placed into a 1.5 ml eppendorf tube and spun for 2 min at 10,000 x g. 400 µl freshly prepared 0.3 M NaCl was added to the gel samples and the DNA was eluted from the gel by rotation ON at 4 °C. The samples were subsequently spun for 5 min at max speed and the supernatant was transferred to a clean tube. In order to eradicate any remaining gel, the supernatant was spun down again for 5 min and a 10 µl pipette tip was inserted in 200 µl pipette tip to transfer the supernatant to a new tube. The DNA was precipitated ON by adding 1153 µl EtOH and 1 µl glycogen and the next day the DNA was eluted in 10 µl EB and the library was quantified using the HighSensitivity Bioanalyzer kit. The small RNA-seq libraries were additionally quantified by Kapa Library Quantification. The libraries were pooled according to their molecular weight and high-throughput sequencing of all libraries was carried out with single-end protocols on a HiSeq 2000 instrument (Illumina).
Chapter 2. Material and Methods

2.3. Methods

ChIP-seq libraries

ESCs were grown on 15 cm dishes coated with 0.1 % gelatine until they were 80 % confluent. Subsequently cells were were cross-linked with 1 % methanol-free formaldehyde in fresh medium for 10 minutes. To quench the cross-linking, 0.2 M final concentration of glycine was added. ESCs were washed twice with ice cold 1 x PBS (137 mM NaCl, 2.7 mM KCl, 10 mM Na₂HPO₄, 2 mM KH₂PO₄ dissolved in 800 ml distilled H₂O, pH was adjusted to 7.4 with HCl) and harvested using a cell scraper. Cells were then pelleted by centrifugation at 8,000 x g at 4°C for 3 min. Pellets were resuspended in LB1 buffer (50 mM HEPES’ KOH, pH 7.5; 140 mM NaCl; 1 mM EDTA; 10 % glycerol; 0.5 % NP-40; 0.25 % Triton X-100, protease inhibitors) for 10 minutes at 4°C, pelleted and resuspended in LB2 buffer (10 mM Tris/HCl, pH 8.0; 200 mM NaCl; 1 mM EDTA; 0.5 mM EGTA, protease inhibitors) for 10 minutes at 4°C. Cells were pelleted and resuspended in LB3 buffer (10 mM Tris/HCl, pH 8; 100 mM NaCl; 1 mM EDTA; 0.5 mM EGTA; 0.1% Na/Deoxycholate; 0.5% N-Lauroylsarcosine, protease inhibitors). Next the cells were sonicated using Misonix Sonicator 3000. Triton X-100 was added to a final concentration of 1 % and the lysate was centrifuged at 20,000 x g for 10 min to pellet the debris. The bead-antibody complexes were prepared before adding the sonicated DNA. Protein G-coupled Dynabeads (Thermo Fisher Scientific, Cat. No. 10003D) and the primary antibodies in PBS with 5 mg/ml BSA were incubated ON. Subsequently, the bead-antibody complexes were added to the sonicated chromatin and both were incubated at 4°C ON. On the following day, beads were washed extensively with RIPA buffer (50 mM HEPES pH 7.6, 1 mM EDTA, 0.7 % Na deoxycholate, 1 % NP-40, 0.5M LiCl), once with 1x TE buffer (1 M Tris-HCl (pH approximately 8.0), 0.1 M EDTA) and eluted in 200 µl of buffer containing 1 % SDS and 0.1 M NaHCO₃. They were then incubated at 65°C ON for reverse cross-linking. RNase A treatment at 37°C was carried out for 1 h, then Proteinase K treatment at 55°C for 2 h. The DNA was then extracted with phenol/chloroform, followed by ethanol precipitation. ChIP-seq library preparation was performed using MicroPlex Library Preparation kit (Diagenode) following manufacturer’s instructions. Libraries were quantified using the High Sensitivity DNA Bioanalyzer kit and Kapa library quantification. Equal amounts
of each indexed library were pooled for multiplex sequencing on a HiSeq 2500 Illumina system (50 bp single-end).

2.3.15 Sequencing

Sequencing was performed at the Wellcome Trust Sanger Institute and the Babraham next generation sequencing facility. Libraries were sequenced on either an Illumina MiSeq or an Illumina HiSeq using the default RTA analysis software.

2.3.16 Bioinformatics

For all experiments involving next-generation sequencing (NGS), raw sequence reads were trimmed to remove poor quality reads and adapter contamination using Trim Galore (v0.4.1). All data was analysed using SeqMonk (www.bioinformatics.babraham.ac.uk/projects/seqmonk/).

WGBS-seq

After the raw sequences were trimmed, the remaining sequences were mapped using Bismark (v0.14.4) (Krueger and Andrews, 2011) with default parameters to the mouse reference genome NCBIM37 (mm9) using paired-end mapping mode. Following this, the reads were deduplicated and the CpG methylation calls were extracted by running the Bismark methylation extractor (v0.14.4) in paired-end mode. Data were quantitated using SeqMonk (www.bioinformatics.babraham.ac.uk/projects/seqmonk/). Probes were defined to span 50 CpG sites and have a minimum read count of 4. The global methylation level was used for calculation of the bean plots covering genomic features defined as follows: exons (probes overlapping exons), introns (probes overlapping introns), promoters (probes overlapping 1000 bp upstream of genes), CGI promoters (promoters containing or within 250 bp of a CGI), non-CGI promoters (all other promoters), intergenic (probes not overlapping with gene bodies), LMRs (Stadler et al., 2011), H3K27ac and
H3K4me1 enhancers (Creyghton et al., 2010). The imprinted control regions were used from Tomizawa et al., 2011.

**Total RNA-seq**

The trimmed reads were mapped to the mouse NCBIM37 (mm9) genome assembly using TopHat v2.0.12 (Trapnell et al., 2009). Strand specific quantification of the data was quantitated overlapping mRNAs using the RNA-seq quantitation pipeline in SeqMonk software (www.bioinformatics.babraham.ac.uk/projects/seqmonk/). Differentially expressed (DE) genes were determined using DESeq2 (p-value 0.05, with multiple testing correction) and an intensity difference filter (p-value 0.05, with multiple testing correction) (Love et al., 2014). The DE genes with high confidence were defined as the genes which were called as significant by both statistical tests.

Global pervasive transcription, defined as genic antisense transcription, was calculated as following. The antisense transcription overlapping protein coding genes was quantitated for WT samples as well as for each time points after *Dnmt1* KO, using a binomial test with a false discovery rate (FDR) of p < 0.05. Additionally, the raw antisense transcription counts for all samples was calculated and significant differential antisense expression was calculated using DESeq2 with an FDR < 0.05. The overlap of the two quantifications was used to define pervasive transcription. And the difference in antisense transcription between WT and KO samples at each time point was plotted using R.

**Gene Ontology**

Functional annotation enrichment analyses were performed using Panther webtools (Mi et al., 2016).
Small RNA-seq

For small RNA-seq, trimmed sequencing reads were aligned to mouse genome assembly NCBIM37 (mm9) using Bowtie2 \cite{Langmead2012}. For miRNA analysis, probes were generated overlapping CDS (coding DNA sequence) and read count enrichment was calculated normalised by total read counts (RPM = reads per million mapped reads) using Seqmonk. Differentially expressed genes were determined using DESeq2 (p-value 0.05, with multiple testing correction) and an intensity difference filter (p-value 0.05, with multiple testing correction) \cite{Love2014}. The high confidence DE genes were defined as the genes which were called as significant by both statistical tests.

For transposon analysis, small RNA-seq data was analysed using the small RNA-seq pipeline piPipes (https://github.com/bowhan/piPipes) \cite{Han2015}. Trimmed data were mapped using Bowtie against the mm9 genome. Additionally, the trimmed reads were mapped to different annotations: piRNA annotations were defined earlier \cite{Li2013a}. Repeats were defined during the analysis using the mouse repeatmasker annotation (http://www.repeatmasker.org). The plots shown were generated as described below. The distribution of small RNAs was computed by mapping all small RNA-seq reads to the different genomic features. The length distribution was calculated taking all uniquely mapped small RNAs into account. Small RNA reads were then pre-filtered as follows: reads mapping to rRNAs and miRNAs were excluded, then reads aligning to the repeat masked mm9 genome (all annotated repeats were masked) were removed. The resulting small RNA reads were mapped to all repeats. The 5’ end nucleotide composition was computed from the uniquely mapped small RNA reads. Similarly, analysis of the position of 5’ to 5’ overlap was performed on the mapped small RNA reads, and the length distribution and strand orientation of small RNAs shown was generated using uniquely mapped small RNA reads.

For consensus sequence mapping, small RNAs were mapped to TE consensus sequences from repeatmasker using Bowtie (v1.0.1; default parameters).
ChIP-seq

For ChIP-seq, trimmed sequencing reads were aligned to mouse genome assembly NCBIM37 (mm9) using Bowtie2 with default parameters. For calling enrichment, read count quantitation in Seqmonk was used. Probe trend plots and probe alignment plots were generated by calculating average CpG methylation levels of 1 kbp overlapping probes from 5 kbp upstream of the transcriptional start site through gene bodies (which were scaled for visualisation) to 5 kbp downstream of the transcriptional end site.

Repeats Analysis

Repeat locations for all repeat classes of interest were extracted from the pre-masked repeatmasker libraries (mouse, repeatmasker v4.0.3, library version 20130422). Instances of repeats within 2 kb of an annotated gene (defined by Ensembl) were removed to prevent mixing signals from genic expression with expression of repetitive sequences. We excluded all TEs which had less than 1000 calls in each dataset. Bisulfite-seq libraries were processed and mapped as described above. The methylation levels at the instances of repeats were calculated as the sum of the percentages of methylated Cs over all Cs for all methylated and non-methylated calls of each repeat class. The methylation retention was plotted as scatterplot or line graph using R. Total RNA-seq sequences were processed and mapped as described above. Non-directional overlaps were quantitated between the mapped RNA-seq reads and the instances of repeats. Subsequently, the counts of all instances of a repeat class was summed and the sum was corrected for the total length of all repeats and the size of the individual libraries to generate RPKM (reads per kilobase per million mapped reads) expression values. The heatmaps were then generated using the R pheatmap library. For the sense/antisense transcription over repeats, the percentages of sense and antisense transcription over repeats were calculated. Additionally, the bias of sense transcription was calculated as the percentage of antisense transcription subtracted from the sense transcription. Furthermore, the overlap of TEs with genes, as defined by
Ensembl, was calculated.
For ChIP-seq analysis, the sequences were processed and mapped as described above. The counts of all instances of repeat classes were summed and the sum was corrected for the total length of all repeats and the size of the individual libraries to generate RPKM expression values. The heatmaps were then generated using the R pheatmap library.
Chapter 3

Acute demethylation upon $Dnmt1$ conditional KO

3.1 Introduction

During early mammalian development, DNA methylation and other repressive epigenetic marks are erased in mammalian primordial germ cells (PGCs) and the early embryo. Therefore, this represents a critical period of genome defence as transposable elements (TEs) can potentially mobilise, presenting a danger to genome integrity. As such during this period alternative silencing pathways are needed to limit the activity of TEs in the absence of DNA methylation (Allis and Jenuwein 2016).

A recent study has shown that during early developmental epigenetic reprogramming the DNA maintenance methylation machinery is the key driver of $in$ $vivo$ DNA demethylation (von Meyenn et al. 2016). A $Dnmt1$ KO may therefore serve as a good $in$ $vitro$ system to recapitulate $in$ $vivo$ demethylation dynamics.

In this chapter, I used $Dnmt1$ conditional knockout (KO) mouse ESCs to study the alternative regulatory mechanisms involved in TE regulation during DNA hypomethylation.

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3.1. Introduction

Chapter 3. Acute demethylation upon Dnmt1 conditional KO

In order to characterise the in vitro system and to be able to compare it to in vivo epigenetic reprogramming, I analysed the effects of global demethylation upon Dnmt1 on the whole genome using whole genome bisulfite sequencing (WGBS-seq), long and short RNA sequencing as well as chromatin profiling of repressive histone marks (Figure 3.1).

I compared differentially methylated regions (DMRs) which resisted global hypomethylation in our in vitro system with variably erased CGIs (VECs) during epigenetic reprogramming (Seisenberger et al., 2012).

Furthermore, as several genes and miRNAs have been described to play an important role during PGC development, I investigated whether the acute deletion of Dnmt1 resulted into transcriptional changes of these genes.

Earlier research has shown that H3K9me2 becomes lost in PGC development, while H3K27me3 is globally enriched in the genome during the formation of the future germ cells (Hajkova et al., 2008, Seki et al., 2007). Additionally, a remodelling of H3K9me2/3 and H3K27me3 is also found in preimplantation embryos (Santos et al., 2005, Liu et al., 2016, Zheng et al., 2016). Accordingly, I looked at the distribution of repressive histone marks - H3K27me3, H3K9me2 and H3K9me3 - upon global demethylation induced by Dnmt1 deletion.

Figure 3.1: Schematic of in vitro Dnmt1 deletion time-course in mouse ESCs.

Having characterised my Dnmt1 KO in vitro system I began to study the effects of Dnmt1 KO induced hypomethylation on TE elements. It is especially crucial during early phases of development to protect the genome from highly active TE classes as this could lead to germline mutations which can be passed on to future generations.
DNA methylation is essential for TE suppression, however this epigenetic modification is globally erased during epigenetic reprogramming \cite{Walsh_1998}. Further, it has been shown that specific TE classes become upregulated during this global hypomethylation \emph{in vivo} PGC development \cite{Molaro_2014} upon these are the young classes of ERVs - like IAPs and ETns - that still possess the ability to retrotranspose and whose transcriptional activity therefore could result into retrotransposition. Thus, I wanted to investigate whether additional epigenetic modifications play a role in TE transcriptional silencing in the absence of DNA methylation during epigenetic reprogramming.

Earlier research has discovered increased pervasive transcription over TEs in ESCs in comparison to somatic cells \cite{Kelley_Rinn_2012}. It was shown in yeast, that low level of genome-wide pervasive transcription, antisense to genic transcription, can initiate RNAi as a defence mechanism against TEs \cite{Cruz_Houseley_2014}. The regulatory function of an antisense RNA, sense to a protein-coding RNA, has long been suggested \cite{Derrien_2012}. Sense/antisense transcription allows the production of double stranded RNA (dsRNA), which is a target for RNAi. dsRNA viruses are controlled by RNAi in plants, insects, lower eukaryotes as well as mammalian cells (reviewed in \cite{Ding_2010, Li_2013b, Maillard_2013}). Furthermore, RNAi can prevent TE mobility in mammals \cite{Yang_Kazazian_2006, Babiarz_2008}. However, in neither of the studies the source of the dsRNA has been identified and therefore it remained ambiguous how the host genome defence pathway identify TE expression to control their activity.

In this study, I wanted to investigate whether pervasive genome-wide transcription, could feed into an RNAi pathway to defend against TE transcriptional activity, similarly to the mechanism that has been found in yeast. Consequently, I tested whether small RNAs were getting enriched at transcriptionally active TE classes upon global hypomethylation by \emph{Dnmt1} KO.

In addition to DNA methylation, and small RNAs, chromatin has been studied as a mechanism to control TE activity. Knockout of the H3K9me3 histone methyltransferase ESET led to transcriptional upregulation of IAP elements in PGC development.
3.2 Results

3.2.1 Acute deletion of Dnmt1 as a model for global demethylation dynamics

To follow DNA demethylation dynamics upon acute deletion of Dnmt1 in ESCs, I used WGBS-seq, sampling DNA at different time-points after Dnmt1 KO induction (list of WGBS-seq datasets in appendix). I mapped the WGBS-seq libraries to the mouse genome (mm9) using the Bismark alignment tool (Krueger and Andrews, 2011) and did the following analysis using the Seqmonk interface. I defined 50 CpG windows and used the bisulfite pipeline in Seqmonk to analyse the methylation level of the probes in the different libraries. Confirmed knockout of Dnmt1 (Dnmt1$^{-/-}$) compared to wildtype (WT) (Dnmt1$^{fl/fl}$) ESCs led to global demethylation in all genic and intergenic features (Figure 3.2). Dnmt1 KO induced hypomethylation started at day 3 and was only completed on day 6 (Figure 3.4). In general CGIs were always hypomethylated in WT and Dnmt1 KO ESCs in comparison to the rest of the genome. Furthermore, low methylated regions (LMRs) along with enhancer regions, as defined by occupancy of H3K4me1 and H3K27ac histone modifications, followed whole genome demethylation dynamics (Figure 3.3).

I used the Seqmonk binomial test for 100 CpG regions which showed unusually high
Figure 3.2: Genome demethylation upon Dnmt1 deletion. CpG methylation levels measured by WGBS of WT (grey) and conditional Dnmt1 KO ESC induced for 1 day (dark red), 3 days (light red), 6 days (light pink), 9 days (light blue), 11 days (dark blue). Methylated cytosines were counted for each rolling 50 CpG window genome-wide and are expressed as percent of total cytosines per window.

Figure 3.3: Genome demethylation in LMRs and enhancer regions. CpG methylation levels measured by bisulfite sequencing of WT (grey) and conditional Dnmt1 KO ESC induced for 1 day (dark red), 3 days (light red), 6 days (light pink), 9 days (light blue), 11 days (dark blue). Methylated cytosines were counted for each rolling 50 CpG window genome-wide and are expressed as percent of total cytosines per window. Enhancer regions are defined by H3K4me1 and H3K27ac marks.
3.2. Results  Chapter 3. Acute demethylation upon Dnmt1 conditional KO

or low methylation in all of the WT to Dnmt1 KO comparisons. Despite global hypomethylation with similar kinetics in all genomic features, I identified 773 differentially methylated regions (DMRs), which consistently changed after Dnmt1 KO - 20 DMRs were hypomethylated and 726 DMRs were hypermethylated (List of top 50 DMRs in appendix).

