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Self-assembly of a fluorescent virus-like particle for imaging in tissues with high autofluorescence†

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Virus-like particles (VLPs) are engineered nanoparticles that mimic the properties of viruses-like high tolerance to heat and proteases-but lack a viral genome, making them non-infectious. They are easily modified chemically and genetically, making them useful in drug delivery, enhancing vaccine efficacy, gene delivery, and cancer immunotherapy. One such VLP is Q β , which has an affinity towards an RNA hairpin structure found in its viral RNA that drives the self-assembly of the capsid. It is possible to usurp the native way infectious Q β self-assembles to encapsidate its RNA to place enzymes inside the VLP's lumen as a protease-resistant cage. Further, using RNA templates that mimic the natural self-assembly of the native capsid, fluorescent proteins (FPs) have been placed inside VLPs in a "one pot" expression system. Autofluorescence in tissues can lead to misinterpretation of results and unreliable science, so we created a single-pot expression system that uses the fluorescent protein smURFP, which avoids autofluorescence and has spectral properties compatible with standard commercial filter sets on confocal microscopes. In this work, we were able to simplify the existing "one-pot" expression system while creating high-yielding fluorescent VLP nanoparticles that could easily be imaged inside lung epithelial tissue.

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Introduction

Virus-like particles (VLPs) are genetically engineered nanoparticles that resemble infectious viruses but lack a viral genome and are not infectious.^{1–3} VLPs typically comprise many identical coat proteins whose self-assembly is templated by the mRNA generated in their recombinant expression. This assembly mimics the properties of the native virus, starting with the nucleation phase and the production of oligomers around the nucleic acids host.^{4–6} The exterior of VLPs displays various amino acids that can be used to fuse peptides or chemically conjugate different molecules through amide coupling, diazonium addition, and click chemistry.^{7–11} Their packing ability has been used to protect RNA by expressing and packing it inside or delivering the scaffold proteins using "one pot" expression systems.^{12,13} Their biocompatibility and easy

chemical and genetic modification have made them useful in drug delivery, enhancing vaccine efficacy,^{14–17} gene delivery,^{18–20} and cancer immunotherapy.^{21–27}

One such representative VLP is Q β , derived from a bacteriophage in the *Leviviridae* family. The VLP forms an icosahedral capsid comprising 180 identical coat protein (CP) subunits.²⁸ Self-assembly of the natural Q β has been thought to be driven by a specific RNA hairpin structure that has an affinity toward the native CPs and is found in the Q β 's viral RNA, positioned at the 5' end, making possible its interaction with CPs during protein synthesis on the bacterial ribosome.^{29,30}

In addition to chemical and genetic modification to the CP—which leads to new covalently bound surface functionality—work on VLPs has focused on using the ample void space as a haven for biomacromolecular cargo. For instance, encapsidating enzymes within the interior lumen of VLPs helps serve as a protease-resistant cage that protects the more sensitive enzymes.^{31,32} Two common ways to pack proteinaceous cargo inside VLPs are through an iterative disassembly-reassembly process or a "one pot" expression system.³³ The former disassembly-reassembly process is best understood as a "supramolecular assembly" process, where self-assembly to reform the capsid is aided by proteins and RNAs that act as templates. For instance, we have previously shown it is possible to encapsidate the small Ultra-Red Fluorescent Protein (smURFP) within Q β by reducing the capsid's

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disulfide bonds and forcing the disassembly of the capsid at low pH and high salt concentration. This is followed by a supramolecular assembly process aided by a 10× fold excess of the smURFP cargo at neutral pH.³⁴ In contrast to previous reports, we found that the charge matching of the interior of the Q β coat protein, which is positively charged, with the anionic charge of smURFP promotes assembly without the need for osmolytes. The 68% yield proved sufficient for small-scale experiments but was cumbersome when significant quantities—i.e., hundreds of milligrams—were required. We were thus drawn to the encapsidation of fluorescent proteins (FPs) inside VLPs *via* cloning, which allows the bacteria to do the time-consuming work of placing cargo within the capsid in high yields producing larger quantities of material.^{35,36} Previous work by Finn and colleagues have shown that proteins or enzymes can be packaged inside VLP Q β using an RNA template that mimics the putative natural way the native capsid self-assembles. This requires engineering recognition elements into the transcribed mRNA to direct fluorescent proteins inside the capsid.^{37,38} As illustrated in Scheme 1A and B, proteinaceous cargo is tagged with an arginine-rich rev peptide tag that recognizes and binds to an RNA aptamer upstream of the ribosome binding site.³⁹ An RNA hairpin downstream of the stop codon binds to the interior of the CP monomers directing the rev-tagged protein in the interior of VLP. Conceptually, this expression system uses the mRNA as a rope that electrostatically ties cargo to the CP. To the best of our knowledge, this method yielded two fluorescent VLPs—one containing the green fluorescent protein (GFP) and one containing the near-infrared proteins (iRFP and mIFP). GFP and iRFPs have outstanding photophysical properties but have some practical and physical limitations in the lab. Most

