A peptide derived from the N-terminus of charged multivesicular body protein 6 (CHMP6) promotes the secretion of gene editing proteins via small extracellular vesicle production

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ABSTRACT
Extracellular vesicles (EVs) are a promising new therapeutic platform. However, the low cargo-loading efficiency limits their clinical translation. In this study, we developed a high-yield EV cargo-loading device and explored its ability to encapsulate gene editing proteins. A series of fusion protein-based systems were constructed and their cargo loading efficiencies were compared by a NanoGlo luciferase assay. A myristoylated (Myr) peptide tag cloned from the N-terminal region of charged multivesicular body protein 6 (CHMP6), termed Myr(CHMP6), outcompeted CD9, ARRD1, and other short polypeptides as an active packaging device. As determined by nanoparticle tracking analysis and transmission electron microscopy, the overexpression of Myr(CHMP6) increased small EV (sEV) production in Lenti-X 293T cells without altering sEV morphology. The high passive packaging efficiency of Myr(CHMP6) was also elucidated for unmodified cargo loading. Western blotting revealed that Myr(CHMP6) facilitated the loading of Cre and Cas9 into sEVs without the generation of packaging device-cargo fusion proteins. Furthermore, Myr(CHMP6)-modified sEVs loaded with Cre or Cas9 promoted gene-editing in recipient cells, as observed using a fluorescence reporter system. Subsequent investigation demonstrated a dose-dependent effect of Myr(CHMP6) tag-induced cargo-loading. Mechanistically, N-myristoylation alone was necessary but not sufficient for the effective packaging of proteins into EVs. Thus, our results indicated that Myr(CHMP6) induces sEV production and may be effective in loading gene editing proteins into sEVs for therapeutic purposes.

1. Introduction
Extracellular vesicles (EVs) are cell-derived vesicles capable of mediating intracellular communication. They are promising next-generation nanomedicines with intrinsic cargo-loading ability, nano-scale particle sizes, high biocompatibility, and targeting potential [1–3]. However, the limited productivity and cargo-
loading efficiency of native EVs impede their application as a therapeutic platform.

EVs have recently been modified by re-engineering various EV-enriched proteins (EPPs), where EEPs are covalently connected to or selectively interact with cargoes, mediating their assembly into EVs. For example, an ARRDC1-p53 fusion structure could encapsulate p53 into EVs [4]. CD9 has also been fused to a blue light-controlled protein–protein interaction module for soluble protein loading [5]. Despite progress in the development of vesicle-mediated cargo delivery platforms, few studies have compared the cargo-loading efficiencies of different fusion protein-based methods, namely, active packaging devices [6–9]. Furthermore, active packaging requires design and manufacturing of new recombinant structures for each cargo of interest, limiting the practical application of engineered EVs.

Another widely used strategy for EV cargo loading is passive packaging. Cargoes are loaded into EVs in a dose-dependent, nonselective manner by overexpression. Despite relatively low yields, passive packaging remains the simplest method, as it does not require the design and construction of new recombinant proteins. This also allows us to encapsulate wild-type cargoes into EVs without fusing them to exogenous constructs, thus, avoiding changes in protein structure and subcellular distributions, and/or impaired cargo function caused by the fusion protein-based strategy.

Various post-translational modifications (PTMs) show regulatory effects on EV protein component sorting [10,11], prompting us to investigate whether polypeptides with PTMs can be re-engineered and used as an EV cargo-loading device. For example, myristoylation sites are found in essential components of viruses or endosomal sorting complexes required for transport (ESCRT), such as in the N-terminal arm of Tacaribe virus (TCRV) Z protein [12], human immunodeficiency virus (HIV) Gag protein [13], and ESCRT-III components CHMP3 [14] and CHMP6 [15]. Prenylation at C-terminal regions of the Rab GTPases is known to mediate their regulatory effect on vesicular transport [16–19]. In this study, after evaluating various candidate proteins with PTMs, we developed a novel packaging device, a 12 amino acid-long polypeptide from the N-terminus of the CHMP6 protein, termed Myr(CHMP6). We aimed to investigate whether Myr(CHMP6) is more efficient than traditional packaging methods using CD9- or ARRDC1- fusion proteins, and to explore its ability in delivering functional gene-editing proteins.

