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Establishment of heterochromatin in domain-size-dependent bursts

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Methylation of histone H3K9 is a hallmark of epigenetic silencing in eukaryotes. Nucleosome modifications often rely on positive feedback where enzymes are recruited by modified nucleosomes. A combination of local and global feedbacks has been proposed to account for some dynamic properties of heterochromatin, but the range at which the global feedbacks operate and the exact mode of heterochromatin propagation are not known. We investigated these questions in fission yeast. Guided by mathematical modeling, we incrementally increased the size of the mating-type region and profiled heterochromatin establishment over time. We observed exponential decays in the proportion of cells with active reporters, with rates that decreased with domain size. Establishment periods varied from a few generations in wild type to >200 generations in the longest region examined, and highly correlated silencing of two reporters located outside the nucleation center was observed. On a chromatin level, this indicates that individual regions are silenced in sudden bursts. Mathematical modeling accounts for these bursts if heterochromatic nucleosomes facilitate a deacetylation or methylation reaction at long range, in a distance-independent manner. A likely effector of three-dimensional interactions is the evolutionarily conserved Swi6HPT, H3K9me reader, indicating the bursting behavior might be a general mode of heterochromatin propagation.

Histone-modifying enzymes covalently attach small polypeptides or chemical groups to nucleosomes, thereby controlling gene expression. Some modifications propagate from nucleation sites over large chromosomal distances. They can be maintained clonally upon cell division or overridden to allow for cell differentiation or adaptation to changing environments. Among the many histone modifications that define chromatin structure, the methylation of histone H3K9 maintains large chromosomal regions in a silent heterochromatic state.

Modeling has pointed to key parameters in chromatin formation and maintenance, for instance, the importance of cooperativity in the recruitment of modifying enzymes to chromatin or how various types of positive feedback would contribute differentially to the properties of chromatin (1–4). Nucleosome interconversion models developed with one organism, fission yeast, have been applied to distant species such as Arabidopsis, Drosophila, or mice, transposed to other modifications, for example, from H3K9 to H3K27 methylation (2, 5–7), and merged with other forms of modeling (8). In Arabidopsis, such models have uncovered fundamental aspects of vernalization by showing how epigenetic memory is digitally acquired and stored in cis at the FLC locus for long-term gene silencing and by anticipating the sequence of events taking place at that locus (2, 4, 6). In Drosophila, nucleosome-based models shed new light on the concerted action of PRC complexes (7). Recently, molecular models have been also complemented by an operational model of chromatin-mediated silencing, which supports the view that silencing is a stochastic event that occurs in an all-or-none fashion in individual cells (9).

A central element of nucleosome interconversion models is that they accommodate feedbacks arising through the direct or indirect recruitment of enzymes by modified nucleosomes. In nature, self-recruitment applies to many histone-modifying enzymes and enzymatic complexes (10–12), some of which might be able to modify nucleosomes brought into proximity by chromatin folding (13–15), while others would uniquely modify immediate neighbors. Hence, among the fundamental questions that quantitative modeling can address are the impacts of short- and long-range nucleosomal interactions occurring in the nucleus on chromatin dynamics. This is deeply connected to the overall expectation that heterochromatin stability depends on the size of the region in question.

To explore these questions, we used here the mating-type region of the fission yeast Schizosaccharomyces pombe. In this system, heterochromatin formation entails methylation of histone H3K9 by a single Su(var)39-family enzyme, Clr4, in combination with nucleosomal deacetylation by various histone deacetylases (HDACs), but no DNA methylation. The region contains the silent mat2-P and mat3-M loci used in mating-type switching (16) and other elements (Fig. 1), including a central cenH element with homology to centromeric and subtelomeric repeats (17, 18).

