Expanding the Versatility of Phage Display I: Efficient Display of Peptide-Tags on Protein VII of the Filamentous Phage

Geir Åge Løset1,2*, Bjarne Bogen1,3, Inger Sandlie1,2*

1 Centre for Immune Regulation, University of Oslo, Oslo, Norway, 2 Department of Molecular Biosciences, University of Oslo, Oslo, Norway, 3 Institute of Immunology, University of Oslo, Oslo, Norway

Abstract

**Background:** Phage display is a platform for selection of specific binding molecules and this is a clear-cut motivation for increasing its performance. Polypeptides are normally displayed as fusions to the major coat protein VIII (pVIII), or the minor coat protein III (pIII). Display on other coat proteins such as pVII allows for display of heterologous peptide sequences on the virions in addition to those displayed on pII and pVIII. In addition, pVII display is an alternative to pIII or pVIII display.

**Methodology/Principal Findings:** Here we demonstrate how standard pIII or pVIII display phagemids are complemented with a helper phage which supports production of virions that are tagged with octa FLAG, HIS6, or AviTag on pVII. The periplasmic signal sequence required for pIII and pVIII display, and which has been added to pVII in earlier studies, is omitted altogether.

**Conclusions/Significance:** Tagging on pVII is an important and very useful add-on feature to standard pIII and pVIII display. Any phagemid bearing a protein of interest on either pII or pVIII can be tagged with any of the tags depending simply on choice of helper phage. We show in this paper how such tags may be utilized for immobilization and separation as well as purification and detection of monoclonal and polyclonal phage populations.

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* E-mail: g.a.loset@imbv.uio.no (GAL); inger.sandlie@imbv.uio.no (IS)

Introduction

Phage display is a platform for selection of binders with affinity for specific target molecules, and also exhibits high versatility with respect to target discovery [1]. In both cases, libraries of polypeptides are created as fusions to phage coat proteins that are solvent exposed [2,3]. The wt filamentous phage virions M13, fd and f1 have about 2,700 copies of the major coat protein pVIII, and in addition, express approximately 3–5 copies each of pIII, pVI, pVII and pIX; pIII and pVI on one virion tip and pVII and pIX on the other [4]. Polypeptides have been fused to and displayed on all five structural proteins, but only pIII and pVIII display have gained widespread use. With the exception of pVII display, which has been evaluated for use with cDNA libraries [5], it is common to the majority of phage display protocols that the heterologous peptide is placed in-frame between an N-terminal signal sequence and the mature form of the viral capsid protein. Alternatively, the heterologous peptide can be combined with a modified version of pII in the periplasm, but both components are still dependent on signal sequence-directed periplasmic targeting [6,7].

Fusion coat proteins may be encoded either in a phage genome or by a phagemid, and in the latter case, complementation by a helper phage is needed to support virion production. It is also possible to combine multiple display of more than one type of fusion protein per virion by using both genome-based vectors [8,9,10,11,12] and phagemid systems [13]. The resulting bifunctional phage particles have been utilized in a number of areas such as life sciences [9], as therapeutics [8,10,11] and in solid state material sciences [12,13].

Since pVII, like pIII, is found at the virion tip in 3–5 copies, peptide display on pVII might be an attractive alternative to pIII display, and indeed, genomic pVII display has been reported [14]. In that study, an N-terminal signal sequence targeting the fusion protein to the periplasm was held as a component critical for successful display. However, of the five structural capsid proteins that build the filamentous phage virion, only pIII and pVIII are synthesized as precursors containing N-terminal signal sequences [4].

