The RNA-induced transcriptional silencing complex targets chromatin exclusively via interacting with nascent transcripts

Yukiko Shimada,1,2 Fabio Mohn,1,2 and Marc Bühler1,2

1Friedrich Miescher Institute for Biomedical Research, 4058 Basel, Switzerland; 2University of Basel, 4003 Basel, Switzerland

Small RNAs regulate chromatin modification and transcriptional gene silencing across the eukaryotic kingdom. Although these processes have been well studied, fundamental mechanistic aspects remain obscure. Specifically, it is unclear exactly how small RNA-loaded Argonaute protein complexes target chromatin to mediate silencing. Here, using fission yeast, we demonstrate that transcription of the target locus is essential for RNA-directed formation of heterochromatin. However, high transcriptional activity is inhibitory; thus, a transcriptional window exists that is optimal for silencing. We further found that pre-mRNA splicing is compatible with RNA-directed heterochromatin formation. However, the kinetics of pre-mRNA processing is critical. Introns close to the 5′ end of a transcript that are rapidly spliced result in a bistable response whereby the target either remains euchromatic or becomes fully silenced. Together, our results discount siRNA–DNA base pairing in RNA-mediated heterochromatin formation, and the mechanistic insights further reveal guiding paradigms for the design of small RNA-directed chromatin silencing studies in multicellular organisms.

Keywords: heterochromatin; RNAi; noncoding RNA; epigenetics; nascent transcript

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RNAi broadly refers to silencing pathways that depend on conserved Argonaute family proteins to repress gene expression (Höck and Meister 2008). A unifying feature of all RNAi pathways is that Argonaute-bound small RNAs guide protein effector complexes to complementary targets to mediate gene silencing. Although very diverse in composition, these are commonly termed RNA-induced silencing complexes [RISCs] [Pratt and MacRae 2009]. Besides mediating sequence-specific degradation or translational repression of target mRNAs (PTGS), small RNAs are also implicated in chromatin modification and transcriptional gene silencing [TGS] in ciliates, fungi, plants, and the germline of animals (Malone and Hannon 2009; Mochizuki 2010; Castel and Martienssen 2013). Pioneering studies in plants and the fission yeast Schizosaccharomyces pombe have provided a conceptual framework for studying the molecular mechanisms of these conserved small RNA-directed chromatin silencing pathways (Moazed 2009; Matzke and Mosher 2014). However, due to technical limitations, some of their most fundamental mechanistic aspects have remained obscure. For most systems studied so far, it is unclear whether the siRNA-loaded RISC targets the DNA or the nascent transcript of a complementary locus. To address this, we made use of the well-understood fission yeast RNAi pathway.

The single S. pombe Argonaute protein (Ago1) is at the core of the S. pombe RISC, known as the RNA-induced transcriptional silencing (RITS) complex [Verdel et al. 2004]. RITS is loaded predominantly with small RNAs complementary to centromeric repeats and is essential for heterochromatin formation at centromeres (Verdel et al. 2004). Upon target recognition, RITS recruits the RNA-dependent RNA polymerase (RDRC), which synthesizes dsRNA using the targeted transcript as a template. This results in additional substrate for siRNA processing by the ribonuclease Dcr1, establishing a positive feedback loop that confers stability to constitutive heterochromatin [Motamedi et al. 2004; Colmenares et al. 2007]. Methylated histone H3 Lys9 [H3K9] is a conserved hallmark of heterochromatin [Allis and Jenuwein 2016]. This chromatin modification is installed by the Clr4 complex [CLRC] (Hong et al. 2005; Horn et al. 2005; Jia et al. 2005), which is recruited to the target locus by RITS (Bayne et al. 2010; He et al. 2013). Clr4 is essential for setting up a stable RNAi-induced heterochromatin domain and is notably the sole S. pombe histone H3K9 methyltransferase.

