Effect of DNA copy number on genetic stability of phage-displayed peptides

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A small model peptide, the FLAG® epitope, was cloned into two filamentous phage display vectors, f88-4 and fd88-4, creating phages f88-FLAG and fd88-FLAG, respectively. Both vectors have a gene VIII display cassette (in addition to their normal phage gene VIII) and display the cloned peptide on a few percent of the virion’s 3000–4000 pVIII (major coat protein) subunits. Vector f88-4 has a replication defect and attains low DNA copy number in infected cells, while vector fd88-4 has no replication defect and attains the normal, high DNA copy number characteristic of wild-type filamentous phage. Almost no loss of displayed peptide was observed during six rounds of propagation of low copy number f88-FLAG phage. In contrast, when high copy number fd88-FLAG phage was similarly propagated, variant clones that did not display the FLAG epitope accumulated gradually. The loss of displayed peptide from the high copy number vector is undoubtedly slow enough to be overcome by even weak affinity selection, and high copy number vectors have important advantages that make their use worth considering, at least when the displayed peptides are small.

INTRODUCTION

Filamentous bacteriophage, the basis of most phage display constructs (1), contains a circular single-stranded DNA (ssDNA) encased in a tubular capsid composed primarily of thousands of copies of the major coat protein pVIII. They infect strains of Escherichia coli that harbor the conjugative F episome and therefore display the F pilus on their surface. The pilus mediates the infection process, which culminates in the penetration of the viral ssDNA (the plus strand) into the cell, where a complementary DNA strand (the minus strand) is synthesized by host polymerases to form the double-stranded replicative form (RF). Minus-strand synthesis is initiated with high efficiency by host RNA polymerase at a special minus-strand origin in the intergenic region (2) of the plus strand but can occur at low efficiency in the absence of the minus-strand origin (3). Rolling-circle replication of the double-stranded RF produces progeny plus strands, a process that requires the phage replication protein pII acting at a plus-strand origin that also lies in the intergenic region. Early in infection, progeny plus strands serve as a template for minus-strand synthesis. Balanced plus- and minus-strand synthesis thus results in the accumulation of double-stranded RF molecules to a copy number of 100 or more. Late in infection and in chronically infected cells, however, the phage ssDNA-binding protein pV sequesters nearly all progeny plus strands into a filament-shaped complex. These ssDNAs are extruded through the cell envelope, concomitantly shedding pV and acquiring the virion coat proteins, including pVIII, from the inner membrane to emerge as completed virions. The extrusion of progeny virions does not kill the cell; chronically infected cells continue to divide, although at a slower rate than uninfected cells. It is the slowing of cell division, not cell lysis, that explains plaque formation by these phage.

Both filamentous phage-display vectors used in this work are of type 88, meaning they have two copies of gene VIII (encoding pVIII). One is the normal, wild-type gene, and it encodes most of the virion’s thousands of major coat protein subunits. The other is a gene VIII display cassette (Figure 1) that has restriction sites that allow for foreign peptide coding sequences to be fused to the gene VIII coding sequence. The bottom part of Figure 1 shows how the coding sequence for the FLAG® peptide is cloned into these sites to create a fusion protein with the FLAG peptide at the N terminus of the mature form of the coat protein (after cleavage by signal peptidase). The display cassette is expressed from an isopropyl-β-D-thiogalactopyranoside (IPTG)-inducible tac promoter operator; under fully induced conditions, approximately 5% of the pVIII subunits derive from the cassette and display the foreign peptide.