Significance

How repressive heterochromatic states propagate along chromosomes and are subsequently maintained for many cell divisions remain important unanswered questions in biology. Combining mathematical modeling and single-cell measurements, we find that heterochromatin does not propagate in a purely linear manner as often assumed. Rather, sudden transitions can affect large domains globally at once. We suggest this is due to long-range interactions that bring distant nucleosomes close to each other to facilitate their modification. This supports the notion that the compartmentalization of chromatin components within the nucleus and the folding of chromatin into loops dynamically control heterochromatin propagation in three dimensions by bringing nucleosomes and their modifiers into close proximity.
Fig. 1. Domain-size-dependent establishment of heterochromatin. (A) Mating-type region of *S. pombe* showing two reporters used to monitor the establishment of heterochromatic silencing. YFP at the *Kint2* site within the nucleation center *cenH* and *mCherry* at the (EcoRV) site. The heterochromatic domain is delimited by the IR-L and IR-R boundary elements and contains two Atf1 binding sites, s1 and s2. The region was extended by inserting, at the indicated *Kint3* site, DNA originating from the *leu1*+ chromosomal region, respectively, a 4.5-kb, two distinct 5.9-kb, and a 7.8-kb fragment (see also SI Appendix, Fig. S1) generating mating-type regions of, respectively, 27.5 kb, 29 kb (SH4), and 31 kb. (B and C) The *clr4* gene was reintroduced into *clr4*Δ cells with extended mating-type regions (B) or original length (C) through genetic crosses, and the establishment of (EcoRV)::*mCherry* silencing was monitored in cultures maintained in exponential growth for 19 d, taking one time point per day. Overlays of brightfield and red fluorescence micrographs are presented for days 2, 4, 8, 12, and 17 as indicated. Histograms of cell fluorescence intensities are shown for the same cultures, each compiling >15,000 cells. Individual channels including YFP are shown in SI Appendix, Figs. S2–S4. (D) Two examples of clonal populations derived from the strain with the 31-kb mating-type region at day 17 and propagated for approximately 40 generations. Each micrograph shows a field 86.5 × 66 µm.
Time course experiments have found that the nucleation of heterochromatin occurs at cenH, stochastically and at a high rate (3). It is driven by RNA interference (RNAi) (19, 20) as at centromeres (21–23). Outward expansion relies on positive feedback by several chromodomain proteins that bind H3K9me. Thus, in Ctr4 chromodomain mutants catalytically proficient but incapable of binding H3K9me (20, 24), or in mutants lacking the HP1 homolog Swi6 (19, 24, 25), H3K9me heterochromatin is restricted to cenH. At centromeres or when heterochromatin is induced by artificially tethering Ctr4, the Ctr4 chromodomain is also critical to the epigenetic heterochromatin state (23, 26, 27). For Swi6, the positive feedback might be exerted by interacting HDAC complexes (28–30) that foster H3K9 methylation (25, 31), and the Swi6 paralog Chp2 is likely to function in a similar way (28, 29, 32–35). These H3K9me/chromodomain feedbacks are induced by artificially tethering Clr4, the Clr4 chromodomain protein, to the DNA, less likely to introduce nucleotide biases or changes in DNA sequence (SI Appendix, Methods). At centromeres or when heterochromatin is induced by artificially tethering Clr4, the Clr4 chromodomain protein, to the DNA, less likely to introduce nucleotide biases or changes in DNA sequence (SI Appendix, Methods).

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The short mating-type region produced exclusively dark-staining colonies starting at early time points, but longer regions showed a delay correlated with the slow silencing of \((\text{EcoRV}):\text{mCherry}\) in them, and distinct patterns of staining were observed (Fig. 2D). The 27.5-kb region produced more heavily streaked colonies than the longer regions, reflecting more frequent successful establishment events during colony growth. For the longest region examined, the 31-kb region, colonies with wild-type iodine staining coexisted side by side with very light colonies for the entire length of the experiment. In this assay, uniformly brown staining shows stability of the heterochromatic state at the timescale of colony formation. The brown colony phenotype was also stable upon growth in culture and replating, and cells in the brown colonies had silenced both \(\text{leu}1^+\) and \((\text{EcoRV}):\text{mCherry}\), while cells in the light colonies showed unstable \(\text{leu}1^+\) and \((\text{EcoRV}):\text{mCherry}\) expression (SI Appendix, Fig. S5). These phenotypes reflect a long-lasting epigenetic inheritance of alternate chromatin states over the mating-type region and denote the stability of the heterochromatic state once it is established. They also show that \((\text{EcoRV}):\text{mCherry}\), the mating-type cassettes, and \(\text{leu}1^+\) all tend to be euchromatic or heterochromatic in individual cells in a correlated manner.