fluorescent microscopes do not have filter sets to image NIR proteins, and GFP has considerable spectral overlap with biomolecules typically found in tissues.^{40,41} The kidney, lung, liver, and spleen are notorious for intense autofluorescence, which makes imaging them with green fluorophores challenging.^{42–44} This often leads to misinterpretation of the results and unreliable science. The solution to overcoming autofluorescence is selecting a spectrally shifted FP that fluoresces in the red channel.

smURFP was evolved from light-harvesting phycobilisomes yielding a photostable protein.^{45,46} The photostability of this protein is higher than other proteins like eGFP, mCherry, and iRFP 1.4. Red fluorescent proteins like iRFPs, iFPs, and smURFP need the fluorophore biliverdin (Bv), which binds to the protein *via* carboxylate groups on Bv to fluoresce. The inside of the smURFP is positively charged, allowing attraction and interaction with the carboxylate groups of fluorophore Bv. Following binding, the fluorescence increases 1400× and shows spectral properties similar to Cy5; however, Bv hydrochloride is expensive, and incorporating Bv post synthetically can produce inconsistent results (*vide infra*).^{36,47–50} Heme oxygenase 1 (HO-1) is an enzyme that breaks heme down into biliverdin, carbon monoxide (CO), and ferrous (Fe²⁺) ions.⁵¹ HO-1 is found in humans but not *E. coli*; however, a recombinant version that can be expressed in bacteria exists.^{52–54}

In this report, we show that the co-expression of HO-1, along with the VLP coat protein and the FP, smURFP, produces a very bright encapsidation complex (s@Q β) both efficiently and in high yield. We demonstrate that the self-assembly of s@Q β is sufficiently robust that neither hairpin recognition motifs nor RNA-based aptamers are necessary to induce high-yielding



Scheme 1 The schematic presentation of previous and current systems described in this work used to co-express proteins for encapsidation inside Q β VLP. (A) Prior work used an RNA aptamer (pink) that binds to the positively charged Rev tag (green) on one end and, on the other end, the hairpin sequence from infectious Q β (purple) binds to the expressed coat protein (light blue), which helps template capsid assembly. (B) This system assembles to form an intact capsid encapsidating cargo. (C) Our simplified approach uses the positively charged Rev tag (green) to interact electrostatically with the negatively charged RNAs (black). This system exploits the known hairpins produced in the coat protein sequence (blue) to template the assembly of the full viral capsid. (D) Our approach also results in cargo loading within the VLP.

s@Q β (Scheme 1C and D). Finally, we prove that we can administer s@Q β via pulmonary lavage, extract lung tissue, and subsequently image s@Q β within the cells of the lung without autofluorescence as a confounding issue.

Results and discussion

Cloning

As a starting point in the cloning process, we used the plasmid pJF51-CP-mIFP (gifted by the Finn laboratory), which contains the sequence for both the CP of Q β and the near-infrared fluorophore mIFP—the mIFP will be digested and replaced with smURFP in this work. The induction of proteins on this plasmid is triggered by the addition of the isopropylthio- β -galactoside (IPTG)—a molecular mimic of lactose. The plasmid was digested with two enzymes XhoI and EagI (New England Biolabs), allowing us to remove the mIFP. The digested backbone was isolated and cleaned from the gel using a NucleoSpin gel and PCR clean-up kit (Macherey-Nagel). Co-expression of the s@Q β with the HO-1 gene was done based on prior reports³⁶ that incorporation of exogenous commercially obtained Bv during expression produces inconsistent results. We decided to test if the co-expression of HO-1, CP, and smURFP could overcome this issue. To eliminate the need for synthetic biliverdin, we used the pBAD-smURFP-HO-1 plasmid, which expresses smURFP and HO-1 under the *L*-arabinose operon. smURFP and HO-1 expression induction is triggered by adding *L*-arabinose to the media. The only restriction enzymes we could use would cause the removal of the smURFP sequence and the HO-1 sequence. Thus, we decided to digest both and reinsert the HO-1 fragment by first amplifying the HO-1 sequence from the pBAD plasmid. Next, the pBAD vector was digested by BamHI and PmeI (New England Biolabs) restriction enzymes. The digested backbone of pBAD was isolated and cleaned with the Macherey-Nagel, NucleoSpin gel, and PCR clean up. The smURFP sequence was ligated with the digested backbone of pJF51-CP (Fig. S1, ESI†). The pBAD backbone and amplified HO-1 were ligated using Gibson assembly (Fig. S2, ESI†). We confirmed the proper ligation by Sanger sequencing. Finally, a double transformation of both plasmids in BL21(DE3) *E. coli* yielded fluorescent s@Q β VLPs.

