Intrinsic retroviral reactivation in human preimplantation embryos and pluripotent cells

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Endogenous retroviruses (ERVs) are remnants of ancient retroviral infections, and comprise nearly 8% of the human genome. The most recently acquired human ERV is HERVK(HML-2), which repeatedly infected the primate lineage both before and after the divergence of the human and chimpanzee common ancestor. Unlike most other human ERVs, HERVK retained multiple copies of intact open reading frames encoding retroviral proteins. However, HERVK is transcriptionally silenced by the host, with the exception of in certain pathological contexts such as germ-cell tumours, melanoma or human immunodeficiency virus (HIV) infection. Here we demonstrate that DNA hypomethylation at long terminal repeat elements representing the most recent genomic integration, together with transactivation by OCT4 (also known as POU5F1), synergistically facilitate HERVK expression. Consequently, HERVK is transcribed during normal human embryogenesis, beginning with embryonic genome activation at the eight-cell stage, continuing through the emergence of epiblast cells in preimplantation blastocysts, and ceasing during human embryonic stem cell derivation from blastocyst outgrowths. Remarkably, we detected HERVK viral-like particles and Gag proteins in human blastocysts, indicating that early human development proceeds in the presence of retroviral products. We further show that overexpression of one such product, the HERVK accessory protein Rec, in a pluripotent cell line is sufficient to increase IFITM1 levels on the cell surface and inhibit viral infection, suggesting at least one mechanism through which HERVK can induce viral restriction pathways in early embryonic cells. Moreover, Rec directly binds a subset of cellular RNAs and modulates their ribosome occupancy, indicating that complex interactions between retroviral proteins and host factors can fine-tune pathways of early human development.

Given the substantial contribution of transposons to the human genome and their emerging roles in shaping host regulatory networks, understanding the dynamic expression and function of these genomic elements is important for dissecting both human- and primate-specific aspects of gene regulation and development. We used published single-cell RNA-sequencing (RNA-seq) data sets to analyse the expression of major transposon classes at various stages of human preimplantation embryogenesis, a developmental period associated with dynamic changes in DNA methylation and transposon expression. This analysis revealed two major clusters, one primarily consisting of repeats that begin to be transcribed at the onset of embryonic genome activation (EGA), which in humans occurs around the eight-cell stage, and a second cluster of repeats, whose transcripts can be detected in the embryo before EGA, indicating maternal deposition (Extended Data Fig. 1a). Within each cluster, more discrete stage-specific changes in repeat transcription could be observed, such that analysis of the repetitive transcriptome alone was able to distinguish pre- and post-EGA cells, as well as eight-cell/morula cells from blastocyst cells (Extended Data Fig. 1a). For example, HERVK and its regulatory element, long terminal repeat (LTR)SHS, were both induced in eight-cell stage embryos, morulae, and continued to be expressed in epiblast cells of blastocysts (Fig. 1a–c and Extended Data Fig. 1a). We further observed that although HERVK was expressed in blastocyst outgrowths (passage 0 human embryonic stem (ES) cells), it was downregulated by passage 10 (Fig. 1d). In contrast, transcripts of another HERV, HERVH, and of its regulatory element LTR7, were detected before EGA and throughout preimplantation development, including in all blastocyst lineages and human ES cells (Extended Data Fig. 1a–c).

Recent studies have reported conditions for capturing a human naive pluripotent state in vitro, and we used RNA-seq to analyse the repetitive transcriptome of ELF1, a cell line derived from an eight-cell-stage human embryo under naive culture conditions, and compared it to the repeat expression in ELF1 cells matured in vitro into a primed state. Surprisingly, although many transposon classes (for example, HERVK and LINE1-HS) were highly expressed in both cell states, only a few showed differential levels between the two (Fig. 1e). In particular, transcripts corresponding to HERVK proviruses and their regulatory elements, LTR5HS (but not the older LTR5a or LTR5b), were among the most strongly induced in naive versus primed ELF1 cells (Fig. 1e and Extended Data Fig. 1d). Similar results were obtained by analysing available transcriptomes of primed H1 human ES cells and naive 3iL cells derived from them, as well as of primed H9 human ES cells and those ‘reset’ to the naive state by NANOG and KLF2 transgene expression (Fig. 1e). Therefore, naive-state-specific upregulation of HERVK is consistent across multiple genetic backgrounds, derivation methods or culture conditions.

From an evolutionary perspective, HERVK is especially interesting, as it is the most recently acquired HERV from which multiple insertions have retained protein-coding potential. While HERVK is present in all Old World primates, nearly a third of its proviruses in the human genome represent human-specific insertions, and 48% of those show polymorphisms in the human population, suggesting that HERVK was active within the last 200,000 years. All human-specific and human-polymorphic HERVK elements are regulated by a specific LTR subgroup, LTR5HS, whereas insertions representing older integrations typically have regulatory elements of the LTR5a or LTR5b subtype (Extended Data Fig. 2a). Interestingly, during human preimplantation development and in the naive state, transcripts originating from LTR5HS, but not LTR5a or LTR5b, are preferentially expressed.

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(Fig. 1e), and we observed an upregulation of human-specific provirus sequences compared to evolutionarily older elements (Fig. 2a). We hypothesized that this differential regulation can be explained by cis-regulatory change in LTR5HS. Indeed, sequence analysis uncovered an OCT4 motif at position 693–699 base pairs (bp) of LTR5HS, which was conserved across diverse LTR5HS sequences, but not present in LTR5a/LTR5b, despite their overall high (~88%) sequence homology with LTR5HS (Fig. 2b and Extended Data Fig. 2a). To test whether OCT4 binding contributes to the transcriptional activation of LTR5HS, we used pluri-potent NCCIT human embryonic carcinoma (EC) cells, which express OCT4, but, in contrast to human ES cells, are permissive for HERVK expression.19 (Extended Data Fig. 2b–d). Chromatin immunoprecipitation with quantitative polymerase chain reaction (ChIP-qPCR) analysis of human EC cells showed preferential occupancy of OCT4, p300 and histone marks of active chromatin at LTR5HS elements, as compared to LTR5a/LTR5b (Fig. 2c). In contrast, we did not detect OCT4 or p300 binding at LTR5HS in primed human ES cells (Extended Data Fig. 2f). Consistent with a functional role in HERVK activation, knockdown of OCT4 or SOX2, but not of NANOG, led to a significant decrease in viral transcripts in human EC cells (Extended Data Fig. 2e and Extended Data Fig. 3a). Furthermore, the activity of transcriptional reporters driven by LTR5HS was impaired by mutations in the OCT4 motif (Fig. 2d and Extended Data Fig. 3b).