**Temporally Correlated Silencing of Genes outside \(\text{cenH}\).** To assess the relative timing of silencing of reporters at a finer scale, we performed a \(\text{clr}4^+\) reintroduction experiment in a strain where the \(\text{YFP}\) gene, instead of being at \(\text{cenH}\), was at a location between \(\text{cenH}\) and \((\text{EcoRV}):\text{mCherry}\), approximately 5 kb from \((\text{EcoRV}):\text{mCherry}\) (Fig. 3). In that setup, \(\text{YFP}\) and \(\text{mCherry}\) were silenced in a highly correlated fashion. Throughout the experiment, with few exceptions, cells either expressed or had silenced both reporters (Fig. 3). Hence, despite being 5 kb apart, the two reporters were silenced at the same time, or nearly. This was confirmed by time-lapse microscopy of dividing cells during the establishment period (examples shown in Movies S1–S6). In cases where a short delay separated silencing of the two reporters, either \(\text{mCherry}\) (Movies S3 and S4) or \(\text{YFP}\) (Movies S5 and S6) could be silenced first. The fact that \(\text{mCherry}\) could be silenced before \(\text{YFP}\) is interesting, as it argues against a linear spreading of heterochromatin from \(\text{cenH}\). It does not exclude that linear spreading might occur from elements located between the \(\text{YFP}\) and \(\text{mCherry}\) genes, such as \(\text{Atf1}\) binding sites (Fig. 1). However, even in this case, spreading would be catalyzed at a distance by \(\text{cenH}\), as, in the absence of \(\text{cenH}\), heterochromatic silencing at \(\text{EcoRV}\) occurs once in 2,000 cell divisions, on average, whereas, here, in the presence of \(\text{cenH}\), it occurred two orders of magnitude more efficiently. The possibility of linear spreading from \(\text{Atf1}\) binding sites facilitated by physical contacts with \(\text{cenH}\) through looping was investigated as part of our modeling.

**Modeling Heterochromatin Formation in the Mating-Type Region.** To model heterochromatin establishment, the wild-type and three expanded mating-type regions were represented by 153, 182, 191, and 203 nucleosomes, respectively, each containing a special 31-nucleosome region to mimic the ability of the \(\text{cenH}\) element to recruit the \(\text{Clr4}\) methyltransferase through RNAi. In addition to explaining the observed heterochromatin establishment kinetics of the differently sized mating-type regions, we demanded that the model reproduce bistability of a 76-nucleosome system lacking the special region, to mimic the phenomenon observed with \(\Delta K\) mutations that show a stochastic switch from euchromatin to heterochromatin and vice versa every 2,000 cell generations, approximately (41). Therefore, a small system with 76 nucleosomes lacking the special region was used for simulations to model the observed bistability. As in ref. 1 and depicted in SI Appendix, Fig. S1, the
model considers three nucleosome states, with S representing a silent nucleosome methylated at histone H3K9, U representing an unmodified nucleosome, and A representing an acetylated nucleosome.

The model considers two types of reactions, recruited conversions, where an enzyme is recruited by a nucleosome and modifies another nucleosome in the system, and direct conversions that do not result from nucleosome-based recruitment processes. Direct conversions encompass nucleosome state changes due to nucleosome turnover, to the binding of enzymes to DNA or transcription factors, or to chromatin contacts from outside the mating-type region, among others. The reactions were simulated using an event-based Gillespie algorithm in which each discrete event with an assigned rate parameter represents one modification event (presented in SI Appendix). The algorithm was executed by first selecting an event, either a direct conversion or a recruitment. When a direct conversion attempt was selected, one nucleosome in the system was chosen randomly, and its state was changed accordingly. In case of a recruited conversion attempt being chosen, a recruiting nucleosome was chosen first. If this nucleosome was either in an A state or an S state, a second nucleosome in the system was chosen, and, if this second nucleosome was in a state that permitted a conversion, the state of this second nucleosome was changed according to the action of the recruited enzyme.

In ref. 3, the necessity of a combination of nonlocal (global) and local interactions was highlighted, and several possible models were analyzed and discussed. Here, the consideration of different system sizes restrained the possible model versions further. Fig. 4 shows simulation results of a model version where the S-mediated feedback on A- to U-state conversions is global, and the remaining three feedbacks act only locally. Other models, including looping models, are discussed farther down and presented in SI Appendix, Figs. S7–S19.

