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Mammalian retrovirus-like protein PEG10 packages its own mRNA and can be pseudotyped for mRNA delivery

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Abstract

Eukaryotic genomes contain domesticated genes from integrating viruses and mobile genetic elements. Among these are homologs of the capsid protein (known as Gag) of long terminal repeat (LTR) retrotransposons and retroviruses. We identify several mammalian Gag homologs that form virus-like particles (VLPs) and one LTR retrotransposon homolog, PEG10, that preferentially binds and facilitates vesicular secretion of its own mRNA. We show the mRNA cargo of PEG10

Competing interests: M.S., B.L., and F.Z. are co-inventors on a U.S. provisional patent application filed by the Broad Institute related to this work. F.Z. is a cofounder of Editas Medicine, Beam Therapeutics, Pairwise Plants, Arbor Biotechnologies, and Sherlock Biosciences.

Supplementary Materials Materials and Methods Supplementary Text 1 to 7 Figs. S1 to S16 Tables S1 to S4 Data S1 References (34–56)

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M.S. and F.Z. conceived of the project. M.S., B.L., and F.Z. designed the experiments. M.S., B.L., J.S. A.L., X.J., and C.L. performed the experiments. M.S., B.L., J.S., and F.Z. analyzed the data. M.S., S.M., and E.V.K. performed bioinformatics analysis of Gag protein diversity. F.Z. supervised the research and experimental design with support from R.M. M.S., B.L., R.M., and F.Z. wrote the manuscript with input from all authors.

can be reprogrammed by flanking genes of interest with *Peg10*'s untranslated regions. Taking advantage of this reprogrammability, we develop <u>Selective Endogenous eN</u>capsidation for cellular <u>Delivery (SEND)</u> by engineering both mouse and human PEG10 to package, secrete, and deliver specific RNAs. Together, these results demonstrate SEND as a modular platform suited for development as an efficient therapeutic delivery modality.

One-Sentence Summary:

The LTR retrotransposon-derived Gag protein PEG10 can be harnessed to facilitate efficient and specific intercellular delivery of cargo mRNAs in mammalian cells.

More than 8% of the human genome is composed of sequences derived from LTR retroelements, including retroviruses, that have integrated into mammalian genomes throughout evolution (1–5). Retroviruses and retrotransposons have many common mechanistic features including the core structural gene (known as gag), however, while retrotransposons replicate intracellularly, the acquisition of the envelope (env) gene by retroviruses has enabled intercellular replication (6). Most endogenous retroelements have lost their original functions, but some of their genes have been recruited for diverse roles in normal mammalian physiology. For example, the fusogenic syncytins evolved from retroviral env proteins (7). The gag homolog, Arc, which forms capsids and has been reported to transfer its own mRNA (8-10), is involved in memory consolidation and regulates inflammation in the skin (11, 12). Another gag homolog, the LTR retrotransposonderived protein PEG10, which has been reported to bind RNA and also forms capsids (13), is involved in mammalian placenta formation (14, 15). These examples raise the possibility that additional retroelement-derived proteins encoded in the mammalian genome transfer specific nucleic acids, providing a potentially programmable mechanism for intercellular communication.

Computational survey of mammalian capsid-forming gag homologs

To identify genes with the potential to transfer specific nucleic acids, we focused on homologs of *gag* containing the core capsid (CA) domain, which protects the genome of both retrotransposons and exogenous retroviruses (16, 17). Previous genome analyses identified many endogenous *gag* homologs in mammalian genomes (18), and experimental efforts have validated the ability of some of these proteins including MmArc and MmPEG10 to form capsid-like particles that are secreted within extracellular vesicles (EVs) (10, 13). To ensure a complete list of candidates, we searched the human and mouse genomes for *gag* homologs. This search identified 48 *gag*-derived genes in the human genome and 102 *gag* homologs in the mouse genome; for 19 human genes, an orthologous relationship between human and mouse was readily traced (in several cases, with additional mouse paralogs), whereas the remaining ones appeared to be species-specific (Table S1 and S2).

