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High-resolution Xist binding maps reveal 2-step spreading during X-inactivation

Matthew D. Simon^{#1,2,*}, Stefan F. Pinter^{#1,3}, Rui Fang^{#2}, Kavitha Sarma^{1,3}, Michael Rutenberg-Schoenberg², Sarah K. Bowman¹, Barry A. Kesner^{1,3}, Verena K. Maier^{1,3}, Robert E. Kingston^{1,*}, and Jeannie T. Lee^{1,3,*}

¹ Department of Molecular Biology, Massachusetts General Hospital, and Department of Genetics, Harvard Medical School, Boston, MA 02114

² Dept. of Molecular Biophysics and Biochemistry, and Chemical Biology Institute, Yale University, West Haven, CT, 06516

³ Howard Hughes Medical Institute

[#] These authors contributed equally to this work.

Abstract

The Xist long noncoding RNA (lncRNA) is essential for X-chromosome inactivation (XCI), the process by which mammals compensate for unequal numbers of sex chromosomes¹⁻³. During XCI, Xist coats the future inactive X (Xi)⁴ and recruits Polycomb Repressive Complex 2 (PRC2) to the X-inactivation center (*Xic*)⁵. How Xist spreads silencing on a 150 Mb scale is unclear. Here we generate high-resolution maps of Xist binding on the X chromosome across a developmental time course using CHART-seq. In female cells undergoing XCI *de novo*, Xist follows a two-step mechanism, initially targeting gene-rich islands before spreading to intervening gene-poor domains. Xist is depleted from genes that escape XCI but may concentrate near escapee boundaries. Xist binding is linearly proportional to PRC2 density and H3 lysine 27 trimethylation (H3K27me3), suggesting co-migration of Xist and PRC2. Interestingly, when the Xi is acutely stripped off Xist in post-XCI cells, Xist recovers quickly within both gene-rich and -poor domains on a time-scale of hours instead of days, suggesting a previously primed Xi chromatin state. We conclude that Xist spreading takes distinct stage-specific forms: During initial establishment, Xist follows a two-step mechanism, but during maintenance, Xist spreads rapidly to both gene-rich and -poor regions.

Xist RNA is a prototype lncRNA with global epigenetic function^{1-3,6}. The initiation of XCI depends on Xist⁷ and loading of the Xist-PRC2 complex at a nucleation site within the Xic^8 . Thereafter, Xist RNA forms a "cloud" over the X-chromosome, signaling the initiation of

AUTHOR CONTRIBUTIONS

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^{*} Correspondence should be addressed to M.D.S. (matthew.simon@yale.edu), R.E.K. (kingston@molbio.mgh.harvard.edu), and J.T.L. (lee@molbio.mgh.harvard.edu).

R.E.K., J.T.L., M.D.S. and S.F.P. designed the experiments; S.F.P., K.S. and R.F. performed cell culture; R.F. and M.D.S. established CHART conditions and performed CHART; S.K.B. prepared CHART libraries; K.S. performed LNA knockoffs; S.F.P. and V.K.M. performed RNA-seq; S.F.P. performed bioinformatic analyses with M.D.S., M.R.S., R.F. and B.A.K.; M.D.S., S.F.P., R.F., J.T.L., and R.E.K. interpreted the results; J.T.L, M.D.S, R.E.K, S.F.P. and R.F. wrote the manuscript.

DATA DEPOSITION

CHART-seq and RNA-seq datasets have been deposited in GEO (accession number GSE48649).

The authors declare no competing financial interests.

chromosome-wide silencing⁴. Concurrently, PRC2 accumulates broadly along the X-chromosome⁹. Although Xist RNA coats the Xi at cytological resolution, whether and where Xist binds at molecular resolution remains unknown. In one model, Xist targets PRC2 to the *Xic*, but outward spreading of PRC2 does not involve Xist. Alternatively, both nucleation and spread involve Xist, in which case Xist and PRC2 would co-migrate at a molecular scale.