One of the type 88 vectors used in the present study, f88-4, is derived from phage fd-tet (4), which has a replication defect and therefore reaches low DNA copy number in infected (or transfected) host cells. As diagrammed in the upper part of Figure 2, the replication defect results from a disruption of the minus-strand origin by a long (2.8-kb) tetracycline-resistant determinant (Tet), and the gene VIII display cassette replaces a few hundred noncoding base pairs within the determinant. Plaques formed by fd-tet derivatives are so small that it is impractical to quantify infection in terms of plaque-forming units (pfu). However, because infection transduces the infected cell to tetracycline resistance, infectious units can be effectively quantified as transducing units by spreading infected cells on tetracycline-containing nutrient agar and counting colonies. These phages can be propagated independently of infection, even in an uninfected F− host, by culturing the phage-bearing cells in medium containing the antibiotic. The replication defect reduces infectivity (transducing units per physical particle) about 10-fold relative to wild-type phage (typical infectivity of 0.5 pfu/particle). That is because minus-strand synthesis is greatly delayed in a newly infected cell, thus delaying the expression of the tetracycline-resistant determinant and subjecting the unpaired plus strand to degradation. When the cells are finally challenged with a high concentration of the antibiotic on agar plates or in liquid medium, only a small minority survives. The yield of physical particles in fd-tet-derived vectors such as f88-4 is also reduced to about 25% of normal levels. The net effect of low infectivity and low particle yield is an overall 40-fold reduction in the yield of infective units when compared with wild-type high copy number phage.
The obvious disadvantages of the replication defect in f88-4 and other fd-tet-derived vectors are offset by a key advantage: it largely averts a complication called “cell killing.” When phage assembly is fully or partially blocked, intracellular phage DNA and gene products accumulate to toxic levels, and the host cell is killed without releasing progeny phage (5). Cell killing is nearly absent in fd-tet derivatives such as f88-4 because of their low RF copy number, and even severe morphogenetic defects are readily tolerated (6). Display vectors based on fd-tet therefore accommodate recombinant coat proteins that partially impair phage assembly or that are directly toxic in their own right. Because there is less selection pressure against foreign peptides, we expect displayed peptides to be more stable genetically in low copy number than in high copy number vectors.

The low copy number vector f88-4 has already been used extensively for both random and natural peptide libraries (7–9). But is it necessary to incur the disadvantages of a low copy number vector, especially when the displayed peptides are short? To address this question, we constructed a high copy number type 88 vector, fd88-4, which has the same gene VIII display cassette as f88-4. As shown in the lower part of Figure 2, the cassette is inserted into a nonessential part of the intergenic region of wild-type fd, where it does not disrupt either the minus-strand origin or the nearby stem loop that serves as a packaging signal during the extrusion of progeny virions. A short model peptide, the FLAG epitope recognized by commercially available monoclonal antibodies, was cloned into both the high and low copy number vectors as diagrammed in the lower part of Figure 1, creating clones fd88-FLAG and f88-FLAG, respectively. Each construct was subjected to six rounds of propagation without selection in favor of the FLAG epitope. Then, after the final round of nonselective propagation, several dozen individual phage subclones were tested for retention of the FLAG epitope. Most, but not all, of the fd88-FLAG subclones had lost the displayed peptide, while all but one of the f88-FLAG subclones retained it. These results therefore corroborate the supposition that displayed peptides are more genetically stable in low copy number vectors than in high copy number vectors. At the same time, the rate of epitope loss from the high copy number construct is not so high as to make fd88-4 unappealing as a vector for small peptide libraries.

MATERIALS AND METHODS

Solutions and Media

Tris-buffered saline (TBS) is 50 mM Tris-HCl, pH 7.5, 0.15 M NaCl. TBST is TBS with 0.5% (v/v) Tween® 20. Dialyzed bovine serum albumin (BSA; Sigma, St. Louis, MO, USA) was dissolved at 50 mg/mL in water and filter sterilized. TTDB is TBST supplemented with 1 mg/mL in water and filter sterilized. TBS/gelatin is TBS supplemented with 1 mg/mL gelatin and autoclaved. NZY and NZY/Tet liquid medium and nutrient agar Petri dishes have been previously described (10). Soft agar contains 1 g Bacto™ tryptone (Fisher Scientific, St. Louis, MO, USA), 0.5 g NaCl, and 0.75 g Bacto agar (Fisher Scientific) per 100 mL and is stored at room temperature after autoclaving and melted in a microwave oven before use. X-gal stock solution contains 2% X-gal in dimethyl formamide and is stored at -20°C, protected from light.

Phage Constructs and E. coli Bacterial Host Strains

All constructions were accomplished by standard recombinant DNA techniques and confirmed by DNA sequencing. The low copy number vector f88-4 has been previously described (GenBank® accession no. AF218363). The high copy number vector fd88-4 was constructed by inserting the gene VIII display cassette (upper part of Figure 1) into the wild-type phage fd. Starting at fd position 5605 (GenBank accession no. J02451), the plus-strand sequence immediately preceding the first position of the cassette is 5′-GC-CACGTT-3′. Ending at fd position 5624, the plus-strand sequence immediately following the last position of the cassette is 5′-CGGCTTTCC-3′. The coding sequence for the FLAG peptide
was installed in both vectors, as shown in the lower part of Figure 1. 