The aforementioned observations are consistent with transactivation by OCT4 being a driver of LTR5HS regulatory activity, but do not explain the differential transcriptional status of HERVK in primed versus naive human ES cells and in human EC cells, as all three express OCT4. We hypothesized that DNA methylation may contribute an additional layer of regulation, and indeed we observed HERVK hypomethylation of solo and proviral LTR5HS (but not the Gag open reading frame (ORF)) in human EC cells and naive ES cells, as compared to primed human ES cells and human induced pluri-potent stem cells (iPSCs) (Fig. 2e and Extended Data Fig. 3c, d). Strong and preferential demethylation of LTR5HS was also observed in recently published DNA methylation maps from human preimplantation embryos, whereas HERVK coding sequences remained more highly methylated.11 Importantly, treatment of primed human ES cells with a

DNA methylation inhibitor, 5-aza-2′-deoxycytidine, for 24 h induced HERVK transcription, with 8–12-fold upregulation of an early transcript encoding an accessory protein, Rec (Fig. 2f). In addition, inhibition of DNA methylation, together with overexpression of OCT4 and SOX2, jointly facilitated HERVK transcription in HEK293 cells (Fig. 2g and Extended Data Fig. 3e), indicating that DNA hypomethylation and transactivation by OCT4 synergistically promote HERVK expression.

A defining characteristic of HERVK is that multiple proviruses have retained ORFs encoding full-length retroviral proteins. Consequently, HERVK reactivation in pathological conditions has been associated with the presence of HERVK proteins3–7, prompting us to examine whether retroviral proteins are also present in human embryos. We used a well-characterized monoclonal antibody recognizing the HERVK Gag precursor and its proteolytically processed form Capsid, which detects cytoplasmic signal with a characteristic punctate pattern in human EC cells and in a subset of naive ELF1 cells, but shows no staining in primed human ES cells and loss of signal in human EC cells after gag short interfering RNA (siRNA) knockdown (Extended Data Fig. 4a–d). In human blastocysts, Gag/Capsid staining was also detected in dense cytoplasmic puncta resembling those seen in human EC cells and naive ELF1 cells (Fig. 3a and Extended Data Fig. 4a, d, e), with all analysed blastocysts (n = 19/19) showing a robust signal. Several HERVK-positive human EC cell lines have been shown to produce viral-like particles (VLPs).20 Remarkably, heavy metal staining transmission electron microscopy (TEM) of blastocysts revealed the presence of cytoplasmic, electron-dense particles of approximately 100 nm in diameter—the reported size of reconstructed HERVK VLPs—with electron-lucent cores21,22 (Fig. 3b). Additionally, human blastocyst cells also contained cytosolic vesicles enclosing 50 or more more highly electron-dense particles of approximately 75 nm in size, which resembled the immature VLPs also seen in human EC cells (Fig. 3c and Extended Data Fig. 5a). The presence of HERVK-derived particles in human blastocysts was further supported by immuno-gold TEM staining of blastocysts, which detected VLPs (or vesicles with multiple VLPs) labelled by Gag/Capsid antibodies either within embryonic cells or on the cell surface, similar to those seen in immuno-gold TEM staining of human EC cells (Fig. 3d, e and

Figure 1 | Transcriptional reactivation of HERVK in human preimplantation embryos and naive human ES cells. a, Schematic of human preimplantation development. b, HERVK expression in single cells of human embryos at indicated stages. Solid line indicates mean. Oocyte (n = 3), zygote (n = 3), 2-cell (n = 6), 4-cell (n = 11), 8-cell (n = 19), morula (n = 16). b–d, Data are taken from ref. 10. *P value < 0.05, non-paired Wilcoxon test. RPKM, reads per kilobase per million. c, HERVK expression in single cells of human blastocysts, grouped by lineage. Solid line indicates mean. Trophoderm (TE; n = 18), primitive endoderm (PE; n = 7), epiblast (EPI; n = 5). d, HERVK expression in single cells of blastocyst outgrowths (passage (p)0) or human ES cells at passage (p)10. Solid line indicates mean. p0 (n = 8), p10 (n = 26). e, Analysis of the repetitive transcriptionomes of three, genetically matched naive/primed human ES cell pairs. Left, naive/primed ELF1 human ES cells (data from this study) (n = 3 biological replicates for both conditions). Middle, 3i/L/C primed H1 human ES cells (data are taken from ref. 12) (n = 3 biological replicates for both conditions). Right, naive/primed H9 human ES cells (data are taken from ref. 15) (n = 3 biological replicates for both conditions). Significant repeats indicated in red at false discovery rate (FDR) < 0.05. DESeq. hESC, human ES cells.
Extensive data Fig. 5b); control blastocyst staining showed no signal from secondary antibody (Extended Data Fig. 5c). Altogether, these data demonstrate that human preimplantation development proceeds in the presence of retroviral proteins and VLPs (summarized in Extended Data Fig. 5d).

Recent studies highlight the ability of TEs to contribute regulatory sequences to mammalian genomes\(^2\),\(^3\),\(^4\). For example, MERV-L elements in mouse have been reported to function as alternative promoters, driving expression of many two-cell stage-specific chimaeric transcripts\(^5\). However, we did not detect robust evidence for HERVK-associated chimaeric transcription (Extended Data Fig. 6a, b and Supplementary Table 1), suggesting that LTR5HS is unlikely to contribute promoter activity to nearby host genes. Alternatively, LTR sequences derived from ERVs could be co-opted to act as long-distance enhancers for the host\(^24\). In agreement with such a possibility, LTR5HS elements were marked by p300 and H3K27ac (Fig. 2c), while genes located in their vicinity showed a strong bias for naive-state-enriched expression, regardless of their upstream or downstream position in relation to the LTR5HS (Extended Data Fig. 6c–e). However, we cannot rule out that this result could be a consequence of preferential HERVK integration near genes active in the naive state.

HERVK encodes a small accessory protein, Rec, homologous to HIV Rev, which binds to and promotes nuclear export and translation of viral RNAs\(^6\). rec, an early viral transcript derived through alternative splicing of the env gene (Extended Data Fig. 2a), is expressed in naive ES cells and human blastocysts, and is rapidly induced in primed human ES cells exposed to 5-aza-2′-deoxycytidine (Extended Data Fig. 7a and 7f). We hypothesized that Rec-mediated nuclear export of viral RNAs into the cytoplasm might ultimately lead to the induction of innate antiviral responses, which typically rely on cytosolic detection of viral RNA/DNA and protein. We noted a striking induction of messenger RNA encoding an interferon-induced viral restriction factor IFITM1 (ref. 26; also known as FRAGILIS2) in human epiblast messenger RNA encoding an interferon-induced viral restriction factor IFITM1 (ref. 26; also known as FRAGILIS2) in human epiblast cells\(^26\), as well as upregulation of IFITM1 transcripts and surface protein levels in human naive versus primed human ES cells (Extended Data Fig. 7b, c, f and Supplementary Table 6). Furthermore, expression of a rec transgene in human EC cells was sufficient to elevate surface-localized IFITM1 protein levels (Fig. 4a). This was at least in part mediated through an effect on IFITM1 mRNA transcription or stability, as Rec overexpression or knockdown had, respectively, increased or decreased IFITM1 mRNA levels (Extended Data Fig. 7d).

