A tractable method for simultaneous modifications to the head and tail of bacteriophage lambda and its application to enhancing phage-mediated gene delivery

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ABSTRACT

There is considerable interest in the use of bacteriophage vectors for mammalian cell gene transfer applications, due to their stability, excellent safety profile and inexpensive mass production. However, to date, phage vectors have been plagued by mediocre performance as gene transfer agents. This may reflect the complexity of the viral infection process in mammalian cells and the need to refine each step of this process in order to arrive at an optimal, phage-based gene transfer system. Therefore, a flexible system was designed that allowed for the introduction of multiple modifications on the surface of bacteriophage lambda. Using this novel method, multiple peptides were displayed simultaneously from both the phage head and tail. Surface head display of an ubiquitinylation motif greatly increased the efficiency of phage-mediated gene transfer in a murine macrophage cell line. Gene transfer was further increased when this peptide was displayed in combination with a tail-displayed CD40-binding motif. Overall, this work provides a novel system that can be used to rationally improve bacteriophage gene transfer vectors and shows it may be possible to enhance the efficiency of phage-mediated gene transfer by targeting and optimizing multiple steps within the viral infection pathway.

INTRODUCTION

Bacteriophage lambda has appealing characteristics as a gene and vaccine delivery vector, which include a high degree of physical stability, compatibility with rapid and inexpensive production/purification methods, genetic tractability and inherent biological safety in mammalian cells (1–3). In addition, the dimensions of the lambda phage particles are broadly similar to those of many mammalian viruses and recent structural evidence points to a shared ancestry between tailed bacteriophages and mammalian DNA viruses (4).

Lambda phage vectors have been successfully used to transfer exogenous genes to mammalian cells, following surface modification of either the phage coat protein (gpD) or the major tail protein (gpV) (5–8). Phage heads contain between 405 and 420 copies of gpD (9), while the phage tail consists of 32 rings, each containing six subunits of gpV (10–12). Thus, both of these proteins can be used to display foreign proteins or peptides at high copy numbers on the surface of lambda phage particles.

Most attempts to enhance lambda phage-mediated gene transfer to mammalian cells have concentrated on optimizing the binding of phage particles to mammalian cells (7,8,13,14). However, the eukaryotic cell poses numerous barriers to phage-mediated gene transfer. After receptor binding and internalization, phage must gain access to the cytoplasm, uncoat and deliver their DNA payload to the nucleus. Thus, the ultimate success of phage-mediated gene transfer depends on the ability to overcome multiple intracellular barriers. This is likely to require the use of combination strategies that increase the efficiency of each step involved in phage-mediated gene transfer, including cell attachment, cytoplasmic entry, endosomal escape, uncoating and nuclear import (7,15,16).

This article reports the design and development of a novel and tractable lambda-based vector that allows for the facile generation of phage particles that display multiple peptides or proteins of interest on their surface. We hypothesized that this system could surmount current...
obstacles to efficient phage-mediated gene transfer, by generating phage vectors that display a combination of exogenous peptides each intended to circumvent a separate barrier to efficient phage gene delivery. This article presents the first example of single lambda phage constructs incorporating multiple surface modifications that, collectively, enhanced in vitro gene transfer to mammalian cells.

**METHODS**

**Lysogens**

The λD1180(luc) lysogen was a gift from Dr Mahito Nakasai and DAVEC Corporation (6); λD1180(luc) contains a firefly luciferase gene under the transcriptional control of the major human cytomegalovirus (CMV) immediate-early promoter. Lysogen λD1180 (Dam15 del EcoRI-SacI CIs857 nin5 Sam100) is deficient in gpD, the S lysis gene, and contains a temperature-sensitive mutation in the cI repressor. The *E. coli* lysogen host (Top10, Invitrogen) is *sup0*, ensuring that neither *Sam100* nor *Dam15* are expressed.