**A Combination of Three Local and One Distance-Independent Silencing Recruitment Reactions Can Explain the Profiles of de Novo Heterochromatin Formation.** For the model presented in Fig. 4, the rates of A-mediated conversions of U to A nucleosomes and S to U nucleosomes were assigned a fixed rate of 100 conversion attempts per nucleosome per cell generation, whereas the remaining two recruitment reactions were varied as shown. In addition to the spontaneous conversions indicated by the connecting straight black arrows (each having a rate of one conversion attempt per nucleosome per generation), the special cenH region was assigned a higher rate of direct U- to S-nucleosome conversions (300 conversion attempts per nucleosome per generation). To ensure that the ability of the model to reproduce the observed establishment kinetics is not restricted...
Fig. 4. A combination of three local and one distance-independent silencing recruitment reaction can explain the domain-size-dependent kinetics of de novo heterochromatin establishment. (A) The model considers four types of recruited conversions based on positive feedback, where enzymes are recruited by the same histone modifications that they deposit. Only the S-mediated A- to U-nucleosome conversions are global, whereas the remaining three reactions act locally on the nearest-neighbor nucleosomes. Spontaneous conversions between states with a higher rate of U-state to S-state nucleosomes within the cenH region are also included in the model. (B) Scanning revealed that this model can recapitulate the observed bistability of the ∆K mutants (indicated by cyan, where the system remains in an active and a silent state for more than 100 consecutive generations), and blue squares, where the system remains in an active and a silent state for more than 500 consecutive generations) and observed the kinetics of the systems with different sizes (23, 27.5, 29, and 31 kb). Dark blue squares highlight that, in addition to the ΔK mutants being bistable, the average lifetime of the 29-kb system in the expressed system is at least twice as long as the lifetime of the 23-kb system. (C) Time–space plots for the four differently sized systems show the region state of one representative cell from 0 to 100 cell generations with parameter values indicated by a gray arrow. The plots illustrate a near-system-size-independent first stochastic switch to heterochromatin at cenH and a system-size-dependent time-delayed second stochastic switch from euchromatin to heterochromatin in the remaining region. (D) Profiles obtained in simulations with the same parameter values as in C, compiling 10,000 cells. The proportion of “ON” cells is shown over time, illustrating that larger system sizes result in substantially slower establishment of heterochromatin. SI Appendix, Fig. S6 depicts how profiles were obtained.

to the relatively low direct conversion rates, we also performed simulations with higher direct conversion rates and were still able to obtain good fits (SI Appendix, Fig. S8). In order to examine whether the model can recapitulate the experimentally observed kinetics, the S-mediated U-to-S and A-to-U nucleosome conversion rate constants were varied as shown in Fig. 4B. A total of 400 pairs of rate constants were tested, 1,000 simulations were performed for each pair, and the operation was repeated for each size of the mating-type region. Bistability is illustrated by cyan and blue squares, where simulations of the 76-nucleosome system are started with either only A- or S-state nucleosomes, and the system remains in each state for more than 100 (cyan) or 500 (blue) generations, respectively. From experimental kinetic measurements performed in five independent experiments plotted in Fig. 5, we could estimate, for each system, the average lifetime (b) of (EcoRV)::mCherry expression following reintroduction of the clr4+ gene (Figs. 2A and S5). The average lifetime (b) for the simulated systems was determined by fitting exponential functions to the simulation results. The measure $Gap = b_{29Kb} - 2 * b_{23Kb}$ (with $b_{23Kb}$ and $b_{29Kb}$ being the average lifetime for the 23-kb system and 29-kb system) was defined to quantify the system size dependency of the model. Dark blue squares indicate simulation results where the 76-nucleosome system was bistable (blue) and where $Gap > 0$. We considered a fit to be good if the $b$ value theoretically determined for a given system within a given parameter regime did not deviate more than 50% from the experimentally determined $b$ value. Since the rate constants were varied in relatively large steps of 10, we reasoned that a reasonably tolerant range around the experimentally measured $b$ values would be suitable to spot a parameter region with a potentially good fit of the data that could be later refined. Thus, we demanded the average lifetime (b) of 1,000 simulations representing 1,000 reporter states in individual cells to be between 2.35 and 7.0 generations for the 23-kb system, between 8.5 and 25.6 generations for the 27.5-kb system, between 24.5 and 73.6 generations for the 29-kb system, and between 59.5 and 178.5 generations for the 31-kb system. For simulations, we considered the system to be switched to heterochromatin when half of the nucleosomes within the (EcoRV)::mCherry reporter became S nucleosomes. In Fig. 4B, correct timing of the 23-kb system is shown by green circles. Red circles indicate correct kinetics of both the 23- and 27.5-kb systems, and silver circles...
indicate correct timing of the 29-kb system in addition to the smaller systems. Golden dots with a smaller black circle in the middle indicate correct timing of all systems. Simulation profiles and the experimental data with fits are shown in Fig. 5. Each simulation profile consists of 200 data points where the proportion of “ON” cells among 10,000 cells was calculated after each generation before the simulation of DNA replication (see SI Appendix, Fig. S6 for a more detailed explanation). In these profiles, the switch-off dynamics of the outer (EcoRV) mCherry reporter results are plotted together with the switch-off dynamics of their respective systems are plotted in gray to highlight the differential switching of cenH and the outer region of the systems. Overall, the model can correctly recapitulate the switching dynamics of the 23-, 27.5-, 29-, and 31-kb systems.

