A LINE1-Nucleolin Partnership Regulates Early Development and ESC Identity

Graphical Abstract

Highlights

- LINE1 RNA is abundant and nuclear in mouse ESCs and pre-implantation embryos
- LINE1 knockdown inhibits ESC self-renewal and induces transition to a 2C state
- LINE1 RNA recruits Nucleolin/Kap1 to repress Dux and activate rRNA synthesis
- In embryos, LINE1 inhibition causes persistence of the 2C program and impairs ZGA

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In Brief
RNA transcribed from LINE1 elements acts as a nuclear scaffold to direct gene expression programs essential for ESC self-renewal and pre-implantation embryo development.
A LINE1-Nucleolin Partnership Regulates Early Development and ESC Identity

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SUMMARY

Transposable elements represent nearly half of mammalian genomes and are generally described as parasites, or “junk DNA.” The LINE1 retrotransposon is the most abundant class and is thought to be deleterious for cells, yet it is paradoxically highly expressed during early development. Here, we report that LINE1 plays essential roles in mouse embryonic stem cells (ESCs) and pre-implantation embryos. In ESCs, LINE1 acts as a nuclear RNA scaffold that recruits Nucleolin and Kap1/Trim28 to repress Dux, the master activator of a transcriptional program specific to the 2-cell embryo. In parallel, LINE1 RNA mediates binding of Nucleolin and Kap1 to rDNA, promoting rRNA synthesis and ESC self-renewal. In embryos, LINE1 RNA is required for Dux silencing, synthesis of rRNA, and exit from the 2-cell stage. The results reveal an essential partnership between LINE1 RNA, Nucleolin, Kap1, and peri-nucleolar chromatin in the regulation of transcription, developmental potency, and ESC self-renewal.

INTRODUCTION

Only about 1.5% of mammalian genomes are comprised of single-copy protein-coding sequences, whereas approximately half of their DNA derives from transposable elements (TEs). Despite their abundance, the roles and regulation of TEs have been understudied, in large part because of the difficulty in mapping repetitive sequences to the genome. Nevertheless, TEs are now accepted as key players in genome evolution by rewiring gene regulatory networks, including in the human genome (Bourque, 2009).

The retrotransposon long interspersed element 1 (LINE1) makes up the largest proportion of TE-derived sequences, and is the only class of autonomous TEs still active in human (Magiorkinis et al., 2015). LINE1-induced mutations have been linked to a growing number of diseases, including hematopoietic and neurological disorders as well as several types of cancer (reviewed in Burns [2017]). For this reason, LINE1 is generally thought to be silenced in differentiated cell types to avoid uncontrolled mutagenesis. However, the view of TEs such as LINE1 as strictly detrimental to cells may be too simplistic. LINE1 is expressed in normal neural progenitor cells, where it has been proposed to promote neuronal diversity (Muotri et al., 2005). LINE1 is also expressed in the pre-implantation embryo (Fadloun et al., 2013) and in the fetal germline (Ohno et al., 2013; Percharde et al., 2017b). A TE of a different class, mouse endogenous retrovirus type L (MERVL), is also expressed in cleavage-stage embryos, where it drives the expression of many transcripts specific to zygotic genome activation (ZGA) and totipotency (Kigami et al., 2003; Macfarlan et al., 2012; Svoboda et al., 2004). Importantly, the rate of LINE1 retrotransposition in embryos and germ cells in vivo is low given the high levels of LINE1 RNA expression (Kano et al., 2009; Richardson et al., 2017; Newkirk et al., 2017). These observations raise the possibility that LINE1 RNA has as yet undefined cellular roles, independent of retrotransposition.

We set out to test the hypothesis that LINE1 plays essential functions in mouse pluripotent cells. Our data point to a model whereby LINE1 and its chromatin partners are essential to orchestrate developmental progression during pre-implantation and for the self-renewal of embryonic stem cells (ESCs).

RESULTS

LINE1 RNA Is Nuclear Localized and Promotes ESC Self-Renewal

We first set out to investigate the expression and localization of LINE1 in mouse ESCs. LINE1 RNA is detected at high levels
in a punctate pattern in the nuclei of ESCs, but not in the cytoplasm (Figure 1A), the opposite localization pattern of the open reading frame (ORF) 1 protein (Orf1p) encoded by LINE1 (Figure S1A). LINE1 RNA is associated with euchromatin and generally excluded from heterochromatic foci (Figure 1A). These results are in agreement with the general localization of Cct-T repeat RNA, which includes LINE1 RNA (Hall et al., 2014). A similar nuclear localization of LINE1 RNA is detected in mouse 2-cell embryos and blastocysts (Figure 1B) (Fadloun et al., 2013). These results raised the possibility that LINE1 RNA plays a role in transcriptional regulation in ESCs.

Next, we developed a LINE1 RNA knockdown (KD) strategy using antisense oligos (ASOs) (Figure 1C). LINE1 ASOs lead to a significant reduction in nuclear LINE1 fluorescence in situ hybridization (FISH) signal, unlike control reverse complement (RC) ASO-treated ESCs (Figures 1D, 1E, and S1B). We validated these results using two small interfering RNAs (siRNAs) as an independent KD method (Figures 1C and S1C). The lower level of KD using RNAi is to be expected given the greater ability of ASOs...
to KD nuclear RNAs (Lennox and Behlke, 2016). Surprisingly, KD of LINE1 results in a dramatic decrease in ESC self-renewal (Figure 1F), a result validated by LINE1 RNAs (Figure S1D). In agreement, LINE1 KD ESCs exhibit a drastically reduced cell expansion rate and an altered cell-cycle profile, with a significant decrease in the proportion of cells in S phase and an increase in cells in G2/M (Figures S1E and S1F). This is accompanied by only a modest increase in cell death (to ~1% of the total population) (Figure S1G) and no changes to the overall levels of Oct4 or Nanog proteins (Figure S1H). Taken together, these data indicate that LINE RNA is required for the efficient propagation of ESCs.

**LINE1 Represses the 2C Transcriptional Program in ESCs**

To analyze the transcriptional impact of LINE1 RNA KD, we performed RNA sequencing (RNA-seq), which revealed that LINE1 KD ESCs are reproducibly distinct from controls (Figures 2A and S2A). Genes with a LINE1 element situated within or nearby show no evidence of downregulation, arguing for a direct effect of KD of LINE1 RNA itself (Figure S2B). Interestingly, LINE1 KD induces a significant upregulation of 414 transcripts (log2-fold change > 0.7, false discovery rate [FDR] < 0.05), with only overall mild downregulation of single-copy genes (Figure 2B). Upregulated genes do not include markers of the three germ layers (Figures S2C and S2D), confirming that LINE1 KD does not induce precocious differentiation. Instead, there is a striking upregulation of genes transiently expressed at the 2-cell (2C) stage upon LINE1 KD (Figures 2C, 2D, and S2E). The 2-cell stage encompasses the switch from maternal control to zygotic genome activation (ZGA) and is associated with a sharp, transient upregulation of many genes, which are collectively called “2C genes” or the “2C program.” Several 2C genes contain promoters originally derived from the TE MERVL, which is also sharply induced at this stage (reviewed in Schoorlemmer et al. [2014]). MERVL and 2C genes are rapidly repressed after the 2-cell stage, and in most ESCs (Macfarlan et al., 2011; Wu et al., 2016). 2C genes upregulated upon LINE1 KD include well-known markers, such as Zscan4, Dubl1, Gm4340, Tcstv1/3, and Ztp352 (Figures 2E and 2F; Table S1), along with the 2C-specific transposon MERVL (Figures 2G–2I). 2C gene upregulation was additionally confirmed using LINE1 RNAi (Figure S2F). Inhibition of LINE1 retrotransposition using antiretroviral drugs (Jones et al., 2008) does not phenocopy LINE1 RNA KD, indicating that its role in ESCs is independent of retrotransposition (Figures S2G–S2I). Moreover, LINE1 KD does not induce MERVL/2C expression in mouse embryonic fibroblasts (Figure S2J). We examined whether LINE1 KD induces conversion of ESCs to a 2C-like fate, using an ESC line in which endogenous 2C-like cells are marked by GFP (Ishiuchi et al., 2015; Macfarlan et al., 2012) (Figure S2K). Depletion of LINE1 significantly increases the percentage of 2C-like cells that display the expected features: loss of chromocenters and lack of Oct4 protein (Figures 2J and 2K) or Nanog protein (data not shown). These results indicate that LINE1 acts to repress MERVL and the 2C transcriptional program in ESCs.

**LINE1 Represses Dux, a Master Activator of the 2C Program**

Next, we explored how LINE1 might repress the 2C state in ESCs. LINE1 KD induces a similar phenotype whether ESCs are grown in serum/LIF or 2i/LIF conditions (Ying et al., 2008) (Figures S3A and S3B). The expression of known repressors of the 2C state is not altered in LINE1 KD ESCs (Figure S3C). Interestingly, we found that the transcription factor Dux is significantly upregulated upon LINE1 KD (Figure 3A; Table S1). Recently, Dux was shown to bind directly to many 2C gene promoters and to be necessary for 2C gene upregulation in ESCs and for pre-implantation development (De Iaco et al., 2017; Hendrickson et al., 2017; Whiddon et al., 2017). Moreover, CRISPR/Cas9-mediated deletion of Dux in zygotes impairs pre-implantation development, suggesting a potentially critical role in early development and ZGA (De Iaco et al., 2017). Analysis of RNA-seq data revealed that Dux target genes are among the most highly activated genes upon LINE1 KD (Figure 3B). Moreover, the repressive chromatin mark H3K9me2 is reduced at Dux and its downstream targets in LINE1 KD ESCs (Figures S3D–S3F). Next, we performed control or LINE1 KD using ASOs, with or without simultaneous KD of Dux using siRNAs (Figures 3C and S3G). RNA-seq

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**Figure 2. LINE1 KD Causes Upregulation of 2C Genes and MERVL**