Here, for the first time we demonstrate that signal sequence directed periplasmic targeting of the fusion is not necessary for functional pVII display. We explore display through pVII and engineer helper phage genomes that combined with standard pIII or pVIII phagemid display support the production of bispecific virions. Defined tags were added to virions with pIII or pVIII fusions. Thus, we used the modified helper phages to effectively FLAG-, HIS6- and AviTag-tag phage particles with pIII or pVIII phagemid-encoded fusions. We also show pIII/pVII phage-
M13 Helper Phages with Tags Fused to pVII

A series of mutants were constructed based on the two genetically different, but functionally identical helper phage genomes, M13K07 and VCSM13. We inserted either an octa-FLAG, hexa-His (HIS6) or a BirA recognition sequence (AviTag) N-terminally to the pVII open reading frame (ORF) such that the critical single nucleotide spacing to the upstream pV ORF and the start codon was maintained [15] (Fig. 1A). The M13K07 modified helper phage genomes were then introduced into host cells, propagated as normal helper phages and the culture supernatants titered for phage content as kanamycin resistant colony forming units (cfu\(^{\text{kanR}}\)), which was found to be comparable to the non-pVII modified helper phage (Fig. S1A). Identical results were obtained with the pVII modified VCSM13 (data not shown). We then tested the modified M13K07 for their ability to rescue pIII display phagemids encoding either a single chain Fv (scFv) or a single chain T cell receptor (scTCR) [16]. The three pVII tagged helpers performed essentially as the untagged helper. Thus, the phage clearly tolerated peptides that differed in length, pI and charge fused N-terminally to pVII (Fig. S1B - D). We then absorbed virions to tag-specific supports (Fig. 1B-E). The AviTag-virions were absorbed to streptavidin (SA) coated beads, and bound virions detected by an anti-M13-HRP monoclonal antibody (mAb). Initial experiments demonstrated inefficient virion capture, presumably due to low endogenous BirA activity as the standard phagemid packaging protocol is conducted at 30°C (data not shown). This was overcome by making a new F\(^+\) E. coli host strain, AVB100FmkII, which over-expresses birA. Phagemid rescue using this strain led to markedly improved SA-specific binding of the virions (Fig. 1B, left part). Approximately 70% of the input phages were captured on beads estimated by titering the phage content in the waste (as cfu\(^{\text{ampR}}\)) after extensive washing of the bead-virion complexes (Fig. 1B, right part). These data demonstrate that the AviTag-pVII fusion is surface exposed and functional as a BirA substrate. The FLAG-tagged phages bound strongly to two different anti-FLAG mAbs, M2 and M5 (Fig. 1C). M5 binds the FLAG peptide only when located at an unprocessed N-terminus, and the strongest reactivity was seen with M5, nearly reaching the

Figure 1. pVII tag-modified helper phages are structurally and functionally identical to normal helper phages, and donate a defined phenotype to pVII. (A) Schematic illustration of the pV, pVII, pIX genomic junctions in the M13 genome framed by the unique BsrGI/SnaBl RE sites. The three tag modifications, their site of insertion and physical characteristics are given. Isoelectric point (pl) as well as average charge ($) was computed using ProtParam (http://ca.expasy.org/). (B) AviTag-pVII functionality as BirA substrate assessed by virion binding to magnetic 5A beads and detection with an anti-M13 Ab (BirA enzyme activity provided by E. coli XL1-Blue or AVB100FmkII). (C) FLAG-pVII functionality as assessed by virion binding to immobilized anti-FLAG M2 and M5 mAbs and detection with anti-M13 Ab. (D) HIS6-pVII functionality assessed by virion binding to magnetic IMAC beads and detection with an anti-M13 Ab. Virion binding to the beads was done without (gray bars) or with (black bar) 300 mM imidazole in the binding buffer. (E) HIS6-pVII functionality assessed as in D, employing HIS6-pVII tagged pVIII phagemid display (pGALD8mFN, unpublished) virions (left part). Specific IMAC bead capture efficiency assessed by titration of imidazole-eluted pVIII phagemid virions after bead capture, shown as percent of in-put (% recovery = out-put (cfu\(^{\text{ampR}}\))/in-put (cfu\(^{\text{ampR}}\)) x100), right part. doi:10.1371/journal.pone.0014702.g001