Of note, although the minimal components of the JAK/STAT interferon pathway are present in human EC cells, many other interferon-induced genes are not upregulated or expressed, indicating that HERVK triggers a precise antiviral response in host cells (Supplementary Table 2). To test whether HERVK expression provides viral resistance, we infected control wild-type human EC cells, control human EC cells expressing a green fluorescent protein (GFP) transgene, or two independent clonal Rec human EC cell lines (Rec-hECCs) with influenza H1N1 (PR8) virus. Interestingly, the Rec-hECs exhibited substantially attenuated infec-
iCLIP-seq in human EC cells expressing Flag–GFP (GFP) or Flag–GFP-tagged rec transgene (Extended Data Fig. 8a, b). We did not detect associated RNA in the control Flag–eGFP purifications, indicating low nonspecific RNA recovery of our assay (Extended Data Fig. 8b). In contrast, parallel Rec purifications from two Flag–GFP Rec expressing clones yielded ultraviolet-crosslinked RNAs, sequencing of which demonstrated that in vivo, Rec robustly binds LTR5HS, but only in the region previously defined as containing the highly structured Rec-responsive element.25–28 (Fig. 4c and Extended Data Fig. 8b, c). In addition, Rec directly interacts with ~1,600 host mRNAs, preferentially in their 3′ untranslated regions (UTRs), a positional preference analogous to that observed in the viral RNA (Fig. 4d, e, Extended Data Fig. 9a and Supplementary Table 3). We did not detect specific RNA sequence motifs enriched at Rec-bound sites; however, multiple examined Rec iCLIP targets were predicted to fold into stable secondary structures (Extended Data Fig. 9b). This is reminiscent of the interaction of Rec with its HERVKLTR response element, which is mediated by RNA secondary structure, rather than a discrete specific binding site.28 We also observed Rec association with mRNAs encoding surface receptor molecules and ligands (for example, FGFR1, FGFR3, IFITM1, KLRG2, Igf1R, Fzd7, Gdf3) and chromatin regulators (for example, Dnmt1, Chd4) (Extended Data Fig. 9a and Supplementary Table 3).

Given that Rec binding to viral RNAs promotes their nuclear export and translation, we next examined if endogenous mRNAs bound by Rec are also more efficiently targeted to ribosomes.22–23. Ribosome profiling of Rec-hECCs, in comparison to wild-type human EC cells, revealed both increases and decreases in ribosomal occupancy, with differential enrichment of 941 mRNAs, of which 134 were also Rec iCLIP targets, representing a significant overlap (P value < 0.05, hypergeometric test) (Fig. 4f and Supplementary Table 5). Notably, mRNAs bound by Rec in 3′ UTRs or coding sequences were more likely to be upregulated in their ribosomal occupancy than expected by chance (P value < 0.05, hypergeometric test), but we did not observe such enrichment for mRNAs bound in their 5′ UTRs. We also noticed that several Rec-bound transcripts (for example, Rpl22, Rpl31, Rps13, Rps20, Eif4g1) encoding ribosome components and translation regulators had increased occupancy in Rec-hECCs,
potentially contributing to additional indirect translational effects of Rec overexpression (Fig. 4e, f and Supplementary Table 5).

Altogether, our results demonstrate that early human development is accompanied by the stage-specific transcriptional activation of HERVK, translation of its ORFs, and assembly of VLPs (Extended Data Fig. 10a). Beyond preimplantation development, we predict that HERVK reactivation occurs in human primordial germ cells (PGCs), which are also characterized by the presence of OCT4 and genome-wide DNA hypomethylation. HERVK protein products have the potential to engage host machinery, as exemplified here by modulation of cellular mRNAs by Rec, which are also characterized by the presence of OCT4 and genome-wide DNA hypomethylation. HERVK protein products have the potential to engage host machinery, as exemplified here by modulation of cellular mRNA synthesis by Rec. This fine-tuning of cellular functions by HERVK proteins may contribute to human-specific or even individual-specific aspects of early development, as the retroviral ORFs are preferentially expressed from the human-specific proviruses, many of which are polymorphic in the human population. Finally, our data raise the intriguing possibility that HERVK provides an immunoprotective effect for human embryos against different classes of viruses sensitive to the IFITM1-type restriction. Although IFITM family members were first described as interferon-induced genes, they are also classical naive-state and PGC markers in the mouse, which nonetheless appear to be dispensable for development. These observations suggest that IFITM1-mediated restriction may be a evolutionarily conserved mechanism protecting both embryos and germ cells from either reinfection from infectious ERVs or exogenous viral infection (Extended Data Fig. 10a).

Online Content Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

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Supplementary Information is available in the online version of the paper.

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Author Contributions E.J.G. and J.W. conceived the project, designed experiments and wrote the manuscript, with input from all authors. E.J.G. carried out most of the experiments and data analyses. S.L.C., M.W. and E.J.G. performed human biopsy/tissue handling and immunofluorescence with expertise and resources provided by R.A.R.P., R.A.F., L.M. and H.Y.C. performed and analysed (CLIP) experiments. R.A.F. provided assistance with ribosome profiling experiments and analysis. N.L.B. and C.A.B. contributed influenza infection experiments. C.B.W. provided human naive cell lines and reagents. D.J.W. performed expression analysis of LTREHS-associated genes.

Author Information Sequencing data sets generated for this study are deposited under the Gene Expression Omnibus under accession number GSE63570. Reprints and permissions information is available at www.nature.com/reprints. The authors declare no competing financial interests. Readers are welcome to comment on the online version of the paper. Correspondence and requests for materials should be addressed to J.W. (wysocka@stanford.edu).
**METHODS**

DNA and RNA isolation at reverse transcription. Genomic DNA was isolated using phenol:chloroform:isoamyl (100:100:1) (Invitrogen). Briefly, cells were digested in 10 mM Tris-HCl (pH = 8.0), 0.1 M EDTA, 0.5% SDS for 37 °C for 1 h, then proteinase K was added to final concentration of 100 μg/ml and then incubated for 3 h at 50 °C. DNA was PCR extracted, ethanol precipitated, and resuspended in TE. RNA was extracted using Trizol (Invitrogen) according to the manufacturer’s instructions. DNase treatment with Turbo DNase (Ambion) was performed for 30 min at 37 °C, PCR extracted, ethanol precipitated, and resuspended in water. Reverse transcription was performed with SuperScript III (Invitrogen) using ~500 ng of DNAase treated total RNA following the manufacturer’s instructions. No reverse transcriptase controls were performed where necessary.