We mapped genome-wide binding locations of Xist RNA by performing CHART-seq (Capture Hybridization Analysis of RNA Targets with deep sequencing), a technique to localize lncRNAs on chromatin using complementary oligonucleotides to enrich for DNA targets¹⁰ (Extended Data Fig. 1a). We designed a cocktail of 11 complementary oligonucleotides for Xist CHART based on conserved or functional Xist domains^{7,11-13} and RNase H mapping for accessibility (Extended Data Fig. 1b,c; Extended Data Table 1). Allele-specific CHART-seq was performed at four developmental stages (Extended Data Fig. 1d): Before XCI in undifferentiated female mouse embryonic stem (ES) cells (d0; <1% of nuclei XCI positive, showing an Xist cloud or H3K27me3 focus); early-XCI (d3; <10% positive); mid-XCI (d7; 40-50% positive); and post-XCI (mouse embryonic fibroblast (MEF) clone, >95% positive). About 600,000 sequence polymorphisms between the M. musculus (mus) and M. castaneus (cas) X-chromosomes enabled ~35% allele-specific mapping to Xi and Xa (active X), respectively⁹. Disabling the mus *Tsix* allele in the female ES cells ensured that the mus X will be Xi¹⁴. We validated results by comparing two independent capture oligo submixtures and an alternative 40-oligo cocktail targeting across the length of Xist (Extended Data Fig. 2a-e, Extended Data Table 1). Regions with significant Xist enrichment localized almost exclusively to Xi (>99% X-linked, p<0.001; >90% Xi-skewed, p<0.05, Extended Data Fig. 2f,g,i). On autosomes, binding was minimal and of questionable significance. Enriched segments were not complementary to captureoligos and showed minimal enrichment on Xa of d0, d3, d7 and MEF cells. Enrichment was not observed using sense control oligos (Extended Data Fig. 2a,c). These experiments excluded artifactual enrichment, validating Xist CHART-seq specificity.

The dominant CHART peak lay in Xist exon 1 and was specific to Xi (Fig. 1a). A developmental time course demonstrated a progression in Xist density, with enriched segments increasing from <0.1% coverage of the X in pre-XCI cells, to ~20% in early- and mid-XCI, and ~54% in post-XCI cells (Fig. 1b,c, Extended Data 2h). Thus, Xist RNA not only forms a cytological cloud but also binds broad swaths of the Xi at molecular resolution. Xist could either spread uniformly along the Xi or target specific regions. Intriguingly, in cells undergoing XCI (d3, d7), Xist preferentially targeted multi-megabase domains (Fig. 1c). In post-XCI MEFs, Xist spread into intervening gene-poor regions throughout the Xi. The d3 and d7 patterns were more similar to each other than to MEF patterns (Fig. 1d, e, Extended Data Fig. 3a). Furthermore, comparative analysis identified MEF-specific domains not found during XCI (Fig. 1e). Despite heterogeneity in the onset of XCI in the ex vivo ES differentiation system, the highly similar d3 and d7 distributions show that Xist targets generich domains first. Extension of ES differentiation to d10 showed statistically significant filling in of gene-poor domains (Extended Data Fig. 3b,c), though not to the extent observed in somatic cells (MEFs). We infer that full spreading across Xi may only be achieved later in development, once differentiation into somatic lineages occurs. Thus, during de novo XCI in the embryo, Xist likely follows a two-step pattern of spreading, first targeting gene-rich clusters (hereafter, "early" domains) and eventually spreading to intervening gene-poor regions ("late" domains). Throughout the process, gene bodies of escapees^{15,16} were depleted of Xist, but occasionally demonstrated Xist enrichment in flanking regions (Fig. 1f, Extended Data Fig. 4), suggesting boundaries that sequester Xist and prevent spreading into neighboring privileged escapee loci.