Display of Tags on pVII

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level observed with polyclonal anti-pVIII antibodies. The latter was surprising, since phages contain thousands of pVIII epitopes and a maximum of only five pVII-FLAG epitopes. Thus, the observed binding clearly underscores strong and specific M5 phage absorption. The HIS6-tagged virions bound IMAC matrix beads in pH dependent manner (Fig. S2), and the binding could be specifically abolished using imidazole (Fig. 1D). Increasing the spacer length between pVII and the HIS6-tag did not increase the efficiency of bead capture (Fig. S2). Moreover, similar IMAC bead capture properties was also seen with pVIII display phagemid virions rescued using this HIS6-pVII helper phage system (Fig. 1E, left part), and based on the in-put to out-put titers after elution, there was a 150-fold differential recovery depending on the absence or presence of the HIS6-modification (Fig. 1E, right part). Imaging analysis by electron microscopy showed that the pVII modified virions were morphologically identical to the unmodified virions (data not shown), ruling out that the pVII modification introduces any gross physical alterations of the virions as compared to their unmodified counterparts.

**pVII-tagging differentiate mixed virion populations**

The ability to produce distinct virion populations with different pVII tags may allow their physical separation from complex mixtures. As proof of concept, we blended a small amount of *E. coli* AVB100FmkII in *in vivo* biotinylated AviTag-virions into a large amount of untagged “background” virions. We then did a single SA bead capture, and the bead-virion complexes used for direct infection of fresh host cells after extensive washing. We found that the procedure led to a 5 x 10^9-fold enrichment of tagged virions with an untagged background of 0.002%. (Fig. 2).

**Bispecific multivalent pIII and pVII display**

The results above clearly show that bispecific phages are obtained by the combined use of pIII (phagemid encoded) and pVII (helper phage encoded) display. To test whether virions are assembled when pIII and pVII fusions are encoded within the same phage genome, we introduced the AviTag-pVII modification and the scTCR-pVII fusion into fUSE5 [17], thereby creating the bispecific pIII-pVII phage genome display vector f37AviTag. Functional display on both distal tips of the virions was indeed demonstrated (Fig. S3).

**Discussion**

Heterologous peptide display on pIII or pVIII is based on signal sequence dependent translocation of the fusion from cytosol to the periplasm. Here, we show efficient display on another capsid protein, namely pVII. In contrast to earlier reports, this was achieved without leader mediated periplasmic targeting of the fusion [14,18]. Despite the fact that the pVII protein lacks any known signal sequence, it is yet inserted into and spans the inner membrane prior to virion incorporation without post-translational processing [4,19]. Although the mechanism of both wt and modified pVII membrane insertion is unknown, the lack of a signal sequence signature suggests a different transport route than either of the SRP-, SEC, TatABC or YidC-dependent pathways of the *E. coli* secretory machinery [20]. The discrepancy between the current and an earlier report regarding the feasibility of genomic pVII fusion protein display without an N-terminal leader sequence may at least partly be due to the nature of the chosen fusion protein [19]. The former study used glutathione-S-transferase, which is a globular protein that readily folds and dimerizes in the cytosol, which may have hampered periplasmic targeting [21]. The lack of an N-terminal leader sequence may alleviate unpredictable heterogeneity in functional peptide display due to incomplete or lack of leader peptidase signal sequence cleavage [22].

All three pVII-tag fusions reported herein were encoded on helper phage genomes, which in turn were used to support production of virions with heterologous fusions to pIII or pVIII encoded on phagemids. Thus, the resulting virions displayed a tag on pVII as well as larger fusions on pIII or pVIII. In contrast to earlier reports on bispecific display that requires reformatting to new dedicated vectors, these novel helper phages significantly improves the versatility and ease of use with existing systems [8,9,13,18]. Any pIII or pVIII display phagemid encoding single clones or a library can be used with one of the pVII modified helper phages described, and bispecific display with a pVII-tag of choice is obtained after standard phagemid rescue. Importantly, the tag on pVII does not interfere with the function of the other fusion. Furthermore, the tag-peptides reported herein, namely AviTag, FLAG and HIS6, which are among the most commonly used tags for detection, immobilization and purification, may be very easily exchanged with another tag of interest in a quick helper phage mutagenesis step. As a very high affinity SA tag, AviTag was chosen, rather than alternatives such as the Strep-tag, despite its need for enzymatic modification to gain functionality. This potentially labor intensive step was easily integrated into the normal phage propagation cycle merely by exploiting an F positive version of the *bchA* over-expressing *E. coli* strain AVB100. In *in vivo* biotinylation of the virion was thereby achieved without adding extra processing steps.

