Rapid, High-Level Transient Expression of Papillomavirus-Like Particles in Insect Cells

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ABSTRACT

Empirical scanning of natural or engineered peptide sequences for functional residues is inherently dependent upon efficient expression of large numbers of individual sequence variants to assay their relative functional potency. The insect baculovirus system has been widely used for expression of viral coat proteins, but it generally requires prior isolation and expansion of a plaque-purified recombinant viral stock to generate useful quantities of self-assembled virus-like particles. In search of a more rapid means of expression of analytical levels of the L1 coat protein of cotton-tail rabbit and human type 11 papillomaviruses, we found that even brief transient cotransfection of insect cells with baculovirus plasmid transfer vectors and viral DNA yielded assembled particles that were immunologically indistinguishable from particles obtained with plaque-purified viral stocks. Within six days of plasmid/viral DNA cotransfection of SF9 cells, at least 1–2 µg of assembled L1 particles/100-mm plate could be demonstrated, which proved more than sufficient to assay functionality. Transient cotransfection of insect cells should provide general utility for rapid high-level expression of sets of protein sequence variants, as well as other sequence-scanning applications such as sequence optimization in protein engineering.

INTRODUCTION

Many aspects of protein structural analysis depend upon the expression of a large number of properly modified or folded variant sequences to screen for function (see, for example, Reference 20). In some cases, sufficient quantities of full-length peptide can be recovered by in vitro transcription/translation, but proper folding cannot be controlled (1,16). Short sequences, such as single-chain Fv antibodies, can be expressed and “displayed” in bacteria or in filamentous phage in partially or even completely folded form for screening purposes (23). However, larger proteins of mammalian origin often require expression in eukaryotic cells for proper folding and assembly. Furthermore, a high local concentration of expressed protein monomers can be important in proper folding, particularly in the case of assembly of viral coat proteins. Because transient transfection of mammalian cells such as Cos or 293 is relatively inefficient and low-yielding, screening large numbers of variant sequences by these means can prove quite laborious (8,9). One system that overcomes some of these difficulties is the insect cell baculovirus gene expression vector system (19), which has, for example, proven very popular for generation of virus-like particles (VLPs) (7,22). Nevertheless, a significant drawback of this system is the relatively long time required to obtain plaque-purified, expanded recombinant baculoviral stocks with which to express useful quantities of VLPs. Even the most time-efficient baculovirus system reported to date, the bacmid system, requires two weeks from initial transfection to recovery of screenable product (15).

We have an interest in identifying the amino acid residues that confer antigenic type-specificity on human papillomavirus (HPV) subtypes (2,4). Many of the type-specific epitopes are conformationally dependent and are detectable only upon VLP assembly. The L1 structural coat protein of several animal and human papillomaviruses efficiently self-assembles when expressed in insect cells by means of recombinant baculovirus (13). The time and labor involved in the generation of pure recombinant virus precludes the use of this method to screen a large number of VLP variants produced through site-directed mutagenesis. However, we previously observed that, when expressed in the baculovirus system, recombinant proteins such as anti-stasin and trigrammin were detectable as secreted products in µg/mL quantities within 5–7 days of the initial transfection of insect cells with plasmid and viral DNAs (6,10). Based upon this observation, we examined whether sufficient quantities of papillomavirus L1 protein would accumulate to allow self-assembly into VLPs upon transient expression, particularly if a more efficient baculovirus transfection system, such as the BaculoGold™ system, were utilized. Employing a rapid 6-day transient transfection protocol, we report the production of L1 coat protein of cotton-tail rabbit papilloma virus (CRPV) and HPV11, both properly assembled into VLPs. Extracts prepared from transiently transfected cells contained immunogenic material recognized by type-specific and VLP-dependent monoclonal antibodies generated against either CRPV or HPV11 VLPs. The transiently expressed material was not cross-reactive with other type-specific antibodies, and recognition was
sensitive to alkaline denaturation. The insect-cell-based transient expression system we describe should prove useful in many other high-throughput protein sequence screening applications where high-level expression, coupled with high-fidelity folding and assembly, are required.

MATERIALS AND METHODS

Transient Expression of VLPs in SF9 Cells

The HPV11 L1 structural gene was cloned from clinical isolates using the polymerase chain reaction (PCR) with primers designed from the published L1 sequence (5,12). The CRPV L1 structural gene was cloned by PCR from viral genomic DNA and was the generous gift of Drs. Kathrin Jansen and Loren Schultz (Merck Research Laboratories, West Point, PA, USA). The L1 genes were subcloned into pVL1393 (Stratagene, La Jolla, CA, USA) for expression in SF9 cells.