I carried out a gene ontology (GO) analysis of the DMRs resistant to demethylation, using the Panther webtool (Thomas et al., 2006). I found significant enrichment of biological terms such as neuronal development and cell differentiation (Figure 3.5).

The FGF signalling pathway also retained more DNA methylation in comparison to the rest of the genome. Nevertheless, I did not find any overlap between resistant regions (VECs) in in vivo PGC reprogramming and the Dnmt1 KO ESC system, other than IAPs, which resisted global demethylation as I will expand on later in this chapter.

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Figure 3.5: Regions resisting global demethylation upon Dnmt1 KO. GO term analysis of all 773 regions which resisted global demethylation from day 6 after Dnmt1 KO onwards. Only 258 genes were found with a mapped GO term.
Chapter 3. Acute demethylation upon \textit{Dnmt1} conditional KO

3.2. Results

To permit a comparison between gene expression and global demethylation, I carried out total RNA-seq analysis of the same time-points after \textit{Dnmt1} KO, as I did for WGBS-seq (list of total RNA-seq datasets in appendix). Total RNA-seq was performed to allow for transcriptional analysis of all TE classes, also those classes that lack a poly adenylation (poly-A) tail.

3.2.2 \textit{Dnmt1} KO leads to transcriptional upregulation of imprinted loci

I did a pairwise comparison of genes upregulated between \textit{Dnmt1} KO and WT ESCs at each time-point and checked for differentially expressed genes. For this, I used the Seqmonk intensity difference filter with Benjamini and Hochberg correction for multiple testing with a p-value threshold of < 0.05 and overlapped them with the genes called differentially expressed by DESeq2 with a p-value threshold of < 0.05 and multiple testing correction (Love et al., 2014). Apart from \textit{Dnmt1} itself, none of the other genes showed any real change at the first time-point after KO induction (Day 1: 3 differentially expressed (DE) genes). However, an increased regulation with time thereafter (Day 3: 43 DE genes, Day 6: 357 DE genes, Day 9: 269 DE genes, Day 11: 50 DE genes) was found (List of differentially expressed genes upon \textit{Dnmt1} KO in appendix). Furthermore, I called significantly differentially expressed genes between WT and \textit{Dnmt1} KO from day 3 onwards. I identified 85 differentially expressed genes by using the overlap of genes identified by DESeq2 and Seqmonk intensity difference with a p-value of < 0.05. The genes were consistently up or down regulated in response to \textit{Dnmt1} KO.

There was a large bias for genes to be upregulated in response to the \textit{Dnmt1} induced DNA demethylation (n = 80), but there were also a small number of genes (n = 5) which were downregulated in response to global hypomethylation (Figure 3.6).

Consequent to \textit{Dnmt1} deletion, \textit{Dnmt1}, \textit{Igfbp2} and \textit{Grb10} were downregulated, whereas the imprinted genes \textit{Xlr3a}, \textit{Miry} and \textit{Rian} were upregulated (Figure 3.7). Furthermore, genes important for PGC development - \textit{Dazl}, \textit{Lefty2}, \textit{Eif2z3y}, \textit{Zscan4d/f} - were up-
regulated after global DNA hypomethylation.

**Figure 3.6: Genes respondent to Dnmt1 driven demethylation.** Scatter plot of total RNA-seq data mapped to the whole genome at day 9 after Dnmt1 KO compared with WT ESCs. Genes significantly differentially expressed from day 6 after Dnmt1 deletion onwards (black), genes which did not change expression levels significantly (grey) across the whole time-course.

**Figure 3.7: Highest respondent genes upon Dnmt1 deletion.** Bar graph of RNA-seq of differentially expressed genes between WT (grey) and conditional Dnmt1 KO ESCs induced for 0 days (black), 1 day (dark red), 3 days (light red), 6 days (light pink), 9 days (light blue), 11 days (dark blue). Data shows RNA-seq data of 2 biological replicates of each time-point.

The pluripotency network *Oct4, Nanog, Klf4, Esrrb* and *Sox2* - was normally expressed after knockout of the DNA maintenance methylase (Figure 3.8).
Chapter 3. Acute demethylation upon Dnmt1 conditional KO

3.2. Results

I next investigated the correlation between gene transcription and promoter methylation throughout the time-course of global DNA hypomethylation. To do this, I grouped the genes in the WT ESCs into not expressed (RPM < 0), low expressed (2.5 < RPM > 0), expressed (5 < RPM > 2.5), high expressed (10 < RPM > 5) or very high expressed (RPM > 10) and then looked at their methylation level throughout the time-course after Dnmt1 deletion. Genes highly expressed in WT ESCs (RPM > 10) had much lower initial promoter methylation than other genes, in spite of this the expression of all other genes was independent of their promoter methylation level. Additionally, I compared the DNA methylation level of genes which were Dnmt1 dependent, but they
also demethylated with the same kinetics as the rest of the genome (Figure 3.9).

3.2.3 Small RNAs from Dlk and X-chromosome locus Xlr3 become upregulated upon Dnmt1 KO

In order to analyse whether Dnmt1 deletion had any effect on the expression of small RNAs, I carried out small RNA-seq at each time-point after Dnmt1 KO and compared it to the respective WT ESCs (list of small RNA-seq datasets in appendix). I mapped the small RNA-seq libraries to the whole genome, and by using the Seqmonk small RNA-seq quality control plot found that over 90% of reads mapped to miRNAs. The small RNAs were 20-24nt long (Figure 3.10A) and endogenously expressed miRNAs in ESCs were expressed throughout the whole time-course after Dnmt1 deletion (Figure 3.11). Moreover, reads for both the 3’ arm as well as the 5’ arm of the mature miRNA mapped to known endogenously transcribed miRNAs in ESCs (Figure 3.10B).

![Figure 3.10: Small RNA-seq libraries are 90% made up of miRNAs.](image)

(A) small RNA size distribution as well as classification of different small RNA classes in Dnmt1 KO and WT ESCs, miRNAs (grey), rRNA (green), small nuclear RNAs (snRNAs) (violet), miscellaneous other RNAs (misc RNAs) (red), small nucleolar RNAs (snoRNA) (orange) and tRNA (light blue). (B) genic location of miRNA 200c with reads mapped in Dnmt1 KO and WT ESCs, each line representing one read.

Next, I wanted to test whether any of the miRNAs were differentially expressed upon global hypomethylation. To address this question, I carried out pairwise comparison
Chapter 3. Acute demethylation upon Dnmt1 conditional KO

3.2. Results

Figure 3.11: Not significantly altered expression of endogenous miRNAs in ESCs. small RNA-seq of miRNA expression in WT (greys) and conditional Dnmt1 KO ESC induced for 1 day (dark red), 3 days (light red), 6 days (light pink), 9 days (light blue), 11 days (dark blue). Data shows small RNA-seq data of 3 biological replicates of each time-point.

of Dnmt1 KO and WT samples of each time-point and looked at the differentially expressed miRNAs. To call differentially expressed miRNAs, I overlapped the differentially expressed miRNAs using DESeq2 and Seqmonk intensity difference filter with a p-value of < 0.05. Two distinct miRNA loci were upregulated in Dnmt1 KO in comparison to WT ESCs (Figure 3.12). The first locus was a miRNA cluster of the imprinted Dlk locus (Edwards et al., 2008, Charalambous et al., 2004). This locus has been previously shown to drive the expression of noncoding RNAs in ESCs (Labialle et al., 2014 Cavaillé et al., 2002 Seitz et al., 2004). The second locus was part of the imprinted Xlr3 locus on the X Chromosome (list of differentially expressed miRNAs upon Dnmt1 KO in appendix).

In order to verify the small RNA-seq results, I carried out small RNA quantitative real time PCR (qRTPCR) of some of the differentially expressed miRNAs confirming that miRNAs from the Dlk locus, mmu-miR-367 and mmu-miR-543, were significantly upregulated in Dnmt1 KO ESCs in comparison to WT ESCs (Figure 3.13).
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Chapter 3. Acute demethylation upon Dnmt1 conditional KO

Figure 3.12: miRNAs from the Dlk and Xlr3 locus are dependent on Dnmt1. (A) Scatter plot of total RNA-seq data mapped to the whole genome of day 9 after Dnmt1 deletion (y-axis) compared to wildtype (x-axis). miRNAs significantly upregulated upon Dnmt1 KO from the Dlk locus (black), miRNAs significantly upregulated upon Dnmt1 KO from the Xlr3 locus (green). (B) Bargraph of small RNA-seq of differentially expressed miRNAs - of the Dlk and Xlr3 locus - between WT (grey) and conditional Dnmt1 KO ESCs induced for 1 day (dark red), 3 days (light red), 6 days (light pink), 9 days (light blue), 11 days (dark blue). Data shows small RNA-seq data of 3 biological replicates of each time-point. Differences between conditions that are statistically significant are denoted by (* p-value<0.05, ** p-value<0.005, (Student’s t-test))

Figure 3.13: Upregulation of mmu-miR-367 and mmu-miR-543 upon Dnmt1 KO. (A) small RNA qRT-PCR on mature mmu-miR-367 and mmu-miR-543 at day 9 after Dnmt1 deletion (light blue) and wildtype (grey). Each qRT-PCR was done in 3 technical replicates. Differences between conditions that are statistically significant are denoted by (* p-value<0.05, ** p-value<0.005, (Student’s t-test))
In summary, WGBS-seq together with long and short RNA-seq analysis upon Dnmt1 KO allowed me to identify imprinted genes as the highest respondent genes. As an example, I analysed the Dlk locus in more detail and in the WGBS-seq the imprinted control region (ICR) became demethylated upon Dnmt1 deletion. This led to transcriptional upregulation of genes of the Dlk locus as well as upregulation of small RNAs in this imprinted region (Figure 3.14).

3.2.4 No genome-wide chromatin changes upon global demethylation in ESCs

I wanted to know whether any global chromatin rearrangements were taking place subsequently to induction of global DNA demethylation in vitro by Dnmt1 KO.

During global DNA demethylation in vivo PGC development H3K9me2 becomes depleted while H3K27me3 becomes enriched genome-wide (Hajkova et al., 2008; Seki et al., 2007). Furthermore, a remodelling of H3K9me2/3 and H3K27me3 is also found.
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Figure 3.15: Enrichment of repressive histone marks in the genome. Pie chart of enrichment of H3K27me3, H3K9me3 and H3K9me2 in repeats (dark violet), genic regions (light violet), promoters (dark green), CGIs (middle green), intergenic regions (light green) in wildtype ESCs.

in preimplantation embryos (Santos et al., 2005, Liu et al., 2016, Zheng et al., 2016).

To study the connection between global DNA demethylation and chromatin organisation in ESCs, I performed ChIP-seq of three repressive histone marks - H3K9me3, H3K9me2 and H3K27me3.

I prepared ChIP-seq libraries from ESCs at day 4 and day 8 after Dnmt1 KO induction, as well as the respective WT samples (list of ChIP-seq datasets in appendix). Those time-points were chosen because at day 3 post Dnmt1 deletion global hypomethylation was not yet complete, while at day 8 the whole genome was totally hypomethylated. Simultaneously, the transcriptional changes were mostly seen at day 6 after Dnmt1 deletion. To permit ChIP-seq analysis, I carried out a read count quantitation in Seqmonk and normalised to total read counts.

H3K27me3 was more highly abundant in the genome (n=50599 enriched sites) than H3K9me3 (n=30184) and H3K9me2 (n=2541). Additionally, H3K27me3 was mostly enriched in genic regions, CGIs, promoter sites as well as repetitive elements, whereas H3K9me3 was depleted from CGIs, intergenic regions as well as promoters and most of the enrichment of this histone mark could be found in repeats. H3K9me2 was evenly distributed across the whole genome with a slight enrichment at genes and repeats.
Figure 3.16: **Histone marks over gene body and TSS.** Probe enrichment of H3K9me3 (green), H3K9me2 (yellow) and H3K27me3 (blue) over gene body and TSS.

![Histone marks over gene body and TSS](image)

Figure 3.17: **Histone mark enrichment in the genome.** Chromosome view of ChIP enrichment of H3K9me3 (green), H3K27me3 (blue) and H3K9me2 (yellow) over a 500kbp region in Chromosome 12. Intensity of the enrichment on the y-axis.

![Histone mark enrichment in the genome](image)
3.2. Results

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(H3K27me3) was specifically enriched at the transcription start site (TSS) with a depletion at the gene body (Figure 3.16), while H3K9me2/3 were depleted in genic regions. The same trend could be seen in a specific example of a 500 kilobase (kb) region on chromosome 2: H3K9me3 was specifically enriched in the active class IAPs (IAPEZ), while H3K27me3 was highly enriched at TSSs and H3K9me2 was ubiquitously found across the whole genome (Figure 3.17).

Figure 3.18: Minor redistribution of global histone marks upon Dnmt1 KO. Scatter plot of repressive histone marks overlapping genes in wildtype (y-axis) versus Dnmt1 KO (x-axis) ESCs.

Subsequently, I tested whether any of the histone marks changed enrichment at any regions in the genome upon Dnmt1 deletion. However, only minor changes in H3K9me3, H3K9me2 and H3K27me3 peaks was measured globally upon DNA demethylation induced by Dnmt1 deletion (Figure 3.18).

To investigate the correlation between histone marks and gene expression in ESCs, I compared the expression levels of genes in WT ESCs and overlaid them with their histone enrichment across the time-course after Dnmt1 deletion. In WT ESCs H3K27me3 enrichment was higher over TSS of lowly expressed genes than highly expressed genes (Figure 3.19) in keeping with its repressive role (Marks et al., 2012). However, neither of the histone marks changed enrichment over genes throughout the time-course of global hypomethylation, nor did any of the repressive marks change enrichment at genes whose expression changed upon global hypomethylation (Figure 3.20).

Finally, I wanted to study the correlation between enrichment of repressive histone
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**Figure 3.19:** H3K27me3 strong enrichment at TSSs of lowly expressed genes. Aligned probe plot of H3K27me3 enrichment surrounding 5kb of TSS.

**Figure 3.20:** No change of histone marks in expressed genes across *Dnmt1* KO. H3K9me3, H3K27me3 and H3K9me2 read enrichment over genes which showed different expression level in WT ESCs: not expressed (RPM < 0, red), low expressed (2.5 < RPM > 0, violet), expressed (5 < RPM > 2.5, pink), high expressed (10 < RPM > 5, dark blue) or very high expressed (RPM > 10, light blue) as well as genes which were *Dnmt1* dependent (orange) across the time-course after *Dnmt1* deletion.
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Figure 3.21: Histone marks are not enriched on regions dependent on DNA maintenance methylation. Line graph of H3K9me3, H3K27me3 and H3K9me2 read enrichment over very high (dark red) (100-75%) , high (orange) (75-50%), medium (light blue)(25-50%) and non methylated (blue)(0-25%) promoter regions before and after Dnmt1 deletion.

marks and global methylation state of promoters after Dnmt1 deletion.

To approach this, I categorised the promoter methylation levels at day 11 after Dnmt1 deletion into four categories: (1) hypermethylated (75-100% methylation), (2) methylated (75-50% methylation), (3) low methylated (25-50% methylation) and (4) hypomethylated (0-25% methylation).

This allowed me to compare the histone enrichment specifically at promoters which resisted global demethylation. But despite the fact that some promoters resisted global demethylation, and still had 75% of methylation at day 11 after Dnmt1 deletion, neither of the repressive histone marks was enriched at those promoters in comparison to the low level methylated promoters (0-25% methylated) (Figure 3.21).

In our in vitro system of Dnmt1 conditional KO I was able to induce global demethylation and found mostly imprinted loci that became transcriptionally activated, while the pluripotency network continued to be expressed normally. Therefore, the conditional KO ESCs proved to be a good model to study the regulation of TEs during global demethylation dynamics in vitro.

3.2.5 Mapping of TEs in next generation sequencing libraries

In order to analyse DNA methylation, transcription and histone enrichment across TEs, with the help of Simon Andrews we mapped all reads to the mouse genome allowing only unique mapping of reads. Subsequently, we ignored all reads that were within 2kb
of genes. TEs are frequently integrated into genes or lie within close proximity of genes. This leads to wrongly annotated 3'UTRs and therefore reads which are derived from genic expression would be wrongly ascribed to TE transcription (Figure 3.22). These reads would also be seen in consensus sequence mapping. Due to this conservative approach to mapping TEs we are likely to lose information on recently integrated TEs, as we only allow unique mapping. We will probably underestimate TE methylation and transcription levels but on the other hand we can be confident about the TE transcription to be real. For small RNAs, we did consensus sequence mapping to TE classes, as not many genic reads are expected in small RNA-seq libraries.

**Figure 3.22: Mapping of TEs in WGBS-seq, RNA-seq and ChIP-seq libraries.** Sequencing reads overlapping genes and transposons sitting within 2kb of genes.

### 3.2.6 TE classes become demethylated upon Dnmt1 KO in ESCs

The Dnmt1 conditional KO system allowed me to study DNA methylation, small RNA contribution as well as chromatin changes as potential regulators of TE transcriptional activity. First I analysed the demethylation of TEs upon Dnmt1 KO.

Acute Dnmt1 KO led to hypomethylation of all transposon classes (Figure 3.23). Nevertheless, some TEs had altered demethylation kinetics compared to the rest of the genome (Figure 3.24).