(A) PCA plot for all genes across all samples, showing that LINE1 KD ESCs have distinct gene expression profiles and are separated from controls along PC1. (B) MA plot showing log2-fold changes in the expression of each gene following LINE1 KD. Horizontal red or blue lines indicate FDR < 0.05 and log2 fold change (FC) of > 0.7 or < –0.7, respectively. Select upregulated 2-cell (2C) genes are labeled in black. (C) Heatmap showing expression changes of 142 2C genes as defined in Macfarlan et al. (2012), upon LINE1 KD. (D) Gene set enrichment analysis (GSEA) for 2C genes like in (C) showing a preferential upregulation of nearly all genes upon LINE1 KD. (E) Browser RNA-seq screenshots of 2C genes Zscan4d and Dub1 in RC or LINE1 ASO samples. (F) qRT-PCR validation of 2C gene upregulation following LINE1 KD with ASOs. Data are mean ± SEM, n = 3 biological replicates. (G) Distance analysis performed on the indicated sets of genes, calculating of log2 absolute distance in base pairs (bp) to the nearest MERVL element. 2C/ASO intersect, S2 2C genes from Macfarlan et al. (2012) also significantly upregulated with LINE1 KD; ASO upregulated, all significantly upregulated genes upon LINE1 KD. *p < 0.01, two-sided Wilcoxon rank-sum test, calculated between 2C/ASO intersect and all expressed genes. (H) MA plot showing log2-fold changes in repeat expression following LINE1 KD. Upregulated MERVL repeats and downregulated rRNA repeats are indicated. (I) qRT-PCR validation of MERVL expression following LINE1 KD. Data are mean ± SEM, n = 3 biological replicates. (J) Percentage of 2C-like cells and representative micrographs in 2C-GFP reporter ESCs 48 hr after nucleofection with ASOs. Data are mean ± SEM of 2 independent experiments. (K) Immunofluorescence analysis of LINE1 KD-induced 2C-like cells. Graph depicts the percentage of GFP+ cells that have the expected features (loss of chromocenters and Oct4 protein). Scale bar, 10 μm; n, number of cells.

See also Figure S2 and Table S1.
revealed that *Dux* depletion significantly reduces the upregulation of *Dux* targets upon LINE1 KD (Figure S3H). Moreover, hierarchical clustering of *Dux* targets revealed that *Dux* KD rescues the effect of LINE1 KD (Figure 3D). Similarly, MERVL is no longer upregulated upon simultaneous KD of LINE1 and *Dux* (Figures 3E–3G and S3I). These data reveal that the upregulation of the 2C-program in LINE1 KD ESCs is *Dux* dependent, raising the question of whether LINE1 directly or indirectly acts to restrict *Dux* expression in ESCs (see below).
LINE1 Promotes ES Self-Renewal Independently of Its Role in Repressing Dux

We subsequently investigated whether the induction of Dux is related to the decrease in self-renewal of LINE1 KD ESCs. We found that Dux KD does not rescue the self-renewal deficit observed upon LINE1 KD (Figure 4A), indicating that LINE1 plays additional Dux-independent roles in ESCs. Transcripts upregulated upon LINE1 KD include p53 targets (Figure S4A). However,
LINE1 KD in p53⁻/⁻ ESCs (Sabapathy et al., 1997) both activates 2C gene expression (Figure S4B) and significantly decreases ES self-renewal (Figure S4C), similar to wild-type ESCs. P53 activation is therefore not a primary cause of the self-renewal deficit of LINE1 KD ESCs.

While downregulation of single-copy genes is mild in LINE1 KD ESCs (Figure 2B), functional annotations related to ribosomal biogenesis and translation are significantly enriched among downregulated genes (Figures 4B and S4D; Table S1). Moreover, ribosomal protein genes are reduced in expression upon...
LINE1 KD (Figures S4E and S4F) and remain downregulated upon simultaneous Dux depletion (Figure 4C; Table S2). Interestingly, ribosomal RNA (rRNA) is significantly downregulated upon LINE1 KD (Figure 2H). In agreement, total RNA levels per cell are reduced upon LINE1 KD (Figure 4D), along with nascent transcription, rRNA synthesis and nascent translation (Figures 4E–4H). Direct chemical inhibition of rRNA synthesis (Haddach et al., 2012) leads to a similarly sharp reduction in the self-renewal of ESCs (Figures S4G–S4I). These results suggest that LINE1 promotes high levels of rRNA synthesis typical of and required by rapidly growing pluripotent cells (Guzman-Ayala et al., 2015). The self-renewal deficit of LINE1 KD ESCs is likely due to a combination of the reduction in nascent transcription, including rRNA, and the cell-cycle defects (Figures S1E and S1F).

**LINE1 RNA Binds Dux and rDNA Loci in ESCs**

We sought to reconcile the seemingly unrelated roles of LINE1 RNA in Dux repression and rRNA synthesis. The high abundance of nuclear LINE1 RNA (Figure 1A), as well as previous reports of its close association with chromatin (Hall et al., 2014) suggest that it may act similar to nuclear long non-coding RNAs (lncRNAs) to regulate gene expression. We therefore performed LINE1 chromatin immunoprecipitation by RNA purification (ChIRP) (Chu et al., 2011). ChIRP probes designed against the length of L1s can successfully capture LINE1 RNA, but not unrelated RNAs (Figure SS). To assess the specificity of LINE1 RNA ChIRP, we performed parallel Malat1 ChIRP and verified its specific association with the Malat1 and Neat1 loci (Figure S5B) (Engreitz et al., 2014). LINE1 RNA is robustly detected at LINE1 DNA (Figure SSC), as might be expected from the previously reported close association of LINE1 RNA with chromosome domains from which it is transcribed (Hall et al., 2014).

Interestingly, we found that LINE1 is significantly enriched at both Dux and rDNA loci compared to Malat1 RNA, but not at control regions (Figures 5A and SSB). The LINE1 ChIRP signal is RNA dependent, as RNase treatment largely prevents the recovery of LINE1 RNA-bound sites (Figure 5A). The repeated nature of LINE1 and the likelihood that unmapped LINE1 insertions exist in the genome may confound the source of the LINE1 ChIRP signal. Nevertheless, these data indicate that LINE1 RNA interacts with the Dux and rDNA loci, and raise the possibility that it cooperates with factors that regulate chromatin activity at these sites.

**Nucleolin Depletion Recapitulates LINE1 KD**

Studies in the retrotansposition field have uncovered a number of proteins that interact with LINE1 RNA. One protein that caught our attention is Nucleolin, a well-known rDNA and rRNA binding protein required for rRNA synthesis and processing (Ginisty et al., 1998). Nucleolin was recovered in a search for factors that interact with mouse LINE1 RNA, and this interaction appears to be conserved in human (Moldovan and Moran, 2015; Peddigari et al., 2013). Intriguingly, Nucleolin was also reported to bind to and repress the human DUX4 genomic repeat, D4Z4, in HeLa cells (Gabellini et al., 2002). Using RNA immunoprecipitation (RIP)-qPCR, we verified that Nucleolin strongly associates, directly or indirectly, with LINE1 RNA in ESCs, at levels similar to that of pre-rRNA (Figure 5B). The Nucleolin RIP-qPCR signal is DNA-independent, indicating that the association of Nucleolin with LINE1 RNA detected by RIP is not secondary to binding to DNA (Figure SSD). In contrast, Nucleolin does not bind other RNAs such as the spliceosomal-RNA U1 (Figures 5B and SSD), despite its abundant nuclear expression (Figure S5E).

Next, we tested whether Nucleolin may be a repressor of Dux and the 2C program in ESCs. A targeted RNAi screen for putative 2C regulators revealed that KD of Nucleolin, but not several other candidates tested, causes dramatic upregulation of Dux, MERVL, and 2C genes (Figures 5C and S5F–S5H). RNA-seq analysis revealed that Dux targets and MERVL repeats are among the most highly upregulated transcripts upon Nucleolin KD (Figures 5D, 5E, and S5I; Table S3). Moreover, there is a remarkably high similarity between the transcriptomes of ESCs depleted for LINE1 or Nucleolin (Figure 5F, Spearman’s ρ = 0.6). These results indicate that Nucleolin KD largely recapitulates LINE1 KD at the transcriptional level. Nucleolin KD also mimics LINE1 KD ESCs with regard to inducing significant decreases in ribosomal protein gene transcription (Figure S5J), total RNA levels (Figure 5G) and ESC self-renewal (Figure 5H). Similarly, Dux KD rescues 2C gene upregulation, but not the self-renewal deficit induced by Nucleolin KD (Figures S5K–S5L). Together, our findings indicate that LINE1 RNA and Nucleolin protein interact and have overlapping functions in ESCs, including promotion of rRNA synthesis and repression of the Dux/2C program.

**Nucleolin and Kap1 Bind to Dux and rDNA in a LINE1 RNA-Dependent Manner**

The only factor to date shown to directly bind and repress Dux in mouse ESCs is the co-repressor tripartite motif-containing protein 28 (TRIM28/Kap1) (De Iaco et al., 2017). We found that Nucleolin interacts with Kap1 in ESCs, suggesting that they may function together in a complex (Figure 6A). Using inducible mutant CreERT2;Kap1 fl/fl ESCs (Rowe et al., 2010), we confirmed that Kap1 deletion induces a significant increase in Dux and 2C genes and MERVL (Figures 6B and 6C), and that Kap1 or Nucleolin KD both lead to over 10-fold increases in 2C-like cells in vitro (Figure 6D). Analysis of published RNA-seq data from Kap1-deleted ESCs (Ecco et al., 2016) also revealed a similarity in transcriptional changes to LINE1 KD, albeit less striking than the Nucleolin KD-LINE1 KD similarity (Figures 5F and S6A, Spearman’s ρ = 0.34). We found that both Nucleolin and Kap1 bind Dux loci, and the levels of both proteins at Dux are significantly reduced upon LINE1 KD (Figure 6E). Moreover, efficient Nucleolin binding to rDNA is also dependent on LINE1 (Figure 6F). Surprisingly, Kap1 is also robustly recruited to rDNA in a LINE1-dependent manner (Figure 6F), and Kap1 deletion leads to similar reductions in nascent rRNA transcription as LINE1 or Nucleolin KD (Figure S6B). LINE1 KD does not affect the levels of Nucleolin or Kap1 RNA or protein (Figures S3C and S6C–S6E), nor their interaction (Figure S6F). Collectively, these data indicate that LINE1 is essential for the efficient chromatin binding of a Nucleolin-Kap1 complex that represses Dux and promotes rRNA expression.
Figure 6. LINE1 Promotes Binding of Nucleolin and Kap1 to Dux and rDNA