In the examples above, the pIII and pVII fusions are found on separate genetic elements (a phagemid and a helper phage genome); hence only low valence pIII is achieved. Using the genome display vector f37, we also show that pIII and pVII fusions are equally well tolerated when encoded in the same vector rendering multivalent bispecific display. This genomic system also offers the potential advantage of simplifying and speeding up the turn over time during propagation as no helper phage super

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**Figure 2. Tag-pVII mediated enrichment of virions.** In *in vivo* biotinylated phagemid (ampR) virions displaying a scTCR on pIII and AviTag on pVII were spiked into VCSM13 helper phage (kanR) at a ratio of approximately 3 x 10^-7. After a single round of SA bead capture and extensive washing, the amount of phagemid (cfu<sub>ampR</sub>) and helper phage (cfu<sub>kanR</sub>) virions captured were determined using the beads for direct infection as described in Methods. The results are shown as percent of in-put retrieved (NA, not applicable). doi:10.1371/journal.pone.0014702.g002
infection step in conjunction with controlled bacterial growth and processing steps are needed.

Finally, we show that one virion population can be retrieved after mixing with a different virion population, using the pVII fusion tag. Importantly, the separation of the virion particles takes place in solution before any selective propagation step. Differential tagging of e.g. two separate pIII libraries would allow library against library selection [23] and physically distinguish between two or more populations of virions independently of their POIs displayed.

Although not explored here, it is highly conceivable that type 7 offers an attractive alternative to current type 3 display for construction of and affinity selection from large peptide libraries [24]. Since pIII is of particular importance for early events in E. coli entry, a complete lack of infectivity interference is expected when pIII is wt [25]. Indeed, in a direct comparison between signal sequence-dependent genomic pIII and pVII display, Kwanskiowski et al. reported superior antigen reactivity with pVII display [14]. It may well be that signal sequence independent peptide libraries offer an additional advantage due to lack of heteronomous leader peptidase processing [22]. An additional advantage to pVII display, leaving pIII unaltered, is that virion rescue following a library selection step may be performed without breaking the virion-target bond, as elution may be done by infection directly on the solid phase. Especially for retrieving high affinity binders, when the strong virion-target interaction may be resistant to a variety of elution strategies, this simplifies rescue and may well increase the successful isolation of such binders [26].

In summary, we have demonstrated display of three peptide tags on pVII that are up to 17 aa in length and have different charge and pI. Hence, despite the large heterogeneity in the physio-chemical properties between these three fusions, no apparent influence on wt pVII function was observed. A summary of the different phagemid and phage genome display methods are shown in Figure 3. We believe this pVII-tagging approach by use of modified helper phages will significantly expand the versatility of existing pIII and pVIII display.