2. Materials and methods

2.1. Cell culture

Lenti-X 293T cells (632,180; Clontech, Mountain View, CA, USA) were maintained in Dulbecco’s modified Eagle medium (DMEM) with high glucose (HyClone, Logan, UT, USA), 10% fetal bovine serum (FBS; Gibco, Waltham, MA, USA), 100 units/mL penicillin G sodium, and 100 µg/mL streptomycin sulfate (Gibco).

2.2. Plasmids and transfection

Nluc-3×FLAG containing SalI and BsrGI site was synthesized (Genewiz, Guangzhou, China) and inserted into pLV-mCherry (#36,084; Addgene, Watertown, MA, USA) using In-Fusion (Takara) to generate pLV-mCherry-Nluc-3×FLAG. Genes encoding CD9, ARRDC1, mono-ubiquitin (Ub), and sonic hedgehog (SHH) were cloned from the Lenti-X 293T cDNA library. Genes encoding polypeptides with myristoylation (Myr) or prenylation (Prenyl) sites were synthesized (Genewiz) or PCR-amplified from primer pairs complementary at the 3′ ends to generate primer dimers and then inserted at the 5′ end or 3′ end of mCherry-Nluc-3×FLAG with In-Fusion to generate recombinant plasmids encoding fusion protein-based cargo-loading systems. GFP was cloned from pGreenFire1-NF-kB (TR012PA-1; System Biosciences, Palo Alto, CA, USA) and the HA tag was synthesized (Genewiz). Overlap extension PCR was performed to generate GFP-HA, followed by In-Fusion insertion into pLV-mCherry (cut with SalI and BamHI) to generate pLV-GFP-HA or into Myr(CHMP6)-mCherry-Nluc-3×FLAG (cut with SalI and Spel) to generate pLV-Myr(CHMP6)-GFP-HA. Cre-3×FLAG was synthesized and inserted into pLV-mCherry (cut with SalI and XbaI) to generate pLV-Cre-3×FLAG. A G2A mutation was introduced in the Myr(CHMP6) tag using
mutagenic primers. A fluorescence reporter system segment loxp-target-mCherry-stop-target-loxp-GFP-stop containing BsmBI and SalI sites was synthesized (Genewiz) and inserted into pLV-mCherry with T4 DNA ligase (Thermo Scientific, Waltham, MA, USA) to generate pLV-Cre/Cas9-reporter. A segment encoding a U6 promoter and a sgRNA, termed sg-1, was synthesized (Genewiz) and In-Fusion-inserted into pLV-Cre/Cas9-reporter (cut with Clal) to generate pLV-Cre/Cas9-reporter-sg-1. Inserted DNA fragments were confirmed by sequencing. Cells were transfected using Lipo293 Transfection Reagent (Beyotime, Shanghai, China) at 60–80% confluency according to the manufacturer protocol. Primers used to construct plasmids are listed in Table S1.

2.3. Microscopy for cell imaging

Transfected cells were seeded on 35 mm confocal dishes (Biofil, Guangzhou, China), cultured overnight, and examined using a LSM 780 laser scanning microscope (ZEISS, Oberkochen, Germany). GFP was excited with a 488 nm laser while mCherry was excited with a 561 nm laser.

2.4. Luciferase activity assay

Cells were seeded into 24-well plates. Culture medium was changed 24 h after transfection and collected after another 24 h, followed by sequential centrifugation at 300 × g for 10 min, 2000 × g for 10 min, and 10,000 × g for 30 min at 4°C to remove large debris. The supernatant was loaded onto a white 96-well plate (Corning, NY, USA), and 50 μL of Nano-Glo Luciferase Assay Substrate (Promega, Madison, WI, USA) diluted 1:50 with the provided buffer was added to each reaction. Cells in 24-well plates were lysed with 300 μL of 1× Passive Lysis Buffer (Promega) per well and centrifuged at 12,000 × g for 10 min. Nluc activity was measured in 50 μL of cell lysate supernatants diluted 1:1000.