Corresponding author: marc.buehler@fmi.ch
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Central to the understanding of RNAi-directed heterochromatin formation is how RITS is targeted to chromatin. The current view is that siRNA-programmed RITS initially recognizes the nascent transcript of its target locus. However, this has not been demonstrated directly. It is supported by the observation that both RITS and the RDRC can be cross-linked to centromeric RNA as well as to DNA [Motamedi et al. 2004]. This suggests that RITS is targeted to chromatin through base pairing between siRNA and pre-mRNA followed by recruitment of RDRC and histone-modifying enzymes [Bühler et al. 2006; Bühler and Moazed 2007]. However, it does not rule out base pairing of siRNAs also with the target locus DNA. Moreover, it has remained elusive whether pre-mRNAs are targeted by siRNA-programmed RITS and what impact, if any, mRNA splicing has on the kinetics of heterochromatin formation.

In wild-type S. pombe cells, siRNAs do not trigger the formation of stable heterochromatin at euchromatic genes in trans [Sigova 2004; Bühler et al. 2006]. This has hampered a systematic mechanistic dissection of siRNA-mediated chromatin silencing. We recently discovered that siRNA-directed heterochromatin formation in S. pombe is inhibited by the RNA polymerase-associated factor 1 complex [Paf1C]. Paf1C mutant strains are highly susceptible to de novo assembly of heterochromatin and stable gene silencing by synthetic, trans-acting primary siRNAs complementary to the coding sequence of protein-coding genes [Kowalik et al. 2015]. This has provided us with a unique tool to address these remaining fundamental mechanistic questions. In this study, we demonstrate for the first time that RITS cannot target DNA in S. pombe. We also found that transcription above a minimal threshold is obligatory for small RNA-directed heterochromatin formation but that high transcriptional activity at an siRNA target locus counteracts heterochromatin assembly. We further show that pre-mRNA splicing is compatible with RNAi-directed heterochromatin formation and that intronic sequences can serve as binding sites for siRNAs that are acting in trans. Together, our results provide the first direct evidence for the nascent transcript model of small RNA-directed epigenetic gene repression and reveal guiding paradigms for the design of small RNA-directed heterochromatin silencing experiments in other organisms.

Results

siRNAs complementary to intronic sequences trigger formation of heterochromatin

Current models of RNAi-directed heterochromatin assembly propose that RISC/RITS is targeted to chromatin via base-pairing interactions with nascent or chromatin-bound RNAs [Motamedi et al. 2004; Nakama et al. 2012; Holoch and Moazed 2015]. However, it has not been demonstrated that truly nascent transcripts (i.e., unspliced pre-mRNAs) can be targeted by siRNA-programmed RITS for the formation of heterochromatin in fission yeast.

To test this directly, we used S. pombe strains in which the leo1+ gene was deleted. Leol is a protein subunit of Paf1C, which inhibits siRNA-directed assembly of heterochromatin at euchromatic genes [Kowalik et al. 2015]. In addition, we used a synthetic RNA hairpin construct that is expressed from the nmt1+ locus, driven by an adh1+ promoter [Fig. 1A; Iida et al. 2008]. This construct contains a 355-nucleotide (nt) intron from the cox4+ gene, separating inverted repeats that are complementary to ura4+ and encoding the stems of the hairpin [Fig. 1A]. Besides siRNAs originating from the double-stranded ura4+ sequence in the hairpin, comparable amounts of siRNA reads are generated from the cox4+ intron in the loop of the hairpin construct [Fig. 1A; Yu et al. 2014; Kowalik et al. 2015]. The hairpin-derived small RNAs are, on average, 23 nt in length and predominantly start with a uridine, confirming that they are bona fide siRNAs [Fig. 1B; Supplemental Fig. S1].