Materials and Methods

Plasmids, bacterial strains, phage and materials

The phOx-BSA and pSEX81 phagemid vector harboring a phOx-BSA specific affinity matured human scFv was kindly provided by Affitech Research AS (Oslo, Norway). The pSEX and pFKPDN phagemids harboring the scTCR V\22C2 and pVII have been described previously [16,27]. The E. coli strains K91K (thi, lacZ::Tn9, K12) and MC1061 (hisD mcrB araD139 M15 transcat [Tn10 (Tet)] and CJ236 (F [lacD139K, rpsL, hsdS (K1- strains XL1-Blue ( recA1 endA1 ugrA96 thi-1 hsdR17 supE44 relA1 lac [F proAB lacZAM15 Tn10 (Tet)] and C236 [F (ThiIII)]-cot (Thy+PcrCam+)/ ung-1 relA1 dut-1 thi-1 spo1T1 mcrA) were purchased from Stratagene (LaJolla, CA, USA) and New England Biolabs (Ipswich, MA, USA), respectively. The E. coli MC1061-derived AVB100 strain that harbors a L-arabinose inducible biaA cassette was purchased from Avidity LLC (Aurora, CO, USA). To obtain the F-positive AVB100FmikII (Strr/Tetr), the AVB100 parent was mated with XL1-Blue using standard methodology [2]. Likewise, the F-positive AVB100F (Strr/Camr) was obtained by mating of AVB100 with CJ236. M13K07 and VCSM13 helper phages were purchased from GE Healthcare (Uppsala, Sweden) and New England Biolabs (Ipswich, MA, USA), respectively. All restriction enzymes (RE) were purchased from New England Biolabs (Ipswich, MA, USA). DNA oligos were purchased from MWG Biotech AG (Ebersberg, Germany). Pfu Ultra and Phusion DNA polymerases were purchased from Stratagene (LaJolla, CA, USA) and Sigma-Aldrich (Oslo, Norway), respectively. The anti-M13-HRP Ab and anti-FLAG M2 and M5 mAbs were purchased from GE Healthcare (Uppsala, Sweden) and Sigma-Aldrich (Oslo, Norway), respectively. The F23.2 mAb was a kind gift from Dr. Uwe D. Staerz (Department of Medicine, National Jewish Medical and Research Center, Denver, USA) and the GB113 mAb was purified from cell supernatant on protein G-sepharose (GE Healthcare,

![Figure 3. Tag-pVII phage display routes.](image-url) (A) The M13K07 helper phage modified with an N-terminal pVII tag assembles into virions displaying the tag of choice on all 3–5 pVII copies on the virion tip. In principle, this modified helper phage genome is therefore analogous to a pII phage genome vector. (B) Used in phagemid rescue of any standard pII phagemid in a 3-3 system, the Tag-pVII modified helper phage yields defined, bispecific virions. Normally, such pII phagemids render low valence display on pII, whereas there will be multivalent Tag-pVII display. (C) Used in phagemid rescue of any standard pVII phagemid in an 8+8 system, the Tag-pVII modified helper phage yields bispecific display where all pVII carry the tag, whereas the many thousand pVIII copies of the virion body will be a heterogeneous blend of POI-pVIII dispersed among wt pVIII. (D) When the Tag-pVII modification is integrated into a pII phage genome display vector, a system is made in which both virion tips are fully modified with a fusion peptide on all copies of both pIII and pVII. doi:10.1371/journal.pone.0014702.g003
Design and construction of the pVII modified phage genomes

Codon-optimized versions for prokaryotic expression of Avi-TagTM (N-MSGLDNIDEFAQKIEWHE-C), FLAG-tag (N-MDYKDDDDKK-C) and the HIS6-tag (N-MHHHHHHHHH-C) peptide sequences were attached 5′-terminally to the pVII ORF in M13K07 by QuikChange™ in vitro mutagenesis according to the manufacturers’ protocol (Stratagen, LaJolla, CA, USA). Primer design was based on a DNA sequence alignment of M13K07, VCSM13 and iUSE5 (Fig. S4). The modifications were verified by DNA sequencing (in-house ABI lab DNA sequencing core facility, Dept. Molecular Biosciences, University of Oslo). To ensure a clean vector background, a BosGI/StuI RE fragment containing the modified pVII was moved into unmodified genomes on compatible RE sites using standard techniques. The DNA constructs were introduced into the *E. coli* hosts XL1-Blue or MC1061 by electroporation. Primer sequences and GenBank accession numbers for the resulting constructs are listed in Table S1.