MMERVK10C always remained more methylated than the rest of the genome and IAP
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Figure 3.23: TE family demethylation upon Dnmt1 deletion. CpG methylation levels measured by WGBS of wild type (grey) and conditional Dnmt1 knockout ESC induced for 1 day (dark red), 3 days (light red), 6 days (light pink), 9 days (light blue), 11 days (dark blue). Methylated cytosines were counted for each rolling 50 CpG window genome-wide and are expressed as percent of total cytosines per window.

elements resisted demethylation until day 6 and then followed genome demethylation kinetics (Figure 3.24). In regard to IAP elements this recapitulates in vivo demethylation dynamics, as IAPs are also resistant to global demethylation during primordial germ cell development (Seisenberger et al., 2012).

3.2.7 Dnmt1 KO leads to transcriptional activation of ERV elements

To analyse whether any classes of TEs were dependent on DNA maintenance methylation for their transcription, I mapped the total RNA-seq data uniquely to the genome and checked for transcriptional activation of TEs. Although all TEs became demethylated only a small number of TE classes were transcriptionally upregulated (Figure 3.25). I grouped the TE elements into 3 categories: (1) members of the “white” category were constantly expressed, (2) members of the “black” category increased in transcription upon Dnmt1 KO and (3) members of the “grey” category were transcriptionally upregulated and subsequently resilenced (in the absence of DNA methylation). In summary, regardless of global TE hypomethylation, with the aforementioned exceptions,
Figure 3.24: Resistance to demethylation of some TEs. (left) Scatter plot of DNA demethylation at day 11 after Dnmt1 deletion with red dot indicated TE family methylation level, (right) genome background model of demethylation dynamics set to zero (grey line) and demethylation dynamics of TE classes (black line) above.
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only a very specific class of endogenous retroviruses (ERVs) became transcriptionally upregulated upon Dnmt1 KO (white category not shown) (Figure 3.25).

Figure 3.25: TE classes upregulated upon Dnmt1 KO. Heatmap of RNA-seq data mapped to TE classes. Only TE classes which were intergenic and within 2kb of genes were considered. Mean of TE classes with at least 1000 integration sites were regarded. RNA-seq was done in duplicates at each time-point with day 0 being the compiled mean of the control RNA-seq datasets.

Figure 3.26: Unique TE elements upregulated upon Dnmt1 KO. RNA-seq data of day 0, day 6 and day 9 upon Dnmt1 KO. Average transcription is shown as read line. Differences between conditions that are statistically significant are denoted by (* p-value<0.05, ** p-value<0.005, (Student’s t-test))

LINE and SINE elements were constantly expressed irrespective of global hypomethylation and fell into the “white” category. MMERVK10C increased in expression upon DNA hypomethylation and made up the “black” category. Members of all 3 families of
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ERVs were expressed and silenced upon Dnmt1 KO, and were therefore members of the “grey” category: ERV1 elements - ORR1B1, RLTR1B, RLTR9E, RLTR45, ERV2 elements - IAP-d, IAPLTR3, IAPEZ, ETnERV2, MMERVK9C and ERV3 elements - MERVL.

I examined single intragenic TE insertions of each category - 1: L1MdGf; 2: MMERVK10C; and 3: IAPs and MERVL; and found that their transcriptional profiles recapitulated the mean profiles of the respective classes, which confirmed that transcriptional upregulation was not driven by a minority of elements but was rather a class wide response (Figure 3.26).

3.2.8 Sense/antisense transcription of repeat families feed into the RNAi pathway

Earlier studies in yeast have shown that low level of genome-wide pervasive transcription, antisense to genic transcription, can initiate RNAi as a defence mechanism against TEs (Cruz and Houseley, 2014).

With the help of Simon Andrews I performed an analysis of antisense transcription in the genome. In general, there was increased pervasive transcription - defined as genic antisense transcription - upon global hypomethylation induced by Dnmt1 deletion (Figure 3.27). Upon further investigation, it transpired that the pervasive genic transcripts that we detected were in fact produced by transcription of TEs that had integrated in antisense direction to these genes. Indeed, 94.9 % of the genic antisense transcripts overlapped with TEs which were upregulated upon Dnmt1 KO (Figure 3.28). A genome-wide study of genic integration of TE elements confirmed this trend, as TEs were generally depleted in genes; however when overlapping they were inserted in the antisense direction to genes (Figure 3.29). To investigate this further, we analysed my total RNA-seq data to determine whether we could observe both sense and antisense transcription at sites of TE integration.
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Figure 3.27: Increased genome-wide pervasive transcription upon Dnmt1 KO. Quantification of genic antisense transcription increases upon Dnmt1 KO. The difference in Dnmt1 KO versus WT ESCs was calculated at each time point after induced Dnmt1 KO.

Figure 3.28: Pervasive transcription overlaps with TEs. Chromosome view of genic transcription with an integrated TE insertion. Sense reads (red), antisense reads (blue).

Figure 3.29: TE orientation antisense to genes. Quantifying TE insertion sense (red) or antisense (blue) to genes as well as percentage of TEs overlapping genes. The distribution and percentage of all repeats served as a control.
Figure 3.30: Sense/antisense transcription in TE families. Barplots of sense/antisense strands of RNA-seq data mapped to TE families in conditional Dnmt1 knock-out ESC induced for 0 days (black), 1 day (dark red), 3 days (light red), 6 days (light pink), 9 days (light blue), 11 days (dark blue), 17 days (dark green) and 25 days (light green). Sense (filled), Antisense (hatched).

To do this we mapped both sense and antisense reads to the different TE classes and showed that TE antisense transcription was found in all of the TE classes, while only the sense transcripts of members of the “grey” TE category were upregulated (Figure 3.30).

As discussed previously, TE antisense transcription is a result of sense transcription of the genes in which the TEs are integrated. As such we were interested in whether genic sense transcription, could actually be serving as a trap for TE transcriptional activity, in order to control expression of TEs. This system may function as follows: the production of sense and antisense transcripts over TEs leads to the production of dsRNAs, which subsequently feed into the RNAi pathway thus silencing TEs post transcriptionally (Figure 3.32). To test this hypothesis, I performed small RNA-seq at...
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time-points after Dnmt1 deletion.

![Model of TE regulation by RNAi](image)

Figure 3.31: Model of TE regulation by RNAi. Sense and antisense transcription of TE elements with genic sense transcription working as a trap of TE sense transcriptional upregulation. This leads to the production of dsRNA and allows for the production of small RNAs which can in turn regulate transcription of TEs.

3.2.9 Small RNAs are produced from TEs upon loss of Dnmt1

To evaluate whether small RNAs played a role in resilencing of the TE elements in our system, I carried out small RNA-seq at different time-points after Dnmt1 deletion. I mapped the small RNA-seq libraries to the consensus sequences of TE classes defined by repeatmasker (www.repeatmasker.org/). A significant increase of small RNAs that mapped to IAPLTR1a, MERVL, MMERVK10C and ETn elements upon Dnmt1 KO was detected. No increased amounts of small RNAs were detected mapping to any of the other classes of TEs, such as L1MdGf and RLTR45, upon KO of the DNA maintenance methylase Dnmt1 (Figure 3.32).

In order to better characterise the nature of the small RNAs which mapped to TEs, I concentrated on small RNAs which mapped to the IAP consensus sequence. The small RNAs mapped across the full length of the IAP transcript. There was no specific small RNA enrichment at the 3’UTR, a characteristic for miRNAs (Figure 3.33). Consequently, I wanted to next check whether or not the small RNAs were bona fide players of the RNAi pathway.
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Figure 3.32: Small RNAs map to TEs. Bar graph of Small RNA-seq libraries in wild type (grey) and conditional Dnmt1 knockout ESC induced for 1 day (dark red), 3 days (light red), 6 days (light pink), 9 days (light blue), 11 days (dark blue), 25 days (light green). Small RNA-seq libraries were done in 3 biological replicates for each time-point. Each qRT-PCR was done in 3 technical replicates. Differences between conditions that are statistically significant are denoted by (* p-value<0.05, ** p-value<0.005, (Student’s t-test))

Figure 3.33: Small RNAs map to the IAP consensus sequence. Chromosome view of small RNA-seq library reads of Dnmt1 KO and WT ESCs at each time-point mapped to the IAPEZ consensus sequence. Each line represents one read.
3.2.10 Small RNAs are actively loaded into the RNAi machinery

An essential component of the RNA induced silencing complex (RISC) is one of the four ARGONAUTE proteins (AGO). While all AGOs can bind small RNAs, only AGO2 has the ability to complete endonucleolytic slicing of the target mRNA, which is characteristic of endosiRNAs (Meister et al., 2004, Liu et al., 2004, Song et al., 2004, Okamura et al., 2004, Rand et al., 2005, Matranga et al., 2005).

In order to test whether the small RNAs mapping to IAPs are bona fide siRNAs, I carried out an AGO2 immunoprecipitation (IP) and analysed the small RNA content of the pulldown by small RNA-seq (Figure 3.35A).

I mapped the small RNAs to the whole genome as well as repeat classes using the piPipes small RNA-seq pipeline (Han et al., 2015).

![Figure 3.34: AGO2 small RNA-seq libraries.](image)

(A) Pie chart of AGO2 libraries mapped to genome and repeatome. miRNAs (black), 5'UTR bound small RNAs (light green), 3'UTR bound small RNAs (yellow), intron bound small RNAs (dark blue), repeats bound small RNAs (dark green), rRNA and tRNA (grey), unannotated small RNAs (white). (B) Bar graph of size distribution of sense and antisense reads mapped to all repeats.

The AGO2 pulldown was successful, as 90% of small RNAs mapped to the whole genome were miRNAs for both KO and WT sample (data not shown). In the Dnmt1 KO ESCs at day 9 after induction, 40% of the remaining small RNAs mapped to repeats (Figure 3.34A, WT data not shown).

The small RNAs that mapped to repeats were 22 nucleotides (nt) long and mapped to
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Figure 3.35: AGO2 siRNAs map to TE classes. (A) Graphical depiction of AGO2 pulldown of small RNAs. Ago2 antibody is used to pulldown AGO2 protein with small RNAs bound. These small RNAs are released subsequently and subjected to small RNA-seq. (B) Bargraph of small RNA-seq libraries mapped to TE classes. AGO2 pulldown were done in 4 biological replicates at day 9 after Dnmt1 KO. Differences between conditions that are statistically significant are denoted by (* p-value<0.05, ** p-value<0.005, (Student’s t-test))

both sense and antisense strands of all repeat classes (Figure 3.34B).

I found small RNAs that mapped to all TEs in WT ESCs, but they significantly increased upon Dnmt1 KO in IAPEZ, ETn, MMERVK10C and L1MdGf (Figure 3.35B). The small RNAs mapping to TEs had the specific characteristics of endosiRNAs, as they were 22 nt long, had a 5’T overhang at nucleotide 20 and formed 5’-5’ overlaps to each other at nucleotide 20 (Figure 3.36). Small RNAs mapped sense and antisense to all TE elements (Figure 3.37). There was no strand bias for small RNAs mapping to LINE-1 elements, while for IAPs and ETn elements there was a sense strand bias.

3.2.11 EndosiRNAs and not miRNAs play a critical role in resilingencing of IAPs

To address the question whether endosiRNAs or miRNAs were playing the primary role in IAP resilingencing I did siRNA knock down (KD) of different components of the RNAi pathway in Dnmt1 KO ESCs. DGCR8 works upstream of DICER in the miRNA pathway, but not of the siRNA pathway, and siRNAs are exclusively bound by endonucleolytically active AGO2. I did a KD of Ago2, Dicer as well as Dgcr8 in the Dnmt1
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Figure 3.36: Analysis of AGO2 bound small RNAs that mapped to repeats. (A) Bargraph of 5’5’ overhang of small RNAs mapped to repeats, (B) Bargraph of nucleotide contribution along the stretch of the small RNAs mapped to repeats.

Figure 3.37: Sense/antisense small RNAs map to TE families. Bargraph of sense (blue) and antisense (red) small RNA reads mapping to TE families at day 9 after Dnmt1 deletion with WT in transparent. Mapping of unique reads to repeatmasker repeat elements using piPipes small RNAseq pipeline.
KO at day 12 after KO induction and checked for IAP expression to identify whether the miRNA or endosiRNA pathway played the primary role in IAP silencing.

**Figure 3.38: IAP is upregulated upon Dicer and Ago2 KD.** (A) Graphical depiction of knockdown study, IAP expression (red), small RNA expression (grey). (B) qRTPCR of IAP expression after siRNA knockdown (KD) of Dicer, Ago2, Dgcr8 in comparison to non-targeting control. Every siRNA KD was done in 3 technical replicates. Differences between conditions that are statistically significant are denoted by (* p-value<0.05, ** p-value<0.005, (Student’s t-test)).

**Figure 3.39: siRNA knockdown of RNAi pathway.** qRTPCR of Dicer, Ago2 and Dgcr8 with (+) or without (-) siRNA. Every siRNA KD was done in 3 technical replicates. Differences between conditions are statistically significant are denoted (* p-value<0.05 (Student’s t-test)).

I performed the RNAi experiment at day 9 after Dnmt1 KO to allow for transcriptional upregulation of IAPs. The expression of Dgcr8, Ago2 and Dicer was downregulated to 20% of their normal expression level by siRNAs (Figure 3.39). I measured the
expression of IAPs by qRTPCR at day 12 after Dnmt1 KO (Figure 3.38A). RNAi against Dicer8 and the scrambled non-targeting control siRNA resulted in a resilingencing of the IAPs, whereas in the Dicer and Ago2 KDs resilingencing of IAPs was impaired (Figure 3.38 B).

I therefore propose that endosiRNAs are primarily involved in IAP resilingencing.

### 3.2.12 Histone modifications may account for different behaviour of TE families upon acute Dnmt1 deletion

In ESCs IAPs are tightly guarded by H3K9me3 methylation (Mikkelsen et al., 2007). The KO of the histone methyltransferase ESET in PGCs leads to a loss of H3K9me3 on ERV elements - specifically IAPs (Leung et al., 2014). H3K27me3 has been shown to be highly enriched in MERVL elements (Hayashi et al., 2016). Global demethylation by transition from serum to 2i cultured ESCs has shown that repressive histone marks were redistributated to potentially silence TEs (von Meyenn et al., 2016, Walter et al., 2016).

In order to check whether this redistribution of repressive histone marks also played a role in resilingencing TE classes in our system, I carried out ChIP-seq of H3K9me3, H3K9me2 and H3K27me3 before and after Dnmt1 deletion (Day 4 and Day 8). H3K9me3 peaks were constantly found over IAPs, and H3K27me3 was enriched at MERVL elements at day 8 after Dnmt1 deletion in comparison to day 4 (Figure 3.40, 3.41).

Redistribution of repressive histone marks H3K9me2/3 and H3K27me3 could indeed
partially explain the resilencing of certain TE classes, as follows. The TEs could be subdivided into the ones which acquired H3K27me3 or H3K9me2, while some had enrichment of both marks after day 8 of \textit{Dnmt1} deletion. MMERVK10C of the “black” category showed high enrichment of all three histone marks, which could explain the low expression level throughout the course of \textit{Dnmt1} deletion, as these ERVs are known to be highly dependent on H3K9me3 \cite{Maksakova2011}. TE elements of the “grey” category showed varied enrichment of repressive histone marks, as follows. MERVL showed high enrichment of H3K27me3 but depletion of H3K9me2 upon \textit{Dnmt1} KO, which could explain the transcriptional upregulation and resilencing of those ERV elements upon global demethylation (Figure 3.42). H3K27me3 enrichment but loss of H3K9me3 occurred at IAPLTR3 and IAP-d elements. H3K9me3 and H3K27me3 were lost over ORR1B and RLTR14 and those elements were expressed until day 11 after \textit{Dnmt1} deletion, when potentially other mechanisms are being put in place for long term silencing. H3K9me2 switched to H3K9me3 over RLTR1B upon \textit{Dnmt1} KO, while H3K9me2 became depleted and H3K9me3 became enriched at RLTR45, RLTR9E, IAPEZ and MMERVK9C upon \textit{Dnmt1} KO. Differing repressive histone marks also controlled TEs of the “white” category. While LTRSI2 and RMER16 did show increased enrichment of H3K9me2 and me3 upon \textit{Dnmt1} KO, L1MdGf and ERVB4 showed high enrichment of H3K27me3, RLTR14 lost H3K27me3 and H3K9me2 but gained H3K9me3 modifications.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure3.41.png}
\caption{H3K27me3 and H3K9me3 enrichment over MERVL elements. Scatter plot of H3K27me3 and H3K9me3 enrichment over all MERVL elements in the genome at day 4 (dark red) and day 8 (dark blue) \textit{Dnmt1} KO ESCs.}
\end{figure}
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Figure 3.42: Repressive histone marks control TEs upon Dnmt1 KO. Heatmap of ChipSeq data of H3K9me3, H3K9me2 and H3K27me3 mapped to TE families at day 4 and day 8 after Dnmt1 deletion. Read count enrichment. Each library was done with no replicate. Enrichment of histone marks was calculated relative to WT enrichment.

Altogether, the repressive histone marks I analysed after Dnmt1 deletion may explain the repression of some TE classes after transcriptional upregulation. But also TEs, which were already targeted by small RNAs, got enriched by repressive histone marks after Dnmt1 deletion.

In summary, I found demethylation of nearly all TE classes after Dnmt1 deletion. Subsequently, I classified TE elements dependent on their expression following Dnmt1 deletion.