Canonical genomes of both LTR retrotransposons and retroviruses encode a long polyprotein consisting of several conserved domains: The matrix (MA), CA, and nucleocapsid (NC) form the *gag* subdomain and are responsible for membrane attachment, capsid formation, and genome binding, respectively. The *pol* subdomain contains the protease (PRO),

responsible for cleaving the polyprotein, the reverse transcriptase (RT), which converts retroelement RNA into DNA, and the integrase (IN) domain, which integrates the genome into that of the host. Some families of endogenous *gag*-containing proteins, such as the PNMA family, contain only the CA and NC subdomains of *gag*, whereas others, such as RTL1 and PEG10 (also known as RTL2 or Mart2), additionally include subdomains of *pol*, namely a PRO domain and a predicted RT-like domain (Fig. 1A, Table S1). Phylogenetic analysis of *Peg10* and its homologs supports the origin of this gene from LTR retrotransposons (fig. S1, A and B).

Among these genes, *Arc* is the most well studied. *Drosophila Arc1* (*darc1*) is a *gag* homolog which contains the MA, CA, and NC domains. It has been shown to form capsids, bind its own mRNA, and transfer it from motor neurons to muscles at the neuromuscular junction (9). *darc1* mRNA binding is dependent on its own 3' untranslated region (UTR), and fusion of this sequence to heterologous mRNAs can initiate their export and transfer as well. *Mus musculus Arc* (*MmArc*), by contrast, contains only the CA domain and has also been shown to form capsids and transfers its own mRNA across synapses (10), but thus far this transfer has not been shown to be specific to *Arc* mRNA, likely due to the lack of a NC domain.

To narrow down the scope of our analysis, we focused on CA-domain containing proteins that are conserved between human and mouse and have detectable levels of mRNA in adult human tissues, reasoning that such proteins were most likely to have been co-opted for important physiological roles in mammals (fig. S2). We produced mouse versions of the selected CA-containing proteins in *E. coli* and found that a number of these formed higher molecular weight oligomers identified by size exclusion (Fig. 1B, and fig. S3A), as previously noted for some of these proteins, such as MmArc (10). Electron microscopy of these aggregated proteins showed that MmMOAP1, MmZCCHC12, MmRTL1, MmPNMA3, MmPNMA5, MmPNMA6a, and MmPEG10 self-assemble into capsid-like particles, many of which appear spherical (Fig. 1, C and D, and fig. S3, B and C).

MmPEG10 binds and secrets its own mRNA

To determine if these proteins are secreted within an EV, we overexpressed an epitope tagged mouse ortholog of each CA-containing gene in HEK293FT cells and harvested both the whole cell lysate and the virus-like particle (VLP) fraction by clarification and ultracentrifugation of the culture media (Fig. 1E). We found that MmMOAP1, MmArc, MmPEG10, and MmRTL1 were all present in the VLP fraction (Fig. 1F, and fig. S4A), but MmPEG10 was the most abundant protein in the VLP fraction (Fig. 1G). Additionally, endogenous MmPEG10, but not MmMOAP1 or MmRTL1, was readily detectable in cell-free adult mouse serum (fig. S4B).

We next tested whether any of the capsid-like particles formed by Gag homologs contained specific mRNAs using RNA sequencing. To avoid the possibility of transfected Gag homolog expression plasmids contributing to high background signal during sequencing, we used CRISPR activation (19) to induce expression of endogenous genes in mouse N2a cells (Fig. 2A and S5A). We performed mRNA sequencing on whole cell lysate and the

VLP fraction (following nuclease treatment to remove any residual, unencapsidated RNA) to identify RNA species in the VLP fraction. We found *MmPeg10* transcriptional activation led to accumulation of significant amounts of full length *MmPeg10* mRNA transcripts in the VLP fraction (Fig. 2, B and C). Previous work on MmPEG10 demonstrated that it binds a number of mRNAs inside of trophoblast stem cells including itself (13), however, here we further show that MmPEG10 binds and secretes its own mRNA into the VLP fraction. An important caveat of this experiment is that some of these proteins, particularly MmArc, are subject to regulation at the level of translation, so lack of enrichment in the VLP fraction could be due to low protein expression (20).