**Plasmid design**

*pTrc:gpD-Fusion* and *pTrc:gpD*. The plasmids *pTrc:gpD* and *pTrc:gpD-Fusion* were derived from *pTrcHis* (Invitrogen) with the addition of a synthetic, codon-optimized D gene as described (17). The D gene was inserted as an NcoI to BamHI fragment, with subsequent loss of the NcoI site. A linker sequence [G(SGGG)2 SGGT] was then added (BamHI to KpnI), to permit insertion of DNA sequences encoding exogenous peptides of interest between the sites KpnI and HindIII. The plasmid *pTrc:gpD* was constructed with the addition of a stop codon immediately after the codon optimized D gene.

*pTrc:gpD-UBHA*. A ubiquitinylation motif derived from the Hepatitis A Virus (HAV) 3 protease [LGVKDDW LLV; (18)] was constructed by overlapping PCR using the primers: UBHAfor (5'-AACCTGGGTACCTAGGC GTAAAAGATGACTGGTTGCTG-3') and UBHArev (5'-CAGGCTAAGCTTCAACACGCAACCAGTCA TCTTTAAC-3'; underlining denotes the translational stop codon). The PCR product was digested with KpnI and HindIII and cloned into *pTrc:gpD-Fusion* to create *pTrc:gpD-UBHA*.

*pTrcRSF:gpV-Fusion*. The gpV expression plasmid was constructed from *pTrcHis*. First, the truncated gpV gene was PCR amplified from the phage genome using the primers: pVForA (5'-AGCTCTCCATGGCGCTTACC AAATCTCAATG-3') and pVRevLink (5'-AGCTG GATCCCCCCTTCCACCCAGGTGC-3'). The PCR product was digested with BamHI and NeoI and cloned into *pTrcHis* at the corresponding sites. A linker with the sequence G(SGGG)8T was synthetically constructed by GeneART (Regensburg, Germany) and inserted as an NcoI to BamHI fragment, with subsequent digestion of the corresponding DNA fragment from *pTrcRSF:gpV-CD40*. A DNA sequence encoding the CD40 receptor-binding peptide, ATYSEFPGLNLK (19), was subcloned from the parent plasmid *pAT-043-CD40* using KpnI and HindIII digests and then cloned into *pTrcRSF:gpV-Fusion* at the corresponding restriction sites to create *pTrcRSF:gpV-CD40*.

**Production of polyclonal antiserum**

The production of the polyclonal anti-gpD antiserum has been described (17). Polyclonal anti-gpV antiserum was produced by transforming the plasmid *pBAD:gpVtrunc* into Top10 *E. coli* (Invitrogen). Transformed bacteria were grown to mid-log phase and then induced for 3h with 0.002% L-arabinose. After expression, bacteria were lysed with BugBuster (Novagen), treated with benzonase and lysozyme and cleared by centrifugation. The lysate was purified on a Ni–NTA matrix and fractions containing the purified gpV protein were identified on immunoblot with an anti-His antibody. SDS-PAGE purified fractions containing the truncated gpV protein were submitted to Sigma Genosys for the production of rabbit anti-gpV antiserum. Antiserum reactivity was verified by immunoblot and aliquots were stored at −80°C.

**Phage production**

Chemically competent lysogens of λD1180 were transformed with purified DNA plasmids of interest, and transformants were then selected using media containing the appropriate antibiotic. Single colonies were grown overnight at 32°C with antibiotic selection, and the resulting cultures were then used to inoculate 11 of fresh antibiotic-containing medium the next day (at a dilution of 1:100). Cultures were grown at 32°C with vigorous shaking (300 r.p.m.) until an OD600 of between 0.3 and 0.4 was reached. Lysogens were induced by transferring the bacteria to a water bath set between 51 and 53°C, followed by incubation with gentle shaking for 20 min. After thermal induction, the cultures were vigorously shaken for an additional 3h at 38°C. Bacteria were then pelleted and resuspended in phage suspension media (SM: 50 mM Tris-HCl, pH 7.5, 100 mM NaCl, 10 mM MgSO4, 0.01% gelatin) and lysed with the addition of 10% chloroform. After chloroform treatment, bacterial DNA was lysed with...
DNaseI at a final concentration of 10 μg/ml. The lysate was then cleared of cellular debris by low-speed centrifugation and phages were pelleted by ultracentrifugation at 110 000 g. Pelleted phage particles were subsequently purified by cesium chloride equilibrium density gradient centrifugation and phage were dialyzed against dialysis buffer (10 mM NaCl, 50 mM Tris-Cl, pH 8.0, 10 mM MgCl₂) prior to storage at 4°C. Typical titers were 1 x 10¹⁰ p.f.u./ml.