I found that small RNAs of the endosiRNA class mapped to certain TE classes, irrespective of their transcriptional upregulation after Dnmt1 KO. Additionally, I found repressive histone marks being targeted to specific demethylated TE classes with the potential to silence them, some with no major contribution of endosiRNAs, whilst others had abundant amounts of endosiRNAs. Hence it is possible that the small RNA silencing pathway is independent of histone modification pathways in the control of TEs during global demethylation.
3.3 Discussion

In this chapter, I first presented and evaluated the Dnmt1 KO ESCs as a mechanistic system appropriate to dissect in vivo demethylation dynamics. I used this hypothesis in an in vitro system by acute deletion of Dnmt1 in mouse ESCs, carried out genome-wide methylome and transcriptome analysis at specific time-points after deletion. Furthermore, I studied the dynamics of repressive chromatin changes during the induced global hypomethylation.

The analysis of genomic demethylation upon Dnmt1 deletion by WGBS-seq showed that most genomic elements became demethylated at a similar rate, nevertheless I was able to identify DMRs that were relatively resistant to demethylation and which overlapped loci implicated in FGF signalling as well as neuronal development. The resistance to hypomethylation of genes important for neuronal development could indicate that these genes need to be kept silenced in order to prevent transcriptional activity and hence differentiation of the ESCs into neuronal lineages.

I found that a minority of transcripts became upregulated upon Dnmt1 deletion; the genes that were consistently upregulated from day 6 onwards were mostly imprinted loci. Reassuringly, genes important for PGC development also became upregulated. Interestingly, I could only confirm a general link between transcription and methylation levels for genes that were highly expressed in ESCs, as their promoters also had the lowest methylation level in WT ESCs, while no further correlation between transcription and DNA methylation was found.

Most known miRNAs (which are also expressed during PGC development) were unaffected by Dnmt1 deletion. I found two imprinted loci harbouring clusters of miRNAs which were upregulated upon global hypomethylation. More detailed analysis of the Dlk locus confirmed that the demethylation of the ICR was associated with the transcriptional upregulation of genes and miRNAs in this locus.

Repressive histone modifications (H3K9me2 and H3K27me3) are significantly remodelled during PGC development and in preimplantation embryos (Hajkova et al., 2008, Seki et al., 2007). However, H3K9me2/3 and H3K27me3 becomes enriched in preim-
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plantation embryos (Santos et al., 2005; Liu et al., 2016; Zheng et al., 2016). In order to ascertain if there was also global loss of H3K9me2 and enrichment of H3K27me3 in hypomethylated ESCs it would be necessary to check the levels of these histone marks by western blotting. Nevertheless, I saw that the enrichment of these marks throughout the genome confirmed earlier studies (Mikkelsen et al., 2007; Marks et al., 2012).

The enrichment of H3K27me3, H3K9me3 and H3K9me2 did not change upon Dnmt1 deletion. This also explains the missing connection between repressive histone marks and transcription of genes in ESCs. At the same time, regions which resisted global hypomethylation throughout the time-course of Dnmt1 deletion did not change in histone mark occupancy. I consider it most likely that resistance to demethylate reflects targeting of these regions by de novo methyltransferases DNMT3a and DNMT3b.

After I confirmed that the Dnmt1 conditional KO in ESCs served as a good system to recapitulate global demethylation dynamics in vivo, I tested the hypothesis that pervasive transcription, antisense to genes, across TEs may work as a trap of transcriptional activation of TEs.

The key conclusion from this work is that global demethylation in ESCs leads to sense strand transcriptional activation of certain TE classes, while the antisense transcription across TE classes was constant. I demonstrated that overlapping sense/antisense transcription was indeed found in TEs and that this can feed into an endosiRNA pathway working through AGO2 to potentially silence TE classes by PTGS.

The Dnmt1 KO system enabled me to uncover TE classes potentially controlled by endosiRNAs and repressive histone marks as well as other TE classes, that were solely enriched in repressive histone marks to potentially re-silence TEs during phases of transcriptional activation through acute DNA demethylation.

In more detail, I found specific ERV elements, IAPs and MMERVK10C, RLTR45 and RLTR1B, which partially resisted global demethylation. In the case of IAP elements, this recapitulates demethylation kinetics in vivo in preimplantation embryos and in PGC development (Seisenberger et al., 2012; Kobayashi et al., 2012).
Additionally, the Dnmt1 conditional KO system allowed me to dissect the transcriptional response of TE classes after acute DNA demethylation and test my hypothesis that gene derived pervasive transcripts can sense TE transcription. Interestingly, only a subset of TEs became transcriptionally upregulated upon Dnmt1 deletion, among which were IAPs and MERVL elements; these TEs have been reported before to become transcriptionally active in vivo during PGC development and in preimplantation embryos (Molaro et al., 2014, Friedli et al., 2014).

Furthermore, I found the same classes of TE elements upregulated in my system as during global waves of demethylation during the transition from serum to 2i grown ESCs (Walter et al., 2016). However, while in serum to 2i transition also LINE elements became transcriptionally upregulated in my global hypomethylation system by deletion of Dnmt1 LINE elements were expressed independent of DNMT1 across the whole time-course. Yet, with my system I created a KO of the DNA maintenance methylase machinery, which recapitulates more closely in vivo demethylation dynamics (von Meyenn et al., 2016).

Sense/antisense transcription has been described to occur in TEs (Svoboda et al., 2004) and in the oocyte pseudogenes provide the antisense strand to TEs to feed into an RNAi pathway (Tam et al., 2008). In the case of ERV elements, the insertion of forward and reverse promoter sequences (LTRs) can produce sense/antisense reads (Svoboda et al., 2004). Here I identified pervasive genic transcription antisense to TEs as a trap of TE sense transcriptional activation.

Interestingly, earlier studies have shown that TEs are mostly found in intergenic regions but if found in genic locations, they are integrated in an antisense direction to the genes (Medstrand et al., 2002, Zhang et al., 2008). This suggests a strong negative selection bias by the genome as the integration of TEs is random (Brady et al., 2009, Lander et al., 2001).

In this study, I present a model that the antisense insertion of TEs into genes may represent a mechanism used by the genome to control the TE transcriptional upregulation. I suggest that genic sense transcription in antisense direction to TEs may work
as a trap of TE sense transcription, this would increase the production of dsRNAs and feed into an endosiRNAs pathway to control TEs post-transcriptionally.

The small RNAs that mapped to different TE elements were shown to be bound by AGO2. Although, I could detect sense and antisense reads which mapped to all repeat classes, I found a sense bias in IAP and ETn mapped small RNAs.

This may be explained by the action of AGO2, which endonucleolytically cleaves its target, and could thereby lead to TE RNA degradation products. However, I cannot rule out the possibility that there could be a mixed class of small RNAs that play a role in TE silencing. Furthermore, the increased amount of AGO2 bound small RNAs mapping to IAPs, ETnERV2 and MMERVK10C upon Dnmt1 KO could be explained by the higher abundance of IAP transcript produced upon global hypomethylation. This interpretation does not mean that the small RNAs I find to be bound by AGO2 are not important but rather that their enrichment upon Dnmt1 KO could be overestimated.

In chapter 4, I will address the role of these small RNAs in TE silencing upon Dnmt1 KO in more detail.

To date, several histone modifications have been implicated a role in TE silencing. Knockout studies in ESCs of H3K9 histone methyltransferases ESET and G9a have shown that members of the ERV families are highly repressed by H3K9me2/3 (Maksakova et al., 2013). H3K9me3 is highly enriched on IAPs in ESCs potentially to prevent IAPs from expression upon DNA hypomethylation (Mikkelsen et al., 2007; Hutnick et al., 2010). The knockout of ESET in PGCs leads to a loss of H3K9me3 on ERV elements - specifically IAPs (Leung et al., 2014; Hutnick et al., 2010). Additionally, H3K27me3 has also been found to be enriched in TEs during epigenetic reprogramming in PGCs at E13.5 (Ng et al., 2013) but in PGCs at E11.5 no specific enrichment of H3K27me3 has been found on IAPs (Kim et al., 2014). Therefore, H3K27me3 could play a role in certain TE classes but potentially not in the silencing of IAP elements.

My observations show a similar redistribution of repressive marks as reported in ear-
lier studies during demethylation induced by 2i (Walter et al., 2016). H3K27me3 was also identified in our system to be a key potential driver of TE silencing of MERVL elements, as previously reported (Walter et al., 2016).

The present study shows that MMERVK10C, as the only element, had enrichment of H3K9me3 upon Dnmt1 deletion. It is also notable that this element is the one most resistant to Dnmt1 deletion induced demethylation consistent with H3K9me2/3 marked regions being more resistant to global demethylation (von Meyenn et al., 2016).

Sharif et al., (2016) proposed a model of UHRF1 control of IAP activity through a combinatorial effect of DNA methylation together with H3K9me3 (Sharif et al., 2016), however, in my system upon Dnmt1 deletion H3K9me3 became depleted at IAPs, with no subsequent enrichment, in agreement with a recent report using ESCs serum to 2i transition (Walter et al., 2016). Nevertheless, I cannot exclude the fact that potentially H3K9me3 might play a role at later stages after Dnmt1 KO, as I only looked at day 8 after deletion.

I suggest a model in which endosiRNAs play a key role as a first response after transcriptional activation of IAPs. It could well be the case that the small RNAs target the histone marks to the TE classes - in order to test this KOs of histone methyltransferases in the background of Dnmt1 deletion would need to be carried out.

In summary, this study presents a mechanistic dissection of epigenetic modifications on silencing of TE elements. Previous studies have shown that TEs are silenced by DNA methylation (Walsh et al., 1998, Arand et al., 2012, Sharif et al., 2016), histone marks (Karimi et al., 2011, Walter et al., 2016, Leung et al., 2014) as well as small RNAs (Flemr et al., 2013, Svoboda et al., 2004, Tam et al., 2008, Aravin et al., 2007). This study took the unique approach in analysing all three epigenetic marks together as means to preserve genome integrity by controlling TE activity. As such, I was able to identify endosiRNAs as key players of TE silencing as well as the importance of several epigenetic modifications to work in parallel in order to keep the genome intact during global hypomethylation in ESCs.
Figure 3.43: Schematic of endosiRNAs becoming expressed to control TEs upon \textit{Dnmt1} deletion in ESCs. TEs in mouse ESCs are methylated (filled lollipops) and get demethylated (open lollipops) upon \textit{Dnmt1} KO. Sense transcription of TEs (red) is inhibited by DNA methylation and gets upregulated upon \textit{Dnmt1} KO. Genic sense transcription (grey) is constant antisense to TEs. This results in the production of dsRNA and allows the production of endosiRNAs bound by AGO2 (black) which silence TEs via post transcriptional gene silencing. Additionally, histone marks (blue hexagons) become enriched at TEs which get transcriptionally upregulated and may be involved in the subsequent transcriptional silencing.
Chapter 4

TE regulation upon \textit{Dicer} KO and \textit{Dicer}/\textit{Dnmt1} KO

4.1 Introduction

The transcriptional activity of TEs induced by \textit{Dnmt1} KO potentially exposes an achilles heel for TEs, as it activates small RNA-based mechanisms that may serve to restrain the TEs. Indeed small RNAs have been described to play a role in TE control with miRNAs as well as endosiRNAs having been found to be involved in TE silencing in the oocyte (Heras et al., 2013, Flemr et al., 2013, Stein et al., 2015).

In the previous chapter I showed that endosiRNAs bound to TEs became enriched at TEs during induced global demethylation in mouse ESCs.

In this chapter, I examine whether these endosiRNAs play a role in TE repression after TE transcriptional activation upon \textit{Dnmt1} KO. Additionally, I aim to test the hypothesis that lowly expressed LINE-1 elements throughout the time-course of \textit{Dnmt1} deletion are post transcriptionally repressed by small RNAs, similarly to LINE element repression by endosiRNAs in human cancer cell lines (Goodier and Kazazian, 2008).

I constructed \textit{Dicer} constitutive and conditional double KOs in the background of \textit{Dnmt1} KO ESCs - (DKO) and (cDKO) respectively, to compare the long term effect
of Dicer KO with an acute deletion of this cellular endonuclease.

Eukaryotic small RNAs are generated through the RNase-III activity of DICER, which is the central player of the RNA interference (RNAi) network mediating gene silencing on a transcriptional or post transcriptional level, TGS or PTGS, respectively (Bartel, 2004; Hammond, 2005). RNAi plays a role in early development as the knockout of crucial players of the RNAi pathway - Ago2, Dicer and Dgcr8 - lead to early embryonic lethality (Bernstein et al., 2003). Nevertheless, it is possible to obtain Dicer KO mouse ESCs that show normal stem cell properties but an inability to differentiate in vitro and in vivo (Kanellopoulou et al., 2005). Dicer KO in mouse ESCs leads to a loss of the RNAi machinery by complete depletion of miRNAs (Kanellopoulou et al., 2005, Murchison et al., 2005). Further to the depletion of small RNAs, Dicer KO in ESCs has been shown to also have an effect on DNA methylation levels by controlling Dnmt expression through the miR-290 cluster (Sinkkonen et al., 2008, Benetti et al., 2008). In this chapter, I characterise the transcriptome of my Dicer/Dnmt1 double KO (DKO) and compare them to previously studied Dicer KO ESCs.

After this genome-wide characterisation of my Dicer/Dnmt1 double KO (DKO), I examine the role of DICER in TE transcriptional re-silencing, after global hypomethylation.

Dicer knockdown in mouse preimplantation embryos leads to transcriptional upregulation of MERVL and IAP elements through the production of dsRNAs (Svoboda et al., 2004). EndosiRNAs have been found to regulate TE transcription in the growing oocyte (Tam et al., 2008, Watanabe et al., 2006) and Dicer KO in human cancer cell lines leads to upregulation of LINE elements (Goodier and Kazazian, 2008). I follow TE transcriptional activation by total RNA-seq and qRTPCR in the Dicer/Dnmt1 double KO (DKO) and conditional double KO (cDKO). To understand whether TE transcriptional re-silencing, after global hypomethylation, is achieved by repressive histone marks or by PTGS through small RNAs I also perform ChIP-seq of H3K27me3, H3K9me2 and H3K9me3 in Dicer/Dnmt1 DKO ESCs.
Recent work has shown that miRNAs as well as endosiRNAs can play a major role in TE silencing in mouse ESCs (Flemr et al., 2013) and I present an interesting miRNA binding to IAPEZ and investigate the potential for translational gene silencing of TEs in ESCs.

This chapter sheds light onto the interplay of small RNAs and DNA methylation in ESCs. Further it uncovers the role of DNA methylation and small RNAs in TE regulation in constant Dicer/Dnmt1 DKO and acute cDKO ESCs.

## 4.2 Results

### 4.2.1 Constitutive KO of Dicer by CRISPR-Cas9

To assess whether AGO2 bound small RNAs had a mechanistic role in TE silencing during global hypomethylation in Dnmt1 KO ESCs, I generated a Dicer KO via CRISPR-Cas9 in the Dnmt1fl/fl ESCs. From this, I was able to create Dicer KO as well as Dicer/Dnmt DKO ESCs.

The Dicer KO was constructed with Cas9 guide RNAs targeting Exon 23 and 24 - encoding the RNase III domain and the RNA binding domain, respectively (Bernstein et al., 2003). The resulting genomic deletion resulted in a catalytically inactive protein (Figure 4.1).

Dicer mRNA levels in Dicer KO ESCs (Dicer−/−/Dnmt1fl/fl) were reduced compared to WT (Dicer+/+/Dnmt1fl/fl) ESC (Figure 4.2 left). To confirm the loss of enzymatic activity in the Dicer KO ESCs, I checked for the expression of endogenously expressed miRNAs in Dicer KO compared to WT ESCs, by small RNA specific qRTPCR.

The expression of mmu-miR93 in Dicer KO compared to WT ESCs was abolished. However, small nuclear RNAs (snoRNAs), which are DICER independent were still being expressed (Figure 4.2 right).

After I confirmed that the engineered Dicer KO genes encoded for an enzymatically inactive DICER protein, I characterised the transcriptome of those ESCs.
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Chapter 4. TE regulation upon Dicer KO and Dicer/Dnmt1 KO

Figure 4.1: CRISPR-Cas9 KO of Dicer in Dnmt1^{fl/fl} ESCs. KO strategy for Dicer in mouse ES cells constructing gRNAs against Exon 23 and 24 of Dicer mRNA. gRNA protospacer adjacent motif (PAM) sequences (red).

Figure 4.2: Characterisation of Dicer KO in Dnmt1^{fl/fl} ESCs. Dicer mRNA is downregulated in 1 clone of Dicer KO ESCs (blue) versus WT ESCs (black). (C) endogenously expressed mmu-miR-93 in mouse ES cells is downregulated upon Dicer KO, relatively to snoRNA expression. Differences between conditions that are statistically significant are denoted by (* p-value<0.05, ** p-value<0.005, (Student’s t-test))
4.2.2 Transcriptional changes in Dicer KO and Dicer/Dnmt1 double KO ESCs

To study the effect of DICER on the mRNA transcription, I carried out total RNA-seq of Dicer KO ESCs (list of total RNA-seq datasets in appendix). Additionally, I performed RNA-seq in the Dicer KO at day 1 and day 11 after CRE mediated deletion of Dnmt1, to analyse the effect of depletion of small RNAs as well as global hypomethylation on self-renewal properties of ESCs. First, I compared the transcriptional profiles of Dicer KO and WT ESCs I called differentially expressed genes using the Seqmonk intensity difference filter with Benjamini and Hochberg for multiple testing correction with a p-value threshold of < 0.05. Most of the differentially expressed genes were transcriptionally silenced upon deletion of Dicer (n = 90). The downregulated genes comprised Lin28 as well as Dnmt3L. Furthermore, I found a small number of genes (n = 7) which were upregulated upon Dicer KO, among these was the developmentally important gene Lefty1 (Figure 4.3) (list of differentially expressed miRNAs upon Dicer KO in appendix). To finish the analysis of the Dicer KO ESCs, I carried out a GO Term analysis, using Panther webtool, of the differentially expressed genes between KO and WT ESCs. Biological processes which were mostly affected by the KO of Dicer were involved in organism development and angiogenesis (Figure 4.4).
4.2. Results

Chapter 4. TE regulation upon Dicer KO and Dicer/Dnmt1 KO

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Figure 4.4: GO term analysis of differentially expressed genes between Dicer KO and WT ESCs. Significantly expressed genes were analysed by Panther GO Term analysis.