To confirm our observation for *MmPeg10*, we transiently transfected overexpression plasmids of UTR-flanked *MmPeg10* into N2a cells and found only enrichment for *MmPeg10* mRNA in the VLP fraction (Fig. 2D) under this overexpression condition. PEG10 contains two putative nucleic acid binding domains, namely the NC and RT, which are released from the polypeptide upon PEG10 self processing (21) (Fig. 2E, Supplementary Text 1, fig. S5, B–D). We generated deletions of these domains and found that mRNA export depends on the MmPEG10 NC, as loss of the nucleic acid-binding zinc finger CCHC motif (residues 416–429) from the MmPEG10 NC substantially reduced export of its mRNA (Fig. 2F).

To better understand the roles of the nucleic acid binding domains of MmPEG10 in RNA binding, we performed eCLIP in N2a cells after transient transfection with HA-tagged MmPeg10 as well as the NC and RT mutants (fig. S6, A and B). Compared to the control, MmPEG10 strongly bound a number of mRNAs in N2a cells including its own mRNA (Fig. 2G). Importantly, both the NC and the RT domains are required for the binding of these mRNAs by MmPEG10 (Fig. 2H and fig. S6C). To confirm MmPEG10's cellular role in an *in vivo* context, we generated knock-in mice carrying an N-terminal HA-tag on the endogenous MmPEG10 protein (fig. S6D). Expression of *MmPeg10* in cortical neurons has been demonstrated previously (fig. S6E) (22). Endogenous MmPEG10 was also found to bind its own mRNA as well as other transcripts abundant in neurons (fig. S6, F and G); in contrast to previous datasets, we detected strong MmPEG10 binding in the 5' UTR as well as some additional binding near the boundary between the nucleocapsid and protease coding sequences and in the beginning of the 3' UTR (Fig. 2I) (13).

MmPEG10's binding of mRNA has been reported to increase the cellular abundance of target transcripts (13). To confirm this role of MmPEG10 in its native context *in vivo*, we perturbed *MmPeg10* gene expression in the postnatal mouse brain and assessed expression changes of MmPEG10 bound transcripts (Supplementary Text 2). We found that the mRNAs of 49 genes downregulated in the brain upon *MmPeg10* knockout are bound to MmPEG10 in the age-matched mouse brain (fig. S7F), suggesting that one of the functions of MmPEG10 is to bind and stabilize mRNAs with fundamental roles in neurodevelopment.

Pseudotyped PEG10 VLPs can deliver engineered cargo mRNAs bearing RNA packaging signals from PEG10 UTRs

To reprogram MmPEG10 to bind and package heterologous RNA, we tested whether a cargo mRNA consisting of both the 5' and 3' UTR of *MmPeg10* flanking a gene of interest would be efficiently packaged, exported, delivered, and translated in recipient cells (Fig. 3A). This UTR grafting approach has been demonstrated for the Ty3 retroelement and *darc1* (9, 23). We first used a Cre-loxP system, a highly sensitive system for tracking RNA exchange that has been used previously with exosomes *in vivo* (24). We flanked the Cre recombinase coding sequence with the *MmPeg10* UTRs and co-transfected it with *MmPeg10* with and without a fusogen, the vesicular stomatitis virus envelope protein (VSVg) (Fig. 3A). We found that MmPEG10 VLPs pseudotyped with VSVg are secreted within EVs that mediate transfer of Cre mRNA, not protein, into target loxP-GFP reporter N2a cells in a VSVg- and *MmPeg10* UTR-dependent manner (Fig. 3B, C, D, and S8, Supplementary Text 3). This result suggests that addition of the *Peg10* UTRs enables functional intercellular transfer of an mRNA via VLPs and that these VLPs require a fusogenic protein for cell entry.