E. coli K-12 host strains MC1061 [F− hsdR mcrB Δ(araABC–leu)6779 araD139 ΔlacI74 galU galK strA thi] and K91 (Hfr Cavalli thi) have been previously described (10). K91 deploys the F pilus and is therefore infectible by filamentous phage. MC1061 has no F pilus and is uninfectible but supports phage production at normal levels.

**Titering Plaque-Forming and Transducing Units**

The nonreplication-defective phage fd88-4 and fd88-FLAG were titered for blue plaques as follows. Phages were diluted to a suitable range of concentrations in TBS/gelatin, and portions of 25–400 μL were pipetted into sterile 15-mL polypropylene tubes (Fisherbrand®; Fisher Scientific). To each tube, we added 400 μL of a late log-phase culture of K91 cells and approximately 3 mL molten soft agar at 50°C premixed with 60 μL 2% X-gal. Immediately after the addition of the molten agar, the contents of each tube were poured onto an NZY agar plate, which was tipped and rocked to distribute the mixture evenly over the entire surface. After the soft agar had gelled, the plates were incubated overnight at 37°C. Plaques are visible within about 10 h, and blue color is well-developed in positive clones 6–10 h later. Phage variants that have lost a functional lac operator give white (i.e., naturally colored) plaques under these circumstances (see Results and Discussion).

Tetracycline transducing units for the replication-defective phage, such as f88-4 and f88-FLAG, were titered on K91 cells as previously described (10). Each transducing unit is manifested as a colony of phage-bearing tetracycline-resistant cells on agar medium containing that antibiotic.

**Nonselective Propagation of f88-FLAG**

Two well-separated colonies of E. coli strain MC1061 cells bearing phage f88-FLAG (about 10^7 virions/colony) were used to inoculate 35 mL of NZY/Tet medium. After shaking overnight at 37°C, the cells were cleared at 4°C by two successive 10-min centrifugation.

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**Gene VIII display cassette**

![Gene VIII display cassette diagram]

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**FLAG insert in gene VIII display cassette**

![FLAG insert diagram]
tions at 3000× and 7600× g. Virions were precipitated from the resulting culture supernatant by two successive polyethylene glycol (PEG) precipitations as previously described (10). A portion of each clone containing about 1.6 × 10^{10} virions was used to inject 500 μL of an early log-phase NZY culture of *E. coli* strain K91 for 10 min at room temperature, and 400 μL of the infected cells were diluted into 20 mL NZY medium supplemented with 0.2 μg/mL tetracycline (a subinhibitory concentration that suffices to induce expression of the tetracycline resistance gene) in a 125-mL culture flask that was shaken vigorously at 37°C for 1 h. Tetracycline was added to a final concentration of 20 μg/mL (fully inhibitory) and IPTG to a final concentration of 1 mM, and shaking was continued overnight at 37°C. Culture supernatants were cleared of cells by centrifuging at 15,000× g small volumes of culture for 1 min in a microcentrifuge. The propagation cycle described in the previous sentence was repeated four more times, except that the phage inoculum was 1 μL of culture supernatant from the previous propagation cycle (about 5 × 10^8 virions). The final culture supernatants contained virions that had been propagated nonselectively (i.e., without regard to retention of an intact gene VIII display cassette) for a total of six cycles. The phage amplification factor for the first propagation cycle (from the single colonies) was approximately 1.75 million, for the second cycle about 800, and for the remaining four cycles about 25,000.

Nonselective Propagation of fd88-FLAG

Two single blue fd88-FLAG plaques (about 10^7 virions/plaque) were used to inoculate 35-mL portions of an early log-phase culture (about 10^8 cells/mL) of *E. coli* strain K91 in NZY medium, and after shaking overnight at 37°C, the virions were prepared by PEG precipitation from cleared culture supernatant as described in the previous subsection. A portion of each clone containing about 7 × 10^{10} virions was used to inoculate 20 mL of an early log-phase K91 culture that was shaken overnight at 37°C, and culture supernatants were cleared of cells by centrifuging at 15,000× g small volumes of culture for 1 min in a microcentrifuge. The propagation cycle described in the previous sentence was repeated four more times, except that the phage inoculum was 2 μL of culture supernatant from the previous propagation cycle (approximately 4 × 10^9 virions). The final culture supernatants contained virions that had been propagated nonselectively for a total of six cycles. The phage amplification factor for the first propagation cycle (from the single plaques) was about 7 million, for the second cycle about 570, and for the remaining four cycles about 10,000.