**Immunoblot analysis of purified phage**

Phage was denatured and structural proteins separated by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE). Gels for the detection of gpD contained between 15 and 20% polyacrylamide, while gels for gpV contained between 12 and 15% polyacrylamide. Phage proteins were detected by western blotting with the appropriate primary antibody (rabbit polyclonal antiserum specific for either gpD or gpV, which were used at dilutions of 1:1000 or 1:2000, respectively). Detection of bound rabbit antibody was achieved by incubation with a 1:3000 dilution of anti-rabbit HRP (Sigma) for 1 h, followed by detection with ECL-Plus Substrate (Amersham Biosciences).

**Detection of CD40 expression on RAW 264.7 cells**

RAW 264.7 cells were purchased from the American Type Culture Collection (ATCC) and maintained according to ATCC recommendations. For detection of the CD40 receptor 1 x 10⁶ cells were stained for 15 min on ice with a 1:200 dilution of PE-CD40 or an appropriate isotype control antibody (BD Pharmingen). After staining, cells were washed with PBS, resuspended in FACS buffer, and analyzed by fluorescence-activated cell sorting using a FACSCalibur instrument.

**Luciferase assay for phage-mediated gene expression in RAW 264.7 cells**

RAW 264.7 cells were plated at 1 x 10⁵ cells/well in 12-well plates, 12 h before the planned phage transduction. After the 12-h incubation, the media was removed and replaced with lambda phage (1 x 10¹¹ p.f.u.) diluted in 1 ml of serum-containing medium. Infection of cells was centrifugally enhanced [1200 g for 1 h at 37°C; ‘spinoculation’ (20–22)] and cells were subsequently incubated with the phage for 48 h at 37°C. Cells were then either used for quantitative DNA PCR analysis (see below) or used for luciferase assays. For the latter, cells were washed with phosphate buffered saline (PBS) and then lysed in 30 μl of Promega’s passive lysis buffer (PBL) for luciferase analysis. Protein concentration in the lysates was determined by Bradford assay, and equal protein quantities were used in luciferase assays, utilizing a luciferase assay kit (Promega).

**Quantitative DNA PCR assay for phage genome copies in RAW 264.7 cells**

RAW 264.7 cells were transduced with lambda phage particles as described above. At the end of the 48-h incubation, cells were washed four times with PBS and total genomic DNA was extracted using the Wizard Genomic DNA Extraction Kit (Promega). Total genomic DNA was quantitated spectrophotometrically and 10 ng was then used for QPCR analysis using a luciferase-specific Taqman probe (5'-CATTTCGCAGCCTACGGTGGTT-3') together with the reverse primer TaqLucR (5'-TTGCAACCCCTTTTGGAAAA-3') and the forward primer TaqLucF (5'-AAGGTAATTCCTAACAGTATG-3'). The amplification conditions comprised an initial denaturation step at 95°C for 10 min followed by 40 cycles at 95°C for 15 s and 60°C for 1 min. A standard curve for lambda DNA quantitation was generated by using serially diluted phage DNA and samples were analyzed with a Bio-Rad iCycler. All standards and samples were assayed in triplicate.

**RESULTS**

**Incorporation of truncated gpV into phage tails**

We previously designed and evaluated a plasmid-based, trans-complementation system that allowed the incorporation of recalcitrant peptides into the head of phage lambda (17). We reasoned that a similar approach might also permit the display of exogenous peptides on the major tail protein, gpV.