**Cell lines and culture.** NCCIT and HEK293 cells were obtained from ATCC. NCCIT cells were maintained in 10% FBS (Omega), 1× non-essential amino acids (100× stock, Invitrogen), and basal media RPMI 1640 (Hyclone). HEK293 cells were maintained in 10% FBS/0.1% sodium azide and stained with 1:100 IFITM1 antibody (rabbit pAb, Proteintech, #50556193) for 30 min at 4 °C. Washed cells were then incubated with chick, anti-mouse A647 secondary for 30 min at room temperature. Control transfection constructs for Rec expression in NCCIT cells were used with eif1a promoter, N-terminal Flag–eGFP-tagged Rec cloned into a piggy-back construct with a puromycin selectable marker. Control construct using Flag–eGFP alone (vector only control) was also used in parallel. Transgene constructs were cotransfected with piggy-back transposase plasmid to generate stable lines. Clones were selected and expanded. Flag–eGFP–Rec clone #1 has >30x endogenous expression of rec mRNA (as measured by qPCR) and Flag–eGFP–Rec clone #2 has >14x endogenous expression of rec mRNA (qPCR), data not shown.

**siRNA knockdown.** siRNA was generated using baculovirus-produced Gardia DICER as described. Briefly, 1 μg of PCR product was in vitro transcribed using Megascript T7 (Ambion) and digested using Dicer at 37 °C for 16 h. siRNA was purified using Purelink RNA mini Kit (Ambion) and the absence of >22 nucleotides RNA was verified using gel electrophoresis and ethidium bromide staining. NCCIT cells were plated onto Matrigel-coated 24-well plates, transfected using 1.5 μl of RNAi-max (Invitorgen) in optiMEM (Gibco) with 25 nM siRNA concentrations for 4 h before addition of fresh media. siRNA knockdowns were performed for three consecutive days, cells were harvested 24 h after final transfection. Two independent siRNA pools were generated for OCT4, NANOG and SOX2, each for turboRFP (non-targeting control) and rec, which overlaps the env ORF. Primers used to generate double-stranded RNA (dsRNA) templates are listed in Supplementary Table 10.

**Human embryo source and procurement.** Human embryos were obtained as previously described. Approximately 25 supernumerary human blastocysts from successful IVF cycles, subsequently donated for non-stem-cell research, were obtained with written informed consent from the Stanford University RENEW Biobank. De-identification was performed according to the Stanford University Institutional Review Board-approved protocol #10466 entitled ‘The RENEW Biobank’ and the molecular analysis of the embryos was in compliance with institutional regulations. Approximately 25% of the embryos were from couples that used donor gametes and the most common cause of infertility was unexplained at 35% of couples. No protected health information was associated with any of the embryos.

**Human embryo thawing and culture.** Human embryos cryopreserved at the blastocyst stage were thawed by a two-step rapid thawing protocol using Quinn’s Advantage Thaw Kit (CooperSurgical) as previously described. In brief, either cryostaws or vials were removed from the liquid nitrogen and exposed to air before incubating in a 37 °C water bath. Once thawed, embryos were transferred to a 0.5 mol l−1 sucrose solution for 10 min followed by a 0.2 mol l−1 sucrose solution for an additional 10 min. The embryos were then washed in Quinn’s advantage medium with HEPEs (CooperSurgical) plus 5% serum protein (CooperSurgical) and each transferred to a 25 μl microdrop of Quinn’s advantage blastocyst medium (CooperSurgical) supplemented with 10% serum protein substrate under mineral oil (Sigma). The embryos were cultured at 37 °C with 6% CO2, 5% O2 and 89% N2 under standard human embryo culture conditions in accordance with current clinical IVF practice. Embryos used in this study were DPF 5–6.

**Immunoﬂuorescence.** Cells were grown on Matrigel-coated glass coverslips, fixed using EM-grade 4% PFA (Electron Microscopy Sciences) for 15 min at 27 °C, washed three times with PBS, blocked and permeabilized with 1% BSA, 0.3% Triton-X 100 in PBS (antibody buffer) supplemented with 5% serum for 1 h at 27 °C. Primary antibodies were incubated in antibody buffer and incubated at 4 °C overnight. Washes were performed three times using 0.1% Triton-X 100 in PBS, and secondary antibodies were added for 1 h at 27 °C in the dark. Cells were mounted using Prolong-fade gold (Invitrogen) with DAPI and imaged on Zeiss LSM 700 confocal.
For embryo immunostaining, the zona pellucida (ZP) was removed from each embryo by treatment with acidified Tyrode’s solution (Millipore) and ZP-free embryos were washed in PBS plus 0.1% BSA and 0.1% Tween-20 (PBS-T; Sigma-Aldrich) before fixation in 4% paraformaldehyde for 20 min at room temperature. Once fixed, the embryos were washed three times in PBS-T to remove any residual fixative and permeabilized in 1% Triton X-100 (Sigma-Aldrich) for 1 h at room temperature. Following permeabilization, the embryos were washed three times in PBS-T and then blocked in 4% of chicken or goat serum in PBS-T overnight at 4°C. The embryos were incubated with primary antibodies in PBS-T with 1% serum sequentially for 1 h each at room temperature at the following dilutions: 1:200 OCT4, 1:100 Gag/Capsid. Primary signals were detected using the appropriate 488- or 647-conjugated Alexa Fluor secondary antibody (Invitrogen) at a 1:250 dilution at room temperature for 1 h in the dark and subsequently DAPI stained. Immunofluorescence was visualized by sequential imaging, whereby the channel track was switched each frame to avoid cross-contamination between channels, using a Zeiss LSM510 Meta inverted laser scanning confocal microscope. The instrument settings, including the laser power, pinhole and gain, were kept constant for each channel to facilitate semi-quantitative comparisons between embryo.

DNA demethylation treatment. HEK293 cells were plated on Matrigel-coated 24-well plates, and treated with 0.1 or 10 μM 5-aza-2′-deoxycytidine (Calbiochem) freshly prepared every 24 h. Cells were then transfected with 1 μg each of pdNA3.1-1.OCT4 and pdNA3.1-SOX2 expression plasmids. Media was changed 24 h later, and cells were harvested 3 days after transfection for RNA analysis. Human ES cells (H9) were grown as described earlier, except mTeSR was supplemented with Rock inhibitor (y-27632, Sigma) at 5 μM, and treated with 0.1 or 10 μM 5-aza-2′-deoxycytidine (Calbiochem) for 24 h.

RNA-seq library construction. Libraries were constructed as described35, using ~10 μg of total RNA followed by poly-A selection with oligo-dT beads, ligation and ten cycles of PCR with NEBnext kit oligonucleotides, and sequenced using Illumina HiSeq2000 at the Stanford Sequencing Facility or ELIM Bio.