All in all, the model can recapitulate bistability of the丧失nucleosome system and the observed establishment dynamics of all system sizes. Slight deviations of the lifetime between simulations and experiments might be explained by simplifications in the model that could influence the establishment dynamics in the experimental setup such as DNA composition or small temperature fluctuations. Some variation also occurred in experimentally determined average lifetimes, even though this did not change the overall observation that strains with larger mating-type regions showed drastically slower establishment dynamics of heterochromatin. We also observed that the model can recapitulate the observed establishment kinetics with higher noise rates (SI Appendix, Fig. S8).

Moreover, in simulations, we could see that small changes of the global rate had substantial effects on the speed of heterochromatin formation. Interestingly, this effect was stronger in larger system, whereas the wild-type 23-kb system was more robust to these changes (SI Appendix, Fig. S8).

![Fig. 5](https://doi.org/10.1073/pnas.2022887118)
Variations on the Model Show the Need for a Global S-Mediated Feedback to Account for the Observed Kinetics. Fig. 6 shows simulation results of the best fits of other model versions with one global recruitment reaction and three local recruitment moves. Fig. 6A shows the simulated dynamics of the model from Fig. 4 with an altered global reaction, where nucleosomes to be modified are now chosen with a probability inversely proportional to their distance to the recruiting nucleosome. The simulations illustrate that, in addition to an extensive weakening of the model’s ability to reproduce the bistable behavior of the ΔK mutant, the model fails to reproduce the heterochromatin formation dynamics in larger systems (Fig. 6B). Fig. 6C shows the simulation results of a model version where the S-mediated feedback on the U-to-S reaction is global and the other recruited conversions act locally with the shown parameter constants. The simulations show that this model version can also reproduce the observed heterochromatin establishment dynamics but fails when the operating mode of the global reaction is changed to a recruited conversion probability attempt decaying as a power law (1/x), with distance x between recruiting and substrate nucleosome (Fig. 6D), similar to the previously examined S-mediated global A-to-U conversions (Fig. 6A and B). Fig. 6E shows simulation results of a model version where the A-mediated S-to-U reaction is global and all other reactions are local with the indicated parameter constants. Although this model shows similar system-size-dependent heterochromatin establishment dynamics, the bistability of the ΔK mutant model system is rather fragile (see SI Appendix, Fig. S11), and the dynamics of larger systems is faster than observed in experiments. Interestingly, this model shows system-size-dependent heterochromatin establishment dynamics even with a distance-dependent global reaction in power law (1/x) operating mode (Fig. 6F). Nevertheless, a large shoulder is visible during the first generations in simulations of larger system sizes, which was not observed experimentally. Moreover, the smaller 76-nucleosome system does not show any bistability for the model configuration shown (see SI Appendix, Fig. S11). Lastly, Fig. 6G shows a model version where the global feedback is mediated by A nucleosomes that stimulate U- to A-state transitions. This model shows hardly any system-size-dependent heterochromatin establishment dynamics, and the dynamics remain unchanged for a global reaction with power law contact probability (Fig. 6H).

In SI Appendix, Figs. S7–S19, a total of seven model motifs (four motifs with each one global and three local feedbacks, and three motifs with each two global and two local feedbacks) have been investigated. Each motif was examined in two different versions, with or without internal looping between cenH and two Atf1 binding sites (s1 and s2 in Fig. 1). Each standard model version, without specific internal looping, was simulated with four different modes of the global feedback reactions (three distance-dependent modes and one distance-independent mode), and the model with an interaction between cenH and Atf1 was simulated with a distance-dependent global feedback reaction (1/x contact probability). Additionally, an Atf1 looping-model version with only local feedbacks was simulated. Distance-dependent feedback reactions followed a power law-distributed contact probability between substrate and recruiting nucleosome as a function of distance between the two nucleosomes. In total, 36 model versions were examined (described in SI Appendix, SI Text and Figs. S7–S19). Of the 36 models, 7 were able to fit the experimentally observed silencing kinetics and fulfill the bistability requirement of the smaller 76-nucleosome system with the same set of parameter values (SI Appendix, Fig. S19). Six of the “working” models have in common that they all have one or two global feedback reactions that are distance independent or very weakly distance dependent (power law exponent ≤ 0.5).