Isolation of naturally occurring mutations in gpV indicate that up to one-third of the carboxyl terminus is unnecessary and can be deleted without adversely effecting tail assembly (23). Therefore, exogenous peptides have been fused to the C-terminus of either full-length or truncated gpV, and displayed on the tail of lambda particles (24–26). Pilot experiments showed that a truncated gpV protein bearing a hexahistidine tag was efficiently incorporated into our phage particles (not shown). Therefore, the truncated gpV gene was used in all subsequent experiments.

**Selection of peptide motifs for display on the head and tail of bacteriophage lambda particles**

Successful gene transfer by bacteriophage vectors is likely to depend on the circumvention of multiple intracellular barriers to efficient gene expression. Therefore, a vector displaying multiple modifications, each intended to address a distinct barrier to gene delivery, may result in more efficient gene transfer when compared to phage that display only a single modification on their surface. Uncoating represents an important, but little studied, aspect of phage-mediated gene transfer that may be enhanced by the display of specific peptide motifs on the phage surface. We were therefore intrigued by reports that gene transduction by a number of mammalian viruses can be improved in the presence of proteasome inhibitors (27–29). This suggests that some viruses may use ubiquitinylation and proteasomal degradation to trigger head uncoating and genomic release. There is also evidence that ubiquitin, especially mono-ubiquitinylation, plays an important role in vesicular trafficking (30) and that ubiquitin can serve as a signal
for localization to multi-vesicular bodies (31). With this in mind, we set about displaying an ubiquitylation motif from the hepatitis A virus 3C protease (UBHA) on the surface of lambda phage particles. To do this, the ubiquitylation motif (LGVKDDWLLV) was fused to the C-terminus of gpD, and then displayed on phage particles (17).

In order to test the feasibility of simultaneously co-expressing two different peptide motifs on the head and tail of lambda phage particles, we selected a second, receptor-binding peptide for display on the major tail protein, gpV. For this purpose, we elected to display a CD40-binding peptide (ATYSEFPGNLKP) that was previously identified in our laboratory (19). We therefore designed the plasmid pTrcRSF:gpV-CD40 to display this peptide on the major lambda phage tail protein, gpV.

*Figure 1.* Expression plasmids for the production of bi-functional phage. Production of bi-functional phage displaying peptide modifications on both gpD and gpV was achieved by transformation of lambda lysogens with a series of compatible plasmid expression constructs. (A) pTrc:gpD for the expression of unmodified wild-type gpD. (B) pTrc:gpD-UBHA for the expression of gpD displaying a ubiquitylation motif from the hepatitis A virus 3C protease (UBHA). (C) pTrc:gpD co-transformed with pTrcRSF:gpV-CD40 for the expression of wild-type gpD in conjunction with gpV displaying a CD40-binding peptide (CD40). (D) pTrc:gpD-UBHA co-transformed with pTrcRSF:gpV-CD40 for the simultaneous expression of gpD-UBHA and gpV-CD40.