We showed previously that the intronless ade6+ gene is highly susceptible to siRNA-directed assembly of heterochromatin in Paf1C mutant but not wild-type cells [Kowalik et al. 2015]. To assess the impact of splicing on the formation of heterochromatin at the endogenous ade6+ locus, we inserted the cox4+ intron sequence in the middle of the ade6+ ORF by homologous recombination. To discriminate between exonic and intronic targeting, we inserted the cox4+ intron in either forward or reverse orientation (target 1 and target 2, respectively). Insertion of a GFP sequence of similar length but devoid of any canonical splice sites (target 3) served as the control for siRNA specificity [Fig. 1C]. RT–PCR analysis of total RNA revealed effective ade6+ splicing if the cox4+ intron is transcribed in its forward orientation [Fig. 1D]. As expected, the cox4+ intron in reverse orientation or the GFP fragment did not enable ade6+ pre-mRNA splicing.

Consistent with our previous results, we observed methylation of H3K9 at the nonsliced locus only in the presence of complementary hairpin-derived cox4+ siRNAs [Fig. 1E, target 2]. In contrast, in the absence of sequence complementarity to the siRNA in target 3, H3K9 methylation was not induced irrespective of the presence of cox4+ siRNAs [Fig. 1E]. Importantly, we also observed cox4+ siRNA-directed H3K9 methylation when the siRNA targeted region is spliced in target 1 [Fig. 1E]. The level of H3K9+ methylation was the same at both the spliced and nonsliced target loci, and the ade6+ mRNA levels of both targets were strongly reduced upon siRNA-directed H3K9+ methylation [Fig. 1F]. Furthermore, in leo1A but not leo1+ cells, secondary siRNAs were generated at the targeted ade6+ locus [Fig. 1G; Supplemental Fig. S2], similar to our previous findings when providing ade6+ siRNAs [Kowalik et al. 2015]. These secondary small RNAs have the characteristic siRNA signature [Fig. 1H], and their spreading is highly directional, as expected, due to the activity of RDRC. Notably, besides generating abundant secondary siRNAs toward the 5′ end of the ade6+ locus, spreading extends even several kilobases beyond, including the bab1+ gene [Fig. 1I]. Thus, intron-targeting primary siRNAs are sufficient to effectively initiate silencing of ade6+, including recruitment of RDRC. These results unequivocally show that pre-mRNA splicing is compatible with RNAi-directed heterochromatin formation and that intronic sequences can serve as binding sites for siRNAs that act in trans.
Promoter-proximal introns are inferior to distal introns

The above results suggest that RNA splicing does not prevent RNAi-directed heterochromatin assembly per se. To test whether this also applies to splicing events at the very beginning of a transcript that presumably occur immediately after the nascent transcript emerges, we generated strains in which the cox4+ intron was inserted 46 nt after the annotated transcription start site of the ade6+ gene (Fig. 2A). As before, we inserted the cox4+ intronic sequence in both orientations (targets 4 and 5) as well as a GFP fragment of similar length as a specificity control (target 6) and confirmed splicing by RT–PCR (Fig. 2B).

In contrast to target constructs 1 and 2 (Fig. 1), we observed significant differences in the amount of H3K9 methylation and ade6+ mRNA silencing between the spliced and nonspliced targets 4 and 5. Although H3K9 methylation was induced at both loci specifically by the cox4 siRNAs, H3K9me2 levels were inherently lower if the cox4+ target sequence underwent splicing (Fig. 2C, D). Consistent with this observation, ade6+ silencing was cox4 siRNA-specific and very robust without splicing of the cox4+ intronic sequence (targets 5 and 6) (Fig. 2E).