Sequence analysis. For RNA-seq repeat analysis of data from embryo and human ES cell libraries (for Fig. 1 and Extended Data Fig. 1), FASTQ files were aligned to rebase consensus sequences (downloaded from RepBase) with bowtie using the command “bowtie -q -p 8 -s -n 2 -e 70 -I 12 -m 20 800 -k 1 -best.” These bowtie parameters ensure that only the best alignment (highest scores) is reported, further only one alignment per read is reported, that is, these settings do not allow multiple-matching. For Fig. 2a analysis of HERVK proviruses, RNA-seq reads were aligned to hg19 using the same parameters described earlier, and the overlap between the manually curated HERVK provirus data set8 is reported. For ReSeq analysis for RNA-seq libraries generated for this paper (ELF1 naive or primed human ES cells; from human EC siRNA RNA-seq, or Rec-hECC versus wild-type human EC cell experiments), reads were processed using DNAnexus software to obtain read counts and RPKM. Reads were counted and where indicated normalized to repeat length and library size using RPKM. Differential expression in RNA-seq experiments described earlier was performed using DESeq, with reported FDR using Benjamini–Hochberg correction.

Interferon-induced gene set analysis. Genes were defined as interferon induced if they were induced fivefold in interferon-treated cells or tissues for experimentally defined interferon responses, PubMedlisted interferon-induced gene set analysis. These transcript models were then compared using the RPMK and DESeq analysis as described earlier. Differential enrichment of LTRHS-associated transcripts in naive/primed upregulated versus naive/primed downregulated was analysed using non-paired Wilcoxon test, and significance is reported at P value < 0.05. Higher average naive/primed RPMK of LTRHS-associated versus non-LTRHS-associated genes was tested using non-paired Wilcoxon test.

Chimaeric transcript identification. One-hundred base-pair paired-end RNA-seq reads generated with ELF1 naive versus primed human ES cells (see earlier) were analysed using a published pipeline32. Briefly, Cufflinks software was used to perform de novo identification of transcript models. These transcript models were then used to identify splice junctions in which one side of the transcript model overlapped the GTF file (for hg19 from UCSC), cataloguing known genes and long noncoding RNAs (lincRNAs), and the other side of the transcript model aligned to hg19 classified as a repeat (UCSC genome browser, repeat track). Transcripts that fulfilled these criteria were classified as chimaeric transcripts, and are reported in Supplementary Table 1.

Clustering. Hierarchical clustering was performed using Gene-e software (http://www.broadinstitute.org/cancer/software/GENE-E/index.html) using K-means clustering of log2-transformed RPMK.

Statistical tests. A list of the statistical tests, multiple-hypothesis testing corrections, and normality criteria for parametric tests is reported in Supplementary Table 7.

Electron microscopy. Samples were fixed using 4% PFA and 0.01% glutaraldehyde for 15 min at 27°C. Routine heavy metal staining was conducted where indicated. Immuno-TEM with 1:100 dilution of anti-HERVK Gag/Capsid using overnight incubation at 4°C and labelling was visualized using 5 nm gold-labelled anti-mouse secondary antibody. Secondary only controls demonstrated specificity of the antibody for this application. TEM was performed at the Electron Microscopy core at Stanford University using a Joel JEM-1400 electron microscope.

iCLIP and data analysis. The iCLIP method was performed as described before with the specific modifications below36. Flag–GFP–Rec (Flag-Rec) expressing NCC cells were UV-C crosslinked to a total of 0.3 J cm−2. Each iCLIP experiment was normalized for total protein amount, typically 1 mg, and partially digested with RNaseL (Life Technologies) for 10 min at 37°C and quenched on ice. Flag-Rec was isolated with anti-Flag agarose beads (Sigma) for 3 h at 4 °C on rotation. Samples were washed sequentially in 1 ml for 5 min each at 4°C. 2X high stringency buffer (15 mM Tris-HCl pH 7.5, 5 mM EDTA, 2.5 mM EGTA, 1% Triton X-100, 1% Na-deoxycholate, 120 mM NaCl, 25 mM KCl), 1X high salt buffer (15 mM Tris-HCl pH 7.5, 5 mM EDTA, 2.5 mM EGTA, 1% Triton X-100, 1% Na-deoxycholate, 1 M NaCl, 250 mM MgCl2, 0.5% NP-40). Purified Flag-Rec was then eluted from anti-Flag agarose beads using competitive Flag peptide elution. Each sample was resuspended in 500 μl of Flag elution buffer (50 mM Tris-HCl pH 7.5, 250 mM NaCl, 0.5% NP-40, 0.1% Na-deoxycholate, 0.5 mg ml−1 Flag peptide) and rotated at 4°C for 30 min. The Flag elution was repeated once for a total of 1 ml. Flag-Rec was then captured using anti-GFP antibody (Life Technologies, A-11122) conjugated to Protein A dynabeads (Life Technologies) for 3 h at 4°C on rotation. Samples were then washed as described previously in the anti-Flag agarose beads. 3′-End RNA dephosphorylation, 3′-end single-stranded RNA (ssRNA) ligation, 5′ labelling, SDS–PAGE separation and transfer, autoradiograph, RNIP isolation, Proteinase K treatment and overnight RNA precipitation took place as previously described37. The 3′-ssRNA ligation adaptor was modified to contain a 3′ biotin moiety as a blocking agent. The iCLIP library preparation was performed as described elsewhere37,38. Final library material was quantified on the BioAnalyzer High Sensitivity DNA chip (Agilent) and then sent for deep sequencing on the Illumina HiSeq 2500 machine for 1× 75 bp cycle run. iCLIP data analysis was performed as previously described37. For analysis of repetitive noncoding RNAs, custom annotation files were built from the Rfam database. For analysis of endogenous retroviral elements, custom annotation files were built from the rebase database. iCLIP reads were filtered for quality, barcode split, PCR-duplicate removed, trimmed (5′ and 3′ ends), and mapped for unique matches under parameter settings previously described37. Bioinformatic pipeline used for iCLIP data analysis is described previously38. Briefly, RT stops were used to map nucleotide resolution of Rec binding, and only nucleotides supported with three independent RT stops in two replicates (with at least one RT stop in each replicate) were reported as binding events, and are reported in Supplementary Table 3.

Ribosome profiling. Human EC cells (NCCIT) were cultured as described earlier. Total RNA was extracted using Trizol (Life Technologies) and used as input material for the ARTseq Ribosome Profiling Kit—Mammalian (Epicentre) following the manufacturer’s protocol with the following modifications. The 3′ UTR ligation adaptor and cDNA synthesis primers from the iCLIP protocol were for library construction. Final library material was quantified as with the iCLIP experiments and sequenced on the Illumina HiSeq2500 in single 1× 75 bp cycle run. ARTseq reads were then sequenced on the Illumina HiSeq 2500 for 2× 75 bp cycle run. iCLIP data analysis was performed as previously described37. For analysis of repetitive noncoding RNAs, custom annotation files were built from the Rfam database. For analysis of endogenous retroviral elements, custom annotation files were built from the rebase database. iCLIP reads were filtered for quality, barcode split, PCR-duplicate removed, trimmed (5′ and 3′ ends), and mapped for unique matches under parameter settings previously described37. Bioinformatic pipeline used for iCLIP data analysis is described previously38. Briefly, RT stops were used to map nucleotide resolution of Rec binding, and only nucleotides supported with three independent RT stops in two replicates (with at least one RT stop in each replicate) were reported as binding events, and are reported in Supplementary Table 3.