SF9 cells were cotransfected using the BaculoGold Transfection kit (Pharmingen, San Diego, CA, USA) according to the manufacturer’s instructions with the following modifications: 8 × 10^6 SF9 cells were transfected in a 100-mm dish, with 4 µg of BaculoGold viral DNA and 6 µg of test plasmid DNA. Cells were harvested after 6 days, except where otherwise specified, and assayed for VLP production by Western blot or enzyme-linked immunoabsorbtant assay (ELISA) assay (11).

Preparation of SF9 Extracts and ELISA Assays

Six days after transfection, the plates were scraped to resuspend cells, and the cells were collected by low-speed centrifugation. Cells were resuspended in 300 µL of breaking buffer (1 M NaCl, 0.02 M Tris-HCl pH 7.6) and homogenized for 30 s on ice using a Polytron PT 1200 B with a PT-DA 1205/2-A probe (Brinkman Instruments, Westbury, NY, USA) in a Falcon 2059 tube. Samples were spun at 2500 rpm in a GPR centrifuge (Beckman Instruments, Fullerton, CA, USA) for 3 min to pellet debris. The tubes were washed with an additional 150 µL of breaking buffer, and the supernatants were collected in 1.5-mL microcentrifuge tubes, and resuspended for 5 min in an Eppendorf microcentrifuge (Model 5415; Brinkman Instruments). ELISA assays were begun the same day.

Five microliters of extract were diluted in 50 µL of 1% bovine serum albumin (BSA) in phosphate-buffered saline (PBS) solution, aliquoted onto a 96-well Immulon®2 (“U” bottom) microtitration plate (Dynatech Laboratories, Bethesda, MD, USA) and incubated overnight at 4°C. Extracts were removed and the plate was blocked with 5% powdered milk/PBS. All subsequent wash steps were performed with 1% BSA/PBS. The plate was incubated at room temperature with primary antibody for 1 h. Primary monoclonal antibodies CRPV.5A, H6.C6 and H11.F1 were obtained as ascites stock from Dr. Neil Christensen (Pennsylvania State University). CRPV.5A and H11.F1 are VLP-dependent and type-specific antibodies that recognize CRPV and HPV11 VLPs, respectively (3,4). Monoclonal antibody H6.C6, raised against HPV6 VLPs, recognizes a linear outer epitope that is present in both denatured L1 samples and VLPs. Furthermore, the antibody is cross-reactive with HPV11 (N. Chistensenn, personal communication). They were diluted 10^5-fold in 1% BSA/PBS before use. After washing in 1% BSA/PBS, the plates were incubated for 1 h with secondary antibody, horse radish peroxidase-labeled Goat anti-Mouse IgG (γ) (Kirkegaard & Perry Laboratories, Gaithersburg, MD, USA), diluted 10^-3 in 1% BSA in PBS. After a final washing, the 2,2′-azino-di(3-ethylbenz-thiazolone) sulfonic acid substrate (Zymed Laboratories, South San Francisco, CA, USA) was added and absorbance read at 405 nm.

RESULTS

Expression of CRPV L1 is Maximal After Six Days

To determine the time required for maximal expression, plasmid p1393:CRPV was co-transfected with BaculoGold DNA into SF9 cells and plates were incubated at 28°C for 3, 4, 5, 6 and 7 days. Plates were scraped, cells collected by low-speed centrifugation, resuspended in 200 µL of Laemmli loading buffer and heated for 5 min at 95°C. Twenty-microliter fractions were electrophoretically separated under denaturing conditions, transferred to nitrocellulose and probed with a commercially available rabbit polyclonal antiserum to CRPV L1 (Dako Corporation, Carperteria, CA, USA).

Although a number of cross-reactive bands are visible (Figure 1), there is a band, which appears only in the pVL1393:CRPV fractions, that runs at the expected molecular weight of 55 kDa, which is consistent with the intact CRPV L1 protein. The intensity of this band increases to a maximum in the six-day sample. The other detectable bands are also observed with a wild-type baculovirus control extract and undoubtedly represent protein species cross-reactive with the polyclonal serum. We conclude that, in this system, L1 production peaks after six days.

Transfiently Expressed L1 Assembles into VLPs That are Detected by Type-Specific Antibodies

The virus particle is comprised of 72 pentameric capsomers of L1. Several reports have demonstrated that baculovirus expression of L1 from diverse papillomavirus types supports not only L1 production, but also assembly into a higher order structure called a virus-like particle (VLP), which is indistinguishable from native virus in electronmicrograph studies (13). Furthermore, many type-specific epitopes are conditional upon VLP formation for proper conformation and display (3,4). Therefore, it was important to demonstrate that transient expression in SF9 cells supports VLP assembly.