We next asked if there is a minimal UTR packaging signal for mediating efficient packaging and functional transfer. The 3' UTR of *MmPeg10* is approximately 4-kB long, but eCLIP indicates only portions of the 3' UTR are bound by MmPEG10 (Fig. 2I). We created constructs encoding the *MmPeg105*' UTR, Cre, and 500-bp segments of the *MmPeg103*' UTR. We found that the proximal 500 bp of the *MmPeg103*' UTR are sufficient for efficient functional transfer of Cre mRNA into target reporter cells (Fig. 3E). Importantly, no efficient functional mRNA transfer was observed for non-UTR flanked Cre or for Cre without the proximal 500 bp of the 3' UTR. Moving forward, we refer to RNA cargo flanked by the *MmPeg105*' UTR and the proximal 500 bp of the 3' UTR as Mm.cargo(RNA), where "(RNA)" specifies the cargo being flanked (e.g., Mm.cargo(Cre)).

Like the mouse ortholog, HsPEG10 is an abundantly secreted protein in the VLP fraction (fig. S10A). Using the same approach we employed with *MmPeg10*, we identified that the 5' UTR and the first 500 bp of the *HsPEG10* 3' UTR are sufficient to mediate functional transfer of Cre mRNA, hereafter denoted as Hs.cargo(RNA) (Fig. 3F). Interestingly, these functional regions of the UTRs are highly conserved across mammals (fig. S10B). Similar to its mouse ortholog, the human system is specific, requiring *HsPEG10* UTR sequences for functional mRNA transfer whereas non-flanked Cre produced only minimal reporter cell activity (Fig. 3F).

To further boost the packaging of a cargo(RNA) by PEG10, we explored the impact of removing any additional PEG10 *cis* binding elements within the *MmPeg10/HsPEG10* coding sequence. For both human and mouse, transfer was increased as a result of recoding the sequence between the NC and the PRO domains, which corresponds to the MmPEG10-bound region in the eCLIP experiments (Fig. 2I, Supplementary Text 4).

Combining these optimizations, we produced VLPs with the recoded mouse and human PEG10 (rMmPEG10 or rHsPEG10), VSVg, and the optimized cargo(RNA) containing the first 500 bp of the 3' UTR; we refer to this system as \underline{S} elective \underline{E} ndogenous e \underline{N} capsidation

for cellular <u>D</u>elivery (SEND). With SEND we detected a substantial (up to 60%) increase in functional transfer of cargo(Cre) into N2a cells for both human and mouse PEG10 (Fig. 3, G and H). Furthermore, we show that VLPs produced with rMmPEG10 can mediate the functional transfer of H2B-mCherry (fig. S12, A and B). Comparison of SEND to previously developed delivery vectors showed that SEND is 4–5 fold less potent than an integrating lentiviral vector as assayed by digital droplet PCR and functional titration (fig. S12, B, C, D, and E). However, given that SEND delivers mRNA rather than integrating an overexpression cassette, we expect it to perform competitively against other mRNA delivery vehicles.

PEG10 is a modular platform for RNA delivery

To generate a fully endogenous SEND system, we tested whether VSVg can be replaced with an endogenous fusogenic transmembrane protein. Given the overlapping tissue expression of *MmPeg10/HsPEG10* and syncytin genes (Supplementary Text 5), we tested the feasibility of pseudotyping the mouse SEND system with MmSYNA, or MmSYNB compared to pseudotyping with VSVg. Pseudotyped particles were incubated with tail-tip fibroblasts from loxP-tdTomato reporter mice, a cell type which we have found amenable to transduction by these fusogens. Based on previous reports, we added the transduction enhancer vectofusin-1 to the supernatant for MmSYNA and MmSYNB particles to enhance *in vitro* transduction (25). In these primary cells, both VSVg and MmSYNA enabled SEND-mediated functional transfer of Mm.cargo(Cre) while MmSYNB did not (Fig. 4A and B). Again, this packaging was highly specific, as only UTR flanked mRNA (i.e., Mm.cargo(Cre)) was functionally transferred. Together with MmSYNA, SEND can be configured as a fully endogenous system for functional gene transfer.