Only one model version, with an S-mediated A-to-U and an A-mediated S-to-U feedback reaction, could recapitulate the experimental observations with distance-dependent global feedback reactions following a 1/x power law contact probability. Although intriguing, we consider this model version to be problematic, because biological evidence is lacking for a global A-mediated S-to-U feedback reaction, in contrast to the feedbacks used by other models that are well documented experimentally, as discussed farther down. Furthermore, any global feedback toward active nucleosomes (either A(S-to-U) or A(U-to-A)) might lead to repeated attacks by A nucleosomes surrounding the mating-type region and destabilize heterochromatin. Similarly a global S(U-to-S) feedback would allow a silenced mating-type region to destabilize the active surroundings of the mating-type region by converting U nucleosomes to S nucleosomes. A global S(A-to-U) feedback would potentially have less destabilizing effects on active states, since it only slightly increases the proportion of U nucleosomes and thus has an effect similar to DNA replication or higher noise. SI Appendix, Fig. S19 presents an overview and discussion of all working models.

Mimicking Chromosome Looping between cenH and Atf1 Binding Sites near mat3-M Does Not Produce Comprehensively Effective Models. To test how a loop domain might contribute to heterochromatin establishment, we simulated looping between internal Atf1 binding sites (42, 43) (Fig. 1) and cenH. The interaction between cenH and the Atf1 binding sites was simulated as an S(U-to-S) recruited conversion. Whenever a substrate nucleosome was chosen at either position 101 or 42 nucleosomes upstream of the last nucleosome in the region (Atf1 binding sites) and a recruiting nucleosome was chosen within cenH, an attempt to convert the substrate nucleosome to an S nucleosome was made. Conversely, if a substrate nucleosome was chosen inside cenH combined with a recruiting nucleosome at either of the two Atf1 binding sites, an attempt to convert the substrate nucleosome to an S nucleosome was carried out (see SI Appendix for a more detailed description). The modeling results suggest that, although internal looping helps with the timing of the observed system-size-dependent silencing dynamics in some model versions, it always fails to simultaneously meet our primary modeling requirement of conferring bistability to the small 76-nucleosome system and system-size-dependent silencing. All in all, based on our model simulations, we find that, although we cannot rule out the possibility of an interaction between cenH and Atf1 binding sites, looping does not change the need for distance-independent or weakly distance-dependent global feedback reactions.

Discussion

Overall, kinetic measurements performed at the single-cell level and stochastic modeling show how the mating-type region undergoes two stochastic switches during heterochromatin formation where the cenH region always switches to a heterochromatic state before the rest of the region. Moreover, the second stochastic switch occurs at a size-dependent rate. At the molecular level, this means that heterochromatin does not spread gradually from the nucleating region at cenH to the remaining region; rather, heterochromatin propagates in an all-or-none fashion, where whole regions collapse suddenly. This mode of propagation differs radically from the linear propagation proposed in other cases (44).

Our modeling shows that a previously suggested three-state model with a distance-dependent global S(A-to-U) feedback (3) falls short of explaining the size dependence observed here, unless the global reaction is changed to a distance-independent feedback. In general, models with one global feedback and three local feedbacks can only explain the newly observed silencing dynamics and the bistability of ΔK mutants when 1) the global simulation results of a model version where the A-mediated S-to-U reaction is global and all other reactions are local with the indicated parameter constants. Although this model shows similar system-size-dependent heterochromatin establishment dynamics, the bistability of the ΔK mutant model system is rather fragile (see SI Appendix, Fig. S11), and the dynamics of larger systems is faster than observed in experiments. Interestingly, this model shows system-size-dependent heterochromatin establishment dynamics even with a distance-dependent global reaction in power law (1/x) operating mode (Fig. 6F). Nevertheless, a large shoulder is visible during the first generations in simulations of larger system sizes, which was not observed experimentally. Moreover, the smaller 76-nucleosome system does not show any bistability for the model configuration shown (see SI Appendix, Fig. S11). Lastly, Fig. 6G shows a model version where the global feedback is mediated by A nucleosomes that stimulate U- to A-state transitions. This model shows hardly any system-size-dependent heterochromatin establishment dynamics, and the dynamics remain unchanged for a global reaction with power law contact probability (Fig. 6H).

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Discussion

Overall, kinetic measurements performed at the single-cell level and stochastic modeling show how the mating-type region undergoes two stochastic switches during heterochromatin formation where the cenH region always switches to a heterochromatic state before the rest of the region. Moreover, the second stochastic switch occurs at a size-dependent rate. At the molecular level, this means that heterochromatin does not spread gradually from the nucleating region at cenH to the remaining region; rather, heterochromatin propagates in an all-or-none fashion, where whole regions collapse suddenly. This mode of propagation differs radically from the linear propagation proposed in other cases (44).