**Figure 1.** Transcription start site (TSS)-distal pre-mRNA splicing does not affect the stability of heterochromatin. (A) Schematic of the adh1+ promoter-driven uro4 hairpin construct. The double-stranded stem consists of complementary uro4+ ORF sequences (401–679, gray), while the loop encodes the cox4+ intron (pink) (Iida et al. 2008). Normalized 5′ ends of derived siRNAs are depicted below as reads per million (RPM). (B) Length histogram of generated siRNAs from A colored by their 5′ starting nucleotides. (C) Schematic representation of siRNA target constructs. The cox4+ intron sequence was inserted at position +700 nt of the ade6+ ORF in forward or reverse orientation (targets 1 and 2, respectively). A GFP fragment of the same size but without canonical splice sites was inserted at the same position [target 3]. (D) RT–PCR to assess intron removal was performed with primers flanking the exon–exon junction (mb2202 and mb167). (E) Chromatin immunoprecipitation (ChIP) experiments with an antibody recognizing H3K9me2. Fold enrichments were normalized to adh1+ and cendg and are shown relative to the respective leo1+/no siRNA samples. Error bars indicate SD. n = 3 independent biological replicates. Two-tailed Student’s t-test. (F) ade6+ mRNA levels were determined by quantitative RT–PCR. Values were normalized to act1+ mRNA and are shown relative to the respective leo1+/no siRNA samples. Error bars indicate SD. n = 3 independent biological replicates. Two-tailed Student’s t-test. (G) Normalized siRNAs mapping to the spliced (target 1; blue) and unspliced (target 2; red) targets in leo1+ [top] and leo1Δ [bottom] cells. Primary siRNAs produced from the cox4+ intron in the hairpin (shown in A) are shaded in pink. Secondary siRNAs are generated solely at the target locus. (H) Length histogram of target mapping siRNAs, similar to B. (I) Browser screen shot depicting siRNAs mapping to the ade6+ target locus.
However, ade6⁺ mRNA levels were reduced only twofold for the spliced target gene (target 4) [Fig. 2E], indicating that excision of the siRNA target sequence by an early splicing event obstructs heterochromatin formation. Despite the decreased silencing potential, RDRC-dependent secondary siRNAs were generated in a leo1Δ-dependent manner also at this target locus. Concomitantly, low levels of H3K9 methylation spanned the entire locus [Fig. 2D,F, Supplemental Figs. S1, S2]. These observations allow two alternative interpretations: (1) Rapid splicing of the target sequence decreases the overall silencing potential and therefore only mildly reduces target expression in all cells. (2) Alternatively, due to early splicing, the initiation of silencing becomes stochastic, and only a fraction of cells trigger the full response, while others do not initiate silencing at all. Notably, H3K9me2, siRNAs, and RNA expression were all measured at the population level, integrating over millions of cells. Thus, if 50% of cells expressing the spliced target remained in a euchromatic state while the other half became heterochromatic, maximal twofold repression can be expected from such a population average. Therefore, we decided to address this question at
single-cell resolution by growing cells on limiting adenine indicator plates, which leads to red colonies when ade6+ is fully repressed. Indeed, when single cells were seeded, ~50% formed red colonies, indicative of complete ade6+ repression [Fig. 2G, left]. This indicates that the kinetics of pre-mRNA processing is a critical parameter for heterochromatin formation. If the siRNA target is available only briefly, such as our spliced target 4, full assembly of the silencing machinery may occur in only a subset of cells, resulting in a bistable on/off state. Of note, once established, silencing is propagated through mitosis, albeit maintenance fidelity is lower than for targets without early splice events [Fig. 2G, right; Kowalik et al. 2015].

In sum, siRNA-binding sites close to the 5’ end of a transcript perform as well as more distal sites unless removed by RNA splicing. Although heterochromatin is established with low frequency when promoter-proximal introns are targeted, it is less stably maintained through mitosis [Fig. 2G]. These results strongly argue against RITS invading the double-stranded underlying DNA or base-pairing with the ssDNA exposed in an R loop upon transcription [Nakama et al. 2012], as, in both cases, splicing is not expected to affect heterochromatin formation.