Previous papers have shown that the wild-type hairpin (Fig. 1A) is necessary to induce a conformational change of the MS2 CP dimers to form the capsid. We can infer that Q β - and VLPs derived from the same family Leviviridae-use a similar assembly mechanism.⁵⁵ Other data have shown that the Q β hairpin can tolerate significant change and still induce self-assembly, requiring only an eight-base pair stem and unpaired adenosine to induce a CP dimer conformational change, shown as the minimal sequence in Fig. 1B.⁵⁶ Using RNAfold software, which returns several low free energy structures for a given RNA sequence, we discovered that the RNA that codes for the CP always contains hairpin structures (Fig. 1C) consistent with the minimal sequence known to require assembly. This suggests that as long as the CP is expressed, the capsid should be able to assemble. It has been shown through cryo-electron



Fig. 1 The secondary structure of the Q β HP operators. (A) The wild-type operator hairpin. (B) The consensus minimal RNA sequence required for coat protein binding; the letter N in the minimal sequence indicates any nucleotide can be placed in that position, and self-assembly still occurs. In the stem portion, most nucleotides are complementary so that the stem will form. (C) the RNA hairpin with the minimal sequence found in the mRNA used to translate the coat protein. (D) For the largest $r_{\text{threshold}}$ value of 3.0 Å, a picture of the Q β VLP (cutaway view) along with the void locations visualized in yellow.

microscopy experiments on Q β that RNA hairpin interactions are needed for more than just creating the structure; they also stabilize the capsid.⁵⁷ Cryogenic electron microscope EM data shows that fifty-nine stem-loops in the Q β viral capsid appear to contribute to its stability; however, it is not known what the minimum amount of stem-loops are needed for a Q β VLP to form. Another change in our strategy is the removal of the α REV aptamer, which is putatively needed to capture the rev-tag fused to the N-terminus of the FP. In our system, we retained the rev tag—which is strongly cationic but removed the α REV aptamer. We hypothesize that the transcribed anionic mRNAs that direct CP self-assembly would also interact strongly with the rev tag's cationic surface. We also considered the total void volume to determine how many proteins we could load within the VLP. Since the expression strategy (Scheme 1) mainly involves RNA, with the smURFP protein accounting for less than 5% by mass, we used the space occupied by the genomic RNA in the Q β virion (pdb entry 7LHD, see Fig. 1D) to estimate how much RNA would fill the VLP. This estimate predicts the maximum number of smURFPs complexes that fit inside the Q β as 10–17 (computational details in ESI†).

Expression

The dual system with two different operons carries with it the possibility of inhibiting the *L*-arabinose operon by the presence of IPTG. It has been found that at low concentrations of arabinose, the pBAD can be repressed by IPTG-influenced expression. However, oversaturation of the system with a high amount of arabinose can inhibit the repressor effect of IPTG.⁵⁸ To overcome the inhibition effect of IPTG, we increased the amount of added *L*-arabinose to a total of 1% w/v. For protein expression, bacteria were grown in Luria broth (LB) and induced when the OD₄₅₀ ~ 1. To test if our co-expression system

produces superior results, we prepared two samples—one where Bv was added externally ($s@Q\beta$ -Bv) and one where they were co-expressed ($s@Q\beta$ -HO). For the system $s@Q\beta$ -Bv, induction was carried out with 1 mM IPTG and 15 mM Bv as optimized previously³⁶ in the presence of $50\ \mu\text{g mL}^{-1}$ streptomycin.