Supported by our understanding of the minimal requirements for PEG10-mediated mRNA delivery (i.e., UTRs and an endogenous fusogen), we could begin to probe the endogenous role of MmPEG10-mediated *MmPeg10* RNA delivery in neurons. Functional transfer of MmSYNA pseudotyped VLPs carrying the native PEG10 transcript into primary mouse cortical neurons led to upregulation of a number of genes involved in neurodevelopment (Supplementary Text 6). This finding reinforces the notion that one role of endogenous *MmPeg10* delivery is binding and stabilizing specific mRNA transcripts in recipient cells. RNA sequencing of N2a cells receiving Mm.cargo(*Peg10*) revealed substantial gene expression changes upon *MmPeg10* delivery that were largely abrogated with PEG10 mediated Mm.cargo(Cre) delivery (Supplementary Text 7). This suggests transferring a reprogrammed cargo does not have the same impact on recipient cells as transferring *MmPeg10* and indicates that *MmPeg10* transcript delivery rather than the delivery of MmPEG10 protein is responsible for the observed gene expression changes. It remains unclear whether MmPEG10 VLPs are natively pseudotyped by the endogenous fusogen MmSYNA to enable cellular uptake of PEG10 VLPs in the central nervous system.

To further characterize the modularity of the components of this system, we tested different cargoRNAs. Using the same pipeline developed for cargo(Cre), we tested whether SEND could mediate the functional transfer of a large ~5 kb Mm.cargo(SpCas9) into N2a cell lines constitutively expressing an sgRNA against *MmKras* (Fig. 4C). SEND was able to functionally transfer SpCas9, leading to ~60% indels in recipient cells (Fig. 4D); similar

to the results with Cre, SEND is specific and only able to efficiently functionally transfer *SpCas9* flanked by either the full length or optimized *Peg10* UTR sequences.

To create an all-in-one vector for delivery of sgRNA and SpCas9, we first tested if an sgRNA can be efficiently delivered by SEND. We independently packaged an sgRNA targeting *Kras* into rMmPeg10 VLPs by co-expressing rMmPeg10 with VSVg and a U6 driven sgRNA and incubated them with Cas9 expressing N2a cells; we saw very little activity even though direct transfection of the guide showed robust indel formation (Fig. 4E). We found, however, that co-packaging the guide alongside Mm.cargo(SpCas9) by co-expressing Mm.cargo(SpCas9) with a U6 driven sgRNA on a separate plasmid was sufficient to mediate 30% indels (Fig. 4F). To determine the reproducibility of this genome editing approach, we repeated this co-packaging strategy with the human SEND system and were able to generate ~40% indels in HEK293FT cells at the *HsVEGFA* locus (Fig. 4G).

The development of SEND from an endogenous retroelement complements existing delivery approaches using lipid nanoparticles (26), VLPs derived from bona fide retroviruses (27–29), and active mRNA loading approaches in EVs (30, 31). Moreover, SEND may have reduced immunogenicity compared to currently available viral vectors (32) due to its use of endogenous human proteins. Supporting this is gene expression data from the developing human thymus which demonstrates that *HsPEG10* is highly expressed compared to other CA-containing genes in the thymic epithelium (33) (fig. S16), which is responsible for T cell tolerance induction. As a modular, fully endogenous system, SEND has the potential to be extended into a minimally immunogenic delivery platform that can be repeatedly dosed, greatly expanding the applications for nucleic acid therapy.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Data and materials availability:

Expression plasmids are available from Addgene under a uniform biological material transfer agreement. Additional information available via the Zhang Lab website (https://zhanglab.bio). Next generation sequencing data generated are available from NCBI SRA with accession number PRJNA743280. All other data are available in the paper and supplementary materials.