2.5. sEV isolation

The sEVs were isolated via differential centrifugation [20,21]. Cells in 150 mm culture plates were washed and the culture medium was changed to 25 mL of fresh FBS-free DMEM with high glucose (HyClone) 24 h after transfection. After another 24 h, the culture medium was collected and sequentially centrifuged at 300 × g for 10 min, 2000 × g for 10 min, and 10,000 × g for 30 min, followed by filtration with a 0.22 μm syringe filter (Merck Millipore, Billerica, MA, USA). sEVs were pelleted by ultracentrifugation at 34,100 rpm (~120,000 × g) for 70 min at 4°C in an Optima XE-100 ultracentrifuge (Beckman Coulter, Brea, CA, USA) with the type 70 Ti rotor. Pellets were washed with phosphate-buffered saline (PBS; HyClone), re-ultracentrifuged, and resuspended with PBS (HyClone) or RIPA buffer (Thermo Scientific, Waltham, MA, USA).

2.6. sEV characterization

sEVs isolated from 25 mL of the culture supernatant were resuspended in 175 μL of PBS. For a nanoparticle tracking analysis (NTA), sEVs were analyzed using the NanoSight NS300 (Malvern Panalytical, Malvern, UK). For sEV visualization, 5–10 μL of resuspended sEVs was stained with phosphotungstic acid (PTA) on a copper grid and examined under a transmission electron microscope (JEM-1200EX; JEOL, Tokyo, Japan).

2.7. Western blotting

Cells or sEVs were lysed on ice in RIPA buffer (Thermo Scientific) supplemented with protease and phosphatase inhibitors (Beyotime) and then cleared by centrifugation at 12,000 × g for 10 min. The cell lysates were quantified with a Pierce BCA protein assay kit (Thermo Scientific) according to the manufacturer protocol. The lysates were then resuspended in SDS-PAGE sample loading buffer (Beyotime). Cell lysates were loaded into SDS-PAGE gels based on equal protein amount (2 μg), while sEV lysates were loaded based on equal volume (20 μL) to reflect sEV contents from the same number of cells. Samples were separated by SDS-PAGE and transferred onto a PVDF membrane (Merck Millipore). Blots were probed with primary antibodies in Tris-buffered saline containing 0.025% Tween-20 (TBST; Asegene, Guangzhou, China) containing 5%
bovine serum albumin (BSA; Thermo Scientific), followed by incubation with HRP-conjugated anti-rabbit antibody or anti-mouse antibody. Primary antibodies are listed in Table S2. Protein bands were visualized with an ECL detection reagent (Merck Millipore) according to the manufacturer protocol and captured with the Amersham Imager 600 (General Electric), and quantified using ImageJ (NIH, Bethesda, ML, USA).

2.8. Gene editing protein delivery assay

Donor cells were co-transfected with packaging device Myr(CHMP6)-GFP-HA or its control GFP-HA, along with cargo-encoding plasmid pLV-Cre-3×FLAG, LentiCRISPRv2 (#52,961; Addgene), or the control pLV-mCherry-Nluc-3×FLAG. sEVs in the culture medium were isolated as described in Section 2.5. Recipient cells were incubated with isolated sEVs for 12 h and then transiently transfected with pLV-Cre/Cas9-reporter-sg-1. After another 48 h, the transfected reporter cells were examined using a LSM 780 laser scanning microscope (ZEISS).

2.9. Statistical analysis

Results are expressed as means ± standard error of the mean (SEM) and analyzed using SPSS 25.0 (IBM, Armonk, NY, USA). Comparisons between two groups were performed by unpaired two-tailed Student’s t-tests. Comparisons among three or more groups were performed by one-way analysis of variance (ANOVA) and Bonferroni correction. Values p < 0.05 were considered statistically significant.

3. Results

In this study, we aimed to develop a high-yield EV cargo-loading device. The Myr(CHMP6)-Cargo fusion structure was found to have significantly higher cargo-loading efficiencies than those of traditional CD9- or ARRDC1-Cargo loading systems. By co-expression with Myr(CHMP6)-GFP-HA, cargoes can be effectively loaded into sEVs without fusing to the packaging device. Our data indicated that Myr(CHMP6) can facilitate Cre and Cas9 protein-loading into sEVs and promote gene-editing in the recipient cells, supporting further applications in EV-based therapy. Lastly, we demonstrated a dose-dependent effect and the essential role of a Gly² myristoylation site in Myr(CHMP6)-mediated cargo loading.

3.1. Myr(CHMP6) outcompetes various protein tags as an active packaging device

To investigate whether polypeptides with PTMs can be re-engineered as EV cargo-loading devices, we cloned polypeptides from different genes containing potential PTM sites, including Ubi, Myr, Prenyl, or cholesterol (CHOL) attachment sites (Figure 1). Traditional active packaging methods, based on CD9 and ARRDC1, were used as positive controls. A cargo protein composed of mCherry, nanoluciferase (Nluc), and a 3×FLAG tag was designed for visualization and sensitive quantification.