For the co-expression system, smURFP@ $Q\beta$ -HO induction was performed with 1 mM IPTG and 1% w/v L-arabinose in LB media supplemented with $50\ \mu\text{g mL}^{-1}$ streptomycin and $50\ \mu\text{g mL}^{-1}$ ampicillin. Further purification includes pelleting down the bacteria, sonication, salting out, 40% w/v sucrose cushion, and again pelleting down to obtain the blue protein pellet. Protein concentration was determined by Coomassie Plus Protein Reagent, and from this, we determined that we obtained $40\text{--}60\ \text{mg L}^{-1}$ of smURFP@ $Q\beta$ -HO. This is comparable to the $50\text{--}70\ \text{mg L}^{-1}$ yield obtained using the α REV aptamer approach (Scheme 1A) reported by Finn *et al.*³⁶

Characterization to compare our different preparations of $s@Q\beta$ VLPs with native $Q\beta$ VLP was performed. A non-reducing 18% SDS (Fig. 2A) gel showed the CP monomer at 14.2 kDa and a faint CP dimer at 28 kDa for $Q\beta$, $s@Q\beta$ -HO, $s@Q\beta$ -Bv, and the smURFP at 15 kDa, as expected. Agarose electrophoresis

(Fig. 2B) shows that $Q\beta$, $s@Q\beta$ -Bv, and $s@Q\beta$ -HO do not significantly change in mobility, suggesting that both size and overall charge are unaffected by protein encapsidation. Fluorescence gel imaging (Fig. 2C) showed a signal corresponding to smURFP in the Cy5 channels for both $s@Q\beta$ -HO and $s@Q\beta$ -Bv but not for our control, $Q\beta$, further proving we have encapsidated smURFP within the VLP in both systems. Dynamic light scattering (DLS) was used to measure the size of the $Q\beta$ VLPs. We found the encapsidation did not alter nanoparticle size much: $Q\beta$ ($\sim 34\ \text{nm}$, PDI = 0.077), $s@Q\beta$ -Bv ($\sim 32\ \text{nm}$, PDI = 0.087) and $s@Q\beta$ -HO ($\sim 33\ \text{nm}$, PDI = 0.053) (Fig. 2D). Autocorrelation graphs are reported in Fig. S3 (ESI†). $s@Q\beta$ VLP solution in light is blue to the eye and under UV shows a pink luminescence (Fig. 2E). To observe the VLPs morphology, transmission electron microscopy (TEM) was used for $s@Q\beta$ -HO (Fig. 2F), and $Q\beta$ (Fig. 2G) confirming our DLS results, which found no differences in size or changes in morphology as shown in (Fig. S4, ESI†). Following centrifugation, the $s@Q\beta$ VLP pellet appears blue (Fig. 2H). Using gel densitometry, we determined that ~ 12 smURFPs were loaded in $s@Q\beta$ -HO, larger than the ~ 8 smURFPs inside $s@Q\beta$ -Bv (Fig. S4 and S5, ESI†). We obtained similar results when the amount was calculated with fluorescence intensity— ~ 10 smURFPs were packed in $s@Q\beta$ -HO and ~ 8 smURFPs inside $s@Q\beta$ -Bv (Fig. S7, ESI†).

To further compare these different expression methods, more analyses were performed. The fluorescence intensity was measured at the same concentration of VLPs, and $s@Q\beta$ -HO exhibited a higher fluorescence intensity than $s@Q\beta$ -Bv (Fig. 3A). We speculate that this might be because bacteria cells uptake small molecules less efficiently when introduced externally rather than the ones the cells make themselves. It is worth mentioning that the fluorescence of $s@Q\beta$ -Bv declined significantly after some days, which we did not observe for $s@Q\beta$ -HO. The VLPs were incubated for 30 min at a pH of 2–10 to evaluate the photophysical properties



Fig. 2 Characterization of VLPs. (A) 18% SDS gel showing the bands of CP monomer, dimer (CP-D), and smURFP (s) for native $Q\beta$, $s@Q\beta$ -HO, and $s@Q\beta$ -Bv. (B) Agarose gel comparing VLPs using Coomassie staining and (C) fluorescence imaging. (D) DLS showing the size of the VLPs. (E) The $s@Q\beta$ sample under natural and UV light. TEM images of (F) $s@Q\beta$ (G) native $Q\beta$. (H) Blue protein pellet of $s@Q\beta$ after purification showing a visible blue color from smURFP protein.