**Highly transcribed genes resist siRNA-directed H3K9 methylation**

The above results strongly support the hypothesis that RITS is targeted to chromatin via base-pairing interactions with nascent transcripts. However, they still do not completely rule out siRNA–DNA base pairing [Moazed et al. 2006]. According to the latter model, trans-acting siRNAs are expected to assemble heterochromatin also in the absence of transcription at the target locus. Therefore, we investigated to what extent the deposition of H3K9 methylation is dictated by promoter strength of the target locus. For consistency, we chose the endogenous ade6+ gene as the siRNA target locus and first replaced its promoter with the strong but tunable nmt1 promoter [Fig. 3A]. When cells are grown on thiamine-containing medium, the nmt1 promoter is gradually repressed with increasing concentrations yet is not switched off completely [Fig. 3B, leo1+ cells; Forsburg 1993]. As a source of trans-acting primary ade6+ siRNAs, we used a previously described stably integrated synthetic ade6+ RNA hairpin construct that is driven by an adh1+ promoter [Kowalik et al. 2015].

When grown under repressed conditions with high concentrations of thiamine, ade6 siRNAs efficiently triggered H3K9 methylation specifically in leo1Δ cells [Fig. 3C]. As a consequence, ade6+ mRNA levels were reduced up to sevenfold in leo1Δ compared with leo1+ cells [Fig. 3B,C]. However, we observed a gradual reduction in silencing and H3K9 methylation efficiency with increasing expression of the target locus [Fig. 3B,C]. Consistent with this, leo1Δ cells expressing ade6+ driven by the nmt1 promoter formed red colonies on limiting adenine indicator plates supplemented with 20 µM thiamine, indicating full ade6+ repression [Fig. 3D]. We also observed attenuated repression with decreasing thiamine concentrations for the nmt1-driven ade6+ but not for ade6+ driven by its own promoter [Fig. 3D]. In contrast, leo1+ cells formed white colonies, suggesting that high transcriptional activity counteracts RNAi-directed heterochromatin assembly.

**Figure 3.** High transcriptional activity counteracts RNAi-directed heterochromatin assembly. (A) Scheme depicting the endogenous ade6+ gene [left] or the thiamine-repressible nmt1 promoter [right]. The siRNA target region is indicated by dashed lines. (B) ade6+ mRNA levels were determined by quantitative RT–PCR. Values were normalized to act1+ mRNA and are shown relative to leo1Δ cells grown in the presence of 20 µM thiamine. Error bars indicate SD. n = 3 independent biological replicates. (C) ChIP experiments with an antibody recognizing H3K9me2. Fold enrichments over clr4Δ cells are indicated. Error bars indicate SD. n = 3 independent biological replicates. (B,C) ChIP and quantitative RT–PCR experiments were performed with cells from the same culture. (D) Fivefold serial dilutions were spotted on adenine-limited PMG agar plates supplemented with thiamine at the concentrations indicated. Precultures were grown in YES medium, washed, and diluted in H2O before spotting.
demonstrating that repression was siRNA-specific and not caused by high concentrations of thiamine. \textit{nmt1-ade6} expression in wild-type cells is still sufficient even when reduced at high concentrations of thiamine [Fig. 3B].

These results reveal that high transcriptional activity at an siRNA target locus can counteract heterochromatin assembly. This may be due to increased nucleosome turnover, leading to the loss of K9 methylated H3 or to RITS/Clr4 displacement from chromatin. Both scenarios would affect the positive feedback loop and therefore prevent stable formation of heterochromatin. Intermediate to low promoter activity should thus be considered as an important criterion in the design of small RNA-directed chromatin silencing experiments.