After transfection, Ubi-Cargo, Myr(CHMP3)-Cargo, Cargo-Prenyl(RAB5A), Cargo-Prenyl(RAB7A), and Cargo-Prenyl(RAB11) were uniformly distributed in cells, similar to the control (Figure 2(a) and Figure S1). Cargoes fused with CD9 were localized at the plasma membrane and in the perinuclear region. ARRDC1-Cargo, Myr(TCRV)-Cargo, Myr(Gag)-Cargo, Myr(CHMP6)-Cargo, Cargo-Prenyl(KRAS), and Cargo-CHOL(SHH) showed different levels of enrichment at the plasma membrane and punctate regions in the cytoplasm (Figure 2(a) and Figure S1).

A NanoGlo luciferase assay was then performed on culture medium (Figure 2(b)) and cell lysates (Figure S2). The extracellular-to-intracellular ratio (E/I ratio) was calculated by dividing Nluc activity in the culture medium by Nluc activity in cell lysates of EV-producing cells to investigate the cargo loading efficiency (Figure 2(c)). Fusion of CD9, ARRDC1, Myr(TCRV), Myr(GAG), Myr(CHMP6), Prenyl(KRAS), or CHOL(SHH) to the cargo protein induced a >4-fold increase in cargo release and >40-fold increase in the E/I ratio. Myr(CHMP6) was the most effective active-packaging device (Figure 2(b, c)). After a proteinase K treatment on the culture medium, Myr(CHMP6)-Cargo still significantly increased cargo release and the E/I ratio (Figure S3). To further verify the EV-associated cargo
loading, we examined the Nluc activities of CD63⁺ exosomes, which is an important subgroup of EVs. Myr(CHMP6)-Cargo induced a > 85-fold increase in cargo release in CD63⁺ exosomes purified by CD63 capture beads. And a > 578-fold increase in the exosome/cell cargo ratio. This result demonstrates that Myr(CHMP6) can promote cargo loading into CD63⁺ exosomes (Figure S4).

3.2. Myr(CHMP6) promotes sEV production

Lenti-X 293T cells were transfected with Myr(CHMP6)-GFP-HA plasmids or subjected to mock transfection to determine whether the constructed packaging device affects sEV production. sEVs isolated by ultracentrifugation (Figure 3(a)) were examined by NTA, revealing that transfection with Myr(CHMP6)-GFP-HA increases sEV production (p = 0.002, Figure 3(b,c)). Furthermore, NTA and electron microscopy showed that the majority of Myr(CHMP6)-GFP-HA-induced sEVs were identical in size and morphology to those produced by mock control cells (Figure 3(b,d)). These results suggest that Myr(CHMP6) induces cargo protein secretion at least partially by promoting sEV production.
3.3. Myr(CHMP6) facilitated gene-editing protein loading into sEVs

We performed a NanoGlo luciferase assay to compare the efficiency of Myr(CHMP6)-mediated packaging of mCherry-Nluc-3xFLAG (the cargo protein) by active and passive loading (Figure 4(a,b), and Figure S5). The fusion protein Myr(CHMP6)-Cargo was used to examine the efficiency of active packaging, while co-expression of Myr(CHMP6)-GFP-HA and the cargo represents passive packaging. The Myr (CHMP6) co-expression strategy, though not as effective as the fusion-based method, significantly increased the productivity and E/I ratio of cargoes (all p < 0.01), indicating an effective passive cargo-loading device.