Influenza infection experiments. Human EC cells (NCCIT) were plated in duplicate (1.5× 105 cells per well) on a 96-well flat-bottom plate in 100 μl Virus Diluent (DMEM, Gibco, supplemented with 1% BSA, 1 μl antibiotics and 20 mM HEPEs). Cells were incubated at 37°C and 5% CO2 for 1.5 h. Wild-type human EC cells and Rec-hECCs were then infected with virus (influenza A/H1N1/PR8/1934, diluted 1:10 into 100 μl virus diluent, increasing total volume to 200 μl). Cells were incubated at 37°C for 1 h. FBS (Hyclone) was added to the wells to a final concentration of 10% FBS. Cells were incubated at 37°C for 5 h. 20 mM EDTA (20 μl) was added to all wells and mixed thoroughly to stop infection. Cells were washed with 200 μl 1× PBS (Hyclone), re-suspended in

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100 µl 1× BD FACS Lysing Solution (BD Biosciences) and stored at −80 °C for later processing.

For staining and analysis, cells were thawed in 37 °C for 20 min. One-hundred microlitres FACS wash (1× HyClone DPBS with 2% FBS) was added to each well and plate was centrifuged. Cell pellets were re-suspended in 200 µl BD FACS Permeabilizing Solution II (BD Biosciences). Cells were incubated at room temperature in the dark for 10 min. Plate was centrifuged and cells were washed twice with 200 µl FACS wash. Cells were stained with primary antibody (mouse anti-influenza A nucleoprotein, C43 clone, Abcam) diluted to 2 µg ml⁻¹. Cells were incubated in the dark at room temperature for 30 min and washed twice. Cell pellets were resuspended in 2 µg ml⁻¹ of secondary antibody (chicken anti-mouse Alexa647, Invitrogen) in 50 µl FACS wash and incubated in the dark at room temperature for 30 min. Cells were washed twice and cell pellets were resuspended in 1% PFA (Electron Microscopy Sciences). Cells were gated to exclude dead cells and debris. Infection levels were background subtracted using uninfected wells, and normalized to infection levels in GFP-hECC for each run.

**RNA-seq data sets.** Data sets used in this study can be accessed from: Array Express Database (accession number E-MATB-2031)¹²; Gene Expression Omnibus (accession number GSE36552)¹³; Gene Expression Omnibus (accession number GSE44183)¹⁴; Array Express (accession number E-MTAB-2857)¹⁵.

Sequencing data sets generated for this study are deposited under in the Gene Expression Omnibus under accession number GSE63570, and are summarized in Supplementary Table 8.

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Repeat expression (data from Yan, et al. NSMB 2013)

HERV-K expression (Data from Xue, et al. Nature, 2013)

Data from Yan, et al. Cell Stem Cell 2013.

b HERV-H RNA expression

c HERV-H RNA expression

d Data from this study

naive hESC RNA-seq

zoom

HERV-K 108 provirus

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Extended Data Figure 1 | Additional single-cell RNA-seq data analyses from preimplantation human embryos (supporting Fig. 1). a, Heat map and hierarchical K-means clustering of highly expressed (average RPKM > 6 across 89 embryo libraries) repetitive elements in single cells of human preimplantation embryos at indicated developmental stages (top) and HERVK expression (bottom) using indicated data sets. b, HERVH expression (RPKM) in single cells of human embryos at indicated preimplantation stages. Solid line indicates mean. RNA-seq data are taken from ref. 10. c, HERVH expression (RPKM) in single cells of human blastocysts, grouped by lineage. Solid line indicates mean. Oocyte (n = 3), zygote (n = 3), 2-cell (n = 6), 4-cell (n = 11), 8-cell (n = 19), morula (n = 16), TE (n = 18), PE (n = 7), EPI (n = 5), p0 (n = 8), p10 (n = 26). RNA-seq data set was from ref. 10. d, Genome browser snapshot showing 100 bp PE-RNA-seq reads from ELF1 naive human EScells aligning at the HERVK 108 provirus on chromosome 7.
Extended Data Figure 2 | LTR5 alignments, HERVK expression data in cell lines, and control ChIP-qPCR analyses in primed human ES cells (supporting Fig. 2). a, Top, presence of HERVK(HML-2) sequences in Old World primates, but absence in New World primates. Middle, schematic of HERVK proviral genome; all human-specific insertions contain LTR5HS. Bottom, phylogenetic relationship of HERVK LTR subclasses showing high degree of sequence similarity. Pro, protease; Pol, polymerase; Gag, group-specific antigen; Env, envelope. Bottom, ClustLW multiple sequence alignment of indicated HERVK LTR sequences (top), region around OCT4 motif is boxed, phylogenetic tree (bottom) indicating presence/absence of OCT4 motif.
b, HERVK protein expression in human EC cells and human ES cells. Protein extracts from human EC cells (NCCIT) and human ES cells (H9) were analysed by immunoblotting with an antibody detecting HERVK Gag precursor and the processed Capsid (top), or the glycosylated, unprocessed form of the HERVK envelope protein Env (bottom). Tata-binding protein (TBP) was used as a loading control. Shown is a representative result of three independent experiments. c, RT–qPCR analysis of HERVK RNA expression in human EC cell line NCCIT, human ES cell line H9, and HEK293 cells. Three distinct qPCR amplicons, corresponding to env, gag and pro are shown. Samples were normalized to 18S ribosomal RNA levels. *P value < 0.05, one-sided t-test. Error bars are ±1 s.d., n = 3 biological replicates. d, HERVK gag or env expression in male human ES cell lines HSF-1, HSF-8, female human ES cell line H9 and human EC cell line NCCIT. *P value < 0.05, one sided t-test compared to control siRNA, n = 3 biological replicates. Error bars are ±1 s.d. e, RT–qPCR analysis of HERVK transcripts after siRNA knockdown of NANOG, OCT4 or SOX2 in human EC cells (NCCIT). Signals were normalized to 18S rRNA. *P value < 0.05, one sided t-test compared to control siRNA, n = 3 biological replicates. Error bars are ±1 s.d. f, ChIP-qPCR analyses of human ES cells (H9) with indicated antibodies. Signals were interrogated with primer sets for positive control regions (active human ES cell OCT4 and SOX2 enhancers), LTR5HS, or non-repetitive, intergenic negative regions, as indicated at the bottom. Shown is a representative result of two biological replicates.
**a**

control OCT4 siRNA

control NANOG siRNA

control SOX2 siRNA

**b**

Luciferase assays (hECC)