(A) Co-immunoprecipitation (coIP) showing association of endogenous Kap1 and Ncl proteins in ESCs. (B and C) qRT-PCR confirming Kap1 deletion (B) and upregulation of Dux, 2C genes, and ERVs (C). Data are mean ± SEM; n = 3 biological replicates. (D) Representative micrographs (left) and percentage of 2C-like cells (right) in 2C-GFP reporter ESCs 3 days following siRNA KD of Ncl or Kap1; n = 2 independent experiments. (E and F) ChIP assays for Ncl and Kap1 at Dux (E) and rDNA (F), with or without LINE1 KD. Data are shown as a percentage input normalized to enrichment at int-chr11 negative control region and are mean ± SEM; n = 3 independent experiments for Ncl ChIP; n = 2 independent experiments, Kap1 ChIP. (G) Example images and quantification of nuclear LINE1 RNA foci by RNA-FISH in ESCs (2C-GFP−) or 2C-like cells (2C-GFP+), representative of 2 experiments. Scale bar, 10 μm; n = number of cells. (H) Representative images and quantification of distinct localization patterns of Dux loci in ES versus 2C-like cells by immunofluorescence (IF) for Ncl combined with Dux DNA-FISH, using 2C-GFP reporter ESCs. Nucleoli are labeled with Ncl antibodies. Example Dux loci are indicated (white arrows), showing two nucleolar loci in an ESC versus one nucleoplasmic locus in a 2C-like cell. Statistics are calculated by Chi-square test for the indicated number (n) of Dux loci. Scale bar, 2 μm. See also Figure S6.
Subsequently, we sought to interrogate what possible difference in 2C-like cells, compared to standard ESCs, allows for induction of Dux and the 2C program. While the levels or localization of Kap1 and Nucleolin are unchanged (Figure S6G), the nuclear abundance of LINE1 RNA is significantly reduced in 2C-like cells compared to ESCs (Figure 6G). In addition, 2C-like cells have significantly fewer nucleoli per cell compared to ESCs (Figure S6H). DNA FISH revealed that in ESCs the Dux loci are most often located in the perinucleolar or laminar regions (Figures 6H and S6I–S6K), both of which are thought to be transcriptionally repressive heterochromatin environments (Guetg and Santoro, 2012; Reddy et al., 2008). Intriguingly, the transition of ESCs to the nucleoplasm (Figures 6H and S6K). Taken together, these data suggest that a combination of reduced abundance of LINE1 RNA with changes to the interaction of Dux with Nucleolin at peri-nucleolar heterochromatin may facilitate the entry of ESCs into a 2C-like state, a possibility that deserves future exploration.

**LINE1 Promotes Silencing of the Dux/2-Cell Program, rRNA Synthesis, and Developmental Progression Past the 2-Cell Stage**

Lastly, we investigated the role of LINE1 in embryonic development (Figure 7A). Almost no LINE1 KD embryos reach the blastocyst stage and most arrest at the 2-cell stage (Figures 7B and S7A). Furthermore, late 2-cell LINE1 KD embryos display high upregulation of Dux as well as its target 2-cell gene, Zscan4 (Figure 7C). In contrast, the expression of Eif1a, a marker of ZGA (Davis et al., 1996; Zeng and Schultz, 2005), is sharply reduced upon LINE1 KD. 2-cell embryos depleted of LINE1 exhibit significant reductions in chromatin accessibility (Figure S7B) in agreement with (Jachowicz et al., 2017) and display significantly increased levels of heterochromatin (Figure S7C). While control ASO-injected embryos show the expected strong cytoplasmic accumulation of 18S rRNA, in LINE1 ASO-injected embryos 18S rRNA is largely retained in the nucleus (Figure 7D). In agreement, chemical inhibition of rRNA synthesis from either the zygote (Lin et al., 2014) or the 2-cell stage onward (Figure S7D) is incompatible with pre-implantation development.

Next, we performed RNA-seq on samples where zygotes were microinjected with lower concentrations (0.5X) of ASOs to allow some LINE1 KD embryos to progress to the 4-cell stage (Figure 7E). Very few transcriptional changes are detected in LINE1 KD embryos at the early 2-cell stage. In contrast, LINE1 KD 4-cell embryos are transcriptionally distinct from controls (Figure S7E; Table S4). Using published data (Wu et al., 2016), we defined gene clusters associated with Dux/early 2-cell gene expression (cluster 4, Figures 7F and S7F), or with elevated expression maintained up to the 4-cell stage (cluster 1, Figures 7F and S7F), 4-cell LINE1 KD embryos display a significant upregulation of early 2-cell genes and reduced expression of 4-cell genes (Figures 7G, 7H, and S7G). Genes associated with ZGA (Zeng and Schultz, 2005) are significantly decreased in 4-cell LINE1 KD embryos (Figures 7F and 7G), along with ribosomal genes (Figure 7H).

Finally, we performed LINE1 KD at the late 2-cell stage, when Dux has already been silenced and ZGA initiated (Figure 7J). Very few of these LINE1 KD embryos develop past the 4-cell stage (Figures 7J and S7J). Moreover, 4-cell embryos depleted of LINE1 from the late 2-cell stage display upregulation of MERVL and 2-cell genes (Figures 7K and S7J). Overall, these data indicate that there is an ongoing requirement for LINE1 RNA for repression of the 2-cell program and developmental progression during early embryogenesis. Taken together, the data suggest that a failure to repress the Dux/2-cell program and defective ZGA and ribogenesis contribute to embryonic arrest upon LINE1 depletion.

**DISCUSSION**

The expression of TEs such as LINE1 is generally thought to be detrimental to cells because it can cause mutations or apoptosis (e.g., Malki et al. [2014]; Burns [2017]). In contrast, we report here that the expression of LINE1 regulates exit from the 2-cell stage by performing two main functions: repressing the 2C program induced by Dux, and activating rRNA synthesis to support rapid proliferation. These two functions are unified by the interaction of LINE1 RNA with Nucleolin, which we identify as a novel repressor of the Dux/2C program. Thus, rather than being a simple genomic parasite, LINE1 may be best viewed as a symbiont that is an integral part of the transcriptional networks that regulate cellular potency during early mammalian development (Figure 7L).

Nucleolin is most commonly associated with the positive regulation of rRNA synthesis (Ginisty et al., 1998). In recent years, Nucleolin has been shown to also have roles in chromatin remodeling, DNA replication, and DNA repair (reviewed in Jia et al. [2017]). Our work uncovers a novel function for Nucleolin, and it will be of interest to identify the functional elements in the LINE1 RNA by defining the regions that interact with Nucleolin and potentially other factors, like Kap1. Intriguingly, the region of mouse LINE1 RNA previously used to identify Nucleolin as an interactor was the inter-ORF (Peddigari et al., 2013), the same region against which we designed the LINE1 ASO (Figure 1C). It is therefore possible that the ASO disrupts the interaction between LINE1 RNA and Nucleolin, which in turn may contribute to destabilizing LINE1 RNA.

It remains unclear how LINE1 RNA-Nucleolin-Kap1 are targeted to the Dux cluster. LINE1-Nucleolin-Kap1 may have other interacting partners with DNA binding specificity. For example, YY1 has been implicated in targeting a repressive complex containing Nucleolin to the human DUX4 cluster (Gabellini et al., 2002), although we did not find a role for YY1 in Dux repression in ESCs (Figure 5C). Alternatively, KRAB-ZFP transcription factors are known to recruit Kap1 to repress TEs (Lupo et al., 2013; Wolf et al., 2015), a function that might have been coopted for Dux silencing. Finally, our data leave open the possibility that transcribed LINE1 DNA loci contribute in cis to Dux repression or rRNA activation via higher-order chromosome interactions (Figure 7L). Furthermore, while LINE1 is specifically detected at Dux and rRNA, it is likely to have other genomic targets that remain to be discovered.
Our data suggest that the LINE1-Nucleolin-Kap1 complex has both an activating function (at rDNA) and a repressive function (at Dux). While this might seem paradoxical, Nucleolin and Kap1 have been shown to have both activating and repressive effects. In addition to its role as activator of rDNA, Nucleolin has been reported to repress the expression of genes, such as cMyc (González et al., 2009), MMP13 (Samuel et al., 2008), or the D4Z4 repeat (Gabellini et al., 2002). Conversely, the co-repressor Kap1 also has activating functions at both the single gene level (Singh et al., 2015) and globally (Bunch et al., 2014). It is possible that the reduction in nascent rRNA synthesis in Kap1 mutant ESCs contributes their self-renewal defect, in addition to the previously reported de-repression of endogenous retroviruses (Rowe et al., 2010). Moreover, the localization of Dux loci to Nucleolin-positive peri-nucleolar regions in ESCs, but not 2C-like cells, provides a potential rationale at the level of nuclear 3D organization for the coordinate roles of LINE1/Nucleolin/Kap1 in activation of rDNA versus repression of Dux.

LINE1 is expressed throughout pre-implantation development (Figure 1B) (Fadloun et al., 2013), which implies that its presence alone may not suffice to repress Dux and the 2C program. However, analysis of RNA-seq data from early embryos reveals that Nucleolin and Kap1 are sharply induced at the late 2-cell stage, correlating with Dux silencing (Figure S7K). Thus, it is possible that, by promoting ZGA at the 2-cell stage, Dux induces several of its own repressors, which can then bind LINE1 RNA and silence the Dux loci. Moreover, there are large-scale changes to the organization of the nucleolus and its associated heterochromatin during these stages (Borsos and Torres-Padilla, 2016) that may pertain to the silencing of the Dux/2C program.

It has been proposed that the retrotransposition activity of LINE1 is essential for early mouse development, possibly by making cDNA copies of sperm-derived RNAs (Beraldi et al., 2006; Sciamanna et al., 2009). Using inhibitors that block LINE1 mobility (Figure S2G) (Jones et al., 2008), we found that retrotransposition is not involved in the regulation of 2C gene expression nor of ESC self-renewal (Figures S2H and S2I). Moreover, LINE1 ORF1 protein, which is required for retrotransposition, is predominantly cytoplasmic (Figure S1A), in contrast to the nuclear localization of LINE1 RNA. While we cannot at present exclude a non-canonical function for LINE1 ORF1 or ORF2 proteins in the developmental roles reported here, our data indicate that it acts as a chromatin-associated RNA that binds Nucleolin and Kap1 to regulate gene expression.