Figure 4.5: Differentially expressed genes between Dicer KO, Dicer/Dnmt1 DKO and WT ESCs. Heatmap of hierarchical clustering of genes differentially expressed between Dicer KO, Dicer/Dnmt1 DKO ESCs. Per probe normalised reads are depicted of all RNA-seq datasets quantified as RPM. Each dataset is sequenced as n=1. red = not expressed, blue = highly expressed.
Subsequently, I carried out a differential expression analysis between Dicer KO, Dicer/Dnmt1 double KO (DKO) with a deletion of the DNA methyltransferase Dnmt1 induced demethylation for 1 day as well as 11 days and WT ESCs as control. Also for this analysis, I used the Seqmonk intensity difference filter with Benjamini and Hochberg for multiple testing correction with a p-value threshold of < 0.05.

I found 258 significantly differentially expressed genes between the different conditions. Genes in cluster I and IV were up- or downregulated solely upon Dicer KO (Figure 4.5). Genes in cluster II were upregulated solely upon Dnmt1 KO. Genes in cluster III showed upregulation upon Dicer/Dnmt1 DKO (Figure 4.5). These genes were upregulated in an additive way upon knockout of both genes, they were not expressed in WT and day 1 after Dnmt1 KO, but were upregulated at day 11 after Dnmt1 deletion as well as in Dicer KO, with a further upregulation in the Dicer/Dnmt1 DKO - like Fbln2 in Figure 4.6. This cluster comprised a lot of cell adhesion genes, indicating a change in cell morphology upon KO of both Dicer and Dnmt1 (list of genes in Cluster III in appendix).

The pluripotency genes - Sox2, Klf4, Esrrb, Oct4 and Nanog - were not significantly differentially expressed in the Dicer KO as well as the Dicer/Dnmt1 DKO ESCs (Figure 4.6).
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Figure 4.7: Expression of the pluripotency genes in Dicer KO, Dicer/Dnmt1 DKO ESCs. Bargraph of RNA-seq data of Dnmt genes in WT (grey), Dicer/Dnmt1 DKO induced for 1 day (green) and 11 days (light green), Dicer KO uninduced for 1 day (orange) and 11 days (yellow), conditional Dnmt1 KO ESCs induced for 1 day (dark red) and 11 days (dark blue). RNA-seq libraries (n=1).

4.6, 4.7), therefore the self-renewal of Dicer KO ESCs was not affected. As reported before, the de novo methyltransferases Dnmt3a and Dnmt3b were downregulated upon Dicer KO (Sinkkonen et al., 2008, Benetti et al., 2008). However, I observed the strongest down-regulation of expression on Dnmt3l. Dnmt1 was normally expressed in the Dicer KO ESCs (Figure 4.8).

Figure 4.8: Expression of the Dnmts in Dicer KO, Dicer/Dnmt1 DKO ESCs. Bargraph of RNA-seq data of Dnmt genes in WT (grey), Dicer/Dnmt1 DKO induced for 1 day (green) and 11 days (light green), Dicer KO uninduced for 1 day (orange) and 11 days (yellow), conditional Dnmt1 KO ESCs induced for 1 day (dark red) and 11 days (dark blue). RNA-seq libraries (n=1).

In summary, in the Dicer KO ESCs I recapitulated the same transcriptional changes that have been reported before (Sinkkonen et al., 2008, Benetti et al., 2008). I found a downregulation of the de novo methyltransferases, furthermore, I found a repression of organism developmental genes, which could lead to the inability of Dicer KO ESCs to
differentiate. Additionally, I was able to identify a gene groups, that were dependent on DICER as well as DNMT1 and showed an additive transcriptional effect in the Dicer/Dnmt1 DKO.

4.2.3 Genome-wide histone marks upon Dicer KO and Dnmt1/Dicer DKO

After an analysis of the transcriptome I analysed the chromatin changes upon Dicer KO as well as Dicer/Dnmt1 DKO ESCs. I tested for three repressive histone marks - H3K9me3, H3K9me2 and H3K27me3 in the different KO ESCs (list of ChIP-seq datasets in appendix), because during in vivo epigenetic reprogramming these repressive histone marks are also largely exchanged (Hajkova et al., 2008, Ancelin et al., 2006, Liu et al., 2016, Zheng et al., 2016). I aimed to discover whether the acute demethylation by Dnmt1 KO and additionally deletion of the small RNAs by deletion of Dicer KO would also have an effect on the overall chromatin structure in ESCs.

![Figure 4.9: Enrichment of repressive histone marks at genomic features in Dicer KO and Dicer/Dnmt1 DKO. Bar graph of enrichment of H3K27me3, H3K9me3 and H3K9me2 in repeats (dark violet), genic regions (light violet), promoters (dark green), CGIs (middle green), intergenic regions (light green) in WT ESCs, Dnmt1 KO, Dicer KO and Dicer/Dnmt1 DKO.](image-url)
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At the outset, I analysed the general enrichment of the three histone marks across different genomic locations. As expected, H3K27me3 was enriched in genic regions, in CGIs as well as promoters. Additionally, this repressive histone mark was enriched on repeats. H3K9me3 was depleted in CGIs, promoters and genes but highly enriched over repeats. H3K9me2 was generally distributed across the whole genome with some enrichment in repeats and genic regions. I did not observe any changes of histone distribution for either of the repressive histone marks upon Dicer KO or Dicer/Dnmt1 DKO in ESCs. The same regions were still enriched and I did not find any significant redistribution of histone marks upon deletion of the DNA maintenance methylase or DICER derived small RNAs (Figure 4.9 and 4.10).

As a final analysis, I overlaid transcriptional levels of genes in Dicer KO ESCs with chromatin changes upon the single and DKO s of Dicer and Dnmt1. I could not find any correlation between genes expressed in WT ESCs and chromatin enrichment in the different KOs (Figure 4.11).

The analysis of the transcriptome, as well as chromatin changes upon deletion of Dicer in the single KO as well as in combination on Dnmt1 KO in the Dicer/Dnmt1 DKO, showed that there was an additive effect of transcriptional changes in the DKO ESCs. Though I could not find any synthetic redistribution of the three histone marks I
Chapter 4. **TE regulation upon Dicer KO and Dicer/Dnmt1 KO**

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**Figure 4.11:** Transcriptional profile in *Dicer* KO is not explained by histone mark enrichment. Line graph of histone enrichment of genes which showed different expression level in *Dicer* KO ESCs: not expressed (RPM $< 0$, red), low expressed ($2.5 < \text{RPM} < 0$, violet), expressed ($5 < \text{RPM} < 2.5$, pink), high expressed ($10 < \text{RPM} < 5$, dark blue) or very high expressed (RPM $> 10$, light blue) as well as genes which were *Dicer* dependent (orange).

studied at the whole genome level.

#### 4.2.4 LINE-1 and major satellite expression in *Dicer* constitutive KO ESCs

In order to address the involvement of *Dicer* derived small RNAs on TE regulation, I conducted a detailed analysis of TEs in the *Dicer* KO and *Dnmt1/Dicer* DKO s. To test whether endosiRNAs were playing an active role in TE repression, I repeated the *Dnmt1* deletion experiments in the *Dicer* KO background. Over the time-course of *Dnmt1* KO in the *Dicer* KO ESCs, while expression of LINE-1 and major satellites were detectable by qRTPCR, surprisingly no expression of IAPs and ETns was found (Figure 4.12).

This result showed that DICER regulated small RNAs were involved in the silencing of LINE-1 elements and major satellites. Major satellites became transcriptionally upregulated following *Dnmt1* deletion and therefore showed dependency on both endosiRNA and DNA methylation. Despite that, LINE-1 elements were highly expressed upon *Dicer* KO, albeit showed no additional transcriptional activation upon *Dnmt1*
Figure 4.12: LINE-1 and major satellites are upregulated in a Dicer constitutive KO. qRTPCR analysis of TE classes in Dicer KO and Dicer KO with conditional Dnmt1 induced for 1 day (dark red), 3 days (light red), 6 days (light pink), 8 days (light blue), 10 days (dark blue), and 12 days (black). Error bars represent standard deviation of 2 biological replicates with 2 technical replicates. Values were normalized to Atp5b, Hspcb and major satellites to U1. Differences between conditions that are statistically significant are denoted by (* p-value<0.05, ** p-value<0.005, *** p-value<0.0005 (Student’s t-test))
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Surprisingly, although IAPs and ETns were Dnmt1 KO dependant and became upregulated upon global demethylation, both TEs were transcriptionally silenced in the Dicer/Dnmt1 DKO ESCs. Nevertheless, upon Dicer/Dnmt1 DKO IAPEZ and ETn both got transcriptionally expressed to a much lower amount than in the single Dnmt1 KO.

Earlier studies have shown that IAP elements are specifically silenced by H3K9me3 marks in Dnmt1 constitutive KO ESCs ([Hutnick et al., 2010](#), [Mikkelsen et al., 2007](#)). I wanted to test whether any repressive chromatin enrichment could explain the transcriptional silencing of ETns and IAPs in my Dnmt1/Dicer DKO ESCs.

![Figure 4.13: Dicer dependent TE upregulation.](image)

**Figure 4.13: Dicer dependent TE upregulation.** Heatmap of RNA-seq data mapped to TE classes. Only TE classes which were intergenic and within 2kb away from genes were considered. Mean of TE classes with at least 1000 integration sites were regarded and z-score was calculated between KO and WT ESCs. RNA-seq (n=1).

With the help of Simon Andrews, we undertook an unbiased approach to find the TE classes that were solely upregulated in the Dicer constitutive KOs in our RNA-seq data (Figure 4.13). For the analysis, we followed the mapping of TEs as described in chapter 3, and then analysed only the TE classes which were differentially expressed with a threshold of p < 0.05 in Dicer KO compared to WT ESCs. Only seven TE classes were differentially upregulated upon Dicer KO - and therefore defined as DICER dependent TEs. Two classes of IAP, IAPLTR3 and IAP-d, had already been identified in chapter 3 as being dependent on Dnmt1 and also showed a dependency on Dicer KO.
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4.2.5 Histone modifications may explain different behaviours of TEs to acute Dnmt1 KO

In early mammalian development chromatin marks become remodelled during global epigenetic reprogramming (Hajkova et al., 2008, Ancelin et al., 2006, Liu et al., 2016, Zheng et al., 2016). IAPs are highly guarded by H3K9me3 and MERVL elements by H3K27me3, respectively (Hutnick et al., 2010, Maksakova et al., 2013). H3K9me2 follows DNA methylation dynamics and gets lost upon global demethylation in PGC reprogramming (Seki et al., 2005). The same dynamics have been seen globally in the serum to 2i transition in mouse ES cells (von Meyenn et al., 2016, Walter et al., 2016). Therefore, I wanted to test if repressive histone marks are enriched at TEs specifically upon KO of Dicer as well as in the DKO of Dicer/Dnmt1.

I performed ChIP-seq experiments of H3K9me3, H3K9me2 and H3K27me3 in the Dicer
KO and Dicer/Dnmt1 DKO ESCs with Dnmt1 KO and WT ESCs as a control. I analysed the enrichment of these marks on TE classes dependent on DNMT1 as defined in chapter 3 (Figure 3.25) and DICER as defined earlier in this chapter (Figure 4.13).

H3K9me3 was depleted upon Dicer KO in most TE classes, with a higher depletion in the DKO ESCs. MERVL elements showed a 2-fold enrichment of H3K9me3 in the Dicer KO ESCs than in WT ESCs, but were less enriched than in the Dnmt1 KO ESCs. L1MdGf elements showed higher enrichment of H3K9me3 in the Dicer KO ESCs than in Dnmt1 KO or WT ESCs, but a depletion of this mark in the DKO ESCs. ORR1B1 and RLTR14 showed both an enrichment in Dicer KO and Dicer/Dnmt1 DKO ESCs.

H3K9me2 showed the exact opposite pattern and was generally enriched in all TEs upon Dicer KO and Dicer/Dnmt1 DKO. L1MdGf elements were depleted of H3K9me2 in the Dicer KO and even more depleted in the Dicer/Dnmt1 DKO. MERVL elements
Figure 4.16: H3K27me3 enrichment over TEs dependent on Dicer and Dnmt1. Heatmap of ChIP-seq data of H3K27me3 mapped to TE families at day 4 and day 8 after Dnmt1 KO, Dicer KO and Dnmt1/Dicer DKO in comparison to WT ESCs. Read count enrichment depicted as z-score. ChIP-seq libraries (n=1).
had lost enrichment of H3K9me2 in Dnmt1 KO to the same level as in the Dicer KO but were totally depleted of this repressive histone mark in the Dicer/Dnmt1 DKO ESCs. Interestingly, IAPs and MMERVK10C lost H3K9me2 specifically upon Dnmt1 deletion with highest enrichment of this mark in single Dicer KO ESCs.

H3K27me3 became enriched in all TE classes upon Dicer KO and Dicer/Dnmt1 KO. While L1mdGf and RLTR45 elements had the highest enrichment of H3K27me3 upon Dnmt1 KO in comparison to Dicer KO and WT ESCs, in IAP-d, MERVL, IAPLTR3 and ERVBA4IB the enrichment of H3K27me3 was higher than in WT ESCs in all three KO conditions. Interestingly, IAPEZ, ETnERV2 and MMERVK10C had enrichment of H3K27me3 from WT to Dnmt1 KO to Dicer KO with highest enrichment in the Dicer/Dnmt1 DKO.

This is a compelling result, as it suggests that in the Dicer/Dnmt1 DKO IAPEZ, ETns as well as MMERVK10C elements are enriched for the H3K27me3 repressive histone mark, which is likely to prevent those TEs from transcriptional upregulation in the Dicer constitutive KO ESCs.

### 4.2.6 TE response after double conditional KO of Dicer and Dnmt1

I aimed to investigate whether the unexpected transcriptional repression of IAPs and ETn elements was due to long term Dicer deletion, therefore I constructed Dicer<sup>f<sub>l</sub>/f<sub>l</sub></sup> conditional double KO (cDKO) ESCs in the Dnmt1<sup>f<sub>l</sub>/f<sub>l</sub></sup> background ESCs. The loxP sites were introduced in intron 15-16 and intron 20-21 of the Dicer gene (Figure 4.17). The activation of the CRE recombinase by induction with tamoxifen (4OHT) led to recombination between the loxP sequences, which in consequence led to a down-regulation of the Dicer mRNA (Figure 4.18A) as well as abrogation of its enzymatic activity shown by depletion of endogenous mmu-miR93 expression (Figure 4.18B).

Dnmt1 deletion experiments were repeated in the Dicer<sup>f<sub>l</sub>/f<sub>l</sub></sup> background. qRTPCR of time-points after acute Dicer/Dnmt1 cDKO showed significantly higher transcriptional
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Figure 4.17: Construction of Dicer\textsuperscript{fl/fl} in Dnmt1\textsuperscript{fl/fl} cDKO ESCs. Knockout strategy for Dicer in mouse ES cells introducing loxP sites in Intron 15-16 and intron 20-21 of Dicer. Agarose gel of PCR to screen for genomic recombination of 2 Dicer cDKO clones after addition of tamoxin for 3 days. Recombination of Intron 15-16 was tested with primer set 1, recombination of intron 20-21 was tested with primer set 2 and recombination of both introns was tested with primer set 3, LD = 1000 bp DNA ladder.

Figure 4.18: Dicer mRNA and miRNAs expression in Dicer/Dnmt1 cDKO. (A) Dicer mRNA is getting downregulated upon CRE recombination induced by tamoxifen in 1 clone, (B) endogenously expressed miRNAs in mouse ES cells are downregulated upon Dicer KO controlled by snoRNA expression. Each experiment was done in 3 technical replicates. Differences between conditions that are statistically significant are denoted by (* p-value<0.05, ** p-value<0.005, (Student’s t-test))

activation of IAPs on day 10 (Figure 4.19). Additionally, on-going expression of IAPs was found at later time-points (Day 12) with no significant resilinging of the IAPs. This indicates that the small RNAs produced were DICER dependent and played a role in IAP silencing upon Dnmt1 KO.

MERVL elements showed the same transcriptional kinetics upon Dicer/Dnmt1 cDKO as for Dnmt1 KO and became transcriptionally activated and subsequently resiled. While ETn elements were upregulated and subsequently resiled upon Dnmt1 KO,
these elements were not transcriptionally activated in the Dicer/Dnmt1 DKO, nor in the Dicer/Dnmt1 cDKO ESCs. This may be due to their resilingencing by histone modifications, but could also be explained by Dicer independent small RNAs, as I found an increase of small RNAs mapping to the ETn elements in the third chapter.