Figure 2. Myr(CHMP6) outcompetes various protein tags as an active packaging device. (a) Fluorescence images of cargoes fused to different packaging devices. Ubi-Cargo, Myr(CHMP3)-Cargo, Cargo-Prenyl(RAB5A), Cargo-Prenyl(RAB7A), and Cargo-Prenyl (RAB11) were uniformly distributed in cells, similar to the control group. Cargoes fused to CD9, ARRDC1, Myr(TCRV), Myr(Gag), Myr (CHMP6), Prenyl(KRAS), or CHOL(SHH) showed various altered distributions. (b) A NanoGlo luciferase assay was performed to evaluate different fusion protein-based active packaging systems. Nluc activity in the culture medium (CM) represents cargo release. Scale bar represents 10 μm. (c) The E/I ratio was calculated by dividing Nluc activity in the culture medium by Nluc activity in cell lysates (CL) of EV-producing cells. E/I ratio fold change represents fold change values of extracellular-to-intracellular RLU ratio. Bar graphs present fold change values compared to the first group as means ± standard error of the mean (SEM) (N ≥ 3). One-way ANOVA and Bonferroni correction were used for multiple comparisons to obtain adjusted p-values. *, Compared with Cargo group, Bonferroni adjusted p < 0.05; #, Compared with CD9-Cargo group, Bonferroni adjusted p < 0.05; †, Compared with ARRDC1-Cargo group, Bonferroni adjusted p < 0.05; $, Compared with Myr(CHMP6)-Cargo group, Bonferroni adjusted p < 0.05.
Exogenous cargoes for gene editing require nuclear transport, suggesting that passive loading may be an advantageous approach in gene therapy. To evaluate whether the co-expression of Myr(CHMP6)-GFP-HA can facilitate the packaging of proteins with nuclear localization signals (NLS), we further transfected Lenti-X 293T cells with FLAG-tagged Cre or Cas9 with Myr(CHMP6)-GFP-HA. Western blotting showed enrichment of the exosomal markers Alix and CD81 but not the endoplasmic reticulum protein Calnexin in sEVs (Figure 4(c,e)), reflecting high-quality sEV isolation. An increase of Alix, CD81, and GAPDH (one of the top five most common proteins associated with EVs [22]) in sEVs was observed upon overexpression of Myr(CHMP6) (Figure 4(c,e)), supporting the increase of sEV production. Myr(CHMP6)-tagged GFP-HA was enriched in sEVs, along with co-transfected Cre or Cas9 (Figure 4(c-f)), supporting the nuclear protein packaging ability of Myr(CHMP6).

3.4. Functionalized Myr(CHMP6)-sEVs enables efficient gene editing

A fluorescence reporter system was constructed to explore whether the Myr(CHMP6)-engineered-sEVs promotes delivery of functional macromolecular proteins for gene editing (Figure 5(a)). Donor cells were transfected with a packaging
device (or a control plasmid) and a cargo protein. sEVs from donor cells that express packaging device Myr(CHMP6)-GFP-HA are termed Myr(CHMP6)-sEVs. sEVs from donor cells that express GFP-HA are termed control sEVs. Recipient cells were subjected to sEV treatment, followed by transfection of the fluorescence reporter system plasmid pLV-Cre/Cas9-reporter-sg-1. The gene editing events can be observed as the mCherry^+^GFP^−^ recipient cells would be recombined to mCherry^−^GFP^+^ cells (where all mCherry is knocked out) or mCherry^+^GFP^+^ cells (where a proportion of mCherry is knocked out) between indicated LoxP/sg-1 binding sites by bioactive Cre or Cas9. As confirmed by fluorescence microscopy, recipient cells treated with Myr(CHMP6)-sEVs loaded with Cre or Cas9 showed a red-to-green fluorescent shift (Figure 5(b,c)). In contrast, low GFP expression can be found in recipient cells treated with control sEVs loaded with Cre or Cas9 (Figure 5(b,c)). Myr(CHMP6)-sEVs or control sEVs without gene editing proteins are not able to induce the green fluorescent signal in recipient cells (Figure S6), precluding false positive result induced by the packaging device. These results demonstrate the functionality of gene editing proteins in recipient cells delivered by Myr(CHMP6)-sEVs.
3.5. Dose-dependent effect of Myr(CHMP6) tag on cargo loading

Lenti-X 293T cells were transfected at different packaging device-to-cargo ratios to evaluate the relationship between Myr(CHMP6) expression levels and cargo protein transport. In this experiment, Myr(CHMP6)-GFP-HA was used as the passive packaging device for the cargo mCherry-Nluc-3×FLAG. Considering that the total plasmid amount may affect transfection efficiency, we used different doses of GFP-HA or mCherry to equalize the mass of Myr
(CHMP6)-GFP-HA or mCherry-Nluc-3×FLAG, respectively. Interestingly, for an equal amount of cargo transfection, Nluc activity in cell lysates decreased as Myr(CHMP6) transfection increased (Figure 6(a)). The introduction of Myr(CHMP6) to the co-transfection system significantly increased cargo release (all \( p < 0.01 \), Figure 6(b)). The E/I ratio increased constantly with the increase in Myr(CHMP6) (Figure 6(c)). With equal doses of packaging device, cargo release increased with the level of cargo transfection (Figure 6(e)), while the E/I ratio did not significantly fluctuate (all \( p > 0.05 \), Figure 6(f)). These results demonstrated a dose-dependent effect of Myr(CHMP6)-tag-induced cargo loading.