The partnership between LINE1 and Nucleolin in the regulation of rRNA synthesis indicates that LINE1 contributes to ESC hypertranscription (Percharde et al., 2017a). This notion is supported by the fact that LINE1 KD leads to decrease in total RNA levels, nascent transcription, nascent translation and self-renewal. Although we do not detect LINE1 RNA binding to ribosomal protein genes, the synthesis of rRNA and ribosomal protein genes is highly coordinated (Laferté et al., 2006).

In addition, we have recently shown that reductions in translational output in ESCs rapidly induce a decrease in nascent transcription of highly expressed genes, including rRNA and ribosomal protein genes (Bulut-Karslioglu et al., 2018). Our data support a model whereby LINE1-mediated induction of rRNA synthesis leads to global increases in ribosomal biogenesis, enabling rapid growth of the early embryo. We speculate that the LINE1-Nucleolin partnership may play roles in other stem/progenitor cells, where it might not necessarily act to repress Dux but may still promote ribogenesis and proliferation.

Our work raises the question of how a mechanism for control of developmental potency based on TEs might have evolved. Active TEs are under acute surveillance by cellular pathways that minimize transposition, including by Kap1 (Rowe et al., 2010). In part because of this, and in part because of a loss in ability to transpose due to an accumulation of mutations, the sequence of TEs is generally thought to be subject to a rapid...

---

**Figure 7. LINE1 Regulates Dux Silencing, rRNA Synthesis, and Early Development**

(A) Summary diagram of ASO microinjection experiments in (B)–(D).
(B) Developmental progression in the indicated number (n) of embryos following ASO microinjections, collected in 3 independent experiments. χ² p values were calculated for the developmental rate of embryos injected with LINE1 or control ASOs.
(C) qRT-PCR for the indicated genes in late 2-cell (2C) embryos harvested 24 hr post-injection. Data are mean ± SEM, n = 3 technical replicates, showing 2 independent experiments.
(D) 18S RNA FISH in late 2C embryos 24 hr following microinjection with Control or LINE1 ASOs, with nuclei stained by DAPI. Nuclear or cytoplasmic foci were counted in each embryo and plotted below as 18S cytoplasmic/nuclear ratio. Data show mean ± SD from the indicated number (n) of embryos from 2 independent experiments. Scale bar, 20 μm.
(E) Summary diagram of 0.5× ASO microinjection experiments for RNA-seq in (F)–(H).
(F) RNA-seq data from Wu et al. (2016) showing mean fragments per kilobase of transcript per million (FPKM) expression relative to MII oocyte for the indicated gene sets identified by mean fragments per kilobase of transcript per million (FPKM) expression relative to MII oocyte for the indicated gene sets identified by -means clustering or for ZGA genes defined in Zeng and Schultz (2005). Shading denotes mean ± SEM at each time point.
(G) Boxplot depicting the log₂-fold change in 4-cell (4C) embryos upon LINE1 KD for the indicated number (n) of gene sets displayed in (G), compared to all expressed genes. p value, 2-sided Wilcoxon rank-sum test.
(H) Examples of the expression of early 2C genes, Dux and Zscan4d, in the indicated embryo samples. p value, topetable FDR, showing n = 3 samples per condition.
(I) Summary diagram of late 2C ASO microinjection experiments in (J) and (K).
(J) Developmental progression in the indicated embryos following late 2C ASO microinjections. χ² p values were calculated for the developmental rate of the number (n) of embryos injected with 1X RC or LINE1 ASOs.
(K) qRT-PCR analysis of MERVL expression in 4C or 5-8-cell (5-8C) embryos following late 2C microinjections with ASOs. Data are mean ± SEM; n = 3 technical replicates and are representative of 3 (LINE1 ASO) or 2 (RC ASO) independent experiments.
(L) Model for the role of LINE1 in early development and ESCs. LINE1 acts as an RNA-scaffold and binds to rDNA and Dux. LINE1 RNA-Ncl-Kap1 cooperate to turn off the 2C gene expression program and promote high levels of ribosome biosynthesis during early development.

See also Figure S7 and Table S4.
rate of divergence. In fact, some mammalian species may have completely lost all retrotransposition-competent LINE1 elements in their genome, even though they can still express mutated LINE1 RNAs (Cantrell et al., 2008). Our results indicate that chromatin-associated LINE1 RNA regulates gene expression and developmental potency without requiring retrotransposition activity. This role of LINE1 as a chromatin-associated RNA therefore avoids the potential detrimental effects of LINE1 retrotransposition that have been reported in several disease states, including cancer (Burns, 2017). The interaction of LINE1 RNA with binding partners, such as Nucleolin is expected to be mediated by RNA secondary structure, which is less constrained by primary sequence than protein-coding regions. Thus, rather than being a vulnerability, the regulation of early development by TEs may allow both robustness, due to the repeated nature of TEs, and adaptability, due to their rapid evolution and their potential to support transposition in conditions of stress. In this regard, it is interesting that the percentage of the genome occupied by LINE1 elements seems to have sharply increased with development of therian mammals (e.g., Ivancevic et al. [2017]). The exploration of the function of LINE1 in other species should shed light on the role of TEs in shaping the evolution of development.

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STAR★METHODS

KEY RESOURCES TABLE

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Chemicals, Peptides, and Recombinant Proteins

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**Experimental Models: Organisms/Strains**

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**Oligonucleotides**

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| RC ASO: AGACAGGCCACAAGAACAGAGTGCCA | This paper; Gene Tools | N/A |
| LINE1 ASO: TGGCATTCTGTTCTTGTGGCTGTCT | This paper; Gene Tools | N/A |
| siGenome siRNAs: custom and Smartpools (see Table S5) | This paper; Dharmacon | N/A |
| LINE1 RNA smFISH probes Quasar 670 (see Table S5) | This paper; Biosearch Technologies | N/A |
| Gapdh RNA smFISH probes Quasar 570 | Biosearch Technologies | Cat# SMF-3002-1 |
| Dux DNA FISH probes (see Table S5) | This paper | N/A |
| LINE1 RNA ChiRP probes (see Table S5) | This paper | N/A |
| Primers for qRT-PCR and ChiP-PCR (see Table S5) | This paper | N/A |

**Recombinant DNA**

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CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Miguel Ramalho-Santos (mrsantos@ucsf.edu, mrsantos@lunenfeld.ca).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mice

Animal experiments were performed with 5-8-week-old female and 2-6-month-old male C57BL/6 mice. Animals were maintained on 12 h light/dark cycle and provided with food and water ad libitum in individually ventilated units. Animal experiments in USA were in accordance with the guidelines of the UCSF Institutional Animal Care and Use Committee, protocol AN091331-03. Animal experiments in UK were approved by the University of Edinburgh’s Animal Welfare and Ethical Review Board (AWERB) and carried out under the authority of a UK Home Office Project License.

Mouse ES cell culture

Mouse E14Tg2A (E14) ES cells (male) were used for all experiments (Hooper et al., 1987), except for p53−/− ES cells (Sabapathy et al., 1997), which are V6.5-derived (male, a gift from Scott Oakes). ES cells were cultured on 0.1% gelatin-coated plates in ES-FBS culture medium (high glucose DMEM GlutaMAX with sodium pyruvate (Thermo Fisher Scientific), 15% FBS (Atlanta Biologicals), 0.1mM non-essential amino acids, 50 U/mL penicillin-streptomycin (UCSF Cell Culture Facility), 0.1mM 2-Mercaptoethanol (Millipore) and 1,000U/ml LIF supplement (ESGRO, Millipore). Where indicated, ES cells were grown in N2B27/2i/LIF conditions (DMEM/F-12, Neurobasal medium, 1x N2/B27 supplements, 1μM PD0325901, 3μM CHIR99021, LIF as above) according to Ying et al. (2008) for at least 4 passages before being used for experiments. Deletion of Kap1 in CreERT2;Kap1fl/fl ES cells (undetermined sex) (Rowe et al., 2010) was performed in ES/FBS conditions with 1μM 4-OHT overnight and analyzed 4 days later. Routine testing of E14 ES cells revealed absence of mycoplasma contamination. ES cells were not genotyped.

2C-GFP ES cell line

2C-GFP reporter ESCs were generated as in Ishiuchi et al. (2015), using Addgene plasmid 69071, from parental E14 ES cells. 4x10⁶ cells were nucleofected with 4μg linearized 2C-GFP plasmid, and plated at low density in 10cm² plates. Selection was performed with 250μg/mL G418 (Mirus) commencing 36h after nucleofection and maintained for 8 days before individual colonies were picked and expanded. A clone with high, specific upregulation of 2C genes in sorted GFP+ cells (Figure S2K) was used for all experiments.

MEFs

E13.5 primary MEFs derived from pooled CD1 embryos (mixed sex) were cultured in MEF medium (as ES-FBS but without 2-Mercaptoethanol and LIF), and used within 4 passages of initial derivation.

METHOD DETAILS

ASO- and siRNA-Mediated Knockdown

A morpholino ASO targeting the inter-ORF region of LINE1 was designed with software available at Gene Tools LLC. The morpholino chemical backbone was chosen because of its stability, specificity and extensive use in pre-implantation mouse embryos (e.g., Lin et al. [2013]). ASOs have been shown to be potent inhibitors of nuclear RNAs, such as IncRNAs (Lennox and Behlke, 2016). The LINE1 ASO was validated in silico to be perfectly homologous to at least 500 full-length LINE1 elements, using L1Base (Penzkofer et al., 2005). This sequence was verified by Blast as not having homologies to any known Refseq mRNAs. The reverse complement of the LINE1 ASO sequence does not target LINE1 RNA and was used as a control (RC ASO). In the indicated embryo experiments,
a standard non-targeting control ASO (Gene Tools) was also used. Both lissaminated (cell line experiments) and non-lissaminated (embryo experiments) ASOs were utilized. ASOs were introduced into cells by nucleofection, utilizing an Amxana Nucleofector 2b device and ES nucleofection kit (Lonza), according to the manufacturer’s instructions. 4-5 million cells were used per nucleofection together with 5nmol of the indicated ASO. Cells were plated in ES medium immediately following nucleofection and left to recover for 24-48h. For colony formation assays and expression analyses excluding co-transfections with siRNAs, Lissamine-positive ES cells were purified by FACS utilizing a BD FACSAria II (BD Biosciences) to enrich for successfully-nucleofected cells. Nucleofection efficiency was routinely 70% or above. All other experiments were performed on the bulk population of nucleofected cells without prior enrichment for ASO-positive populations. siRNA transfections were performed in ES cells with Lipofectamine 2000 (Thermo Fisher Scientific) as described previously (Percharde et al., 2012). ES cells were plated 5-7h before transfection at a density of 5x10^4 ES cells per 6-well culture area and transfected with 100 pmol siRNA, according to the manufacturer’s standard recommendations. A non-targeting siRNA (siGenome siControl #2, Dharmacon) was used as a control. For LINE1 siRNA knockdown, 2 independent siRNA sequences were designed using online tools available at Dharmacon and verified to have no perfect match to other mRNAs. For combined ASO/siRNA experiments, ES cells were first nucleofected with ASOs, then plated and left to recover for 7h. Next, the medium was exchanged for fresh ES-FBS without antibiotics and siRNA transfections were performed as above. The medium was exchanged the next morning and cells harvested for RNA extraction approximately 48h post initial nucleofection. All ASO and siRNA sequences are available in Table S5.