Transcription activity above a minimal threshold is necessary for heterochromatin formation

Because the \textit{nmt1} expression system does not switch off completely, we aimed to abolish transcription of the endogenous \textit{ade6} gene entirely by deleting its promoter. However, we still observed substantial residual transcriptional activity (data not shown), preventing us from distinguishing between the DNA and nascent RNA targeting models at the endogenous \textit{ade6} locus. We reasoned that the residual \textit{ade6} activity emanated from neighboring gene activities. Therefore, we examined the \textit{S. pombe} genome for a region with no apparent sign of transcription to serve as a neutral landing site where transgenes would be less influenced by transcription occurring in the vicinity. Inspecting RNA sequencing (RNA-seq) and small RNA-seq data generated from wild-type \textit{S. pombe} cells, we found a 17-kb region on chromosome III with no discernible RNA production. Furthermore, H3K9 methylation was not significantly enriched in this area [Fig. 4A]. At that locus, we inserted different \textit{ade6} transgenes: the full-length \textit{ade6} gene with its own promoter and a terminator sequence, \textit{ade6} without the promoter but with the terminator, \textit{ade6} without the terminator but with the promoter, and the \textit{ade6} ORF alone [Fig. 4B].

![Figure 4](image-url)
To determine the absolute number of ade6+ mRNA molecules per cell produced from these transgenes, we used the droplet digital PCR (ddPCR) technology. This revealed that the full-length ade6+ transgene driven by its own promoter gives rise to ~17 RNA molecules per cell. For the transgene lacking the promoter, we counted only 1.8 RNA molecules per cell, while the transgene without transcription termination signals produced 3.6 ade6+ RNA molecules per cell. For the solitary ade6+ ORF [sole ORF] inserted at the same position, we never counted more than one molecule per cell [Fig. 4C]. Thus, as expected, in the absence of transcriptional regulatory elements, expression of ade6+ is very low or highly infrequent.

If RITS targeted DNA, trans-acting ade6 siRNAs would be expected to trigger H3K9 methylation at all four transgenes. We detected H3K9 methylation at the full-length ade6+ gene, at ade6+ without the promoter, and at ade6+ without the terminator [Fig. 4D–F]. However, we did not observe methylated H3K9 at the ade6+ solitary ORF transgene when less than one transcript per cell was present [Fig. 4G]. Together with the sensitivity to splicing [Fig. 2], these results demonstrate that the RITS complex cannot bind to DNA. We therefore conclude that RITS is targeted to chromatin exclusively via base-pairing interactions with nascent transcripts. The absence of any H3K9 methylation despite residual transcription activity at the solitary ade6+ ORF indicates that synthesis of a minimal number of nascent transcripts must be achieved to recruit sufficient amounts of chromatin-modifying enzymes.

Discussion

siRNA–DNA interactions do not initiate de novo assembly of heterochromatin

Originating from initial discoveries in yeast and plants, the role of RNAi-related pathways in epigenetic genome regulation has triggered much excitement. It provides an attractive mechanistic explanation for the largely elusive question of how chromatin-modifying enzymes find their targets. Current models state that the sequence information contained in the loaded small RNAs guides enzymatic activities to complementary targets via base-pairing interactions with nascent RNAs [Grewal and Ekwall 2015; Martienssen and Moazed 2015]. However, sequence-specific interactions involving siRNA–DNA base pairing have not been refuted. The following key observations from the data presented here rule out that targeting RITS to DNA is sufficient to initiate the formation of silent chromatin in S. pombe: [1] Transcription above a certain threshold is an absolute requirement for the formation of heterochromatin. Silencing does not initiate if less than one RNA molecule is present per cell [Fig. 4]. [2] Splicing of siRNA-binding sites close to the 5′ end of a transcript interferes with heterochromatin assembly [Fig. 2]. If siRNAs targeted RITS to DNA, heterochromatin stability between spliced and nonspliced targets would not be different. Therefore, we conclude that RITS is programmed by Dcr1-produced siRNAs to target specific chromosomal regions exclusively via siRNA nascent transcript base-pairing interactions.