Figure 6. Dose-dependent effect of Myr(CHMP6)-tag-induced cargo-loading. Lenti-X 293T cells in 24-well plates were transfected with cargo plasmids and the packaging device or control plasmids at the indicated doses. A NanoGlo luciferase assay was performed 48 h after transfection. (a-c) When cargo was fixed at 500 ng, Nluc activities in CL, culture medium (CM), and extracellular-to-intracellular RLU ratio (E/I ratio) were measured at different packaging device transfection doses. (d-f) With Myr(CHMP6)-GFP-HA fixed at 500 ng, Nluc activities in CL, CM, and E/L ratio were measured at different cargo transfection doses. Data are presented as means ± SEM (N = 3). One-way ANOVA and Bonferroni correction were used for multiple comparisons to obtain adjusted \( p \)-values. \( * \)p < 0.05, \( ** \)p < 0.01, \( *** \)p < 0.001.
3.6. The Gly\textsuperscript{2} myristoylation site is essential for the membrane affinity and packaging efficiency of Myr(CHMP6)

Myr(CHMP6)(G2A) mutants lacking the putative N-myristoylation site were generated to investigate the role of this modification (Figure 7(a,b)). While Myr(CHMP6)-Cargo distributed in the cytoplasm with enrichment at membrane structures (Figure 2(a), Figure 7(c) and Figure S1), the mutant Myr(CHMP6)(G2A)-Cargo was uniformly distributed in cells (Figure 7(c) and Figure S7), indicating that the G2A mutation decreased the membrane affinity of the Myr(CHMP6) tag. When co-transfected with the cargo, Myr(CHMP6)-GFP-HA showed a dot-like distribution in the cytoplasm and enrichment at the plasma membrane. The mCherry of the cargo remained generally freely distributed (Figure 7(d) and Figure S7). Interestingly, at sites with intense GFP signals, mCherry signals decreased and did not overlap with GFP, suggesting active packaging activity at the Myr(CHMP6)-modified locus (Figure 7(d) and Figure S7). In contrast, the localization of GFP-HA and the Myr(CHMP6)(G2A)-GFP-HA mutant was identical to that of the cargo (Figure 7(d) and Figure S7). Additionally, the G2A mutation abolished the ability of Myr(CHMP6) to induce cargo loading (Figure 7(e,f)). Taken together, these data demonstrated that Gly\textsuperscript{2} is essential for Myr(CHMP6)-induced cargo loading.

Figure 7. The Gly\textsuperscript{2} myristoylation site is essential for the membrane affinity and packaging efficiency of Myr(CHMP6). (a) Schematic of the control cargo mCherry-Nluc-3xFLAG, active packaging device Myr(CHMP6)(G2A)-Cargo, and its mutant Myr(CHMP6)(G2A)-Cargo without the myristoylation site. (b) Schematic of GFP-HA, Myr(CHMP6)-GFP-HA, or the Gly\textsuperscript{2}-mutated Myr(CHMP6)(G2A)-GFP-HA, and its representative fluorescence images of control cargo mCherry-Nluc-3xFLAG, active packaging device Myr(CHMP6)(G2A)-Cargo, and its mutant Myr(CHMP6)(G2A)-Cargo without the myristoylation site. Scale bar represents 10 μm. (c) Representative fluorescence images of cargo co-expressed with GFP-HA, Myr(CHMP6)-GFP-HA, or the mutated Myr(CHMP6)(G2A)-GFP-HA. Scale bar represents 10 μm. (e) NanoGlo luciferase assay of Myr(CHMP6) and its mutant Myr(CHMP6)(G2A)-induced active loading. Data are presented as means ± SEM (N = 3). One-way ANOVA and Bonferroni correction were used to obtain adjusted p-values for multiple comparisons. ***p < 0.001.
4. Discussion

We provide the first evidence that Myr(CHMP6), a short polypeptide tag, is an effective ‘EV production booster’ and can be used as both an active and passive cargo-loading device. As an active packaging device, the efficiency of Myr(CHMP6) outcompeted that of conventional methods utilizing CD9, ARRDC1, or other short protein tags. As a passive packaging device, the co-expression of Myr(CHMP6) enabled the encapsulation of unmodified cargoes, including Cre and Cas9, and induced gene editing in recipient cells. In light of the advantages of EV-based therapy, our packaging device may improve the cargo-loading ability and promote the clinical translation of engineered EVs.