**Figure 4.19: TE expression in the Dnmt1fl/fl and Dicerfl/fl/Dnmt1fl/fl cDKO ESCs.** qRTPCR analysis of TE classes in the Dnmt1fl/fl and Dicerfl/fl/Dnmt1fl/fl cDKO ESCs induced for 1 day (dark red), 3 days (light red), 6 days (light pink), 8 days (light blue), 10 days (dark blue), and 12 days (black). Error bars represent standard deviation of 2 biological replicates with 2 technical replicates. Values were normalised to Atp5b, Hspcb and major satellites to U1. Differences between conditions that are statistically significant are denoted by (* p-value<0.05, ** p-value<0.005, (Student’s t-test))

MMERVK10C were only upregulated in Dnmt1 KO ESCs 25 days after KO induction, therefore it is not surprising that they were not upregulated in the Dicer/Dnmt1 cDKO ESCs until day 12. Major satellites as well as LINE elements were not dependent on knockout of the maintenance methylation machinery alone but in the Dicer/Dnmt1 cDKO they became transcriptionally upregulated. In order to check whether major satellites have the ability to produce endosiRNAs, I mapped the total RNA-seq dataset to major satellite consensus sequences (Figure 4.20A). I observed sense and
antisense transcripts in WT as well as \textit{Dnmt1} KO ESCs at day 9 after induced CRE recombination (Figure 4.20B). Additionally, I pulled down AGO2 bound small RNAs and mapped them back to the major satellite consensus sequence (Figure 4.20C).

Figure 4.20: Long and short small RNAs map to major satellites. (A) major satellite (GSAT element) was analysed by unique mapping in the genome and a general expression of this elements, red bar depicts the mean expression level. (B) sense (red) and antisense (blue) reads mapped to major satellite consensus sequence by piPipes small RNA-seq analysis tool. (C) Bargraph of small RNAs pulled down by AGO2 mapped to major satellite consensus sequence, in WT ESCs = grey and Dnmt1 KO ESCs at day 9 = blue. Experiments were performed as 4 biological replicates.

In summary, I was able to identify DICER dependent small RNAs, that controlled LINE-1 elements and major satellites, as well as IAP elements during acute demethylation in ESCs.

Furthermore, I found histone mark enrichment at RNAi independent TE classes and concluded that upon genome hypomethylation, histone modifications and RNAi play a major role to keep TE elements transcriptionally silenced.

**4.2.7 miRNA production against IAPs suggest multiple levels of gene regulation**

In chapter 3, I identified small RNAs perfectly mapping to the IAP transcript, but also some which imperfectly mapped to the consensus sequence (Figure 4.21). I aimed to investigate whether these smallRNAs could be miRNAs, that regulate IAP expression at the level of translation.

Mmu-miR-7081 mapped to the IAP consensus sequence independent of \textit{Dnmt1} deletion
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Figure 4.21: miRNAs map to IAPEZ consensus sequence independent of DNA demethylation. Consensus sequence mapping of small RNA-seq data to IAPEZ consensus sequences allowing 2 mismatches, sense reads = red, antisense reads = blue, arrow indicating miRNA.

(Figure 4.21). As the miRNA is expressed independent of DNA methylation, the potential mechanism of translational gene silencing by this miRNA is already in place before transcriptional activation of IAPs. The mmu-mir-7081 mapped to the 3’LTR of the polymerase of IAPs. As a bona fide miRNA it mapped with its 5’seed region totally complementary to the IAP 3’LTR followed by a bulge sequence and additional posterior binding (Figure 4.22). The pre-miRNA gene of mmu-mir-7081 is encoded in an intron of Dnmt1 and is only conserved in the murine lineage (Figure 4.23). This is curious, as also IAPs are also specific to the rodent lineage. I suggest that the miRNAs could have coevolved with IAPs to control this highly active TE in the mouse germline.

Figure 4.22: Mmu-miR-7081 complementary to IAP consensus sequence. miRNA 7081 binds with 5’ seed region to IAP sequence at nucleotide 6450. The seed region is binding followed by a 3 nt bulge sequence, additionally there is some 3’ binding with additional bulges. Local alignment was done using EMBOSS Matcher alignment tool with default parameters.

I confirmed the constant expression of pre-mmu-miR-7081 in the small RNA-seq libraries across the whole time course after Dnmt1 deletion (Figure 4.24B) by targeted small RNA qRTPCR at day 9 after Dnmt1 and detected mmu-miR-7081 expression throughout the whole time-course (Figure 4.24A).
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Figure 4.23: Mmu-mirR-7081 is conserved in the murine lineage. Evolutionary conservation of mmu-miR-7081 across different species. UCSC genome browser.

Figure 4.24: Expression of mmu-miR-7081 upon Dnmt1 KO. (A) small RNA qRTPCR on mature mmu-miR-7081 at day 9 after Dnmt1 deletion (light blue) and wild-type (grey), (B) small RNA-seq of miRNA expression in wild type (grey) and conditional Dnmt1 knockout ESC induced for 1 day (dark red), 3 days (light red), 6 days (light pink), 9 days (light blue), 11 days (dark blue).

4.2.8 IAP protein production upon Dicer KO

In order to study whether transcriptional activation of IAPs leads to production of IAP protein we undertook immunofluorescence (IF) staining of Dicer\(^{-/-}\) ESCs. Only in the double KO of Dicer and Dnmt1 IAP protein was detected by confocal microscopy (Figure 4.25). In the single Dnmt1 KO and Dicer KO as well as the WT ESCs no IAP protein was observed. This indicates that Dicer dependent small RNAs are needed for translational resilingencing of IAPs and opens the possibility for miRNA-7081 to play an important role.

Altogether, this could suggest that there are two ways of TE silencing through small RNAs in ESCs. The first one is through endosiRNA dependent post-transcriptional gene silencing of TE transcripts and the second one is through translational repression and depletion of the TE protein.
4.3 Discussion

In chapter 3 of this thesis, I found AGO2 bound small RNAs mapping to TEs. Additionally, my KD study in chapter 3 suggested that IAP transcription is dependent on DICER. I concluded that endosiRNAs are likely to be the major RNAs involved in TE regulation in ESCs.

In this chapter, I aimed to test whether DICER derived small RNAs are regulators of TE resilencing after transcriptional upregulation during global hypomethylation.

I constructed the Dicer constitutive KO ESCs by a deletion of the enzymatic activity and could confirm the loss of enzymatic activity by loss of miRNAs normally expressed in WT ESCs.

Due to differing results of phenotypes and transcriptional changes in different Dicer KO ESCs from other labs (Kanellopoulou et al., 2005; Murchison et al., 2005), I wanted...
to characterise my KOs carefully and carried out a transcriptional analysis through RNA-seq. GO term analysis of the differentially expressed genes showed that genes important for organism development were substantially downregulated in Dicer KO ESCs in comparison to WT ESCs, which suggests that the KO ESCs could have a differentiation defect. Indeed, earlier studies have found a defect in differentiation of Dicer KO ESC cells in vitro and in vivo (Kanellopoulou et al., 2005, Murchison et al., 2005).

As reported before, Dnmt3a and Dnmt3b were downregulated in our Dicer KO ESCs, while Dnmt1 was normally expressed (Sinkkonen et al., 2010, Murchison et al., 2005). Interestingly, Dnmt3l was the most downregulated Dnmt, with no expression in Dicer KO ESCs. Additionally, I found an intriguing additive effect in the Dicer/Dnmt1 DKO ESC. After day 1 of induction of Dnmt1 KO a cluster of genes (Figure 4.25) was much more lowly expressed than at day 11 after induction of Dnmt1 KO and the expression increased in the Dicer KO compared to WT ESCs. This could be attributed to the enzymatic effect of DICER and DNMT1 - and therefore to the activity of small RNAs and DNA methylation - respectively. However, this could also be the effect of DICER on Dnmt expression through small RNAs, as has been suggested before (Murchison et al., 2005).

As the RNA-seq results were all done in one Dicer KO clone with no replicate, it is important to prepare another replicate and repeat the results before coming to any conclusions. However, if the additive effect is confirmed, it would be possible to do RNA-seq of the cDKO ESCs of Dicer/Dnmt1 at different time points to study the dynamics of DNA methylation and small RNAs on transcription in mouse ESCs.

After the transcriptional analysis I also studied the chromatin changes upon Dicer KO. No genome-wide redistribution of H3K9me3, H3K27me3 or H3K9me2 was found upon DNA hypomethylation and depletion of Dicer dependent small RNAs.

After the analysis of transcriptome as well as chromatin changes upon Dicer KO in ESCs, I studied the TE response in the Dicer/Dnmt1 DKO and cDKO ESCs.
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However, I could not find any transcriptional upregulation of IAPs nor ETn elements in the Dicer/Dnmt1 DKO ESCs, while LINE elements and major satellites were highly upregulated in the DKO ESCs.

I suggest that that upon Dicer constitutive KO active TEs were silenced through repressive histone marks. In order to test this hypothesis I performed histone ChIP-seq of repressive histone marks H3K9me3, H3K9me2 and H3K27me3 in the Dicer KO ESCs compared to Dicer/Dnmt1 DKO, Dnmt1 KO and WT ESCs.

Interestingly, I saw high enrichment of H3K27me3 on IAPs as well as ETn elements upon Dicer KO. This enrichment was even higher in the Dicer/Dnmt1 double KO ESCs. This could mean that H3K27me3 histone marks are influenced by the existence of small RNAs, and their depletion could result in higher H3K27me3 redistribution in the genome. To assess a relationship between the H3K27me3 enrichment in the Dicer KO ESCs, it would be necessary to test the specificity of the enrichment by deleting the PRC2 complex in the Dicer and Dicer/Dnmt1 KO ESCs.

After this, I constructed the conditional Dicer/Dnmt1 cDKO ESCs and tested for TE activity. I could only find lack of IAP resilencing in the double conditional KO of both Dicer and Dnmt1. This is an intriguing finding, as it suggests that in the DKO in the state of hypomethylation as well as depletion of small RNAs, targeted chromatin changes on very active TE classes, such as IAPEZ, may prevent these elements from transcriptional expression.

The transcriptional upregulation of IAPs could not be followed for longer than 12 days, as the ESCs started to die afterwards. This could be due to uncontrolled retrotransposition events of IAPs. In order to test this hypothesis it is possible to measure IAP protein levels or to carry out retrotransposition assay of this TE. Another way of testing for retrotransposition would be to investigate chimeric transcripts in RNA-seq data of Dicer/Dnmt1 cDKO ESCs over time.

To my surprise, ETn elements were still guarded by unknown mechanisms in the cDKO ESCs throughout the time-course of deletion. ETn elements are highly active in the mouse germline and guarded by DNA methylation; earlier I found AGO2 bound small
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RNAs mapping to ETns. These TEs were transcriptionally silenced in the Dicer KO as well as in the Dicer/Dnmt1 cDKO ESCs. We found high enrichment of H3K9me2 repressive histone marks upon Dicer/Dnmt1 DKO on ETn elements and therefore suggest that ETn elements are not resienced by endosiRNAs primarily but rather by transcriptional gene silencing through repressive histone marks. However ETns could also be guarded by additional repressive histone marks, which I have not tested for, as their histone enrichment after hypomethylation matched with that of IAP elements.

Furthermore, in this study I only dissected the role of DICER dependent small RNAs on TE silencing, although DICER independent AGO2 bound small RNAs could also play a role in TE silencing. DICER independent small RNAs are highly abundant in mouse ESCs (Babiarz et al., 2008) and are important for major satellite silencing (Cao et al., 2009). DICER independent small RNAs could also explain the repression of ETn elements, which had increasing amounts of AGO2 bound small RNAs mapped to them, but were not responsive upon KO of this cellular endonuclease, therefore the produced small RNAs that mapped to ETns might be DICER independent.

Transcription of LINE-1 elements and major satellites is not dependent on DNA methylation, as both repeats were expressed in ESCs with or without Dnmt1 deletion (Chapter 3). On both elements AGO2 bound small RNAs with an endosiRNA signature were found. Dicer/Dnmt1 DKO, as well as Dicer/Dnmt1 cDKO led to transcriptional upregulation of both elements. This suggests that in mESCs the low transcriptional activity of LINEs and major satellites is due to the activity of endosiRNAs keeping them repressed all the time. However, LINEs as well as the pericentric repeat class are known to be silenced by repressive histone marks (Saksouk et al., 2014, Tsumura et al., 2006, Ip et al., 2012, Goodier and Kazazian, 2008) and I also found increased enrichment of histones upon in upon KO of Dicer and Dnmt1.

The differential silencing of LINE-1 and major satellites primarily by endosiRNA while ERV elements are highly guarded by DNA methylation, could be due to the fact that the older TEs are already domesticated in the mouse genome while ERV elements are still highly active in the male germline. IAP elements are then in the second instance primarily guarded by endosiRNAs while the ETn elements are guarded by repressive histone marks.
I also found a potential role of translational gene silencing of IAP elements, through an miRNA which is transcribed in an intron of Dnmt1. This is very intriguing, as the pre-miRNA of the mature miRNA as well as the IAPs are both confined to the murine lineage. This could suggest that the miRNA mechanism of IAP control has coevolved with IAPs in order to guard the genome from activity of this specific ERVK elements. To test this hypothesis, it is possible to deplete ESCs of this specific miRNA by locked nucleic acids (LNAs) and measure translation of IAP protein by western blot. To reveal whether this specific miRNA also plays a role in the first response to feed into an endosiRNA pathway, as it is the case in plants, it would be necessary to perform the miRNA knockdown at the time of transcriptional expression and check by small RNA-seq, whether endosiRNAs still map to IAP transcripts.

Figure 4.26: Schematic of DICER generated endosiRNAs control TEs upon Dnmt1 deletion in ESCs. TEs in mouse ESCs are methylated (filled lollipops) and get demethylated (open lollipops) upon Dnmt1 KO. Sense transcription of TEs (red) is inhibited by DNA methylation and gets upregulated upon Dnmt1 KO. Genic sense transcription (grey) is constant antisense to TEs. This results in the production of dsRNA and allows the production of DICER dependent (white) endosiRNAs bound by AGO2 (black) which silence TEs via post transcriptional gene silencing. Additionally, histone marks (blue hexagons) become enriched at TEs which get transcriptionally upregulated and may be involved in the subsequent transcriptional silencing.
In summary, I analysed the role of small RNAs during the dynamics of global demethylation in mouse ES cells and were able to show that LINEs, major satellites as well as IAP element expression are dependent on DICER derived endosiRNAs among DNA methylation and histone marks. For other TE elements histone marks and DNA methylation seem to play the primary role in silencing. Additionally, I also found miRNAs to potentially play a role in IAP silencing.

This could expand the way we think about epigenetics as major regulators of TE silencing during early mammalian development. DNA methylation is in place at regions which need to be remodelled. Accompanying this methylome remodelling are major chromatin changes. Potentially small RNAs - such as siRNAs, piRNAs and miRNAs - are involved in keeping TE elements under control to preserve genome integrity for future generations.
Chapter 5

Small RNAs as suppressors of TEs upon demethylation

5.1 Introduction

In chapter 3, I found that during acute demethylation by Dnmt1 deletion endosiRNAs bound by AGO2 map to IAPs, LINEs and major satellites. In chapter 4, I presented evidence that these small RNAs are going through an DICER mechanism of repeat silencing.

In this chapter, I examine whether small RNAs could potentially play a role in TE regulation in other systems of global genome demethylation. Accordingly, I compare the small RNAs mapping to TEs in the transition from (1) serum to 2i grown ESCs, (2) early stages of induced pluripotent stem cell (iPSC) reprogramming as well as (3) in vitro and in vivo reprogramming in PGC development.

Likewise, I study the expression of miRNAs in the three demethylation systems. I compare the miRNAs that influence the pluripotency of ESCs and are indispensable during PGC development across the (1) serum to 2i grown ESCs, (2) early stages of induced pluripotent stem cell (iPSC) reprogramming as well as (3) in vitro and in
miRNAs have been reported to control germ cell development. Knock-out of \textit{Dicer} solely in PGCs, results in poor germ cell proliferation (Hayashi et al., 2008). Additionally, the depletion of the whole miR290-295 cluster in mice leads to embryonic lethality through the impairment of PGC development and premature failure in the ovary (Medeiros et al., 2011). The miR290-295 cluster as well as the miR302-367 are highly abundant in ESCs and as both clusters are cell cycle miRNAs, they are invaluable for ESC self-renewal. The miR16 and miR191 clusters are highly expressed in ESCs and target Smad2 and thereby they inhibit mesoderm differentiation. At the same time, miR23 is able to inhibit the endodermal and ectodermal differentiation. However, miRNAs can also promote differentiation - miR421 inhibits BMP signalling and suppresses the pluripotency factor \textit{Oct4} (Zhao et al., 2014, Hadjimichael et al., 2016).

(1) Mouse ESCs have the ability to generate all somatic and germline cells \textit{in vitro} as well as \textit{in vivo} in chimeric embryos (Smith, 2001). To preserve stem cell identity, pluripotency factors have to be expressed (Nichols et al., 1998, Hochedlinger et al., 2005, Hough et al., 2006, Takahashi and Yamanaka, 2006). However, ESCs express these pluripotency factors heterogeneously in culture (Chambers et al., 2007, Toyooka et al., 2008, Niwa et al., 2009).

This could be due to undefined culturing conditions of ESCs in serum. Since 2008, it is possible to culture ESCs in serum-free conditions in the presence of two inhibitors of the FGF and GSK signalling pathways (2i inhibitors, PD0325901 and CHIR99021) (Ying et al., 2008, Kunath et al., 2007, Wray et al., 2010). ESCs grown in 2i culture conditions adopt a ground state of pluripotency, with loss of mosaic expression levels of pluripotency factors (Wray et al., 2010, Ficz et al., 2013, Marks et al., 2012). A key feature of 2i grown ESCs compared to serum grown ESCs is the global hypomethylation of the whole genome. However, IAPs as well as major satellites partially escape this demethylation (Ficz et al., 2013). Additionally, global histone rearrangements have
been found in the more naïve state ESCs in comparison to serum grown ESCs (Marks et al., 2012, von Meyenn et al., 2016). Earlier studies have shown that this global hypomethylation leads to transcriptional activation of IAPs, MERVL as well as LINE elements, and subsequent resilencing of these TEs by histone redistribution (Walter et al., 2016).