Testing Clones for Retention of FLAG Epitope

Suitable dilutions of the final f88-FLAG culture supernatants were titered for transducing units on K91 host cells as described above. Forty-nine well-separated colonies from each clone were picked to grid points on an NZY/Tet agar dish containing 1 mM IPTG (to induce the

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**Figure 2.** Intergenic region of the filamentous phage genome showing positions of the gene VIII display cassette in the f88-4 and fd88-4 vectors. The three known functional elements of the intergenic region are delineated in the lower part. The tetracycline-resistant determinant (Tet) in f88-4 is 2.8 kb long and disrupts the minus-strand origin; the gene VIII display cassette replaces a few hundred noncoding base pairs of the determinant.
display cassette) with sterile wooden sticks, as were 11 negative control clones harboring a phage that does not display the FLAG epitope. After overnight growth of the gridded clones at 37°C, the Petri dish was cooled and blotted with a disk of nitrocellulose. The blot was washed vigorously in tap water to remove all adherent cells, blocked for 1 h at room temperature in 10 μg/mL dialyzed BSA in TBS, reacted for 3 h at room temperature in 5 μg/mL biotinylated mouse monoclonal anti-FLAG antibody M2 (Sigma) in TTDB, washed extensively with TBST, reacted for 30 min in 2 μg/mL alkaline phosphatase conjugate of streptavidin (Jackson ImmunoResearch, West Grove, PA, USA), washed extensively in TBST, and developed for a few minutes in 1-Step™ BCIP/NBT substrate solution (Pierce Chemical, Rockford, IL, USA) until positive purple spots were well-developed. All but one of the 98 f88-FLAG clones were strongly positive. All 11 negative control clones were negative.

Suitable dilutions of the final fd88-FLAG culture supernatants were titered for blue plaques as described above. Of the plaques derived from clone 1, 76% were blue and 24% white, and 9% of the clone 2 plaques were blue and 91% white (white plaques have lost all or part of the lac operator sometime during the six cycles of nonselective propagation, as explained in Results and Discussion). An NZY agar dish was seeded with 400 μL of an early log-phase NZY culture of K91 in 3 mL molten soft agar premixed with 60 μL 2% X-gal. When the soft agar had gelled, 49 blue clone 1 plaques, 42 blue clone 2 plaques, and 11 white clone 2 plaques were picked to grid points on the seeded dish with sterile toothpicks. The gridded dish was then incubated at 37°C for about 20 h to allow the plaques to form and the blue color to develop. All blue primary plaques but none of the white primary plaques gave blue plaques on the gridded dish. The gridded dish was blotted with nitrocellulose, and the blot was washed, blocked, reacted, and developed as described above. None of the 11 white plaques turned out to be positive for FLAG in this test, and 26 of the 49 blue clone 1 plaques were positive, as were 37 of the 42 blue clone 2 plaques.

RESULTS AND DISCUSSION

Properties of fd88-4 Vector and Clones f88-FLAG and fd88-FLAG

When fd88-4 is plated for plaques in a wild-type E. coli host on nutrient agar containing X-gal but no IPTG (or lactose), the plaques are dark blue. The reason is that the lac operator on the high copy number phage genome (Figure 1) titrates the limited supply of Lac repressor molecules, thus derepressing the lac operon on the host-cell chromosome in the absence of IPTG or other lactose inducer. Subclones derived from fd88-4 that give white plaques in these circumstances have presumably lost all or part of the lac operator sometime during propagation. The fd88-4 display cassette (Figure 1) is also presumed to be fully derepressed, even if no IPTG is included in the medium. (Because of its low copy number, f88-4 does not derepress the host-cell lac operon or its own display cassette substantially in the absence of IPTG or other inducers.)

Four liters of fd88-4-infected cells grown to stationary phase yielded a total of 1.6 × 10^16 purified virions with an infectivity of 0.58 pfu/particle and 2 mg of purified supercoiled RF DNA. These statistics are typical for wild-type filamentous phage. The virion yield, infectivity, and RF yield are about 4, 10, and 2 times higher, respectively, than for fd-tet-based phage. All plaques were blue, indicating that the incidence of phage that had lost the lac operator part of the gene VIII display cassette was very low.