**Embryo Microinjection and Culture**

Females were superovulated by administration of 7.5 I.U. of pregnant mare serum gonadotropin (PMSG; National Hormone Pituitary Program (NHPP), Harbor-UCLA Medical Center, CA, USA or Intervet UK) and 48h later with 7.5 I.U. of human chorionic gonadotropin (hCG; Sigma-Aldrich or Intervet UK). Embryo experiments were performed as described previously (Lin et al., 2013, 2014), and embryos were microinjected using a microinjector (FemtoJet 4i, Eppendorf) and an inverted microscope (Leica, DMi8) equipped with micromanipulators (Narishige). Zygotes were microinjected with Standard Control, RC, or LINE1 ASOs. ASO solutions were injected into the cytoplasm from a stock concentration of 1.5mM (1X) or 0.75mM (0.5X), utilizing 2-5 pl of solution per injection. Embryos were cultured in KSOM+AA medium at 37°C in 5% CO2. For qRT-PCR analysis in late 2C embryos, 10-11 embryos were collected approximately 24h post injection for RNA isolation using the PicoPure RNA Extraction Kit (Arcturus), or were cultured for a further 3 days to monitor developmental progression. Embryo qRT-PCR data are normalized to Hprt expression. For embryo RNA-seq experiments, embryos were microinjected with 0.5X ASO and cultured in KSOM+AA medium. Early 2C embryos were collected on approximately 33 h post hCG (~9 h post-microinjection). 4C embryos were collected after 2 days of in vitro culture. ASO injections at the late 2-cell stage were performed after collection approximately 43h post hCG administration, and 1X ASOs were microinjected into the cytoplasm of both blastomeres. Developmental progression to blastocyst was monitored, or 4C embryos were collected on the day following injection for RNA isolation and qRT-PCR analysis, as above. For inhibition of rRNA synthesis after ZGA, late 2C embryos were cultured in KSOM+AA with 1 μM CX-5461 or 0.1% DMSO.

**DNase-TUNEL Experiment**

DNase I-TUNEL assays were performed in late 2C embryos. Embryos were permeabilized by 0.5% Triton X-100 in pre-extraction buffer (300mM sucrose, 25mM HEPES, 1mM CaCl2, 50mM NaCl, 3mM MgCl2) for 5 min before digesting with 0.2U/μl of DNase I (NEB). Embryos were then fixed in 4% PFA. TUNEL Assays (Click-iT TUNEL Imaging assay, Thermo Fisher Scientific) were followed according to manufacturer’s instructions. The nuclear area was defined according to Hoechst DNA staining and the intensity of nuclear TUNEL signal was quantified using Fiji software.

**RNA FISH**

Embryo RNA FISH was performed as previously described (Lin et al., 2014). Embryos were fixed in methanol, permeabilized in 70% ethanol and hybridized at 37°C using 48 single-molecule probes designed to span the length of LINE1 ORF2 RNA, designed from the sequence of L1spa (L1Md_Tf family) and expected to target the majority of transcribed LINE1 RNAs (Naas et al., 1998), or against 18S rRNA at 1:250 dilution. RNA-FISH in ES cells was performed on glass coverslips 48h after ASO or siRNA treatment, according to the manufacturer’s standard protocol (Biosearch Technologies). Fixation was performed in 1% paraformaldehyde (PFA) for 15 min, followed by permeabilization overnight in 70% ethanol at 4°C prior to hybridization. Hybridization was performed as above, using the probes against LINE1 RNA. Co-incubations were performed with Gapdh probes where indicated (purchased from Biosearch Technologies). For +RNase negative controls, coverslips were incubated in 2X saline-sodium citrate (SSC) buffer plus 10 μg/mL RNase A (Thermo Fisher Scientific) for 30 min at 37°C, then washed twice in 2X SSC, prior to LINE1 FISH hybridizations. For LINE1 RNA FISH in ES versus 2C-like cells, GFP+/− populations of 2C-GFP reporter cells were separated by FACS then plated onto separate wells on matrigel-coated Lab-Tek II Chambered Coverglass for 1h before fixation and hybridization as above. All images were collected on a Leica DMI 4000B inverted scope using a 100 × oil-immersion objective, every 0.25 μm using the Z stack function of the MetaMorph software (Molecular Devices). All nuclei were counterstained with DAPI. RNA FISH foci were quantified with StarSearch online software (Raj et al. [2008], Raj Lab, University of Pennsylvania), using identical threshold settings between different samples and images.
Cell-cycle and Self-renewal Assays

24h following nucleofection or siRNA transfection, ES cells were either trypsinized and counted manually (siRNAs) or isolated by FACS according to Lissamine fluorescence (ASOs); 1000 cells were plated per well of a 12-well plate in ES-FBS medium. Colonies were left to form over a period of 5-6 days and then fixed and stained for Alkaline Phosphatase (AP) according to the manufacturer’s recommendations (Sigma-Aldrich). The numbers of AP-positive colonies per well were manually counted. For proliferation assays, 1x10^6 ES cells were plated immediately following ASO nucleofection and then each day for the next four days the cells were trypsinized and counted, replating 1x10^6 cells each time back into culture. Cell-cycle analysis was performed according to Bulut-Karslioglu et al. (2018) and the standard manufacturer’s protocol, incubating ES cells 48h after nucleofection with 10 μM EdU for 1 h, followed by analysis of EdU incorporation and DNA content with the Click-iT EdU imaging kit (Life Technologies) and FxCycle-Violet, respectively.

RNA Extraction and Expression Analysis

For ASO-nucleofected samples, RNA was isolated from 300,000 FACS-purified ES cells using the RNeasy mini kit (QIAGEN), according to manufacturer’s instructions. RNA was treated once on-column with DNase I and once more off-column (RNase-free DNase I from Thermo Fisher Scientific) to remove any residual DNA contamination. cDNA synthesis was performed from 1 μg RNA with the High Capacity cDNA Reverse Transcription kit (Applied Biosystems) and qRT-PCR performed with SYBR green (KAPA) on an ABI-Prism PCR machine. qRT-PCR analysis following siRNA experiments or ASO/siRNA experiments were performed as above, but without prior FACS purifications. All gene expression data were normalized to two independent housekeeping genes (Rpl7, Ubb or H2A).

Retrotransposition Assay

ES cells were nucleofected with 5 μg codon-optimized LINE1 CAG-ORFeus GF-P reporter plasmid (An et al., 2006) or LINE1 5’UTR-ORFeus GF-P reporter plasmid (Newkirk et al., 2017). The following morning, Stavudine (d4T, 20 μM) or Tenofovir disoproxil fumarate (TDF, 4 μM) (Jones et al., 2008) was added and cells cultured for a further 48h, when retrotransposition rates were determined by flow cytometry for GFP-positive cells, or cells harvested for RNA extraction as above. Colony formation assays in the presence of d4T and TDF were as above, with the addition of inhibitors performed 24h after plating and maintained throughout.

RNA-sequencing

Sample preparation for RNA-seq in ES cells was performed as previously described (Percharde et al., 2017b). RNA was extracted utilizing the RNeasy mini kit as for qRT-PCR, then 800ng DNase-treated total RNA was used per library preparation according to the NEN NEXT Ultra Directional Library Prep Kit for Illumina (NEB). Embryo RNA-seq was performed in pools of 14-18 early 2C or 4C embryos per sample, with total RNA isolated using the PicoPure RNA Extraction Kit (Arcturus). Libraries were generated using the SMARTer Stranded Total RNA-seq Kit v2 (Takara) and 500pg total RNA input per sample. In all cases, three replicates were sequenced per condition at the UCSF Center for Advanced Technology on an Illumina HiSeq 2500 or HiSeq 4000, with 50bp single end reads.

Immunofluorescence

ES cells were plated onto matrigel-coated 8-well chambers for 1h, fixed for 10 min in 4% PFA, and blocked and permeabilized in one step in IF buffer (PBS, 10% donkey serum, 2.5% BSA) plus 0.4% Triton X-100. Incubations were performed overnight at 4°C, using the following antibodies and dilutions in IF buffer: GFP (1:100), Oct4 (1:100), Nanog (1:200), Kap1 (1:500), Nucleolin (1:1000 x), Orf1p (Di Giacomo et al., 2014) (1:500). The next day, slides were washed 3 times in PBS, incubated for 60 min in the relevant fluorescently-conjugated secondary antibodies (1:1000 in IF buffer, Life Technologies), and washed again as before. Slides were mounted with Vectashield containing DAPI and imaged on a Leica SP5 upright confocal microscope at 63X magnification under oil immersion. Embryo immunofluorescence (IF) experiments were performed as above, except blocking was performed in 5% BSA and incubation with primary antibodies was at 37°C for 1.5 hr. with antibodies against H3K9me2 or H3K9me3 (1:50), and nuclei counterstained with DAPI. Images of stained embryos were acquired by a spinning disk confocal (CSU-W1, Yokogawa) on an upright microscope frame (BX-63, Olympus) using a 60x silicon oil immersion objective (UPLSAPO 60XS2, Olympus) with additional 2x intermediate magnification. IF signal intensity was quantified using Fiji software.