Virion production

Phagemid rescue from *E. coli* XL1-Blue using M13K07 or VCSM13 helper phages was done essentially as described [30]. Recombinant phages were amplified from *E. coli* MC1061 transformed with iUSE5 essentially as described [31]. Phagemid rescue in AVB100mkII and phage amplification in AVB100F were done essentially as for XL1-Blue and MC1061, respectively, and increased in *in vivo* biotinylation was achieved by supplementing the growth medium with 50 μM d-biotin and 0.4% w/v L-arabinose (final concentrations). Virion assembly was monitored by spot titration as described [25]. Where applicable, the virions were purified and concentrated by PEG/NaCl precipitation as described [32], and resuspended in PBS, pH 7.4. To estimate the *in vivo* biotinylation level in AVB100mkII, the virions were 2x PEG precipitated and captured on SA beads as described below. The biotinylation level was then determined as 100-f, where f is the fraction (in %) of virions in the waste of the in-put.

Phage capture ELISA and bead capture

In ELISA, the various targets (Abs and phOx-BSA) were absorbed to MaxiSorp™ microtiter plate wells (Nunc, Roskilde, Denmark) in concentrations from 2.5 to 5 μg/ml in PBS, pH 7.4 overnight at 4°C. The wells were blocked with either PBS/TM (PBS supplemented with 0.05% w/v Tween 20 and 4% w/v skim milk) or 2% w/v BSA (in PBS) for 1 h at room temperature (RT). Virion preparations where then added and allowed to react for 1 to 2 h at RT, before captured virions were detected with anti-M13-HRP (1:2000) and the tubes the tubes were incubated for 1 h/RT on a rotating wheel. The tubes were washed 3x in PBST as described above. 0.5 ml of PBST containing an anti-M13-HRP Ab (1:2000) was then added to each tube and the tubes were incubated for 1 h/RT on a rotating wheel. The tubes were washed 3x in PBST as described above. 500 μl TMB soluble was added and the reaction stopped by an equal volume of 1M HCl after 5 min at RT. 100 μl of each solution was transferred to Maxisorp ELISA strips (Nunc, Roskilde, Denmark) and the absorbance measured at A450 nm. For IMAC phage bead capture, 50 μl/sample TALON™ beads were prepared in Washing and Binding (WB) buffer, either with pH 7 or 8, essentially as described by the manufacturer. In one experiment, the WB buffer (pH 8) was supplemented with 300 mM imidazole. The appropriate phage titrations were then added in a total volume of 500 μl (supernatant or PEG precipitated) and incubated for 1 h/RT on a rotating wheel. After washing the beads as described above using WB buffer, anti-M13-HRP Ab (diluted 1:5000 in PBST) was added to each tube and the tubes were further incubated 1 h/RT on a rotating wheel. After washing in PBST, 500 μl TMB soluble was added and the reaction stopped by an equal volume of 1M HCl after 5 min at RT. 100 μl of each solution was transferred to Maxisorp ELISA strips and the absorbance measured at A450 nm.

Supporting Information

**Figure S1** Virion assembly and functionality analysis of pVII modified helper phages. (A) Virion assembly efficiency in *E. coli* XL1-Blue. Normal (denoted as wt) and pVII tag-modified M13K07 helper phage production was assessed by infectious titration, and values given as the number of kanamycin-resistant (kanR) colony forming units (cfu) per ml culture supernatant. (B) Phagemid rescue ability of normal and modified M13K07 helper phages. Ampicillin-resistant (ampR) anti-phOx scFv- and scTCR-pIII encoding phagemids were rescued and the virion production capacity given as the number of cfuampR/ml by infectious titration. (C) Phagemid (ampR) to helper phage (kanR) ratio in infectious titration. (D) The integrity of the pII displayed POI (scFv anti-phOx or scTCR) on virions rescued with normal or the modified M13K07 helper phages assessed by antigen-specific phage capture ELISA. Briefly, phOx-BSA or the TCR surrogate Ag mAb GB113 (the cognate Ag for the TCR is the murine pMHC II complex I-Ed/2315) was coated in microtiter wells and interacting virions detected with an anti-M13 mAb as described in Methods. Notably, the GB113 Ab is clonotypic for the 4B2A1 TCR from which the scTCR is derived [28].