Fig. 3 Comparison and characterization of $s@Q\beta$ -HO and $s@Q\beta$ -Bv at the same $Q\beta$ concentration. (A) Emission spectra and (B) pH stability from 2–10. (C) Absorbance spectra of encapsidated FP in $Q\beta$ and free FP. (D) SEC shows the coelution of smURFP with the total protein peak.

of the smURFP. While this experiment says nothing of the stability of the VLP at these pH's we can say that we observed minimal loss of fluorescence, with the maximum being a 10% reduction in intensity at pH 8 meaning that the cargo (fluorescent protein) is still stable (Fig. 3B). The spectral properties of the encapsulated and free smURFPs showed identical meaning_{max}; however, considerable broadening at lower wavelengths was noted (Fig. 3C and Fig. S8, ESI†). To verify the purity of the VLP, size exclusion chromatography (SEC) was used. It showed coelution of the VLPs indicative of the absorbance at 280 and 260 nm for the protein and at 642 nm for smURFP (Fig. 3D). It is worth mentioning that if smURFP had not been encapsulated post-expression, we would observe the free smURFP as a separate peak in the SEC chromatogram.

From these experiments and in agreement with the literature, we conclude that the co-expression system reliably produced a brightly fluorescent encapsulated complex. An added benefit is that the synthesis does not require adding expensive Bv and produces consistent results. Based on these results, we recommend co-expression whenever possible and will use the s@Qβ-HO as we advance in this manuscript. To further explore if removing the REV tag would allow encapsulation of smURFP in Qβ VLP, we digested the REV tag out, performed ligation, and then started expression. No blue color was observed during expression, and the protein solution appeared white/transparent in color indicating smURFP was not packed inside the VLP. These results demonstrate the necessity of the REV tag in the cargo encapsulation. SDS page following purification confirmed no smURFP was encapsulated (Fig. S9, ESI†).

Cell studies

GFP is a well-known FP widely used in imaging owing to its great brightness and stability. However, imaging with GFP can be very challenging as specific molecules found in cells and tissue are bioluminescent at similar wavelengths that GFP emits. This autofluorescence makes data interpretation a difficult process. For this reason, we have benchmarked our system against the previously reported construct GFP@Qβ and studied them both *in vitro* and *in vivo*.

To observe their behavior first in cells, we performed cell viability studies and cellular uptake in the A549 human lung adenocarcinoma cell line. Using a lactate dehydrogenase (LDH) assay, no toxicity was observed at any concentration from 0.5 mg mL⁻¹ to 10 mg mL⁻¹ (Fig. 4A). Cellular uptake was quantified using flow cytometry (Fig. 4B and Fig. S10, ESI†), and we found the VLP uptake to be the same regardless of the FP used. Additionally, laser scanning microscopy (CLSM) images were acquired for cells treated with saline, GFP@Qβ s@Qβ, and, as shown in Fig. 4C, we saw similar localizations within cells regardless of the FP used.

Fluorescence imaging *in vivo*

In tissues, molecules and endogenous fluorophores become fluorescent and absorb at a specific wavelength. The most well-known fluorophores in tissues are flavins, pyridinic acid, and collagen. Imaging experiments must be performed to distinguish the fluorescence when treating tissue with exogenous