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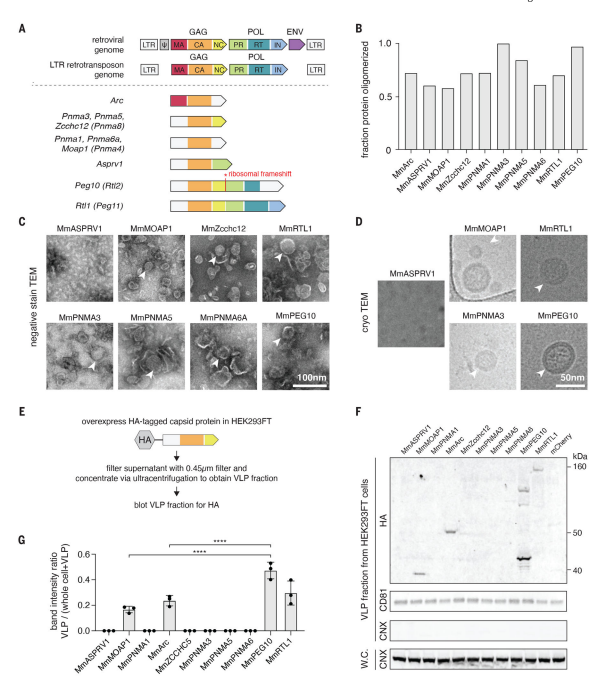
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 $\label{thm:continuous} \textbf{Fig. 1. Identification of mammalian retroelement derived Gag homologs that form capsids and are secreted }$

A. Domain architectures of selected Capsid (CA)-containing mammalian Gag homologs compared to that of typical retrovirus and LTR retrotransposons. Each group of Gag homologs contains a distinct combination of predicted CA, Nucleocapsid (NC), Protease (PR), and Reverse Transcriptase (RT) domains. LTR, long terminal repeat; MA, matrix; IN, integrase.

B. Fraction of the total bacterially-produced protein that forms oligomers (>600 kD), as determined by size exclusion chromatography.

C. Representative negative stain transmission electron micrographs (TEM) of the *Mus musculus* (Mm) orthologues of the CA-domain containing proteins. Scale bar, 100 nm.

- D. Representative electron micrographs using cryogenic electron microscopy (cryoTEM) of a selected subset of the identified CA-domain containing proteins. Scale bar, 50 nm.
- E. Method for detecting extracellular forms of CA-domain containing homologs.
- F. Representative blots of CA-domain containing proteins in the cell-free fraction. CD81 was used as loading control for the ultracentrifuged cell-free fraction. Whole cell (W.C.) and VLP fraction blots for the endoplasmic reticulum marker CALNEXIN (CNX) ensure equal loading of whole cell protein and the purity of cell-free VLP fraction.
- G. Quantification of extracellular CA-domain containing proteins (as in F) based on n=3 replicates.

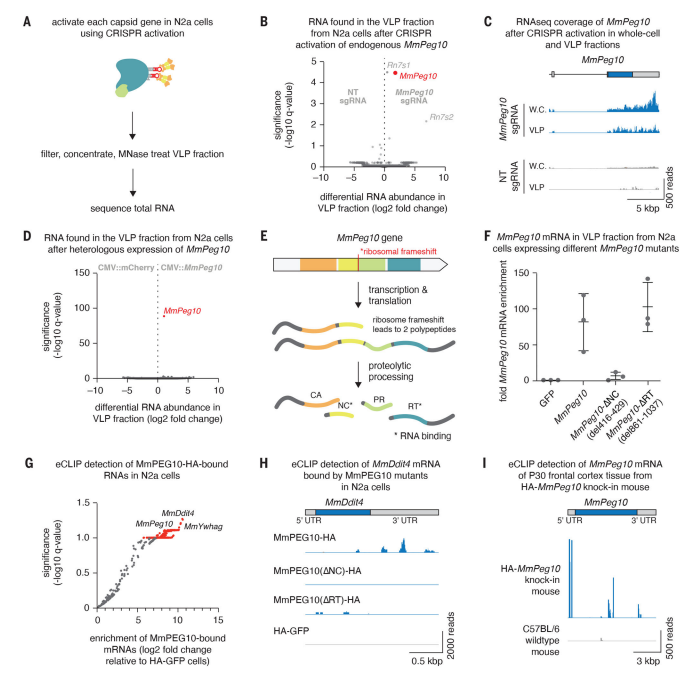


Fig 2. MmPEG10 protein and mRNA is secreted in vesicles by cells in vitro.