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**Figure S2** IMAC bead capture efficiency of HIS6-pVII virions depend on tag-pVII spacer and pH during capture. (A) PEG precipitated VCSM13 helper phage virions or pFKPDN-scTCR 4B2A1 phagemid-derived virions were prepared as described in Methods. The bead capture was done as described by the manufacturer protocol in the recommended sample volumes as described in Methods. The Binding and Washing buffer was used with either pH 7 or 8 and the virion in-put titer was normalized to 1x 1010/ml in PBS for all samples. Binding and subsequent detection of captured virions were done with an anti-M13 Ab as described in Methods. Two different HIS6-pVII modified helper...
phages were used, denoted HIS6 and HIS6-GS2, of which the latter has a 10 aa spacer between the HIS6-tag and the pVII capsid protein (illustration in B). Both versions of the HIS6-modified VCSM13 helper phage virions, but not the unmodified (denoted wt), were captured on the IMAC beads. The HIS6 version bound with higher efficiency than the HIS6-GS2 version. Moreover, virion capture was most efficient when done at pH 8. The latter was also seen with phagemid-derived virions packaged with the HIS6-pVII modified VCSM13 helper phage.

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**Figure S3** Bispecific multivalent scFv-T-III and AviTag-pVII display encoded by a single phage genome. (A) Virion assembly efficiency in E. coli MC1061 from USE5 (harboring a scFv T-III fusing with the fusion partner AviTag). Numbers of tetracycline-resistant (tetR) cfu/ml in culture supernatants and the corresponding PEG precipitated samples are given as assessed by infectious titration in E. coli K91 (B). (C) The integrity of the scFv-T-III fusions on the virions assessed by binding to conformation (F23.2) and TCR clonotypic (GB113) mAbs in ELISA using tier normalized virion in-puts. (C) AviTag-pVII functionality as endogenous BirA substrate (both E. coli XL1-Blue (M13K07) and MC1061 (E3AviTag)) assessed by virion binding to magnetic SA beads detected by an anti-M13 Ab. (D) Increased AviTag-pVII in vivo biotinylation is obtained with E3AviTag using the E. coli AYB100F host strain (see Methods) which over-expresses birA upon L-arabinose induction, as shown by increased virion binding to magnetic SA beads detected by an anti-M13 Ab.

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**Figure S4** Multiple sequence alignment of the M13K07, VCSM13 and USE5 genomes. The M13K07 (New England Biolabs sequence), VCSM13 (GenBank accession no.: AY598820) and USE5 (GenBank accession no.: AF218364) were aligned using ClustalX 2.0.5 and manually annotated using GeneDoc (http://www.psc.edu/biomed/genedoc). Only the relevant parts of the genomes framed by the unique BsRGI and SmaBI RE sites are shown together with the pV, pVIII and pIX ORFs (only partial ORFs for pV and pIX, respectively).

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**Table S1** GenBank accession numbers and QuickChange mutagenesis primers. The M13K07 (New England Biolabs sequence: http://www.neb.com) and VCSM13 (GenBank accession no.: AY598820) is functionally identical both in their normal and pVII-modified versions. Both genomes are 100% identical in the BsRGI/SmaBI defined region where the pVII modification is found. The pVII modifications use were made in the M13K07 genome and verified sequences shuffled into VCSM13 and USE5 (thereby creating E37) on the compatible BsRGI/SmaBI RE sites. The sequences submitted to GenBank were based on the existing VCSM13 and USE5 GenBank entries AY598820 and AF218364, respectively.

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**Author Contributions**

Conceived and designed the experiments: GL. Performed the experiments: GL IS. Contributed reagents/materials/analysis tools: BB. Wrote the paper: GL IS.

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