Various methods, particularly active packaging, have been used to increase the cargo-loading efficiency of EVs. Constitutive proteins of EVs [4,5,23], virus-like particles [24,25], and other vesicle-enriched structures were re-engineered for direct or indirect interaction with modified cargoes. Ubiquitin is recognized by ESCRT [26,27]; however, our mono-ubiquitin tag did not substantially improve the packaging efficiency. Since EVs are largely enriched with cholesterol [28], we also cloned SHH for cholesterol attachment [29], which showed a high active packaging efficiency. The functions of many viral proteins, ESCRTIII components, and endosomal proteins are dependent on N-myristoylation [30–32] or prenylation [17]. Thus, we cloned the N- or C-terminus of EEPs with these lipidation sites as well as a prenylated KRAS, a truncated protein widely used as a membrane anchor [33], and fused these structures to the cargo (Figure 1). Some of these lipidation tags, namely Myr(TCRV), Myr(Gag), Myr(CHMP6), and Prenyl(KRAS), were more efficient than CD9 or ARRDC1 (Figure 2(b,c)). Although Myr(TCRV), Myr(Gag), Myr(CHMP3), and Myr(CHMP6) all included Gly2 myristoylation sites, their packaging efficiencies and membrane affinity differed (Figure 2). This observation demonstrates that downstream amino acid sequences, in addition to the myristoylation site itself, impact the cargo-loading efficiency of the packaging device. The Myr (CHMP6) tag was the most efficient active packaging device evaluated in this study (Figure 2(b,c)). Its packaging ability was further verified in proteinase K-treated culture medium and purified CD63+ exosomes (Figure S3 and Figure S4).

The majority of effective packaging devices interact with membranes. The tetraspanin CD9 has transmembrane domains. The arrestin-like domain of ARRDC1 interacts with the plasma membrane and regulates ARMM budding [34]. Lipidation tags regulate membrane trafficking and protein secretion [35]. In the present study, we found that few active-packaging structures remain in the nucleus (Figure 2(a)). This is an important limitation because therapeutic cargoes for gene therapy usually require nuclear transport. Further, the fusion of these devices inevitably changed the native sequence of cargoes. The packaging device itself as well as interacting molecules would attach to the cargo, potentially disrupting the functionality. For example, ARRDC1-p53 actively packaged in EVs attached to plasma membrane and was ubiquitinated [4], which might reduce the nuclear transport of p53 and preferentially lead to cargo degradation in target cells, thus impairing the overall bioactivity of the engineered EVs. Therefore, structural and functional validation of the cargoes are required for each new design, presenting an obstacle to the implementation of EV-based therapeutic technology. For the above reasons, simple overexpression is a common strategy for the loading of wild-type nuclear or cytosolic cargoes into EVs [36–38], which is rather low-yield, without increasing EV production [4,5,39]. To address these issues, we developed a simple but high-yield passive packaging strategy for unmodified wild-type cargoes. Since the Myr (CHMP6) tag is an efficient active-packaging device, we evaluated its utilization for passive-packing. Transfection of Lenti-X 293T cells with Myr(CHMP6)-GFP-HA significantly increased sEV production, without changing their morphology, demonstrating that it acts as an ‘EV production booster’ (Figure 3). Cargoes co-expressed with Myr(CHMP6) were effectively encapsulated (Figure 4(a,b)).