Conservation of the nascent transcript model

There is accumulating evidence that the nascent transcript model applies to small RNA-directed chromatin silencing pathways also in other organisms. The strongest support comes from studies in Drosophila, where piRNAs in follicle cells originate almost 100% from the antisense strand of transposable elements. These trigger silencing only if the target harbors complementary piRNA sites in sense orientation. Because silencing of reporters and endogenous loci is strictly dependent on the orientation of the piRNA targets, direct DNA targeting as well as targeting of ssDNA exposed in R loops can be excluded also in flies [Sarot et al. 2004; Sienski et al. 2012; Post et al. 2014]. However, Argonaute engagement with the underlying DNA in other systems likewise awaits direct disproof. For example, RNA-directed DNA methylation (RdDM), a major small RNA-mediated epigenetic pathway in plants, requires a specialized transcriptional machinery that comprises the two plant-specific RNA polymerases: polymerase IV [Pol IV] and Pol V [Matzke and Mosher 2014]. Pol IV transcribes precursors that are processed by an RdRP [RDR2] and DCL3 [DICER-LIKE 3] to produce siRNAs that load onto AGO4 [Herr et al. 2005]. Pol V produces transcripts that are widely accepted to act as scaffolds for association of AGO4–siRNA complexes and subsequent chromatin modification [Wierzbicki et al. 2008]. However, although current models depict siRNA base-pairing with Pol V-generated scaffold RNAs, it is still elusive whether guide RNAs bind directly to DNA exposed by Pol V transcription or to nascent transcripts produced from it [Dalakouras and Wassenegger 2014; Matzke and Mosher 2014]. Consistent with our results in S. pombe, 24-nt secondary siRNAs induce DNA methylation in trans at unlinked target sites only if this sequence is transcribed by RNA Pol II. However, methylation can nevertheless occur without detectable transcription at this target in the presence of 21- to 24-nt hairpin-derived siRNAs. This suggests that synthesis of a nascent transcript at some target loci is not essential for RdDM and that methylation of nontranscribed target sequences in Arabidopsis may require multiple size classes of siRNA [You et al. 2013]. The nascent transcript model has also been put forward to explain small RNA-directed chromatin regulation in Caenorhabditis elegans [Grishok 2005; Guang et al. 2010; Burkhart et al. 2011; Buckle et al. 2012; Luteijn et al. 2012; Wedeles et al. 2013]. Whether siRNA–DNA base pairing is negligible in these processes has not been addressed. However, conservation of nascent transcript targeting in fission yeast and Drosophila suggests that this is an evolutionarily conserved mode of inducing chromatin modifications.

Epigenetic silencing is a digital process

Our efforts in triggering the formation of heterochromatin by trans-acting primary siRNAs in S. pombe highlight
a key feature of epigenetic gene regulation. Rather than inducing gradually decreasing mRNA levels, siRNAs initiate gene silencing in an “all or nothing” fashion in Paf1C mutant cells. Once established, the off state is stably propagated (Kowalik et al. 2015) unless the siRNA complementary sequence is situated in an early intron (Fig. 2). Similar digital silencing responses occur upon artificial tethering of various chromatin-modifying enzymes to a transcriptional reporter in mammalian cells (Buntu et al. 2016). This type of silencing is physiologically highly relevant, as exemplified by the regulation of the FLOWERING LOCUS C (FLC) in Arabidopsis. FLC expression is gradually repressed by prolonged cold exposure when measured at the level of a tissue. However, this seemingly “analog” silencing response is the net result of digital silencing events that occur in individual cells of that particular tissue over time. Thereby, a population of cells as a whole can respond quantitatively to an environmental change by simply switching individual cells from an on state to an off state (Berry and Dean 2015).

We infer that the siRNA-induced off switch is triggered as soon as the local concentration of chromatin-modifying activities and their residence time on the target are sufficiently high with respect to the target transcript itself. Only if this ratio is above a certain threshold will the epigenetic changes take effect and be maintained. Thus, future efforts should focus on quantifying such thresholds under physiological conditions for small RNA target interactions as well as for other epigenetic modifiers. This will not only further our understanding of epigenetic gene silencing but hopefully also enable us to generate meaningful predictions of putative targets for epigenetic regulation based on steady-state measurements.