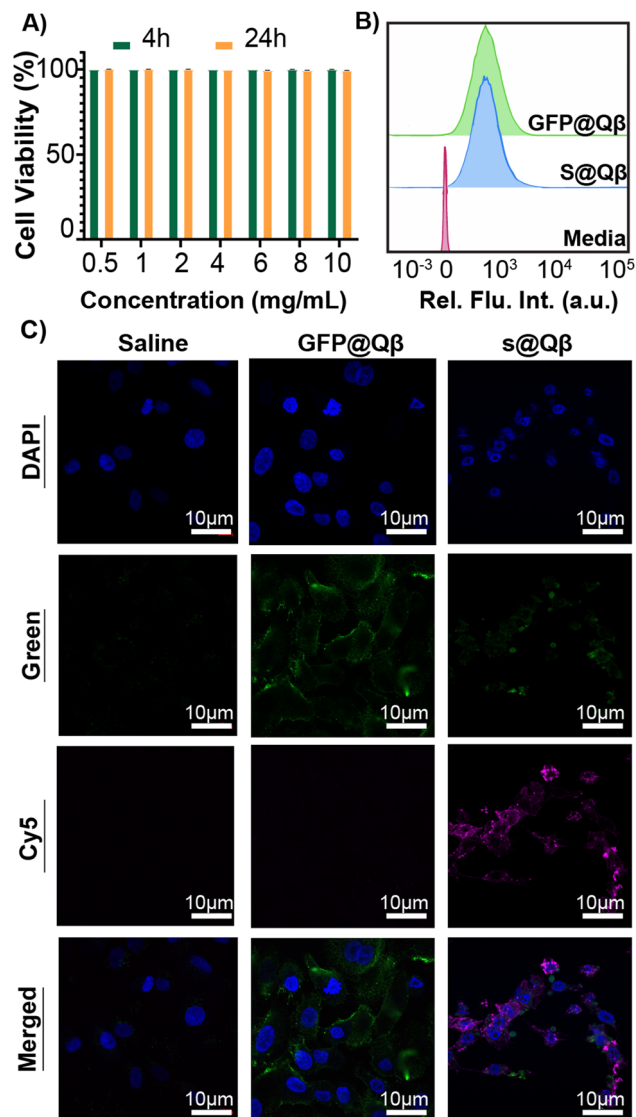


Fig. 4 Cell viability and cellular uptake of A549 cells of s@Qβ and GFP@Qβ. (A) Viability of s@Qβ at 0.5–10 mg mL⁻¹ for 4 h and 24 h. (B) Flow cytometry of A549 cells treated with s@Qβ (blue) compared with GFP@Qβ signal and cells only (pink). (C) Cellular uptake for 4 h imaged with CLSM representing saline, GFP@Qβ, and s@Qβ.

fluorophores. In this experiment, we attempted to observe if s@Qβ VLP would be sufficiently spectrally shifted and bright to be observed in lung tissue. Three groups of BALB/c mice ($n = 3$) were administered with saline, GFP@Qβ, and s@Qβ endotracheally, as shown in Fig. 5A. Four hours post-delivery, the mice were sacrificed, and their lungs were extracted, fixed, paraffin-embedded and thin sectioned using a microtome. Tissue slides were dehydrated and cleaned, and nuclei were stained with DAPI and imaged by CSLM. As illustrated in the micrograph in Fig. 5B, when GFP is used, it is difficult to determine which signal arises because of autofluorescence and what arises from GFP@Qβ. Indeed, green fluorescence is obvious in all the samples, even if they were not given GFP@Qβ. In contrast, the micrographs showing samples where s@Qβ was used is very clear in the Cy5 channel-imaged samples stained with s@Qβ



Fig. 5 (A) Schematic representation of the delivery. (B) CSLM images of BALB/c lung tissues ex vivo for saline (left), GFP@Q β (center), and s@Q β (right).

using a standard Cy5 filter set produced unambiguous results, clearly showing the lung tissue had taken in the VLPs.

Conclusions

This study shows that the mRNA used in translating CPs for the Q β capsid contains hairpin structures sufficient to nucleate capsid formation while also encapsulating cargo. We have also shown that the encapsidation of cargo featuring a REV tag can occur without the complementary α REV aptamer; however, no encapsidation occurs in the absence of the REV tag. We suspect this assembly is driven purely by electrostatic interactions between the cationic REV tag and the anionic backbone of the mRNA. We demonstrated that the genetic co-expression of s@Q β and HO-1 yields consistent results and lowers the cost s@Q β expression compared to the expression of s@Q β with exogenous addition of BV during induction. Finally, we

demonstrated that encapsidation of the smURFP protein inside the Q β VLP drastically allowed for imaging in the Cy5 channel, reducing background auto-fluorescence compared to the spectral window used to image GFP fluorescence.

Ethical statement

Female BALB/c mice were obtained from Charles River Lab (Wilmington, Ma). All animal studies used for endotracheal delivery were performed in strict accordance with the protocol #18-17 approved by the University of Texas at Dallas Institutional Animal Care and Use Committee (IACUC).

Author contributions

Conceptualization: JJG, IT, methodology: IT, OT, YHW, MD, FCH, investigation: IT, OT, YHW, MD, RNE, ACC, CBD, FCH, JG, NDN, SON, JJG, supervision JJG, writing – original draft: IT, JJG, writing review & editing IT, JJG.

Data availability

The raw data associated with this work can be accessed at the Open Science Foundation (OSF) via this link: https://osf.io/cj9r7/?view_only=d06e0579eeb4422f8173663daeb6f706.

Conflicts of interest

There are no conflicts to declare.

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