LTR5_HS wildtype

LTR5_HS oct4 mutant

SIV5 prozone

mock

**c**

**d**

**e**

HEK293 3 days 5-aza-2'-deoxycytidine

transfect: OCT4/SOX2 2 days Measure: HERV-K RNA

OCT4/SOX2 Empty vector

**Letters**

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Extended Data Figure 3 | HERVK regulation by OCT4 and DNA methylation (supporting Fig. 2). a, Transcription factor knockdown in human EC cells (NCCIT). Cells were transfected with siRNA pools targeting indicated transcription factors and protein depletion was measured by immunofluorescence with respective antibodies in comparison with control, mock-transfected cells. DAPI (blue), OCT4 (green, left), NANOG (green, middle), SOX2 (green, right). Shown is one of three representative fields of view at ×20 magnification. b, Dual luciferase assays with indicated reporter constructs in human EC cells (NCCIT) showing that mutation of OCT4 site decreases reporter activity. N = 3 biological replicates, error-bars ±1 s.d. *P value < 0.05, one-sided t-test. SV40 enhancer/promoter construct was used as a positive control. c, Bisulfite sequencing for indicated cell types (WT33 human IPSC) analysing consensus LTR5HS-specific amplicon as in Fig. 2e. d, Bisulfite sequencing analysis of HERVK proviral consensus amplicon containing 3′ end of LTR, primer binding site, and 5′ region of Gag ORF (see Extended Data Fig. 2a) in indicated cell types: ELF1 naive, human ES cell, WT33 human IPSC, NCCIT human EC cell, or H9 human ES cell. e, RT–qPCR analysis of HERVK RNA levels in HEK293 cells treated with indicated concentrations of 5-aza-2′-deoxycytidine for 3 days, followed by transfection with OCT4/SOX2 expression constructs and RNA collection 48 h after transfection. qPCR primer sets were designed to three independent amplicons of HERVK. *P value < 0.05, one-sided t-test. n = 4 biological replicates, error bars ±1 s.d.
a) DAPI OCT4 Gag/Cap merge/zoom
b) hECC siRNA
   | gag #1 | gag #2 | gag #3 | control |
   | 1:2    | 1:2    | 1:2    | 1:2     |
c) GAG TBP
   | dispac/gaG | IPAD | dispac/gaG | 4TCO | EGRE | M |
   | Gag capsid | OCT4 | MERGE | Gag capsid | OCT4 | MERGE |
d) Elf1, naive hESC
   | DAPI | OCT4 | MERGE |
   | Gag capsid | OCT4 | MERGE |
e) Human blastocyst
   | DAPI | OCT4 |
   | Gag capsid | OCT4 | MERGE |
Extended Data Figure 4 | HERVK Gag/Capsid antibody validation and staining (supporting Fig. 3). a, Immunofluorescence analysis of human EC cells (NCCIT) and human ES cells (H9) stained with DAPI (blue), OCT4 (green), Gag/Capsid (red), or IgG control (bottom). White boxes indicate regions shown in higher magnification/merge (right). Shown are representative fields of three independent experiments.
b, Sensitivity of HERVK Gag/Capsid antibody immunoblot signal to HERVK knockdown. Human EC cells were transfected with one of three independent siRNA pools targeting HERVK Gag or with a control, non-targeting pool (synthesized against RFP) and total protein was analysed by immunoblotting with anti-Env and anti-Gag/Capsid antibodies. 1:2 serial dilution of total protein was loaded, as indicated. Blots were stripped and re-probed with TBP as a loading control. Shown is a representative result of two independent experiments.
c, Sensitivity of HERVK Gag/Capsid antibody immunofluorescence signal to siRNA knockdown of Gag/Capsid (top) or control siRNA targeting RFP (bottom). Shown is a representative result of three fields of view. Magnification: 20X.
d, Immunofluorescence of naive ELF1 human ES cells with antibodies against OCT4 (green), HERVK Gag/Capsid (pink), DAPI in blue. Region marked with white box on left is shown with larger magnification (bottom). Magnification = 20x, 40x respectively.
e, Another representative example of immunofluorescence of human blastocysts with DAPI (blue), OCT4 (green) and Gag/Capsid (red) shown (n = 19 blastocysts; DPF 5–6). Original magnification, ×40.
a) heavy metal staining

b) anti-Gag/Capsid

b) anti-Gag/Capsid (zoom)

c) control staining (secondary only)

d) EGA

naive hESC

implantation

primed hESC

in vitro maturation?

DNA methylation

nuclear OCT4

HERV-K expression

EGA

implantation

naive hESC

primed hESC

in vitro maturation?
Extended Data Figure 5 | TEM analyses of human EC cells and control embryo staining (supporting Fig. 3). a, TEM analysis of human EC cells (NCCIT) with heavy metal staining; arrow indicates VLPs. Boxed region is shown with higher magnification in an inset. Scale bar = 500 nm. Shown is a representative example of two independent experiments. b, TEM immuno-gold labelling of human EC cells (NCCIT) with Gag/Capsid antibodies. Shown is a representative example from two independent experiments. c, Secondary antibody only control for immuno-gold labelling of human blastocysts. Shown is a representative example from eight fields of view. d, Model figure summarizing HERVK transcriptional regulation in human embryos and in vitro cultured pluripotent cells. Dashed lines indicate inference of OCT4, DNA methylation and HERVK level changes at implantation from those observed between naive and primed human ES cells, in the absence of data from actual postimplantation human embryos.
a Chimeric transcript identification (Cufflinks analysis ELF1 naive hESC)

b Chimeric transcript identification (Cufflinks analysis ELF1 naive hESC)

c LTR5HS-associated genes

d ELF1 hESC, naive vs primed (This study, Ware, et al. 2013)

e 3iL naive hESC vs primed H1 hESC (Chan, et al. 2013)
Extended Data Figure 6 | Correlation of HERV LTR5HS elements with gene expression (supporting Fig. 4). a, Number of splice junctions identified linking indicated HERV class to annotated ReqSeq genes. Analysis was done using RNA-seq data set from ELF1 naive human ES cells, n = 3 biological replicates. b, Number of reads supporting chimaeric transcripts from indicated HERV class in ELF1 naive human ES cells, n = 3 biological replicates. c, Expression of LTR5HS linked genes plotted as a function of distance to the gene’s transcription start site (TSS). x-axis: distance of TSS to the nearest LTR5HS in kb; y-axis: fold change in expression of the linked gene in ELF1 naive versus primed human ES cells (this study, left) or expression of the linked gene in 3iL versus primed H1 human ES cells (right, ref. 12). d, Top, histograms showing expression of all genes that significantly change in expression between naive and primed ELF1 human ES cells (top histogram, white) or significantly changed genes that are LTR5HS associated (bottom histogram, blue); expression values from naive versus primed ELF1 human ES cell RNA-seq data sets (FDR < 0.05 DESeq). Fischer’s exact test gives stated P value, indicating enrichment of LTR5HS-linked genes in naïve upregulated category. Bottom, quantification of average expression of LTR5HS-linked (blue) or unlinked (white) genes. Non-paired Wilcoxon test with stated P value indicating that genes linked to 1 or more LTR5HS have significantly higher mean expression.