Cre/Loxp and CRISPR/Cas9 editing systems are unique technologies for genetic manipulation [40–42]. Many studies have attempted to encapsulate Cre or Cas9 into EVs by active loading. Cre, Cas9, or sgRNAs have been re-engineered to interact with EEPs or other EV-enriched structures via...
fusion with various modules [25,39,43–45]. Taking advantage of passive-loading, our Myr(CHMP6)-co-expression strategy enabled effective Cre and Cas9 encapsulation without cargo re-engineering (Figure 4). Notably, Myr(CHMP6) induced an increase in exosomal markers Alix and CD81, as well as GAPDH, which is reported to regulate EV biogenesis and binds to EV surface [46], supporting the ability of Myr(CHMP6) in promoting EV production. Furthermore, sEVs loaded with Cre or Cas9 were used to treat recipient cells expressing a fluorescence reporter system to induce gene editing. The gene-editing events in recipient cells were observed via fluorescence microscopy. When pre-treated with Myr(CHMP6)-sEVs, more recipient cells expressed GFP, indicating a higher efficiency of mCherry knock-out (Figure 5 and Figure S6). With these results, we demonstrate Myr(CHMP6) as an efficient tool to increase functional protein loading, even in the presence of NLS, into sEVs, with potential applications for gene therapy.

The dose-dependent effect was examined to investigate how Myr(CHMP6) affects protein trafficking. The results showed that the E/I ratio constantly increased with Myr(CHMP6) transfection, but not the amount of cargo transfected (Figure 6(c,f)), elucidating the role of Myr(CHMP6) in regulating cargo loading efficiency. In contrast, the amount of cargo released into the culture medium is related to transfection dose of both Myr(CHMP6) and the cargo (Figure 6(b,e)). Previous studies have provided evidence that N-myristoylation can target the AMP-activated protein kinase (AMPK) to lysosomes [47]. Our results showed a decrease of Nluc activity in cell lysates as well as the culture medium at the highest dose of Myr(CHMP6) transfection (Figure 6(a,b)). A possible explanation for this might be that the high dose of Myr(CHMP6) tag targeted cargo proteins to lysosomes and thus enhanced cargo degradation.

The Gly7 site of CHMP6 is crucial for its myristoylation, subcellular distribution and interaction with ESCRT components [15]. Our results showed that the G2A-mutated Myr(CHMP6) tag was uniformly distributed in the cytoplasm and nuclei, and it was not efficient in cargo loading (Figure 6 and Figure S7). Additionally, different peptide tags, although all contained N-myristoylation sites, had different packaging ability (Figure 2(b,c)), indicating that N-myristoylation alone was necessary but not sufficient for the effective packaging of proteins into EVs. Interestingly, we observed that cells transfected with Myr(CHMP6) are round and smaller than the control (Figure 6(d) and Fig S7). It was demonstrated that proteins with lipid affinity can regulate membrane shape by altering lipid bilayer force balance [48]. Since Myr(CHMP6)-tagged proteins showed high membrane affinity (Figure 2(a), Figure 7(c,d), Fig S1 and Fig S7), they might also be able to change the cell morphology.

Limitations of this study should also be noted. First, we did not evaluate the specificity of the Myr(CHMP6)-GFP-HA co-expression strategy. A mass spectrometry analysis of highly purified Myr(CHMP6)-induced EVs is necessary to characterize the changes in sEV composition and test the specificity of our system. Second, we did not address the mechanism by which Myr(CHMP6) induced sEV production. Previous studies have suggested that myristoylated proteins have high raft affinity [49,50] and that lipid raft components are involved in EV biogenesis [51]. Studies of the relationship between Myr(CHMP6) and lipid raft behavior may be helpful to understand the mechanism underlying elevated sEV production. Third, our data demonstrated the packaging ability and functionality of Myr(CHMP6)-induced sEVs with Cre or Cas9 proteins. Further in vitro and in vivo studies are needed to investigate the metabolic kinetics and targeting of these types of engineered EVs.

5. Conclusion

Myr(CHMP6) independently boosted sEV production and thus acted as an effective cargo-loading device. Myr(CHMP6) co-expression facilitated the encapsulation of Cre and Cas9 into sEVs, which further promoted gene editing in recipient cells. These finding indicates that Myr(CHMP6), as an effective packaging device, could be further used in gene therapy.

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Author Contributions

W.F., G.W., J.F. and J.P. conceived, designed and developed the experiments; J.F., J.P, X.Z., Y.C., Y.Z., L.H., D.M. and Z.C. performed the experiments; J.F. and J.P. analyzed the data, contributed data visualization and prepared the original draft; W.F. and G.F. acquired funding, supervised the project, reviewed and edited the article. All the authors read, critically revised, and approved the final manuscript.

Disclosure statement

No potential conflict of interest was reported by the author(s).

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