The coding sequence for the FLAG peptide was ligated into the cloning sites of both fd88-4 and f88-4 as shown in the lower part of Figure 1, thus constructing the high copy number clone fd88-FLAG and the low copy number clone f88-FLAG. Similar to their respective parent vectors fd88-4 and f88-4, fd88-FLAG formed blue plaques and f88-FLAG formed tetracycline-resistant transductant colonies when titered as described above.

Genetic Stability of the FLAG Peptide in f88-FLAG and fd88-FLAG

Two clones each of the high copy number fd88-FLAG construct and the low copy number f88-FLAG construct were propagated through six cycles of nonselective propagation—that is, without regard to FLAG peptide expression—as described above. For all rounds of fd88-FLAG propagation and all but the first round of f88-FLAG propagation, the host cell was K91. IPTG was included to induce the expression of the display cassette during f88-FLAG propagation but was unnecessary during fd88-FLAG propagation. A random sample of a few dozen subclones from each of the four final phage populations was then tested for retention of the FLAG peptide and (in the case of fd88-FLAG) for a functional lac operator. Of the 98 f88-FLAG subclones tested, all but one were positive for the FLAG peptide. In contrast, only 76% and 8.8% of the subclones of fd88-FLAG clones 1 and 2, respectively, retained a functional lac operator, and of those that did, only 53% and 88%, respectively, were positive for the FLAG peptide (as expected, none of the operator-negative subclones tested were positive for the FLAG peptide). Overall, therefore, 41% and 7.8% of the subclones derived from fd88-FLAG clones 1 and 2, respectively, retained the ability to express the FLAG peptide after six rounds of nonselective propagation. It is entirely possible that the observed differences between the clone 1 and clone 2 populations are merely stochastic, rather than the result of any intrinsic difference between their starting clones. In any case, there were no sequence differences in their gene VIII display cassettes.

There are at least two types of selective advantage an fd88-FLAG clone might gain by deleting all or part of its gene VIII display cassette. First, the DNA would be smaller and therefore able to replicate faster. However, no comparable advantage would accrue to an f88-FLAG clone because in that case it is initiation of minus-strand synthesis, and not completion of already-initiated plus or minus strands, that is the rate-limiting step in replication. Second, expression of the recombinant pVIII would be eliminated, thus relieving the cell at the least of a slight metabolic burden and at the most of a slightly toxic protein (recall that the fd88-FLAG gene VIII display cassette
Comparison of mRNA gene expression by RT-PCR and DNA microarray

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Few studies have compared the quantification of mRNA by DNA microarray to the results obtained by reverse transcription PCR (RT-PCR). In this study, mRNA was collected from the healing femoral fracture callus of adult and juvenile rats at various times after fracture. Ten samples were measured by both methods for 26 genes. For RT-PCR, mRNA was reverse transcribed, amplified, electrophoresed, blotted, and probed with 32P-labeled internal oligonucleotides, which were quantified. For DNA microarray, the mRNA was processed to biotin-labeled cRNA, hybridized to 10 Affymetrix® Rat U34A microarrays, and quantified. Correlation coefficients (r) for each gene for the agreement between RT-PCR and microarray ranged from -0.48 to +0.93. This variation made the interpretation gene-specific. Genes with moderate expression levels gave the highest r values. Increased numbers of absent calls by the microarray software and increased separation between the location of the PCR primers and the microarray probes both led to reduced agreement. Microarray analysis suggested a floor effect in expression levels measured by RT-PCR for two genes. In conclusion, moderate mRNA expression levels with overlap in the location of PCR primers and microarray probes can yield good agreement between these two methods.

REFERENCES


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INTRODUCTION

Messenger RNA (mRNA) can be quantified by a number of methods (1). Among the more popular methods for genes with relatively low expression levels has been reverse transcription PCR (RT-PCR). This semiquantitative method is labor-intensive and gives data for only one, or a very few, gene(s) for each assay (2). This has led to the search for more efficient methods for studying mRNA levels for multiple genes.

DNA microarrays yield much greater data output since one hybridization results in the measurement of expression of all genes on the array at the same time (3). This allows the study of many gene transcripts of interest as well as the discovery of new genes involved in the system under study. While this new technology holds great promise, relatively few studies have been done to compare the results obtained by DNA microarray...