RNA Immunoprecipitation

RNA immunoprecipitation (RIP) experiments were performed on nuclear extracts according to a standard Abcam protocol, with the following modifications. 1 μg anti-Nucleolin or control IgG antibodies were pre-bound to 30 μL Protein A Dynabeads (Thermo Fisher Scientific) and incubated rotating for at least 3h at 4°C. Beads were next collected on a DynaMag (Thermo Fisher Scientific) and resuspended in RIP buffer (150mM KCl, 25mM Tris pH 7.4, 5mM EDTA, 0.5mM DTT, 0.5% NP40, protease and RNase inhibitors) containing 500ng/μL tRNA (Thermo Fisher Scientific) and 1 mg/mL RNase-free BSA (bioWORLD) to block for 30 min, then collected and used immediately in RNA immunoprecipitation. Where indicated, 40U of Turbo DNase (Thermo Fisher Scientific) was added to nuclear extracts and incubated for 30 min at 37°C, then quenched with 10mM EDTA, before continuing with immunoprecipitations. Prior to immunoprecipitation, ES nuclear extracts were also blocked for 30 min with 20 μL Protein A Dynabeads at 4°C, 30 min. Cleared nuclear lysates were incubated together with antibody-bound blocked beads overnight at 4°C. The next day, lysates were washed...
four times in RIP buffer, once in PBS, and RNA was extracted from beads using Trizol and standard phenol-chloroform extraction. The aqueous phase containing the RNA was loaded onto RNeasy mini columns (QIAGEN) with 2x volume of 100% ethanol and RNA was purified according to the standard protocol. RNA was DNase I treated twice as before and used to generate cDNA for qRT-PCR.

**Nascent Transcription and Translation Assays**

For nascent transcription assays, ES cells were cultured for 45 min in normal ES medium supplemented with 1mM EU. For nascent translation assays, cells were incubated in HPG medium (as ES-FBS medium with the following substitutions: Methionine- and Cysteine-free DMEM, 1mg/mL BSA instead of FBS) for 30 min, followed by HPG medium plus 50 µM HPG for 45 min. Following incubations, cells were collected by trypsinization, fixed, and permeabilized, and nascent RNA or proteins were labeled using the Click-iT Alexa Fluor 488 RNA or Protein Synthesis Assay Kit (Thermo Fisher Scientific). All samples were analyzed on a LSR II Flow Cytometer (BD Biosciences). Relative EU or HPG incorporation was quantified by comparing the EU/HPG median fluorescence intensity between LINE1 and RC nucleofected samples, and carried out 24-48h after nucleofection.

**Nascent RNA capture followed by qRT-PCR**

EU incorporation was performed as above, with the exception that incubations were with 0.4mM EU for 30 min. Where indicated, CX-5461 was added in the morning prior to EU addition, for a total of 8h treatment. ES cells were washed, collected by trypsinization, counted, and 2x10^6 were used to extract RNA. Nascent RNAs was captured according to standard protocols within the Click-iT Nascent RNA Capture Kit (Invitrogen), and used in qRT-PCR assays with the indicated primer sets (see Table S5 for sequences).

**Chromatin Isolation by RNA Precipitation**

ChIRP was performed according to Yin et al. (2015) with some modifications. 59nt DNA probes were biotinylated through terminal transferase (NEB) with Bio-N6-ddATP (ENZO) as substrate. E14 ES cells were harvested by trypsin digestion and crosslinked with 2mM dithiois(succinimidyl propionate), (DSP, Thermo Scientific) in PBS at room temperature for 30 min with gentle end to end rotation. Formaldehyde was added to a final concentration of 3% to crosslink for 10 min further, then quenched with 250mM Glycine at room temperature for 5 min. ES cells were centrifuged and the pellet was washed with ice-cold PBS for 3 times, then snap frozen in liquid nitrogen and stored at −80°C. Crosslinked DNA (−1 x 10^7) were resuspended with 500µl DNase I digestion solution (20mM Tris-HCl, pH7.5, 5mM MgCl2, 0.5mM CaCl2, 0.5% Triton X-100) with 1/20 volume of vanadyl ribonucleoside complex (VRC, NEB), 2.5µl protease inhibitors and 2.5µl 200mM PMSF. Dnase I was added to a final concentration of 12 U/ml; the reaction was rotated for 10 min at 37°C and stopped with 20mM EDTA. Chromatin was pelleted, washed once with nucleic lysis buffer (NLB, 50mM Tris-HCl, pH7.5, 10mM EDTA, 1% SDS, inhibitors) and sonicated in NLB (5 cycles of: 25% amplitude, 6 s on, 15 s off, Vibra-Cell Ultrasonic Liquid Processors). Insoluble material was removed by centrifugation and the supernatant used for ChIRP experiments. For the Rnase treatment control, samples were treated with 10 µg/ml Rnase A/T1 at 37°C for 20 min. For hybridization, samples were incubated with 20pmol probes per 200µl lysate, supplemented with one-fourth volume of 5x hybridization buffer (50mM Tris-HCl, pH7.5, 10mM EDTA, 1.5M NaCl, 50% formamide). The hybridization was conducted at 39°C rotating for 3h. 50 µl pre-balanced streptavidin M280 beads were then added and the incubation continued for an additional 3h. The beads were washed 5 times total with 0.2 x SSC wash buffer (0.1 x SSC, 1% SDS, inhibitors) and sonicated in NLB (5 cycles of: 25% amplitude, 6 s on, 15 s off, Vibra-Cell Ultrasonic Liquid Processors). Insoluble material was removed by centrifugation and the supernatant used for ChIRP experiments. For the Rnase treatment control, after 3 times of washing, the beads were treated once with 10 µg/ml Rnase A/T1 at 37°C in Rnase digestion buffer (50mM Tris-HCl, pH7.5, 75mM NaCl, 1mM DTT), before washing twice more. To elute, the beads were washed once with SDS elution buffer (50mM Tris-HCl, 5mM MgCl2, 75mM NaCl, 1% SDS) at 39°C for 20 min, and once with elution buffer (50mM Tris-HCl, 5mM MgCl2, 75mM NaCl, 0.1% Triton X-100) at 39°C for 5 min. DNA was eluted from the beads by RNase H treatment in two sequential incubations with RNase H (NEB) at 37°C for 20 min, and with SDS elution buffer at room temperature, 2 min, combining all eluents. Crosslinking was reversed by treatment with 0.1 µg /µl protease K, 150mM NaCl, and 10mM EDTA at 65°C overnight and the DNA was purified using the MinElute PCR Purification Kit (QIAGEN). ChIRP enrichments were analyzed by qPCR of the purified DNA. For RNA-ChIRP analysis, beads were boiled in NLB after the 0.1x SSC washes, then further reverse-crosslinked by boiling at 95°C for 30 min in the presence of 1mM DTT. Reverse crosslinked RNA was purified using Trizol and processed for qRT-PCR analysis as for other RIP RNA samples.

**DNA FISH**

Co-staining of Dux DNA-FISH and immunofluorescence followed a protocol modified from what was previously described (Guan et al., 2017). The sequences of 22 ssDNA oligos to tile the 5kb genomic region of one Dux repeat unit were designed via OligoArray2.1 (Rouillard et al., 2003), synthesized by IDT with 5’ amino modifications, pooled together in equimolar ratio, and covalently labeled with Cy3. For immunofluorescence combined with Dux DNA-FISH experiments, 2C-GFP ES cells were seeded onto Lab-Tek II Chambered Coverglass coated with Matrigel for 1h, then fixed 5 min with 4% paraformaldehyde followed by an ice-cold MeOH wash for 5 min on ice. Samples were incubated with 80% (v/v) formamide at 80°C for 10 min followed by a PBS wash for 1 min. Cells were next incubated in IF blocking buffer (2.5% BSA, 0.4% Triton in PBS) at room temperature for 30 min. The hybridization of oligo probes to the Dux repeat region was achieved in 5 min at room temperature in a solution containing 50% (v/v) formamide, 2x SSC, and 1 µM oligo probes followed by a wash of 40% (v/v) formamide at room temperature for 3 min. Samples were next incubated with primary antibodies against Nucleolin and GFP in IF blocking buffer for 30 min at room temperature followed by incubation with secondary antibodies (Alexa Fluor, 1:3000) for 30 min at room temperature, and nuclei counterstained with DAPI. Fluorescence images
were taken on an inverted microscope with 100x (N.A. = 1.4) oil objective using X-Cite LED illumination in DAPI, GFP, RFP, and Cy5 channels, utilizing an Andor Zyla sCMOS camera for detection (Nikon Imaging Center at UCSF). The image stacks were acquired with a spinning disk confocal module with 0.25 μm in z-step. For scoring of Dux localization, all images were analyzed in Fiji, identifying individual z stacks containing one locus, then comparing across Nucleolim or DAPI channels to identify overlap or co-localization with nucleoli, edges of nuclei (lamina), or nucleoplasm. Positive GFP signal was used to identify 2C-like cells and score localization for each Dux locus in each ES versus 2C-like cell. Data were collected and quantified from two separate FISH experiments across multiple individual chambers.