- A. Method for identifying nucleic acids that are secreted in the VLP fraction upon gene activation of CA-domain containing proteins.
- B. Differential RNA abundance and significance in the VLP fraction from N2a cells after CRISPR activation of endogenous *MmPeg10*.
- C. Alignment of sequencing reads showing sequencing coverage of the *MmPeg10* mRNA from (B).
- D. Differential RNA abundance and significance in the VLP fraction from N2a cells after heterologous transfection of *MmPeg10*. n=3 replicates.

E. The four domains of MmPEG10 are translated into two isoforms. These are self-processed by the PEG10 protease into separate domains, of which the NC and RT bind RNA.

- F. Fold enrichment of MmPeg10 mRNA compared to GFP in the VLP fraction from N2a cells transfected with wildtype MmPeg10 or deletions of the predicted nucleocapsid (NC) and reverse transcriptase (RT) domains.
- $G.\ Log_2$ fold change and significance of bound RNAs from eCLIP data comparing HA-GFP to WT MmPEG10-HA.
- H. Representative sequencing alignment histogram of the *MmDdit4* locus generated from eCLIP of N2a cells transfected with wildtype or mutant *MmPeg10*.
- I. Representative sequencing alignment histogram of the MmPeg10 locus generated from eCLIP data of n = 3 HA-PEG10 and n = 3 untagged animals.

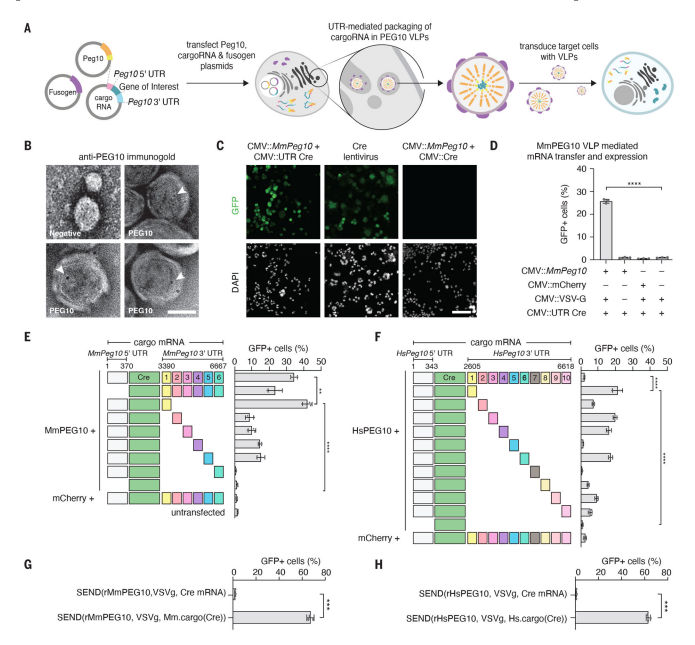


Fig 3. Flanking mRNA with MmPeg10 5' and 3' UTRs enables functional intercellular transfer of mRNA into a target cell.

A. Schematic showing reprogramming MmPEG10 for functional delivery of a cargo RNA flanked with the *MmPeg105*′ and 3′ UTRs (hereafter, "cargo(RNA)")

B epresentative TEM micrographs of VLP fraction immunogold labeled for MmPEG10.

Text labels indicate transfection of cells with MmPeg10 or mock (negative). Arrowheads indicate gold labeling. Scale bar, 50 nm.

C. Representative images of loxP-GFP N2a cells treated with VSVg pseudotyped MmPEG10 VLPs produced by transfecting Mm.cargo(Cre) or Cre mRNA, and a lentivirus encoding Cre. Scale bar, 100 um.

D. Functional transfer of RNA into loxP-GFP N2a cells mediated by VSVg pseudotyped MmPEG10 VLPs. Data quantified by flow cytometry 72 hours after VLP addition, n=3 replicates.