I aim to investigate whether next to histone marks, small RNAs silence TE classes during global hypomethylation during the transition from serum to 2i grown ESCs.

(2) Four transcription factors - cMyc, Klf4, Oct4 and Sox2 - have been identified to reprogram somatic cells into iPSCs (Takahashi and Yamanaka, 2006, Takahashi et al., 2007, Park et al., 2008). In iPSC generation the epigenetic landscape has to be erased to establish a pluripotent state (Maherali et al., 2007, Mikkelsen et al., 2008, Onder et al., 2012, Polo et al., 2012, Papp and Plath, 2013, Lee et al., 2014a). During the generation of iPSCs there is a wave of demethylation (Milagre et al, in preparation). Additionally, it was shown that in iPSC reprogramming endogenous retroviruses (ERVs) become up-regulated (Wissing et al., 2012, Friedli et al., 2014). I generated small RNA-seq libraries during different stages of reprogramming and mapped them to TE classes to investigate whether any small RNAs could be found to control TEs during this event of global hypomethylation.

(3) The precursor of the future gametes - oocyte and sperm - are primodial germ cells (PGCs). A small number of PGCs (n = 40) is first detectable in the epiplast at embryonic day (E)7.25. Afterwards, PGCs migrate to the genital ridges (E8-10.5) and proliferate thereafter extensively. To establish the prerequisite for pluripotency, PGCs have to reset the epigenetic memory in a phase of global epigenetic reprogramming (von Meyenn and Reik, 2015, Reik and Surani, 2015). Due to the limited cell number of PGCs in vivo, in vitro systems to study PGC differentiation was in demand. It has been possible to isolate cells in embryoid bodies (EBs) that express germ cell markers and use them as a proxy for in vitro generation of PGCs (Daley, 2007, Saitou and Yamaji, 2010). However, to properly study PGC development more faithful in vitro systems were demonstrated more recently (Hayashi
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5.2 Results

5.2.1 small RNAs in ESCs during serum/2i transition

I was interested to learn whether the global hypomethylation, that has been reported in the transition from serum to 2i grown ESCs, could lead to an increase of small RNAs that mapped to TEs.

Earlier studies have shown, that the global demethylation happened 72h after switching ESCs serum medium to serum-free 2i medium. Additionally, certain TE classes become transcriptionally up-regulated during serum to 2i transition (Walter et al., 2016). Therefore, I sampled RNA at 24h, 48h and 72h after exchanging the culture conditions, from serum to serum-free 2i medium, and prepared small RNA-seq libraries. I analysed the small RNA composition of the small RNA-seq libraries of the three time-points after adding 2i medium to the ESC using the piPipes small RNA-seq pipeline (Han et al., 2015). Most of the small RNAs were miRNAs (64% miRNAs in 24h 2i, 55% miRNAs in 48h 2i and 42% miRNAs in 72h 2i) (Figure 5.1).
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I observed a compelling increase of small RNAs that mapped to repeats, upon transition of serum to 2i culturing conditions - 7% in 24h 2i, 11% in 48h 2i and 13% in 72h 2i (Figure 5.1). As a consequence, I characterised the small RNA fraction that mapped to repeats further.

![Pie chart of small RNA-seq libraries mapped to 2i grown ESCs for 24h (left), 48h (middle) and 72h (right). miRNAs (black), 5'UTR bound small RNAs (light green), 3'UTR bound small RNAs (yellow), intron bound small RNAs (dark blue), repeats bound small RNAs (dark green), rRNA and tRNA (grey), unannotated small RNAs (white).](image1)

Figure 5.1: Classification of small RNAs mapped to 2i grown ESCs. Pie chart of small RNA-seq libraries mapped to 2i grown ESCs for 24h (left), 48h (middle) and 72h (right). miRNAs (black), 5’UTR bound small RNAs (light green), 3’UTR bound small RNAs (yellow), intron bound small RNAs (dark blue), repeats bound small RNAs (dark green), rRNA and tRNA (grey), unannotated small RNAs (white).

![Bar graph of size distribution of small RNA-seq libraries depleted of small RNAs that mapped to miRNAs in 2i grown ESCs at 24h (yellow), 48h (dark green) and 72h (light green) in comparison to WT ESCs (grey), day 9 Dnmt1 KO ESCs (light blue).](image2)

Figure 5.2: 21nt small RNAs in 2i grown ESCs. Bar graph of size distribution of small RNA-seq libraries depleted of small RNAs that mapped to miRNAs in 2i grown ESCs at 24h (yellow), 48h (dark green) and 72h (light green) in comparison to WT ESCs (grey), day 9 Dnmt1 KO ESCs (light blue).

I studied the size distribution of the small RNA-seq libraries, without the miRNA
fraction, in ESCs at 24h, 48h and 72h after changing into 2i culturing conditions, to analyse whether any distinct size fraction of small RNAs would be present. Indeed, at 24h culturing the ESCs in 2i medium I detected similarly high amounts of 21nt small RNAs as in Dnmt1 KO ESCs at 9 days after KO induction, while the WT ESCs had much less 21nt small RNAs (Figure 5.2). After culturing the ESCs for longer time in 2i - 48h and 72h - the 21nt peak went down.

This is an intriguing finding, as the global demethylation in the transition from serum to 2i has been reported to happen from 48h 2i onwards (von Meyenn et al., 2016, Ficz et al., 2013).

I wanted to know whether the 21nt small RNAs were enriched at a specific TE classes. Therefore, I mapped the 21-24nt fraction of small RNAs, depleted of miRNAs, to LINE, SINE, LTR and DNA transposons. I measured an increase in small RNAs that mapped to SINE elements in 2i grown ESCs. SINE bound siRNAs have been found in mouse ESCs, nonetheless no increase of SINE transcription was observed in earlier studies in 2i ESCs (Babiarz et al., 2008, Walter et al., 2016).

Despite the small RNAs that mapped to SINE elements, the amount of small RNAs that mapped to the other TE families were not change in 2i grown ESCs compared to

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Figure 5.3: Small RNAs map to SINE elements in 2i medium. Bar graph of proportion of small RNAs without the miRNA fraction that mapped to DNA transposons (black), SINEs (dark grey), LINEs (grey), LTRs (light grey).
serum grown ESCs.

This result differs from the response to acute demethylation through *Dnmt1* KO ESCs, where a bigger proportion of small RNAs mapped to ERVs.

To understand whether a change in miRNA profiles was found after the change from serum to 2i medium, I analysed the small RNAs that mapped uniquely to the genome. Reassuringly, I found mostly miRNAs with a size of 21-22nt in the genome in the small RNA-seq quality control plot of Seqmonk (Figure 5.4).

Figure 5.4: miRNAs are the most abundant class of small RNAs mapped to the genome in 2i grown ESCs. Size distribution and classification of small RNAs mapped uniquely to the mouse genome of 2i grown ESCs for 24h, 48h or 72h. miRNAs (grey), miscRNA (red), rRNA (green), tRNA (blue), snoRNA (orange), snRNA (violet).

I compared the expression of known miRNAs, important in PGC development and endogenously expressed in ESCs, across the serum to 2i transition by analysing the miRNAs mapped to the whole genome using Seqmonk (Medeiros et al., 2011, Takada et al., 2009, Hayashi et al., 2008, Zhao et al., 2014, Hadjimichael et al., 2016).

Then, I carried out a hierarchical clustering of the selected miRNAs across the early stages of 2i transition. The clustering was done in Seqmonk using per probe normalised counts and therefore the heatmap depicts changes in miRNA expression across the different time points. miRNAs in cluster I were high expressed at 24h and 48h 2i but became down-regulated in 72h 2i. Cluster I encompassed the miR302 family, which is known to control ESC self renewal by controlling the cell cycle. miRNAs in cluster II were highly expressed in 24h 2i, but down-regulated thereafter. This cluster contained miRNAs, highly expressed in ESCs, like miR200, miR23a, miR421. The third cluster (III) was low expressed at both 24 and 48h 2i, but miRNAs increased expression at
72h 2i. Among these miRNAs were miR-let7g - normally expressed in differentiated cells (Figure 5.5).

![Heatmap of miRNAs expression in 2i grown ESCs.](image)

Figure 5.5: Heatmap of miRNAs expression in 2i grown ESCs. Heatmap of hierarchical clustering of miRNAs expressed in ESCs as well as important in PGC development were compared over the time-course of serum to 2i transition. Per probe normalised.

### 5.2.2 small RNAs in iPSCs reprogramming

In the lab Inês Milagre performed reprogramming of mouse MEFs to iPSCs and assessed the global methylation levels at certain time-points after reprogramming. The global demethylation in iPSCs reprogramming was confirmed on all genic regions. The demethylation went down to 45% with the biggest hypomethylation at passage 12 (data not shown). This global demethylation was also found in all TEs - with resistance to demethylation in IAPs (Seisenberger et al., 2012, Kobayashi et al., 2012) - which opened the possibility for TE transcriptional upregulation and thus another system where small
RNAs could play a role in TE repression (Milagre et al., in preparation).

Figure 5.6: Classification of small RNAs in iPSCs reprogramming. Pie chart of small RNA-seq libraries mapped to MEFs (left), passage 12 (middle) and 17 (right) iPSCs. miRNAs (black), 5′UTR bound small RNAs (light green), 3′UTR bound small RNAs (yellow), intron bound small RNAs (dark blue), repeats bound small RNAs (dark green), rRNA and tRNA (grey), unannotated small RNAs (white).

Figure 5.7: No specific fraction of siRNAs in early stages of iPSCs reprogramming. Bar graph of size distribution of small RNA-seq libraries without miRNAs, mapped to MEFs (yellow), passage 12 (orange) and passage 17 (green) iPSCs.

For this reason, I produced small RNA-seq libraries at four time-points during reprogramming. For the classification I used the piPipes small RNA-seq pipeline (Han et al., 2015). First of all, I classified the small RNAs. 72% of the small RNAs in MEFs were miRNAs, with only 5% repeat mapped small RNAs. However during reprogramming in passage 12 iPSCs, the miRNA fraction went down to 14% miRNAs, while the fraction
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of repeats bound small RNAs increased to 28%. In the iPSCs at passage 17 the miRNAs made up 40% of the total small RNAs with 17% mapping to repeats. I wanted to test whether the small RNAs at passage 12 were siRNAs (Figure 5.6). The intriguing increase of small RNAs that mapped to repeats during iPSCs reprogramming motivated me to study the small RNAs depleted of miRNAs further. However, I only found 22nt small RNAs in MEFs, while during the iPSCs reprogramming no discrete size of small RNAs was detected. I concluded that no siRNAs are mapping to TEs during iPSCs reprogramming at early stages of reprogramming (Figure 5.7).

Even though, the miRNA fraction decreased during iPSCs formation, I observed miRNAs with a mean size of 21-22nt in MEFs, iPSCs passage 12, 17 as well as passage 31 in the small RNA-seq quality control plot of Seqmonk (Figure 5.7) after mapping the small RNA-seq reads to the whole genome. I analysed the expression of miRNAs, known to be important during early mammalian development and essential for ESC pluripotency, throughout iPSCs reprogramming (Medeiros et al., 2011, Takada et al., 2009, Hayashi et al., 2008, Zhao et al., 2014, Hadjimichael et al., 2016).

I performed a hierarchical clustering with per probe normalisation in Seqmonk. This resulted in a division of miRNAs into two main clusters: (cluster I) miRNAs high expressed in MEFs and down-regulated thereafter, (cluster II) miRNAs not expressed in MEFs and up-regulated thereafter. Cluster I was reassuringly comprised of the miR-let7 family, which is known to be highly expressed in differentiated tissues but lowly

Figure 5.8: Classification of small RNAs of iPSCs mapped to the genome. Bar graph of size distribution and classification of small RNA-seq libraries at different passages of iPSCs during reprogramming in comparison to MEFs. miRNAs (grey), miscRNA (red), rRNA (green), tRNA (blue), snoRNA (orange), snRNA(violet).
expressed in pluripotent cells. Cluster II contained the miR290-295 family, which is also known to be important for proper PGC development, as well as the miR200 family which is highly expressed in ESCs (Figure 5.9). Interestingly, the expression of cluster II miRNAs was downregulated again at passage 31.

Figure 5.9: Heatmap of miRNAs expression in iPSCs during reprogramming. Heatmap of hierarchical clustering of miRNAs expressed in ESCs as well as important in PGC development were compared during iPSCs generation.

5.2.3 small RNAs in mouse PGC like cells (PGCLC) and PGCs

piRNAs in vivo and in vitro PGCs

miRNAs as well as piRNAs have been studied in male PGCs at E16.5 in vivo PGCs (Hayashi et al., 2008, Molaro et al., 2014). Here, we carried out a comprehensive study
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of small RNAs in E13.5 to E15.5 male and female PGCs. Additionally, Ferdinand von Meyenn setup a protocol to study in vitro mouse PGC like cells (PGCLC) (von Meyenn et al, 2016, accepted) and I generated small RNA-seq libraries of PGCLCs from two different mouse ESC strains - E14 and C57BL/6 (B6) - and compared them to WT E14 ESCs small RNA-seq libraries, as a control.

I found 54% small RNAs that mapped to repeats with only 9% small RNAs mapping to miRNAs in the PGCLC in comparison to mESCs, where 60% of the library was made up of miRNAs and only 15% of small RNAs mapped to repeats (Figure 5.10).

The small RNAs of the PGCLC, that mapped to repeats were 24-32nt long and had a thymidine overhang at the 5’ ends. Additionally, these small RNAs showed the piRNA profile of a 5’-5’ overlap at nucleotide 10 with reads mapping to sense and antisense strands of repeats (Figure 5.11).

![Figure 5.10: Classification of small RNAs mapping to in vitro PGCs.](image)

I wanted to compare the PGCLC small RNA-seq libraries to the in vivo PGCs. Wendy Dean isolated in vivo PGCs of male and female at E13.5, E14.5 and E15.5 and I prepared small RNA-seq libraries of them. In comparison to PGCLC, I found similar classes of small RNAs in the small RNA-seq libraries of male and female in vivo PGCs, at each developmental stage assessed. I measured 54% small RNAs mapping to repeats and only 9% mapped to miRNAs (Figure 5.12). The size distribution of all of the in vivo PGC small RNAs mapping to repeats, showed that they were comprised of a class of 20-24nt and a class of 24-32nt small RNAs, while in ESCs I only detected a peak of
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20-24nt small RNAs and no signature of piRNAs (Figure 5.13A, B).

Figure 5.11: piRNAs from PGCLCs mapped to TEs. (A) Bargraph of nucleotide contribution along the stretch of the small RNAs mapped to repeats, (B) Bargraph of 5’5’ overhang of small RNAs mapped to repeats, (C) Bargraph of size distribution of small RNA-seq libraries mapped to TE classes of sense (red) and antisense (blue) reads.

Figure 5.12: Classification of small RNAs mapping to in vivo PGCs. (A) Pie chart of small RNA-seq libraries mapped to male E15.5 PGCs in comparison to E14 mouse ESCs. miRNAs (black), 5’UTR bound small RNAs (light green), 3’UTR bound small RNAs (yellow), intron bound small RNAs (dark blue), repeats bound small RNAs (dark green), rRNA and tRNA (grey).

To analyse whether piRNAs map to active TEs in the mouse germline, I mapped the small RNA-seq data of PGCLCs and in vivo PGCs to the consensus sequence of the IAPEZ element. I detected mostly 22nt small RNAs in ESCs (at very low level) and a high number of small RNAs of size 24-29nt mapping to IAPs in PGCLCs. A small number of small RNAs of the size of 22nt could be found in PGCLCs. I mapped the small RNA-seq libraries of in vivo PGCs to IAPEZ consensus sequence and found small RNAs mainly of the size of piRNAs mapping to IAPs. Intriguingly, in E13.5 PGCs I found small amount of piRNAs mostly from male PGCs mapping to IAPs, while at
E14.5 mostly female small RNAs were mapping to IAPEZ consensus sequence, while at E15.5 the piRNAs of the male PGCs are getting very enriched over the female piRNAs (Figure 5.14).

**Figure 5.13: TE bound small RNAs in vivo PGCs and mouse ESCs.** (A) Bar graph of size distribution of small RNA-seq libraries mapped to the whole genome of ESCs and PGCs, miRNAs/siRNAs (light grey), piRNAs (dark grey), (B) Bar graph of nucleotide contribution along the stretch of the small RNAs mapped to repeats in PGCs and ESCs.

A small number of small RNAs of the size of 22nt could be found in PGCLCs as well as during in vivo PGC development. This class of 22nt small RNAs was of interest, as it could represent a class of endosiRNAs acting in vivo during PGC development. For this reason, I examined whether these 22nt small RNAs would also map to other TE classes.

**endosiRNAs in vivo and in vitro PGCs map to TE s**

In order to see whether the 20-24nt small RNAs mapping to all repeats in vivo PGCs, were also endosiRNAs, working potentially through a similar mechanism of TE silencing I presented during global hypomethylation in mouse ESCs (Chapter 3 and 4), I
only mapped 20-24nt small RNAs to all repeats. While most of these small RNAs were miRNAs, I found 10% repeat bound small RNAs (Figure 5.15A). These small RNAs mapped to both sense and antisense of all repeats and had a peak at 21nt (Figure 5.15B).