Our modeling shows that a previously suggested three-state model with a distance-dependent global S(A-to-U) feedback (3) falls short of explaining the size dependence observed here, unless the global reaction is changed to a distance-independent feedback. In general, models with one global feedback and three local feedbacks can only explain the newly observed silencing dynamics and the bistability of ΔK mutants when 1) the global
Modes of heterochromatin propagation suggested by experiments and modeling combined. Positive feedbacks exerted by heterochromatic nucleosomes during heterochromatin formation are represented. A global positive feedback exerted by S nucleosomes on A-to-U deacetylation might be imposed by phase separation of Swi6-associated chromatin with bound HDACs. In contrast, the local positive feedback exerted by S nucleosomes on U-to-S methylation reactions could be due to guided catalysis by nucleosome-bound Clr4. This particular feedback combination was suggested by mathematical modeling and literature to account for the kinetics of heterochromatin formation over the fission yeast mating-type region measured here. During establishment, a first stochastic switch at the nucleation center cenH leads to the fast build-up of a patch of heterochromatin from which heterochromatin expands to the entire mating-type region in a second stochastic switch with size-dependent delays.

reaction is independent of linear distance between substrate and modifying nucleosome and 2) it is toward the silenced state. Three-state models with distance-dependent feedback reactions only work with two global reactions, one toward silenced states and the other toward active states. However, these types of models then require additional mechanisms for isolating the studied region from its surroundings. We discuss these models farther down in light of the positive feedbacks known to operate in the mating-type region.

A key aspect captured by modeling is the role played by the cenH element. Many experiments combined have determined that RNAi acting at cenH facilitates heterochromatin formation over the entire mating-type region (3, 19, 41, 45, 46), but the mechanism has remained obscure. cenH also potentiates the effect of silencing elements at ectopic locations (19, 47, 48). Providing mechanistic insight, modeling shows how the build-up of an H3K9me nucleosome reservoir at cenH could drive the conversion of a larger region to the heterochromatic state via a global feedback by Clr4, as these structures would differ from the dinucleosomes used in vitro to assay the catalytic activity of Clr4 (54). Thus, the possibility of a global feedback by heterochromatic nucleosomes on the methylation reaction will also have to be considered in the future. Moreover, in vivo Clr4 is part of the multisubunit complex ClrC containing, for example, a ubiquitin ligase whose ubiquitylation of histone H3K14 is required for the methylation of H3K9 by Clr4 (61). Potential feedbacks through the whole complex remain to be studied. RNAi reaching outside cenH might also operate over long distances to recruit Clr4 (24, 62). Compared to a global feedback on A-to-U transitions which might be easily counteracted by histone acetyl transferases in euchromatin, the effects of a global feedback on U-to-S transitions could be more difficult to control genome wide. In all cases, the compartmentalization of chromatin components by phase separation and the organization of chromosomes into loop domains would be important factors in limiting the range of global reactions.

In the future, feedbacks can be investigated genetically using chromodomain mutants (e.g., ref. 53) to alter the affinity of H3K9me of the Clr4 chromodomain affecting the S-mediated (U-to-S) feedback, or the Swi6 chromodomain affecting the S-mediated (A-to-U) feedback. In the other direction, the A-mediated (U-to-A) feedback could possibly be perturbed by targeting a positive feedback loop acting on the Mst2 acetyl transferase (63) and the A-mediated (S to U) feedback by targeting the bromodomain protein Bdf2, an interactor of Epe1 (64). Moreover, in an independent approach, varying system size in the absence of the cenH region would allow us to determine the relative effects of cenH with size, and the relative increase in the contribution of global recruitment when considering larger systems.
Materials and Methods

Strain Constructions and Growth Conditions. Standard procedures were used to propagate S. pombe cells and to construct new strains by genetic crosses (65) or by transformation of existing strains with recombinant DNA (66) as detailed in SI Appendix. Strain genotypes are shown in SI Appendix, Table S1, and the oligonucleotides used for strain construction are shown in Table S2. Yeast extract with supplements (YES) was used as rich medium; minimal sporulation agar (MSA) (with 2 g/L arginine as nitrogen source) supplemented with 100 mg/L uracil, 100 mg/L adenine, and 200 mg/L leucine) was used for crosses; MSA (with 2 g/L glutamate as nitrogen source) supplemented with 200 mg/L leucine, when indicated, was used to plate cell cultures during establishment experiments prior to iodine staining; and EMM2 supplemented with 200 mg/L leucine was used as growth medium for the heterochromatin establishment experiments. Cultures were propagated at 30 °C.