We investigated what might target Xist to early domains by comparisons with various chromatin features (see Methods)^{9,17-22}. Interestingly, Xist is more likely to target genes in regions of active chromatin in ES cells. Allele-specific RNA-seq analysis demonstrated Xist's preference for genes that are active (e.g., on the Xa and in d0 and d7 cells) and showed skewed expression in d7 ES cells and in MEFs (Fig. 1c, g, Extended Data Fig. 5a; $p < 2.2 \times 10^{-16}$; Pearson's r=0.46). Furthermore, there were positive correlations with DNase I hypersensitive sites (Pearson's r=0.43), SINEs (r=0.55), and early replicating regions (r=0.45; Fig. 1h, Extended Data Fig. 5b). Strong anti-correlations were observed with LTRs (r=-0.39), LINE1 (r=-0.54), and lamin-associated domains (LADs, r=-0.48)²³. Xist partitioning did not correlate with cytogenetic banding on the X-chromosome (Fig. 1h)^{19,24}. LINE1s have been proposed as spreading elements²⁵, but repetitive reads from Xist CHART-seq aligning to LINE1 were not enriched over input (Extended Data Fig. 5c). Xist's localization showed modest positive correlation with Xic looping contacts inferred from HiC (high-throughput chromosome conformation capture)¹⁸ via an anchor within the Xist locus (Fig. 1h, Extended Data Fig. 5b). Together, these data support a role for open chromatin in guiding Xist, with Xist coming into contact with gene-rich regions (early domains) first, and spreading secondarily to more distal gene-poor inter-regions (late domains).

Given co-nucleation of Xist and PRC2 at the *Xic*⁸, we asked whether Xist continues to associate with PRC2 during spreading. Comparison of Xist, EZH2, and H3K27me3 enrichment revealed strikingly similar chromosome profiles across time (Fig. 2a, Extended Data Fig. 6a-c). By contrast, PRC2 and H3K27me3 densities on Xa did not correlate with Xist, nor did those on Chr13, a representative autosome (Extended Data Fig. 6a). Consistent with the idea that Xist directs PRC2 localization onto Xi⁵, Xist densities demonstrated an extensive linear relationship with EZH2 and its product H3K27me3 across the X in mid-XCI but not pre-XCI cells (Fig. 2b). Correlation with the H3K4me3 control (active mark) was poor. In MEFs, densities of H3K27me3 and Xist remained highly correlated, while reduced densities of PRC2 were observed during maintenance. Interestingly, Xist densities were not necessarily greater at previously defined "PRC2 strong sites"⁹ (Extended Data Fig. 6d,e); instead, Xist densities showed a general correlation with Xi-specific PRC2 enrichment (Extended Data Figs. 6d,h; 5b). This supports the idea that strong sites are Xist-independent (as indeed they are present in d0 cells⁹) and indicate that Xist and PRC2 co-migrate to new regions within the early domains on the Xi.

We then asked if localization mechanisms were inherent to Xist RNA or chromatin context. In perturbation experiments, we stripped away Xist RNA and observed recovery on the Xi of MEFs at 1h, 3h, and 8h. Locked nucleic acids (LNA) directed against Xist RNA's Repeat C prevented nucleation and therefore spreading¹³. RNA fluorescence in situ hybridization (FISH) showed that LNA-4978 did not overtly perturb Xist at 1h, but led to full Xist displacement by 3h, with Xist reassociation at 8h (Fig. 3a). As reassociation requires newly synthesized Xist rather than relocalization of displaced Xist¹³, reassociation must depend on outward spreading of new RNA from the *Xic*, just as during XCI establishment.

Interestingly, however, CHART-seq revealed a pattern not evident cytologically by RNA FISH. At 1h, when Xist was still visualized on Xi (Fig. 3a), CHART-seq demonstrated a relative loss in late domains (Fig. 3b-d), suggesting that Xist binds more weakly to gene-poor than gene-rich regions and consistent with Xist's banded pattern on the metaphase Xi observed cytologically²⁴. At 3h, Xist was strongly depleted from both regions. At 8h, partial recovery was evident in both regions. However, unlike spreading during *de novo* XCI (d3, d7), spreading of Xist during the somatic maintenance phase (MEF) did not follow a two-step process, as Xist reassociation in early and late domains occurred simultaneously (Fig. 3b-d). Therefore, spreading during *de novo* XCI was restricted to early domains and occurred on a time-scale of days in the *ex vivo* system; in contrast, re-covery and

respreading in post-XCI cells occurred more generally in both domains and on a time-scale of hours. This quantitative difference is significant, with accumulation in late domains appearing on the same time-scale as early domains during the recovery period after Xist knockoff (Fig. 3e-g; Extended Data Figs. 7, 8). Similar results were observed using an independent LNA, LNA-C1, targeted to a different sequence in the Repeat C region and in multiple replicates (Fig 3a, Extended Data Figs. 2d, 3a, 7, 8). Despite LNA-C1 being fasteracting¹³ (Fig. 3a), LNA-C1 and LNA-4978 treatment resulted in remarkably similar Xist knockoff and recovery on Xi.