**Figure 5.15:** Contribution of 20-24nt small RNAs mapped to repeats. (A) Pie chart of small RNA-seq with the size of 20-23nt of male E13.5 libraries mapped to all repeats. miRNAs (black), 5’UTR bound small RNAs (light green), 3’UTR bound smallRNAs (yellow), intron bound small RNAs (dark blue), repeats bound small RNAs (dark green), rRNA and tRNA (grey), unannotated small RNAs (white), (B) Bargraph of size distribution of small RNA-seq libraries mapped to TE classes of sense (red) and antisense (blue) reads,
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I mapped the 20-24nt small RNA fraction of the *in vivo* E13.5 and E14.5 PGC libraries to all the repeat classes dependent on *Dnmt1* and *Dicer* defined in Chapter 3 and 4. I measured an increase of small RNAs that mapped to IAPs as well as L1MdGf in E13.5 and E14.5 males, while no small RNAs mapped at MERVL, ETnERV2 and MMERVK10C (Figure 5.16).

**miRNAs in vivo and in vitro PGCs**

To finish the characterisation of small RNAs *in vitro* PGCLC as well as *in vivo* PGCs, I investigated the small RNAs that mapped uniquely to the genome.

I analysed the expression pattern of miRNAs expressed during PGC development, as well as miRNAs which are confined to the ES cell lineage, in the *in vivo* and *in vitro* PGCs small RNA-seq libraries. I performed hierarchical clustering per probe normalised in Seqmonk. Most miRNAs fell into the same cluster. While in *in vitro* PGCs all of the miRNAs were lowly expressed, all of them increased in expression *in vivo* PGCs. Despite this the miRNAs did not form apparent clusters. I could recapitulate
the effect of remethylation of male PGCs earlier than female PGCs from E14.5 onwards, as most of the miRNAs were highly expressed in male E13.5 PGCs but already down-regulated at E14.5 and E15.5 male PGCs. miR-let7g was highly expressed in all *in vivo* samples but not in the *in vitro* PGCLCs.

**Figure 5.17:** Heatmap of miRNAs expression *in vitro* and *in vivo* PGCs. Heatmap of hierachical clustering of miRNAs expressed in ESCs as well as during PGC development in our *in vivo* PGCs cells at E13.5, E14.4 and E15.5 in male as well as female, and *in vitro* PGCLCs.

In conclusion, not in all demethylation systems I found small RNAs mapping to TEs. Although in our demethylation system with *Dnmt1* deletion, as well as during *in vitro* and *in vivo* PGCs and in the serum/2i transition, we detected endosiRNAs next to
piRNAs that map to TE elements, in iPSCs we could not find small RNAs of a distinct size that mapped to TEs.

During the miRNA analysis I confirmed the expression of miR290-295 cluster in pluripotent states, while the let-7 class was higher expressed in differentiated cells like MEFs (Zhao et al., 2014, Hadjimichael et al., 2016, Medeiros et al., 2011, Takada et al., 2009, Hayashi et al., 2008, Jia et al., 2013, Bhin et al., 2015).

5.3 Discussion

In the earlier two chapters, I proposed the role of DICER small RNAs in TE silencing during acute demethylation in ESCs.

In this chapter I wanted to test the hypothesis, that the suggested mechanism could also act in other systems of global hypomethylation.

To begin, I investigated the existence of endosiRNAs in a transition from serum to 2i grown ESCs. The global hypomethylation induced through the addition of FGF and GSK signalling inhibitors, leads to transcriptional upregulation of TE classes (Walter et al., 2016). The highest abundance of 21nt small RNAs were present at 24h after 2i transition. However the DNA demethylation is only apparent after 48h after 2i transition (Ficz et al., 2012, von Meyenn et al., 2016). This suggests, that the small RNAs were responding very quickly to the change from serum to 2i conditions in ESCs, irrespective of DNA methylation. To verify whether 21nt RNAs are increased prior to 24h 2i, small RNA libraries from earlier time-points should be sequenced.

Most of the 21nt small RNAs were mapping to SINE elements and while SINE siRNAs have been found before in mESCs, no SINE up-regulation was detected in serum to 2i transition (Barbierz et al., 2008, Walter et al., 2016). This could mean that the SINE RNAs are guarded by endosiRNAs during serum/2i transition, but are dispensable of DNA methylation, similarly to LINE elements in the acute demethylation system. However, the nature of the 21nt small RNA fraction should be verified by AGO2 IP small RNA-seq.
As a second system of global demethylation, I examined stages of mouse iPSCs reprogramming. I measured 28% and 17% small RNAs binding to repeats in passage 12 and 17 iPSCs - respectively. However, I could not observe any distinct size classes of small RNAs and therefore concluded that at the stages we sampled siRNAs were not mapping to TEs in iPSCs.

This could be because I had very limited amount of RNA (10ng of total RNA) of input material for passage 12 iPSCs. Small RNA-seq library preparation for small input libraries are now available, therefore it would be possible to sample these early time-points during iPSCs reprogramming in a future study.

As a third system, I used in vitro and in vivo PGCs. In the lab we were able to generate PGCLCs as an in vitro system for PGC development. I observed 1.4% of small RNAs that mapped to piRNA clusters as well as 55% small RNAs that mapped to repeats and showed piRNA properties, the small RNA contributions of PGCLCs was comparable to the amounts of piRNAs and repeat mapped small RNAs in vivo male PGC at E15.5. We were able to present the first in vitro model to study mammalian piRNAs in PGCLCs (von Meyenn et al, 2016, accepted). In addition, I produced small RNA-seq libraries from E13.5, E14.5 and E15.5 male and female and also found small RNAs of the size of 24-29nt length mapping to IAPEZ in the E14.5 female PGCs. We also found the typical piRNA features in the fraction of small RNAs that mapped to repeats.

Further to the TE bound small RNAs, I examined the expression of miRNAs in the different demethylation systems. I found miRNAs expressed in naïve ESCs, in iPSCs as well as in PGC development. In the serum to 2i transition miR200, a highly abundant miRNA in ESCs, as well as mir23a, known to suppress ectoderm and endoderm development, were down-regulated in 72h 2i conditions. In addition to this, I measured an increase of the miR-let7 family - known to be expressed in differentiated cell lines in the 72h 2i samples. This suggests that in the early phases of 2i transition, small RNAs were very respondent and changed expression levels quickly. However, it is very com-
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PELLING that miRNAs, that are known to prevent differentiation, were down-regulated after culturing the ESCs for 72h in 2i, as 2i grown ESCs have a transcriptome more closely to ICM stage and have been also described as naïve ESCs. To reveal further insights later time points might explain whether the miRNA profile of 2i ESCs is still changing after 72h of culturing ESCs in 2i medium.

miRNA expression in the iPSCs reprogramming verified the reprogramming of the iPSCs, as miRNAs known to be expressed in differentiated tissues were highly expressed in MEFs but down-regulated thereafter.

Also in PGC development our data supported earlier studies, as we found all the miRNAs important for PGC development to be expressed. Interestingly, I found that miRNAs at E14.5 and E15.5 in the male PGCs were down-regulated more then the same stage female PGCs, which could be explained by the global remethylation of the male PGCs, while the female PGCs are still hypomethylated.

Noteworthy, the presence of endosiRNAs mapping to TE classes during in vivo PGC reprogramming at E14.5 opens the possibility that our in vitro system of Dnmt1 induced global hypomethylation - presented in chapter 3 and 4 - presents a model also for in vivo TE regulation.

During global hypomethylation the activation of TE elements presents a risk for the genome integrity. In order to prevent retrotransposition events, molecular mechanisms need to be in place to suppress TE activity. Here, I present a mechanism of endosiRNAs, that can silence TEs and protect the genome during acute demethylation.
Chapter 6

Conclusions and Outlook

Transposable elements (TEs) refer to DNA sequences which can move from one location in the genome to another; the result of these movements or retrotranspositions can be either detrimental or advantageous to the genome.

On one hand, TEs activity can lead to chromosomal breakage, large-scale genomic rearrangements as well as disruption of protein coding genes and alteration of transcriptional regulatory networks (McClintock, 1951). In addition, TEs are responsible for 10-15% mutations in mice (Kazazian and Moran, 1998, Maksakova et al., 2006).

On the other hand however, TE insertions may also lead to the derivation of protein-coding genes with new and potentially beneficial functions for the host genome. A key example of this is the domestication of the Rag1 gene, which is invaluable for the adaptive immune system and that evolved through TE endogenisation. Another examples is the MERVL transcriptional network which, mostly driven by “solo” LTRs, seems to be implicated in activation of the transcriptional network at the two-cell stage of mouse embryonic development.

These observations lead to the intriguing question of the apparently mutualistic relationship between host genome and TEs. This is sometimes referred to as a “friend or foe?” relationship: while transcriptional activity of TEs is needed to drive evolution, this activity must be tightly controlled by the host genome, since transposons can also disrupt protein-coding genes. Thus, in order to preserve genome integrity, the cellular machinery must guard the host DNA from the activity of TEs. This presents major
challenges for the host, firstly as the genome has to distinguish TEs from protein-coding genes and secondly because a plethora of TE families exist in the genome as they arise through vertical gene transfer. This leads to an “arms race” between genome and TEs: while the genome tries to find ways to control TE activity to preserve genome integrity TEs continuously evolve into different varieties to escape these control mechanisms.

Several epigenetic marks have been implicated in TE control. It has been suggested that DNA methylation actually evolved in order to protect the genome from TE activity and the repressive histone marks H3K9me3 and H3K9me2 are known to be involved in ERV silencing in ESCs. Additionally, small RNAs have been described to play a role in TE control.

While, TE mobilisation is undoubtedly advantageous to TEs, it also presents a disadvantage as this transcriptional activity can activate small RNA-based mechanisms that serve as “guardians of the genome”. For example, TEs that retrotranspose into piRNA clusters are silenced through piRNA pathways during spermatogenesis and feed into this small RNA based immune system. miRNAs and endosiRNAs have also been found to be involved in TE silencing in the oocyte (Flemr et al., 2013, Stein et al., 2015).

It is especially during the global hypomethylation stage of epigenetic reprogramming in the mammalian genome, that genome integrity is in danger. Global hypomethylation allows for transcriptional activation of LINEs and ERVs in oocyte and blastocyst and a burst of IAP transcriptional activity has been found to occur during PGC reprogramming (Molaro et al., 2014). As such the genome is particularly vulnerable for retrotransposition events, with the added risk of germline mutations occurring, and thus it is possible that silencing mechanisms other than DNA methylation have to be put in place to control TE mobilisation at this time.

In this thesis I performed a mechanistic study to follow TE activity after acute demethylation with a system that strongly recapitulates in vivo epigenetic reprogramming: I used Dnmt1 conditional KO ESCs and followed demethylation dynamics after KO induction, by WGBS-seq, total RNA-seq, small RNA-seq and ChIP-seq.
In chapter 3, I show that *Dnmt1* conditional KO leads to genome wide hypomethylation and transcriptional upregulation of ERV elements. The mouse genome presents the perfect model to study TE regulatory mechanisms, as mouse TEs are younger than predominantly inert human TEs, most of them having integrated in the last 25 Myrs containing around 1000 active LINEs and ERVs. Therefore, the mouse genome is still in the process finding ways to protect itself from the mobility of TEs.

I also uncovered a new potential genomic mechanism to identify and target TE transcriptional activity. I observed that in areas where there was genic pervasive transcription, a TE had integrated and that these transcripts can as such serve as an antisense strand to TE sense transcription and feed into an endosiRNA pathway of TE silencing. During a phase of evolutionary expansion TEs do not have a preference for integration in any genomic location, however, TEs are most commonly domesticated in intergenic regions. This may occur through positive selection in order to prevent TE insertions that could disrupt gene expression. For example it has been shown that ERVs are selected to mostly persist intergenically and that, if ERVs are integrated into genic regions they are manifested in antisense direction to the genes ([Medstrand et al., 2002](#)).

In chapter 3 I present evidence that the presence of complimentary TE sense and antisense transcripts could demonstrate a way by which the genome could control the activity of TEs during TE mobilisation, as the genic regions works to constantly produce antisense strand to the TE’s sense transcript. The genic sense strand may thus be regarded as a signal of TE transcription. The TE sense transcript and the genic sense transcript can produce dsRNAs and can feed into an endosiRNA pathway and controlling retrotransposition. This model is supported by findings in the yeast genome ([Cruz and Houseley, 2014](#)). The produced endosiRNA are going through an AGO2 dependent mechanism and are potentially involved in TE regulation subsequent to their initial transcriptional activation.

In chapter 4, I performed *Dicer* KO in the *Dnmt1* KO ESCs. This mechanistic study allowed me to identify DICER dependent pathways that are essential for transcriptional resilencing of certain TE classes. Interestingly, the TEs which were dependent upon DICER activity were IAP elements, which are one of the most active classes of
TEs in the mouse germline.

I also identified an apparently cooperative effect in the defence mechanisms working to silence TE elements. I found IAPs and ETn elements, which were largely dependent on DNMT1 as shown in chapter 3, to be transcriptionally silenced in the double KO of Dicer/Dnmt1 and, as such, by an unknown mechanism. Subsequently, analysis of ChIP-seq data showed an enrichment of H3K27me3 at these sites upon double KO of Dicer/Dnmt1. This is a very intriguing finding, as it could mean that PRC mediated pathways of H3K27me3 deposition may be activated, to preserve TE silencing, upon global hypomethylation in the absence of small RNAs. Future research, possibly with histone methyltransferase knockouts and knockouts of players of the PRC complexes, would allow further insight into this.

Additionally, I found indications that miRNAs might play a role in IAP silencing. The miRNA in the intron of Dnmt1 is only conserved in the mouse genome and thus has potentially evolved in conjunction with the murine-lineage-specific-IAP elements and may play a role in the control of IAPs. miRNAs might play similar roles to silence young TE elements in other lineages. Future experiments involving depletion of this miRNA will allow more insight into this intriguing finding.

In chapter 5, I looked for small RNA based silencing mechanisms that control TEs during the three incidences of global hypomethylation: serum to 2i culturing of ESCs, iPSC reprogramming and in vitro and in vivo PGCs.

In the serum to 2i transition of ESCs I found increased small RNAs binding to SINE elements. This is a very interesting preliminary result and could shed light onto other systems and other TE classes that may become silenced through endosiRNA activity.

Knowledge of TE activity is particularly important in iPSCs as it could result in deleterious effects on the genome and iPSCs are used for regenerative medicine. In chapter 5 I assessed endosiRNAs as a potential mean to control mobility of TEs during iPSC reprogramming. I was unable to find any small RNAs that control TE activity in iPSCs at passage 12, 17 and 31, however future experiments should concentrate on additional time points during iPSC reprogramming, to clarify this result.
Intriguingly, \textit{in vivo}, endosiRNAs were found to bind to TE classes during PGC development. As these small RNAs had characteristics of endosiRNA they suggest the \textit{in vitro} mechanism I uncovered in mouse ESCs upon global hypomethylation to potentially be active during PGC development in the germline. Finally, I was able to detect piRNAs in \textit{in vitro} PGCs, which presents the possibility to mechanistically study those small RNAs and their putative role in control of TEs in greater detail.

In summary, I have presented endosiRNAs to be guardians of the genome, they act to silence TEs by restricting TE expression during acute hypomethylation conditions. This study has also suggested a possible connection between small RNA pathways and histone modifications working in conjunction to keep TEs silenced. My research highlights young classes of TEs, which are still retrotransposition active, to be targeted by several histone modifications and endosiRNA pathways. The different ways of TEs silencing presented in this study indicate the challenge the host genome has to preserve genome integrity by preventing TE retrotransposition events. Several ingenious silencing mechanisms have had to evolve and must be engaged at different stages during early development to allow sufficient protection of the germline DNA from TEs.
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Smalheiser, N. R. and V. I. Torvik

Smallwood, A., P.-O. Estève, S. Pradhan, and M. Carey


Smit, A. F.

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Smith, A. G.


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Sookdeo, A., C. M. Hepp, M. A. McClure, and S. Boissinot


Soper, S. F. C., G. W. van der Heijden, T. C. Hardiman, M. Goodheart, S. L. Martin, P. de Boer, and A. Bortvin

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Speek, M.

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Stocking, C. and C. A. Kozak

Suda, Y., M. Suzuki, Y. Ikawa, and S. Aizawa

Suetake, I., F. Shinozaki, J. Miyagawa, H. Takeshima, and S. Tajima

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Suzuki, M. M. and A. Bird

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Takada, S., E. Berezikov, Y. L. Choi, Y. Yamashita, and H. Mano

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Tang, G., B. J. Reinhart, D. P. Bartel, and P. D. Zamore


Taunton, J., C. A. Hassig, and S. L. Schreiber


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Thomas, J. H. and S. Schneider


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von Meyenn, F. and W. Reik


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Wu, H. and Y. Zhang


Xiong, Y. and T. H. Eickbush


Xu, M., Y. You, P. Hunsicker, T. Hori, C. Small, M. D. Griswold, and N. B. Hecht


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Yang, N. and H. H. Kazazian


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Zhang, T., S. Cooper, and N. Brockdorff

Zhang, Y., I. A. Maksakova, L. Gagnier, L. N. van de Lagemaat, and D. L. Mager

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Zheng, H., B. Huang, B. Zhang, Y. Xiang, Z. Du, Q. Xu, Y. Li, Q. Wang, J. Ma, X. Peng, F. Xu, and W. Xie

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Zhu, J.-K.
# Nomenclature

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**Expression of differentially expressed genes between WT and Dnmt1 KO.**

Expression levels as log2 RPKM values. Upreregulated under Dnmt1 KO.
Expression of differentially expressed genes between WT and Dnmt1 KO. Expression levels as log2 RPKM levels. Downregulated upon Dnmt1 KO.

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**Expression of differentially expressed genes upon Dicer and Dnmt1 KO in cluster III.** Expression values given as log2 RPKM levels.
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Expression of differentially expressed genes upon Dicer and Dnmt1 KO in cluster III. Expression levels as log2 RPKM levels.