Reintroduction of clr4+ Gene into clr4Δ Cells. The S. pombe strains PG3713 or PG3716 were used to reintroduce clr4+ into clr4Δ strains AE14, AE28, SH4, SH5, and PG3873. Mating mixes were prepared on MSA plates supplemented with leucine, uracil, and adenine and incubated for 48 h at 30 °C. To obtain random spores, mating mixes were scraped and resuspended in 1.5 mL of 1% glusulase (Roche) in H2O. The suspensions were incubated overnight at 30 °C and vortexed for 5 min, and random spores were separated by centrifugation. They were washed twice in sterile H2O and resuspended in EMM2 supplemented with leucine for selective germnation and growth. Cultures were set up in 1 mL of medium in 10-mL plastic culture tubes on a rotating wheel or in glass cultures tubes on a floor shaker, at 30 °C, and they were maintained in exponential phase through frequent dilutions for the length of the experiment. At each time point, cells were counted and imaged by fluorescence microscopy. In total, five establishment experiments were conducted, the first with a 23-kb (three technical replicates) and a 31-kb region (two technical replicates); the second with a 31-kb region (three technical replicates); the third with a 23- (two technical replicates) and a 27.5-kb region (three technical replicates); the fourth with a 23- (two technical replicates) and a 27.5-kb region (two technical replicates); the fifth with a 31-kb region (two technical replicates) and a 27.5-kb region (three technical replicates). Cells were imaged with a Nikon Ti Eclipse epifluorescence microscope equipped with a 100× objective, an Andor Luca camera, and a temperature-controlled chamber which was set to 30 °C for the duration of the experiment. Images were taken using the NIS Elements Advanced Research software in the bright-field, YFP, and mCherry channels. Image acquisition times were 800 ms for YFP and 600 ms for mCherry. Images were converted to tiffs and analyzed in MATLAB using custom image-processing software. Cell boundaries were identified automatically using bright-field images, and fluorescence intensity was measured in the middle of the cells where the nucleus is expected in most cases. A threshold for each YFP and mCherry was set to separate ON and OFF cells. Alternatively, double Gaussian functions were fitted to histograms to calculate the proportions of OFF and ON cells. The two methods produced essentially identical results.

Data Acquisition with Xcyto-10 Quantitative Cell Imager. As an alternative to the Nikon Ti Eclipse microscope, fluorescence images were acquired and processed with an Xcyto-10 Quantitative Cell Imager (ChemoMetec) to generate the data presented in Fig. 1 and SI Appendix, Fig. 5. The LED535 lamp and filter 582 to 636 were used for mCherry, LED488 lamp and filter 513 to 555 for YFP, with exposure times of 1 s in both cases. The acquired images were visually inspected and analyzed with the XcytoView 1.0.109.0 software. The YFP and mCherry data were fitted using the curve_fit function of the Scipy.optimize Python package which uses the method of nonlinear least squares to fit the function $f(x) = a + e^{-x}$ to the data points in order to determine the b values of all systems corresponding to the respective parameter configuration. The 95% confidence bands were calculated with the uncertainties Python package.

Fluorescence Microscopy and Image Analyses. Cells were imaged with a Nikon Ti Eclipse epifluorescence microscope equipped with a 100× objective, an Andor Luca camera, and a temperature-controlled chamber which was set to 30 °C for the duration of the experiment. Images were taken using the NIS Elements Advanced Research software in the bright-field, YFP, and mCherry channels. Image acquisition times were 800 ms for YFP and 600 ms for mCherry. Images were converted to tiffs and analyzed in MATLAB using custom image-processing software. Cell boundaries were identified automatically using bright-field images, and fluorescence intensity was measured in the middle of the cells where the nucleus is expected in most cases. A threshold for each YFP and mCherry was set to separate ON and OFF cells. Alternatively, double Gaussian functions were fitted to histograms to calculate the proportions of OFF and ON cells. The two methods produced essentially identical results.

Curve Fitting and Confidence Bands of the Fit. The simulation results and the data shown in Figs. 4–6 were fitted with the curve fit function of the Scipy.optimize Python package which uses the method of nonlinear least squares to fit the function $f(x) = a + e^{-x}$ to the data points in order to determine the b values of all systems corresponding to the respective parameter configuration. The 95% confidence bands were calculated with the uncertainties Python package.


Data Availability. All data generated or analyzed during this study are included in this article and SI Appendix.

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