Taken together, these data argue for distinct mechanisms of Xist spreading during establishment (de novo XCI) in early embryonic cells, when spreading occurs in a two-step fashion (early to late domains), and during maintenance in somatic cells, when Xist spreads more generally into both early and late domains (Fig. 3h). The Xi may retain epigenetic memory of Xist²⁶, enabling more efficient spreading during maintenance. As Xist mostly dissociates from the Xi during mitosis⁴, epigenetic memory could facilitate Xist's resynthesis and re-spreading in G1, and duplication of Xist patterns after DNA replication. Indeed, Xist's continued action is essential for maintenance of XCI²⁷. In summary, we have illuminated the mechanism by which Xist spreads on a 150 Mb scale. Comparing localization dynamics of Xist relative to other lncRNAs (Hasiculeyman and Rinn, personal communication) and 3D conformations²⁸ may prove highly informative for understanding general mechanisms of RNA-directed chromatin change.

METHODS SUMMARY

Cells were grown as previously described⁹. RNA was isolated and sequenced using a modified adaptor ligation protocol, and Xist localization determined by CHART¹⁰ with several modifications as detailed in the extended Methods. LNAs that target Xist were previously described¹³.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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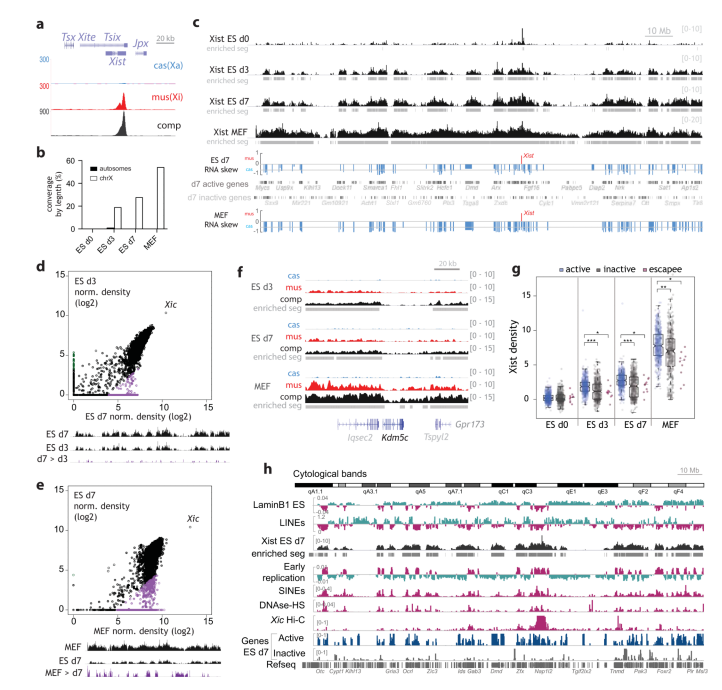


Figure 1. CHART-seq reveals a two-step mechanism of Xist spreading during de novo XCI

a, Xist RNA is enriched on Xi. Normalized read densities displayed in mus, cas, and composite (comp) tracks. **b**, Coverage of enriched segments on chrX and autosomes. **c**, Xist coverage at indicated timepoints relative to gene silencing. Enriched segments shown beneath in gray. Brackets, y-axis scale. Xist peaks at d0 have less amplitude and density, but reflect d3 and d7 patterns, and are Xi-enriched (Extended Data Fig. 2f), consistent with initial Xist spreading to local regions, suggesting initial differentiation in a subfraction of cells. RNAseq of d7 and MEF shown below. Skewed allelic expression consistent with Xi-silencing (value -0.5 = 3-fold expression difference between Xi and Xa). **d-e**, Xist CHART signals (40 kb bins) from d7 correlate with d3 (**d**) and MEF (**e**)(see Extended Data Fig. 3).