**Generation of bi-functional phage particles with modifications to both the head and tail**

The lysogen λD1180(luc) was transformed with either pTrc:gpD or pTrc:gpD-UBHA to produce phage particles that displayed either wild-type gpD (Figure 1A) or a modified gpD head protein fused to a ubiquitylation motif (Figure 1B). To produce phage particles that displayed a modified gpV tail protein fused to a CD40-binding peptide, the λD1180(luc) lysogen was transformed with pTrcRSF:gpV-CD40 and pTrc:gpD (Figure 1C). Finally, bifunctional phage displaying both peptide modifications were produced by dual transformation with pTrc:gpD-UBHA and pTrcRSF:gpV-CD40 into λD1180(luc) lysogens (Figure 1D). Phage particles were then produced following lytic induction of the lysogen, and CsCl density-gradient-purified phage particles were analyzed by western blot with antisera specific for gpD or gpV. As expected, introduction of pTrc:gpD-UBHA into the lysogen resulted in the generation of progeny phage particles that had full replacement of wild-type gpD head protein with gpD-UBHA (Figure 2; top panel). Similarly, introduction of the expression plasmid for gpV-CD40 into lambda lysogens yielded bi-functional phage displaying both peptide fusions. The plasmid pTrc:gpD-UBHA and the plasmid pTrcRSF:gpV-CD40 were co-transformed into lambda lysogens. After lytic induction, phage were purified by CsCl density gradient centrifugation and analyzed. (A) Phage (5 × 10^8 p.f.u.) were loaded on a 12% polyacrylamide gel, subjected to SDS-PAGE and transferred to nitrocellulose. Immunoblot analysis was then performed using an anti-gpD rabbit polyclonal antiserum. This revealed full replacement of gpD with gpD-UBHA in phage produced from lysogens carrying the pTrc:gpD-UBHA plasmid. The molecular weight of gpD is ~12 kDa and of gpD-UBHA is ~16 kDa. (B) Phage (5 × 10^8 p.f.u.) were loaded on a 12% polyacrylamide gel, subjected to SDS-PAGE and transferred to nitrocellulose. Immunoblot analysis was then performed with an anti-gpV polyclonal antiserum. This revealed that the phage expressing the gpV-CD40 fusion protein contained roughly a 1:1 ratio of wild-type gpV and recombinant gpV-CD40. The projected molecular weights are ~30 kDa for wild type gpV and ~27 kDa for recombinant gpV-CD40 (Note that the recombinant gpV fusion protein is based on a truncated form of gpV, and is therefore smaller than its wild-type counterpart).
gpV-CD40 into the lysogen resulted in the generation of progeny phage particles that had ~50% replacement of wild-type gpV tail protein with the gpV-CD40 fusion protein (Figure 2; lower panel). Full replacement of wild-type gpV on the phage tail was not expected, since the lambda lysogen contains a functional wild-type gpV gene. In contrast, the lysogen contains an amber-mutated gpD gene that permits full replacement of gpD by exogenously expressed recombinant gpD, in host cells that lack the amber suppressor tRNA. In addition to the partial replacement of gpV that is evident in Figure 2 (lower panel), it can also be appreciated that the gpV-CD40 fusion protein is smaller than the endogenous gpV protein. This is because we used the truncated form of gpV to construct the gpV-CD40 fusion (see Materials and methods section).

Display of two distinct peptide modifications on the surface of lambda phage results in an enhancement of phage-mediated gene transfer

We hypothesized that phage particles displaying both a CD40-binding motif on the phage tail and an ubiquitylation motif on the phage head would be capable of mediating more efficient gene transfer in CD40-positive mammalian cells, when compared to unmodified phage particles or phage particles that displayed only a single capsid modification.

In order to test this prediction, we performed in vitro gene transfer experiments using the RAW 264.7 cell line. This murine macrophage cell line retains key properties of primary macrophages, including (i) cell surface expression of modest levels of CD40 and (ii) the ability to take up exogenous particles and macromolecules by phagocytosis (32–35). The phagocytic competence of RAW cells may be important since phagocytosis has been proposed to represent the major mechanism by which lambda phage vectors transduce cells in vivo (36).

RAW 264.7 cells were exposed to luciferase-encoding phage particles at high, intermediate and low multiplicities of infection (MOI = 1 x 10⁶, 3 x 10⁵ or 1 x 10⁵, respectively). Forty-eight hours later, the cells were washed extensively and extracts were prepared for analysis of luciferase expression levels.

Unmodified phage particles (gpD) and particles which displayed only the CD40-binding peptide (gpV-CD40) did not mediate detectable levels of luciferase gene expression when compared to control cell lysates (Figure 3; note that there is a modest but non-statistically significant trend towards increased gene transduction in the case of the gpV-CD40 phage, at the high-dose level).