We prepared extracts from SF9 cells...
transiently transfected with either pVL1393:CRPV or pVL1393:11 and evaluated their ability to elicit binding by anti-L1 monoclonal antibodies dependent upon VLP conformation and specific for either CRPV or HPV11 VLPs (Figure 2). Figure 2A shows the results of the ELISA with the CRPV-specific monoclonal antibody CRPV.5A. A strong signal is produced with either purified CRPV VLPs or pVL1393:CRPV extracts. No binding above background level is observed with either purified HPV11 VLPs or pVL1393:HPV11 extracts from transfection.

The results are reversed when the ELISA is performed with HPV11-specific monoclonal antibody H11.F1 (Figure 2B). A strong signal is observed with either purified HPV11 VLPs or pVL1393:HPV11 extracts. No binding above background is observed with either purified CRPV VLPs or pVL1393:CRPV extracts from transfection.

To confirm that binding was VLP dependent, we evaluated binding of monoclonal antibody H11.F1 to a denatured extract (Figure 3). An aliquot of HPV11 SF9 extract was diluted into 0.1 M sodium carbonate, pH 10.5, and incubated at room temperature for 1 h. The extract was coated onto a microplate and allowed to dry. In parallel, untreated pVL1393:HPV11 extract was coated onto a microplate and incubated overnight at 4°C. The presence of L1 was verified with antibody H6.C6. Figure 3 shows that the strong signal elicited by pVL1393:HPV11 extracts is reduced to background level in the alkaline-treated sample. In contrast, alkaline treatment had no effect on recognition by control monoclonal H6.C6.

The results demonstrate that transient expression in Sf9 cells supports production of L1 species that contain type-specific epitopes known to be dependent upon VLP formation for display. We have also successfully expressed HPV6 L1 VLPs in this system (data not shown).

DISCUSSION

In this work, we have confirmed that the level of papillomavirus L1 protein expression obtained in Sf9 cells cotransfected with a pVL1393-based vector and BaculoGold viral DNA is sufficient within 6 days posttransfection to generate analytical levels of self-assembled VLPs. Expression of L1 protein appears to reach about 1–2 µg/100-mm petri dish, easily enough VLP material from a single transfection for epitope expression screening using an ELISA format. We have applied this technique to screen sequence variants produced by site-directed mutagenesis and have succeeded in identifying...
critical residues that comprise a conformationally dependent epitope in human papillomavirus 11 (2).

It had previously been noted that detectable levels of recombinant protein appear within a few days of baculoviral/plasmid DNA cotransfection (6,10,18,24,25). This “transient” expression was used principally to confirm expression of the desired protein. In other work, “transient” expression has been exploited to study baculovirus transcription by linking the promoter to the bacterial reporter gene chloramphenicol transacetylase (CAT) on plasmid-borne vectors (17,21,24,25). However, other applications of the baculovirus expression system, such as the study of viral structural protein self-assembly or various aspects of protein structure/function analysis and engineering, have continued, for the most part, to use infection with plaque-purified virus to produce the desired protein(s).

It is in this context that we have found that the BaculoGold system generates transient expression levels of a viral structural protein high enough to satisfy even the stringent test of self-assembly, a concentration-dependent process. These levels of expressed protein are achieved within 6 days posttransfection, considerably faster than reported for the next quickest baculovirus recombinant viral isolation system, the bacmid system (15). The expression levels are also comparable to the best levels obtainable in mammalian cell transient expression systems. This system is well-suited for inquiries that require only analytical amounts of material that must be properly folded or assembled (2). We believe that it holds promise for investigations into structural parameters on nonviral proteins as well.

The intracellular events that follow DNA cotransfection, recombination between viral and plasmid DNA and expression of plasmid-borne genes, are not well understood. Our results and other published reports that document the gradual kinetics of recombinant protein expression between days 3 and 6 posttransfection are consistent with a hypothesis that both transient expression from transfected plasmid DNA and expression from recombinant viral-borne genes in newly infected cells contribute to the total protein yield. We observed optimal levels 5–6 days posttransfection, when essentially all the cells show signs of infection, and when the concentration of infectious viral particles in the supernatant is over $10^6$/mL (A. Lenny and M. Silberklang, unpublished observations). Because almost all infectious virus produced by the BaculoGold system is recombinant (14), much of the late gene expression (driven by the polyhedrin promoter) observed may be due to infection of
cells by newly released recombinant virus. In this sense, the observed expression is not much different from what would occur with plaque-purified virus. Therefore, the transient system we describe provides a very practical avenue to screen sequence variants with the baculovirus system, and should be applicable to all cases where microgram quantities of expressed protein are sufficient to evaluate results. It additionally offers a key advantage over other transient systems in that further scale-up for production of any particular protein sequence of interest would simply involve expansion by standard methods, from the viral progeny of the transient cotransfection, of a pure recombinant baculoviral stock.

REFERENCES


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