**Chromatin Immunoprecipitation**

Low-cell histone chromatin immunoprecipitation (ChIP) was performed starting from 5x10⁵ Lissamine-positive ES cells purified by FACs 48 h after ASO nucleofection. Cells were fixed in suspension for 8 min in 1% formaldehyde, washed and the pellets snap frozen. Chromatin was prepared utilizing reagents and the protocol from Low Cell Number ChIP Kit (Diagenode). Cell pellets were thawed, lysed and sonicated using a Covaris S220 focused-ultrasonicator for 8 cycles with the following conditions: 60 s per cycle, duty cycle 2%, 200 cycles per burst, intensity 3. IPs and washes were performed according to kit protocols, utilizing 1x10⁶ cells per IP. For transcription factor ChIP, chromatin fixation and preparation of nuclear extracts were performed as described previously (Stock et al., 2007). ES cells were fixed and washed as above, then incubated in swelling buffer (25mM HEPES pH 7.9, 1.5mM MgCl₂, 10mM KCl, 0.1% NP-40) for 30 min on ice with frequent vortexing and passed 5 times through a 25G needle to shear cell membranes. Nuclei were pelleted by centrifugation then resuspended in sonication buffer (50mM HEPES pH 7.9, 140mM NaCl, 1mM EDTA, 1% Triton X-100, 0.1% Sodium deoxycholate, 0.1% SDS) and incubated on ice for 30 min before proceeding to sonication. Swelling and sonication buffers were supplemented with the following inhibitors fresh each time: 1x Halt Protease Inhibitor Cocktail (Thermo Fisher Scientific), 5mM sodium fluoride, 5mM sodium butyrate, 1mM PMSF, along with 10mM N-ethylmaleimide (NEM) (Thompson et al., 2015). Chromatin samples were sonicated using a Covaris S220 focused-ultrasonicator for 14 cycles with the following conditions: 60 s per cycle, duty cycle 5%, 200 cycles per burst, intensity 4. Resultant DNA fragments were of an average length of 200-500bp. Immunoprecipitations were carried out overnight at 4°C with 20 μg chromatin, 3 μg Kap1/control antibody or 0.4 μg Nucleolin antibody, and 30 μL Protein A/G Dynabeads per IP. ChIP washes were performed the following day as follows: twice in low-salt buffer (20mM Tris-HCl pH 8, 0.1% SDS, 1% Triton X-100, 2mM EDTA, 150mM NaCl), once in high-salt buffer (as above but with 500mM NaCl), once in LiCl wash buffer (10mM Tris-HCl pH 8, 250mM LiCl, 0.5% NP-40, 0.5% sodium deoxycholate, 1mM EDTA), once in TE buffer, all for 7 min at 4°C for each wash. Finally, Dynabeads containing immune complexes along with input samples were resuspended in elution buffer (50mM sodium bicarbonate, 50mM Tris-HCl pH 8, 1% SDS, 1mM EDTA) containing 100μg/mL RNase A (Thermo Fisher Scientific) and incubated for 30 min at 37°C. Crosslinks were next reversed by addition of 0.5mg/mL Proteinase K (Ambion) and incubation at 55°C shaking for at least 2h. DNA was purified with Qiaquick PCR purification columns (QIAGEN) and ChIP enrichments analyzed by qPCR. ChIP enrichments were calculated as % input and normalized to enrichment at the negative control region, intergenic chr11, int-chr11 for each experiment. Primer sequences are listed in Table S5.

**Western blotting**

Whole cell extracts were prepared from ES cells in ice-cold RIPA buffer containing protease-inhibitors (as for ChIP but minus NEM). Proteins were separated on 4%–15% Mini-Protean TGX SDS-PAGE gels (BioRad) and transferred to PVDF membranes. Blocking was performed for 45 min in 5% milk/PBS-T buffer followed by incubation overnight with primary antibodies at 4°C. The following day, membranes were incubated with the appropriate anti-mouse/rabbit secondary antibodies conjugated to HRP (Jackson) for 1h, and proteins were detected by ECL or ECL Plus reagent and autoradiography.

**Co-immunoprecipitation**

Co-immunoprecipitation (Co-IP) assays were performed on nuclear extracts. Cell pellets were resuspended in hypotonic buffer (10mM HEPES pH 7.9, 5mM MgCl₂, 0.25M Sucrose, 0.1% NP40, protease inhibitors). Cell suspensions were passed through an 18G needle, incubated for 10 min on ice and centrifuged. The resulting pellet was resuspended in nuclear extraction buffer (10mM HEPES pH 7.9, 1mM MgCl₂, 0.1mM EDTA, 0.5M NaCl, 0.5% Triton X-100, 25% glycerol, protease inhibitors). The nuclear suspension was passed through an 18 G needle, incubated on ice for 30 min, sonicated in a Bioruptor (Diagenode) for 5 min at 30 s on, 30 s off and centrifuged at 4000 rpm for 10 min in a benchtop centrifuge at 4°C. Supernatants containing nuclear extracts were quantified using a Pierce BCA protein assay kit (Thermo Fisher Scientific), and 100 μg nuclear extracts were immunoprecipitated using the indicated primary antibodies, rotating overnight at 4°C. The following day, immune complexes were bound to Protein A/G Dynabeads (Thermo Fisher Scientific), washed 3 times in nuclear extraction buffer at 4°C 10 min, boiled in LDS Sample Buffer (Thermo Fisher Scientific) and used for western blotting analysis.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

**RNA-seq analysis**

RNA-seq reads were quality-checked, trimmed, and aligned to reference genome mm9 using Tophat2 (Kim et al., 2013) and default settings, apart from for TE analysis where reads were aligned using Tophat2 setting g –1 to map each TE to one random location.
Reads were counted for each gene or TE family using the Subread package, FeatureCounts (Liao et al., 2013), and data normalized utilizing DESeq2 in R/Bioconductor (Love et al., 2014). All other RNA-seq analyses and statistics were performed in R/Bioconductor utilizing custom R scripts. For transcriptional analysis of Dux targets, these were identified as the set of genes detected in our dataset that are most highly induced by Dux overexpression in ES cells (Hendrickson et al., 2017), which we verified to be directly bound by Dux in over 70% of cases (Hendrickson et al., 2017).

**Statistical analysis**
All statistical analyses were performed with Graphpad Prism 7.0 software, except genome-wide data analyses which were performed in R/Bioconductor. Details of individual tests are outlined within each figure legend, including number and type of replication performed (n) and the reported error either as standard deviation (s.d) or standard error of the mean (s.e.m). All statistics are * p < 0.05, ** p < 0.01, *** p < 0.001, **** p < 0.0001, #p = 0.056, and calculated by two-tailed Student’s t test unless specifically noted otherwise in the legend. Welch’s correction was applied to t tests whenever the variance was unequal between conditions.

**DATA AND SOFTWARE AVAILABILITY**
The accession number for the sequencing data reported in this paper is GEO: GSE100939.
Figure S1. Knockdown of LINE1 RNA by siRNAs, Related to Figure 1
(A) Representative immunofluorescence staining showing cytoplasmic LINE1 Orf1p in ESCs. Scale bar, 10 μm.
(B) Representative FACS plots showing percentage Lissamine-positive ESCs 48h after nucleofection with Lissaminated RC or LINE1 ASOs
(C) Representative images and quantification of LINE1 RNA FISH in ESCs 48h after transfection with either Control or two independent siRNAs against LINE1. Data are mean ± s.d for the indicated number (n) of cells. Scale bar, 10 μm.
(D) Colony formation assays in ESCs following transfection with control (RC) or LINE1 siRNAs. Data are mean ± s.e.m, n = 3 biological replicates.
(E) Growth curves of ESCs following nucleofection with RC or LINE1 ASOs, shown as mean ± s.e.m, n = 3 biological replicates.
(F) Cell-cycle analysis 48h after RC/LINE1 ASO nucleofection, with representative FACS plots and quantification of n = 3 biological replicates. **p < 0.01, Chi-square test.
(G) Cell viability analysis in Lissamine-positive ESCs, measured by Sytox-blue staining. Dead cells (permeabilized with 4% paraformaldehyde for 30 min) were used as a positive control for Sytox-blue staining. Shown are representative FACS plots plus quantification from n = 3 independent experiments, +/− s.e.m.
(H) Western blot (left) and immunofluorescence (right) analysis showing no overall change to Oct4/Nanog protein levels in ESCs 48h after LINE1 KD. Scale, 10 μm.
Figure S2. Induction of a 2C-like State upon LINE1 Knockdown, Related to Figure 2
(A) LINE1 KD ESCs show a distinct gene expression profile, as evidenced by unsupervised hierarchical clustering of samples using Pearson Correlation coefficients.
(B) Boxplot showing log$_2$FC in LINE1- versus RC ASO-nucleofected samples of the indicated number (n) of genes. Note that categories are non-inclusive, i.e., L1 < 100kb refers to genes < 100kb but > 10kb away from a full-length LINE1 element. P values are calculated by two-sided Wilcoxon rank sum test.

(C) Expression analysis of pluripotency genes and markers from all 3 germ layers, taken from RNA-seq data following LINE1 KD. Data are mean ± s.e.m, n = 3 biological replicates in each condition. Endo, endoderm, Meso, mesoderm, Ecto, ectoderm, plurip, pluripotency.

(D) qRT-PCR validation of germ layer marker expression following ASO treatment. Data are mean ± s.e.m, n = 4 biological replicates

(E) Venn diagrams indicating the overlap between top upregulated genes with LINE1 KD (log$_2$ fold-change > 1.0) and the indicated datasets. Datasets are as follows, ‘Up in 2C-like cells’: significantly upregulated genes in 2C-tdTomato+ cells (log$_2$ fold-change > 1.0), ‘Up with CAF-1 depletion’: overlap of top 200 genes upregulated upon knockdown of both subunits of CAF-1 (p60, p150) (Ishiuchi et al., 2015), ‘Up in Zscan4+/MERVL+ cells’: genes upregulated in Zscan4/MERVL-double-positive cells (log$_2$ fold-change > 1.0) (Eckersley-Maslin et al., 2016). P values were calculated by Fisher’s exact test, assuming a universe of 18,000 expressed genes.

(F) 2C gene expression 48h after transfection of ESCs with Control or LINE1 siRNAs. Data are mean ± s.e.m, n = 3 biological replicates.

(G) LINE1 retrotransposition assay, where retrotransposition mediated by a codon-optimized LINE1 transgene (ORFeus) (An et al., 2006; Newkirk et al., 2017) results in GFP expression. Treatment with LINE1 reverse transcriptase inhibitors, Stavudine (d4T, 20 μM) or Tenofovir disoproxil fumarate (TDF, 4 μM) inhibits LINE1 retrotransposition. Data are number of GFP-positive cells per 300,000 live cells, representative of 2 independent experiments.

(H) Colony formation assay of ESCs treated with reverse transcriptase inhibitors, indicating that inhibition of LINE1 retrotransposition does not affect self-renewal. Data are mean ± s.e.m, n = 4 biological replicates.

(I) 2C gene expression after 48h treatment with reverse transcriptase inhibitors, data are mean ± s.d, n = 2 biological replicates

(J) Expression of 2C genes or TEs 48h after LINE1 KD in MEF cells. Data are mean ± s.e.m, n = 3 biological replicates.