In contrast, phage that displayed the ubiquitylation motif (UBHA) on their surface were able to successfully mediate gene transfer in RAW 264.7 cells, leading to luciferase levels that were roughly 10-fold greater than background, at the high dose level (P < 0.01); gene transfer was not significantly above background at the intermediate dose level. Very similar results were obtained following exposure of cells to a simple mixture of phage particles that displayed either gpV-CD40 or gpD-UBHA on their surface (groups designated as ‘gpD-UBHA + gpV-CD40’ in Figure 3).

Gene transfer was further enhanced (by ~2-fold) when the UBHA motif was co-displayed on the phage surface together with the CD40-binding peptide (groups designated as ‘gpD-UBHA + gpV-CD40’ in Figure 3). At the intermediate dose level, this result achieved statistical significance (P < 0.01) when compared either to gpD-UBHA phage alone or to the mixture of gpD-UBHA + gpV-CD40 phage.

Display of the CD40-binding peptide on the surface of lambda phage results in an enhancement of phage-mediated DNA uptake into cells

To further examine the effect of displaying the CD40-binding peptide on the phage surface, the efficiency of phage genome uptake into cultured RAW 264.7 cells was analyzed. The cells were confirmed to express CD40 by flow cytometric analysis (Figure 4B), and were then exposed to each of the phage constructs at a fixed MOI (1 x 10⁶). Forty-eight hours later, cells were washed thoroughly, and lysates prepared. Analysis of lambda genome copy number in lysates from transduced cells revealed that DNA copy number was slightly, but significantly (P < 0.05), greater in cells which were transduced with phage particles that displayed gpV-CD40 versus unmodified phage (Figure 4A). Lambda genome copy number was further elevated in cells which were transduced with phage particles that displayed gpD-UBHA plus gpV-CD40 versus phage particles that displayed either gpD-UBHA or gpV-CD40 alone (Figure 4A; P < 0.001). This suggests that surface display of the CD40-binding peptide enhances phage-mediated gene transfer at the level of either phage binding or internalization (leading to an increase in the number of phage genome copies in the cell).

Analysis of phage genome copies in transduced cells also revealed that surface display of the UBHA peptide did not lead to an increase in the number of phage genome copies per cell, when compared to unmodified phage particles (Figure 4A; compare gpD and gpD-UBHA). Since surface display of the UBHA peptide resulted in a profound increase in phage-mediated gene expression (Figure 3), we conclude that surface display of the UBHA peptide enhances phage-mediated gene transfer at a post-internalization step. This may reflect an effect on intracellular trafficking of phage particles, on phage uncoating or on another ubiquitin/proteasome-dependent pathway (29).

DISCUSSION

Multiple phage capsid proteins suitable for peptide display have been identified in both lambda and M13. Despite this, simultaneous display from two or more protein platforms has rarely been described, even in the more widely studied filamentous phage display system (37–40). Recently, a bifunctional filamentous phage intended for delivering biological agents was described (40). The phage combined an integrin-targeting moiety at pIII with a
streptavidin-binding sequence at pVIII. The authors tested their phage in vitro for receptor specific cell binding and internalization. In addition, they complexed their phage with quantum dots and then demonstrated tumor-specific accumulation after intravenous injection in mice. Despite these few successes, the design of bifunctional phage remains underexplored and underutilized. In 1996, Dunn proposed a bifunctional lambda phage combining gpD and gpV display, yet, until now, a bifunctional lambda phage has never been described.

The experiments reported here resulted in the simultaneous co-display of two different peptide modifications on the head and tail of lambda phage particles, as translational fusions to gpD and gpV, respectively. In both cases, a high copy number of the displayed peptide was achieved (full replacement, or roughly 400 copies/phage particle in the case of the gpD fusion protein and partial replacement, or roughly 100 copies/phage particle in the case of the gpV fusion protein). Moreover, because the gpD and gpV expression plasmids described here are compatible with the previously described, CDF-origin-based plasmid, pTrcCDF:gpD-Fusion (17), it should be possible to introduce up to three modifications to a single phage particle in the future (two to gpD, and one to gpV).