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**a** rec RNA level (RT-qPCR)

**b** IFITM1 RNA level (RNA-seq)

Data and analysis from Chan, et al. 2013.

Data and analysis from this study

**c** surface IFITM1 staining

**d** IFITM1

**e** hECC

**f** All interferon-induced genes

Data and analysis from this study

Data and analysis from Chan, et al. 2013.

**Note:**
- rec RNA level (RT-qPCR)
- IFITM1 RNA level (RNA-seq)
- Surface IFITM1 staining
- IFITM1
- hECC
- All interferon-induced genes

**Images and Graphs:**
- Bar charts and scatter plots indicating relative expression levels and RNA-Seq results for ELF1 and primed naive hESC.
- Surface IFITM1 staining diagrams with control and treated groups.
- Flow cytometry analysis for IFITM1 expression in hECC.
- Graph showing all interferon-induced genes with log(RPKM) values.
Extended Data Figure 7 | rec and IFITM1 expression in naive human ES cells, and effect of Rec expression on H1N1(PR8) infection (supporting Fig. 4). a, Left, RT–qPCR analysis of HERVK rec expression levels in ELF1 naive human ES cells (n = 3 biological replicates) or H9 primed human ES cells (one biological replicate). Normalized to 18S rRNA. Right, Rec RNA levels in indicated blastocyst lineages. Solid line indicates mean; data are from ref. 10. b, RNA-seq quantification of IFITM1 RNA levels in naive or primed ELF1 human ES cells (left, this study) or 3iL human ES cells versus primed H1 human ES cells from ref. 12 (right). n = 3 biological replicates for each condition, error bars are ±1 s.d. Asterisk indicates significance at FDR < 0.05, DESeq. c, Flow cytometry for surface-localized IFITM1 staining in the indicated H9 human ES cells or naive ELF1 human ES cells (top) or, as a control for IFITM1 antibody specificity, knockdown of IFITM1 with two independent IFITM1 siRNA pools compared to control siRNA-treated cells in Flag–eGFP–Rec-hECCs (bottom). d, Left, IFITM1 expression in control human EC cell versus Rec-hECC (NCCIT) RNA-seq data sets. n = 2 biological replicates. Significance = FDR < 0.05, DESeq. Right, IFITM1 expression in control siRNA versus Rec siRNA-treated human EC cells (NCCIT) RNA-seq. n = 3 biological replicates, error bars are ±1 s.d. Significance = FDR < 0.05, DESeq. e, Flow-cytometry profiles for indicated cell types in H1N1(PR8) infected (top) or non-infected (bottom) wild-type (WT) control human EC cells or Flag–GFP–Rec-hECCs, clone #1. Shown is one representative example of four independent experiments showing a co-plating experiment in which GFP-Rec cells and wild-type control (GFP negative) cells are infected in the same well, stained in the same tube and identified by GFP fluorescence after gating for FSC and SSC. f, Scatterplot of ELF1 naive versus primed human ES cell RNA-seq showing all interferon-induced genes, with differentially regulated genes (FDR < 0.05 DESeq, n = 3 biological replicates each) highlighted in red. There is a significant overlap between differentially regulated genes and interferon-induced genes as measured by a hypergeometric test (P value < 0.05).
Diagram of iCLIP-seq procedure

**a**

- UV-C Crosslinking
- Cell lysis and partial RNaseI digestion
- FLAG IP
  - 1M NaCl + 1% Na-DOC + 1% TritonX100 wash
- GFP IP
  - 1M NaCl + 1% Na-DOC + 1% TritonX100 wash
- FLAG Peptide Elution
- SDS-PAGE, Transfer, and ProtK Digestion
- Library Prep
- Illumina Sequencing

**b**

| Sample       | WT | Flag-eGFP | Flag-eGFP-Rec clones |
|--------------|----|-----------|----------------------|
|              |    |           | #2                   |
| [RNaseI] UV @ 254nm |   |           | #3                   |

**c**

LTR5HS RRE
predicted RNA structure

598 nt → 250 nt

- RNA + Flag-GFP-Rec
- IP: FLAG, peptide elute, IP: GFP
- Blot: P32-labeled RNA
- Input control
  - WB: anti-GFP
  - (FLAG-eGFP-Rec, slow-migrating)
  - (FLAG-eGFP-Rec, predicted=35Kda)
  - (FLAG-eGFP)
- WB: anti-HSP90
Extended Data Figure 8 | iCLIP analysis of Rec-associated RNAs (supporting Fig. 4). a, Diagram of iCLIP-seq procedure (see Methods for details). Briefly, cells are crosslinked using ultraviolet, lysed and digested with RNase to trim RNAs. Sequential immunopurification is performed using Flag M2, peptide elution, and GFP immunoprecipitation (IP). After stringent washing, RNAs are recovered and either radiolabelled (shown in b) or reverse transcribed and prepared for Illumina HTPS libraries. b, Autoradiogram of labelled RNAs (top) recovered from ultraviolet-crosslinked cells using sequential Flag–eGFP immunoprecipitation from: wild-type human EC cells (lanes 1, 2), Flag–eGFP control human EC cells (lanes 3, 4), or two independent Rec-hECC transgenic lines (lanes 5–8), separated on an SDS–polyacrylamide gel electrophoresis (SDS–PAGE) gel. Free Rec protein runs as a ~35 kDa band, while Rec protein crosslinked to RNA molecules show lower electrophoretic mobility. Please note that: (1) Rec-bound RNAs are resistant to even high concentrations of RNaseI, probably indicating extensive secondary RNA structures, and (2) low/no background of contaminating RNAs in control immunoprecipitation from wild-type human EC cells or Flag–eGFP control human EC cells. Western blots with anti-GFP antibody were also performed to confirm the presence of tagged protein in Flag–eGFP control and Flag–eGFP–Rec cells, both in input and immunoprecipitation fractions (middle). HSP90 was used as a loading control (bottom). c, Computationally predicted (using mFold) secondary structure of LTR5HS sequence around the Rec response element (identified experimentally in vitro previously25). Single nucleotide resolution Rec ultraviolet-crosslinking sites determined by iCLIP are shaded in red; n = 2 biological replicates.
Extended Data Figure 9 | Rec target mRNA analysis (supporting Fig. 4). a, Genome browser representations of the Rec iCLIP read (n = 2 biological replicates) distribution at indicated mRNA targets. b, Computationally predicted (using mFold) secondary structures of indicated Rec iCLIP-seq targets. Single-nucleotide resolution Rec ultraviolet-crosslinking sites determined by iCLIP are shaded in red; to orient the reader, browser representation of the folded fragment is shown above each respective cartoon.
Extended Data Figure 10 | Model of HERV-K regulation and function.