(K) Validation of clonal 2C-GFP reporter ESCs, by qRT-PCR analysis performed for Dux and 2C genes in sorted 2C-GFP positive versus negative cells. Data are mean ± s.d n = 2 biological replicates. n.s., not significant, n.d., not detected.
Figure S3. Dux Knockdown Rescues 2C/MERVL Upregulation Induced by LINE1 Knockdown, Related to Figure 3

(A) qRT-PCR analysis showing Dux and 2C upregulation in ESCs cultured in N2B27/2i/LIF conditions upon LINE1 KD. Data are mean ± s.e.m, n = 3 technical replicates, representative of 3 experiments.

(B) Colony formation assay performed in nucleofected ESCs cultured in N2B27/2i/LIF conditions. Data are mean ± s.e.m, n = 3 biological replicates.

(C) Expression analysis of previously reported 2C/MERVL repressors following LINE1 KD, taken from RNA-seq data. Data are mean ± s.e.m, n = 3 biological replicates in each condition.

(D–F) ChIP assays for the histone marks (D) H3K9me2, (E) H3K9me3, or (F) H3K27me3 performed 48h after nucleofection with ASOs and sorting for Lissamine+ cells. Data are mean ± s.e.m n = 2 technical replicates, representative of at least 2 independent experiments.

(G) Principal Component Analysis (PCA) plot for all genes across all samples, confirming ASO/siRNA RNA-seq samples have distinct gene expression profiles.

(H) Boxplot analysis of Dux-target genes following LINE1 KD and co-transfection of either siControl or siDux (n = 100 genes expressed in our data, from the original list of 200 most-upregulated genes). P value is determined by two-sided Wilcoxon rank sum test.

(I) Graph of MERVL repeat expression in RNA-seq samples, showing that simultaneous Dux knockdown rescues MERVL upregulation upon LINE1 KD.
**Figure S4. Effects of LINE1 Knockdown Are Not p53-Mediated and Include Reductions in Ribosomal Protein Gene Expression, Related to Figure 4**

(A) GSEA analysis of RNA-seq data in LINE1 versus RC nucleofected ESCs, showing a preferential upregulation of p53 targets upon LINE1 KD.

(B) qRT-PCR analysis of 2C/MERVL gene expression in p53
- /- ESCs analyzed 24-48h after nucleofection with RC/LINE1 ASOs. Data are mean ± s.e.m, n = 3 biological replicates, normalized to RC ASO 24h. #p < 0.01, LINE1 ASO 24h versus RC ASO 48h.

(C) Colony formation assay in p53
- /- ESCs, showing that LINE1 KD still inhibits self-renewal. Data are mean ± s.e.m, n = 3 biological replicates.

(D) GSEA analysis as in (A), demonstrating a preferential downregulation of ribosomal genes upon LINE1 KD.

(E) Heatmap analysis of n = 86 ribosomal genes in RC or LINE1 ASO RNA-seq samples, with samples grouped by unsupervised hierarchical clustering.

(F) Boxplot depicting downregulation of Rpl/Rps genes compared to all expressed genes upon LINE1 KD. P value calculated by two-sided Wilcoxon rank sum test.

(G) Nascent RNA expression of the indicated ribosomal RNA and protein-coding genes following 8h treatment with CX-5461. Data are mean ± s.e.m, n = 2 biological replicates.

(H) Colony formation assay performed in ESCs plated after 8h treatment with CX-5461. Data are mean ± s.e.m, n = 3 biological replicates.

(I) Growth curves of ESCs with or without 8h CX-5461 treatment. Data are mean ± s.e.m, n = 3 biological replicates.
Figure S5. LINE1 and Nucleolin Repress Dux and the 2C/MERVL Program, Related to Figure 5

(A) Analysis of the amounts of the indicated RNAs pulled down by LINE1 or Malat1 ChIRP probes, expressed as RNA-ChIRP % input. Data are mean ± s.e.m, n = 3 independent experiments.

(B) ChIRP-qPCR analysis for ChIRP positive control, Malat1, confirming that Malat1 RNA, but not LINE1 RNA, is highly associated with Malat1 and Neat1 genomic loci. Data are mean ± s.e.m, n = 3 independent experiments.

(C) ChIRP-qPCR analysis with control or LINE1 primers, indicating that LINE1 RNA is found associated with LINE1 DNA. Data are mean ± s.e.m, n = 3 independent experiments.

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(D) RIP assay performed with or without DNase treatment before immunoprecipitations, indicating that the Nucleolin (Ncl)-LINE1 RNA association is DNase-independent. Data are representative of 2 independent experiments and shown as mean ± s.e.m, n = 3 technical replicates.

(E) Nuclear enrichment of the indicated RNAs taken form RIP input expression data. RNA expression is shown relative to cytoplasmic RNA, Gapdh. Data are mean ± s.e.m, n = 3 independent experiments.

(F-H) qRT-PCR showing knockdown of candidate 2C/Dux repressors alongside Dux and 2C gene expression. Data are representative of two independent experiments and are mean ± s.e.m., n = 3 technical replicates. (G-H) Confirmation of Ncl knockdown, (G) at the RNA level, by qRT-PCR, and (H) at the protein level, by western blotting. Images in (H) were spliced together to remove unrelated lanes.

(I) Expression of MERVL repeats in RNA-seq data upon knockdown of Ncl, showing mean ± s.e.m, n = 3 biological replicates.

(J) GSEA analysis of RNA-seq data upon Ncl knockdown, showing a preferential downregulation of ribosomal genes.

(K) Colony formation assays performed in ESCs plated 24h after transfection with the indicated siRNAs. Data are mean ± s.e.m, n = 3 biological replicates.

(L) qRT-PCR analysis performed in ESCs 48h after transfection with the indicated siRNAs, showing that the induction of 2C genes upon Ncl KD depends on Dux expression. Data are mean ± s.e.m, n = 3 technical replicates, representative of three experiments.
Figure S6. LINE1, Nucleolin, and Kap1 Coordinately Regulate rDNA and Dux, Localized at Peri-nucleolar Regions in ESCs but Not 2C-like Cells, Related to Figure 6

(A) Scatter graph comparing gene expression changes upon LINE1 KD (this study) with changes upon Kap1 deletion (Ecco et al., 2016), with the indicated Spearman’s correlation coefficient. Dux targets are shown in red.

(B) Nascent RNA expression for ribosomal protein coding genes or rRNA upon KD of Ncl, LINE1 or Kap1. Data are mean ± s.e.m, n = 3 biological replicates.

(C and D) qRT-PCR confirmation that KD of LINE1 by (C) ASOs or (D) siRNAs does not alter Ncl RNA expression. Data are mean ± s.e.m, n = 3 biological replicates, n.s., not significant.

(E) Western blot analysis of protein extracts from ESCs treated with RC or LINE1 ASOs, confirming that Kap1 and Ncl protein levels are unaffected.

(F) Co-IP assays performed in ES nuclear extracts 48h after nucleofection with RC/LINE1 ASOs, demonstrating that Ncl and Kap1 still interact in the absence of LINE1 RNA.

(G) Immunofluorescence analysis of Ncl and Kap1 proteins in 2C-GFP cells, showing no change between ES and 2C-like states. Green arrows in (G-H) denote 2C-GFP positive cells. Scale, 10 μm.

(H) Representative immunofluorescence images and quantification of the number of nucleoli per cell in ES versus 2C-like cells. 2C-like cells were identified by positive 2C-GFP signal and absence of Oct4. ****p < 0.0001, Chi-square test. Scale, 10 μm, n = number of cells.

(I) Representative co-immunofluorescence/DNA-FISH images and example scoring for each Dux localization pattern. Each locus was scored as associated with the lamina (L), nucleolus (N), or nucleoplasm (P). Nucleoli were stained with anti-Ncl, and 2C-GFP cells with anti-GFP antibodies. Scale, 2 μm.

(J) Example images of Dux localization in unsorted 2C-GFP cells, showing identification of 2C-like cells by GFP-staining and nucleoli by anti-Ncl. Dux loci are indicated by white arrows, and a 2C-like cell by a green arrow. Scale, 10 μm.

(K) Quantification of the proportion of ES or 2C-like cells containing at least one nucleoplasmic Dux locus versus no nucleoplasmic loci. Statistics are calculated by Chi-square test for the indicated number (n) of cells.
Figure S7. LINE1 Is Essential for Pre-implantation Development, Related to Figure 7

(A) Developmental progression to blastocyst of the indicated Uninjected, Control ASO or LINE1 ASO injected embryos. Data shown are the percent of all embryos from 3 independent experiments at the indicated developmental stage, n = number of embryos.

(B) Analysis of chromatin accessibility in late 2C embryos following zygote ASO microinjections by DNase-TUNEL experiments. Intensities are calculated per nucleus and data shown as mean ± s.d for the number (n) of nuclei, in data combined from 2 independent experiments for each assay. Nuclei are counterstained by Hoechst. Scale, 20 μm.

(C) Immunofluorescence for global repressive chromatin marks H3K9me2/3, quantified as in (C). Scale, 20 μm.

(D) Rates of developmental progression to blastocyst in embryos cultured from the 2C stage with or without CX-5461. P values in (A-B and I) are calculated by Chi-square test, n = number of embryos.

(E) PCA plot of all genes across all samples, showing that samples are separated first by developmental stage, and second by ASO microinjection, according to global gene expression profiles.

(F) FPKM-normalized RNA-seq data from Wu et al. (2016) demonstrating the 6 separate gene clusters identified by K-means clustering. Cluster 4 contains early-2C genes such as Dux itself.

(G) RNA-seq normalized, log2-transformed expression for late 2C genes (which peak in expression in late-2C/4C stage), Ddit4l and Dub1a, in the indicated embryo samples, n = 3 independent batches of embryos were sequenced per condition. *FDR < 0.05, ****FDR < 0.0001, Toptable analysis.

(H) GSEA analysis showing that ribosomal genes are preferentially downregulated in 4C embryos after LINE1 KD.

(I) Developmental progression rates to blastocyst in embryos following late-2C microinjection with ASOs, in embryos collected from 2 independent experiments, n = number of embryos.

(J) qRT-PCR expression analysis of the indicated 2C genes in 4C embryos after late-2C ASO microinjections. Data are mean+/− s.e.m, n = 3 technical replicates, for the indicated independent batches of embryos.

(K) Expression data from Wu et al. (2016) illustrating the expression levels of Kap1/Nci during pre-implantation development. Data are FPKM, normalized to expression level in the zygote.