The long-term goal of our experiments is to develop a bacteriophage lambda vector system that is capable of mediating efficient gene transfer into mammalian cells. As a first step towards this goal, we evaluated phage-mediated gene transfer in a murine macrophage cell line using recombinant lambda phage particles that encoded...
of cells with the CD40-specific antibody. The isotype control antibody, and the unshaded curve represents staining by flow cytometry. The darkly shaded curve represents staining with antibody or an isotype control antibody, and stained cells were analyzed as unmodified phage particles (gpD). Statistical analysis was performed the number of phage genome copies per cell, when compared to the UBHA peptide did not lead to a statistically significant increase in phage displaying gpD-UBHA or gpV-CD40 alone). Surface display of increased lambda genome copy number (\(P < 0.05\)). RAW 264.7 cells were stained with an anti-CD40-PE, and then washed and lysed. Analysis of lambda genome copy number was conducted using 10 ng of total input cellular DNA prepared from the cell lysates. DNA copy number was slightly, but significantly (\(P < 0.05\)), greater in cells which were transduced with gpV-CD40 versus unmodified phage (gpD). The bi-functional phage, combining gpV-CD40 with gpD-UBHA, further increased lambda genome copy number (\(P < 0.001\), when compared to phage displaying gpD-UBHA or gpV-CD40 alone). Surface display of the UBHA peptide did not lead to a statistically significant increase in the number of phage genome copies per cell, when compared to unmodified phage particles (gpD). Statistical analysis was performed using one-way ANOVA with Tukey’s post-test; significance was taken as \(P < 0.05\). RAW 264.7 cells were stained with an anti-CD40-PE antibody or an isotype control antibody, and stained cells were analyzed by flow cytometry. The darkly shaded curve represents staining with the isotype control antibody, and the unshaded curve represents staining of cells with the CD40-specific antibody.

a luciferase reporter gene. These experiments showed that the surface display of an ubiquitinylation motif resulted in a profound enhancement of phage-mediated gene transfer. The ubiquitinylation motif may enhance phage-mediated gene transfer as a result of proteasome-mediated uncoating of the phage particle, or because of effects on the intracellular trafficking of internalized phage particles (30), and possible localization of internalized phage particles to multi-vesicular bodies (31), or due to effects on other intracellular pathways (29). It is noteworthy that certain mammalian viruses such as Vesicular Stomatitis Virus (VSV) rely on multi-vesicular bodies for endosomal escape and efficient infection of host cells (41,42). Attempts to resolve the mechanism by which the displayed ubiquitinylation motif enhances phage-mediated gene transfer were inconclusive, since the well-characterized proteasome inhibitors MG-132 and lactacystin proved toxic to the RAW 264.7 cells, even at the minimum effective dose. Thus, future studies will be needed to investigate the mechanism by which the ubiquitinylation motif enhances phage-mediated gene transfer.

Our proof-of-principle experiments also evaluated whether the simultaneous display of two different modifications on the phage surface may further enhance phage-mediated gene transfer. To do this, we generated luciferase-encoding phage particles that displayed a receptor (CD40)-binding peptide on their major tail protein (gpV) in addition to the ubiquitinylation motif on their major head protein (gpD). These bifunctional phage particles were able to mediate an enhanced efficiency of gene transfer into a cultured murine macro-phage cell line, when compared to phage particles that displayed only a single peptide moiety on their surface. This effect was dose-dependent, with higher levels of gene transfer being detected when larger amounts of phage were added to cells. Furthermore, co-display of both motifs on the same phage particle was required for the observed enhancement of phage-mediated gene transfer; gene transfer efficiency was significantly improved when compared to a simple mixture of gpD-UBHA phage plus gpV-CD40 phage. Thus, the enhanced gene transfer effect by the dual-display construct cannot be attributed to a trans-effect.

Overall, the results reported provide strong support for the notion that it may be possible to rationally improve the efficiency of phage-mediated gene transfer by displaying several different peptides on the phage surface. In the future it should be possible to introduce and test other modifications with the intention of eventually designing a highly efficient phage-based gene delivery vector.

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