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Regions showing >10-fold differences after normalization are colored purple and displayed on the X (lower panels). **f**, Depletion of Xist at a representative escapee. **g**, Xist preferentially targets genes in active chromatin (H3K4me3-marked on d7). Xist densities shown for gene bodies of active (n=532), inactive (n=475), and escapee genes (n=10). Medians are indicated. Individual data points overlaid on boxplot; error bars, 1.5-fold interquartile range. *p<0.05, **p<1×10⁻⁸, ***p<2.2×10⁻¹⁶, Mann-Whitney U tests. **h**, Xist RNA distribution from d7 cells relative to chromatin features. Simon et al.

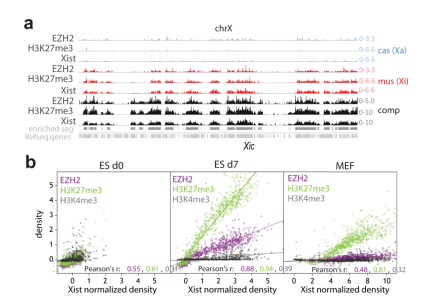


Figure 2. Co-spreading of Xist RNA and PRC2

a, Normalized read densities of Xist, EZH2, and H3K27me3 on chrX in d7 cells. **b**, Xist densities (200 kb bins) correlated with EZH2, H3K27me3 and H3K4me3 signals at different stages of XCI. Pearson's r displayed. EZH2/H3K27me3 R^2 values: 0.3/0.37 for d0, 0.77/0.88 for d7, and 0.23/0.66 for MEFs, respectively. H3K4me3 R^2 values: < 0.15 across all samples.

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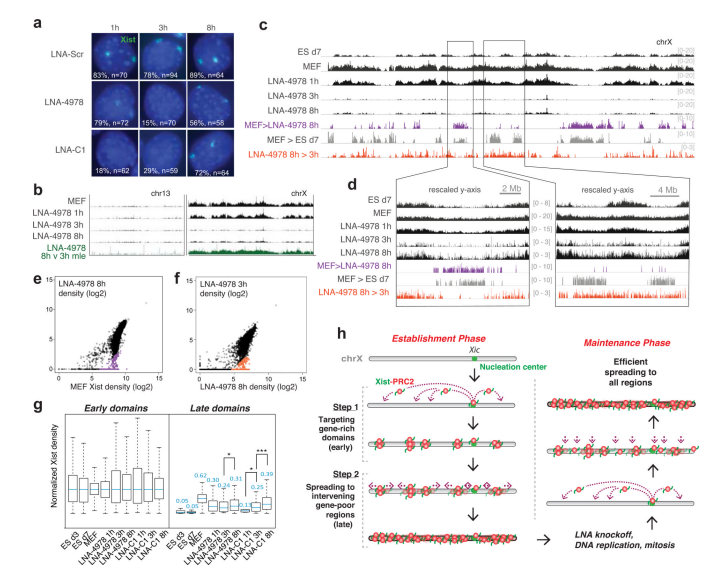


Figure 3. Xist knockoff uncovers a distinct spreading method during the maintenance phase a, RNA FISH shows depletion and recovery of Xist RNA (green) in MEF cells after Xist knockoff. %nuclei with Xist clouds and sample size (n) shown. Scr, scrambled LNA. b, Chromosome-wide recovery of Xist after LNA-4978 knockoff on chrX and chr13. Regions of recovery comparing 8h over 3h LNA-4978 were determined using a maximum likelihood enrichment estimate²⁹. c,d, Xist knockoff and recovery across chrX. Colored regions show >10-fold median-normalized differences between samples. (c) Entire chrX (d) zoom of one region with more late domain recovery (right) than the other (left). e,f, Xist CHART signals (40 kb bins) from LNA-4978 8h correlated with MEF (e); LNA-4978 3h correlated against LNA-4978 8h (f). Regions showing >10-fold differences after normalization are colored as shown in panel d. g, Xist recovery in indicated samples, with 40-kb binned Xist densities normalized to median levels of early domains of each sample, to determine how early and late domains recover from knockoff compared to during de novo XCI. Normalized median values for each sample indicated above box. *, p < 0.05; ***, $p < 10^{-8}$, Wilcoxon test, as in Extended Data Figs. 7c, 3c. h, Model: distinct methods of Xist spreading during establishment and maintenance.