Finally, we note that current state-of-the-art biochemical techniques interrogating cell populations are not well-suited for quantitative analysis of epigenetic gene regulation. As exemplified in this study, epigenetic silencing in individual cells may be easily missed if gene expression is assessed at the population level. Hence, adjusting current and developing new methods at single-cell resolution will be important for future mechanistic studies.

Materials and methods

Strains and plasmids

Fission yeast strains were grown at 30°C in YES medium. All strains were constructed following a PCR-based protocol (Bahler et al. 1998) or by standard mating and sporulation. Strains generated in this study are in Supplemental Table 2.

Silencing assay

To assess ade6+ expression, serial fivefold dilutions of the respective strains were plated on yeast extract plate or PMG plates supplemented with 5.65 mg/L adenine and 226 mg/L each leucine, ura1, histidine, and lysine.

RNA isolation and cDNA synthesis

RNA was isolated using Absolutely RNA miniprep kit (Agilent) as described in Emmerth et al. (2010). cDNA was synthesized using PrimeScript RT Master Mix (Takara).

Quantitative real-time PCR

Real-time PCR on cDNA samples and chromatin immunoprecipitation (ChIP) DNA was performed as described in Emmerth et al. (2010) using a Bio-Rad CFX96 real-time system using SsoAdvanced SYBR Green supermix (Bio-Rad). Primer sequences are in Supplemental Table 1.

Qualitative RT–PCR

PCR on cDNA was performed using a fast-cycling PCR kit (Qiagen). PCR products were analyzed by agarose gel electrophoresis. Primer sequences are in Supplemental Table 1.

ddPCR

Exponentially growing cells were harvested, and the cell number was determined with a hemocytometer (Thoma). Typically, 1.2 × 107 cells were used to isolate RNA using the MasterPure yeast RNA purification kit (Epigenetix). Two-hundred nanograms of total RNA isolated from mouse embryonic stem cells was added to the yeast samples in lysis buffer to estimate the recovery rate of RNA. cDNA was synthesized using the PrimeScript RT Master Mix (Takara). cDNA corresponding to 20 ng of RNA was used as template for ddPCR. The PCR reaction was prepared with QX200 ddPCR EvaGreen supermix (Bio-Rad) and Droplet generation oil for EvaGreen (Bio-Rad). Droplets were generated with a QX200 droplet generator (Bio-Rad). After amplification to the end point, droplets were quantitated by QX200 droplet reader (Bio-Rad).

ChIP

ChIP experiments were performed as described previously in Kowalik et al. (2015) with histone H3K9me2-specific mouse monoclonal antibody from Wako (clone no. MAB0307).

Small RNA-seq

Small RNA libraries were prepared as described previously (Kowalik et al. 2015). In brief, total RNA was isolated, and small RNAs (18–28 nt) were size-selected by PAGE. Libraries were prepared using the Illumina TruSeq small RNA preparation protocol according to manufacturer’s instructions. Libraries were sequenced on an Illumina HiSeq2500 instrument.

Small RNA-seq data analysis

Cutadapt (Cutadapt—a “adapter”–discard-untrimmed -m 18) was used to remove the 3’ adaptor form the raw reads. Reads <18 nt and untrimmed reads were removed. Trimmed reads were aligned to the S. pombe genome (ASM294 version 2.24) using Bowtie (Langmead et al. 2009). No mismatches were allowed, and, for multimapping reads, one random best hit was kept (-M 1 -v 0 –best –strata). For mapping the reads to the hairpin, similar settings were used due to the inverted uorel sequence in the hairpin stem (see Fig. 1A). To map the reads to the targets, only uniquely mapping reads were allowed (-m 1 -v 0 –best –strata). Mapping stats for all samples are in Supplemental Figure S3.
Accession numbers

Small RNA-seq data have been deposited at the NCBI Gene Expression Omnibus (GEO) database and are accessible through GEO series number GSE87672.

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