- E. Functional transfer of RNA into loxP-GFP N2a cells mediated by VSVg pseudotyped VLPs produced with MmPeg10 or mCherry and Mm.cargo(Cre) constructs encoding tiles of the MmPeg103'UTR. Data quantified by flow cytometry 72 hours after VLP addition, n=3 replicates.
- F. Functional transfer of RNA into loxP-GFP N2a cells mediated by VSVg pseudotyped VLPs produced with HsPEG1010 or mCherry and Hs.cargo(Cre) constructs encoding tiles of the HsPeg103'UTR. Data quantified by flow cytometry 72 hours after VLP addition, n = 3 replicates.
- G. Functional transfer of RNA into loxP-GFP N2a cells mediated by VSVg pseudotyped VLPs produced with r*MmPeg10* and Mm.cargo(Cre) or Cre mRNA. Data quantified by flow cytometry 72 hours after VLP addition, n = 3 replicates.
- H. Functional transfer of RNA into loxP-GFP N2a cells mediated by VSVg pseudotyped VLPs produced with rHsPeg10 and Hs.cargo(Cre) or Cre mRNA. Data quantified by flow cytometry 72 hours after VLP addition, n = 3 replicates. For all panels *p<0.05, **p<0.01, ****p<0.001, ****p<0.0001, One-Way ANOVA.

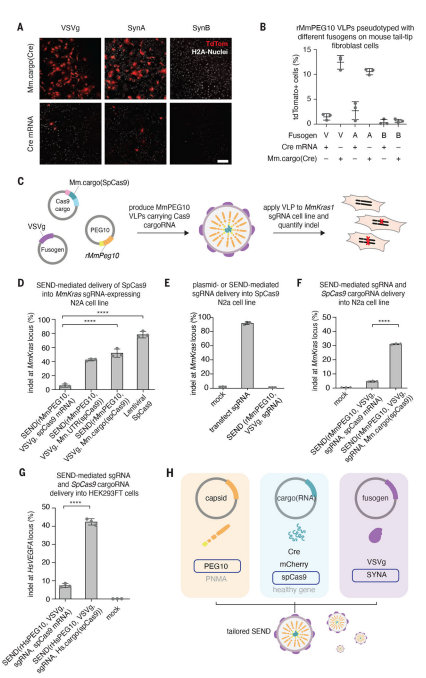


Fig. 4. SEND is a modular system capable of delivering gene editing tools into human and mouse cells.

- A. Representative images demonstrating functional transfer of Mm.cargo(Cre) or Cre mRNA in rMmPEG10 VLPs pseudotyped with VSVg (V), MmSYNA (A), or MmSYNB (B) in Ai9 (loxP-tdTomato) tail tip fibroblasts. Scale bar, $200 \, \mu m$.
- B ercent of tdTomato positive cells out of the total number of H2A stained nuclei from high content imaging of n=3 replicates of (A).
- C. Schematic representing the retooling of SEND for genome engineering.
- D. Indels at the *MmKras* locus in *MmKras1*-sgRNA-N2a cells treated with SEND (VSVg pseudotyped rMmPEG10 VLPs) containing either SpCas9 mRNA, Mm.UTR(SpCas9), or

Mm.cargo(SpCas9) and a lentivirus encoding SpCas9. Indels quantified by NGS 72 hours after VLP or lentivirus addition, n=3 replicates.

E. Indels at the mouse *MmKras* locus in a constitutively expressing SpCas9 N2a cell line either transfected with a plasmid carrying the *MmKras* sgRNA or treated with SEND (rMmPEG10, VSVg, *MmKras* sgRNA). Indels quantified by NGS after 72 hours, n=3 replicates.

F. Indels at the *MmKras* locus in N2a cells treated with SEND (VSVg pseudotyped rMmPeg10 SEND VLPs) containing either SpCas9 mRNA or Mm.cargo(SpCas9) and sgRNA. Indels quantified by NGS 72 hours after VLP addition, n=3 replicates.

G. Indels at the *HsVEGFA* locus in HEK293FT cells treated with SEND (VSVg pseudotyped rHsPEG10 VLPs) containing either SpCas9 mRNA or Hs.cargo(SpCas9) and an unmodified sgRNA. Indels determined by NGS 72 hours after VLP addition, n=3 replicates.

H. SEND is a modular delivery platform combining an endogenous Gag homolog, cargo mRNA, and fusogen, which can be tailored for specific contexts.