Expression of Simian Retrovirus Type D Serotype 2 Envelope in Insect Cell Using Baculovirus Expression Vector System

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Simian retrovirus type-D (SRV) is a causative agent of simian acquired immunodeficiency syndrome in Asian macaques, and can serve as a viral model in understanding of retrovirus infection because of some similarities to human AIDS pathogenesis. Study of infection and pathogenesis of SRV in macaques could be a strategy of vaccine and antiviral development for preventive and therapeutic purposes. We expressed the SRV-2 envelope gene using baculovirus expression vector system and transfected it to Spodoptera frugiferda insect cell line for SRV-2 recombinant protein production. Analysis using PCR and sequencing technique of recombinant in the passage-3 viral stock indicated the occurrence of recombination between SRV-2 envelope and baculovirus genome. Purification using immobilized metal ion affinity chromatography Ni2+-NTA to recombinant protein could minimize the presence non-specific proteins. The SDS-PAGE analysis showed a specific protein for SRV-2 gp70 envelope. Western blot analysis of this purified protein indicated a specific reaction with anti-SRV-2 antibody positive of Macaca fascicularis serum shown as SRV-2 gp70 envelope band.

Keywords: SRV-2, baculovirus, Sf9 cell, Macaca fascicularis

The simian retrovirus (SRV) is a Betaretrovirus capable of causing an AIDS-like disease in Asian macaques. Of the five simian retrovirus neutralization serotypes identified (SRV-1 to SRV-5), three (SRV-1 to SRV-3) have been molecularly cloned and genomically sequenced (Daniel et al. 1984; Marx et al. 1984; Gardner et al. 1988). Disease caused by the more commonly found SRV-2 infection in macaques is characterized by diarrhea, chronic weight loss, anemia and sometimes retroperitoneal fibromatosis (Moazed and Thouless 1993). SRVs have emerged as significant pathogens in captive macaques following recognition of their etiologic role in outbreaks of immunodeficiency disease at several Regional Primate Research Centers in the US in early 1980 (Daniel et al. 1984; Gardner et al. 1988).

Serological studies in Macaca fascicularis, Macaca nemestrina and Pongo pygmeus in Indonesia showed the presence of antibodies to SRV-2, leading to the assumption that these animals have been infected with SRV or similar agent (Iskandriati et al. 1998a; Iskandriati et al. 1998b; Warren et al. 1998). The high prevalence of disease among wild populations pose some problems for breeders providing a population of SRV-free macaques, since macaques are frequently used in biomedical researches (Lerche and Osborn 2003). Regarding potentially active infection and immune abnormality affected by this virus, SRV-2 is a pathogenic agent that should be eliminated in the Macaca breeding colony (Marx et al. 1984; Lerche et al. 1995; Morton et al. 2008).

In this study, we developed SRV-2 envelope (Env) recombinant protein production method using baculovirus expression vector system (BEVS). This recombinant protein could be used as antigen sources for serological test against antibody anti-SRV-2 or as SRV-2 recombinant protein vaccine candidate. Recombinant baculoviruses are widely used to express heterologous genes in insect cells. The BEVS has some advantages, such as the capacity for large inserts of DNA and high yield of recombinant protein. Protein produced in BEVS is very similar to naturally occurring human proteins in terms of post-translational modifications, biological activity and protein stability (Possee 1997; Joshi et al. 2000). For this reason, BEVS is widely used in academia and industry for expression of a variety of recombinant protein in insect cells (Kost and Condreay 1999).

MATERIALS AND METHODS

Determination of Entry Clone SRV-2 Env. The SRV-2 provirus isolated from peripheral blood mononuclear cells (PBMCs) of Indonesian M. fascicularis (provided by PSSP LPPM IPB) was PCR amplified using specific primers SRV-2 BacNTerm 5823U to SRV-2LTR90L that produced the blunt end PCR product. This PCR product was cloned to pENTR/D/TOPO vector (Invitrogen), which facilitated the entry into baculovirus genome. The plasmid pENTR/D/TOPO was then analyzed using sequencing methods with specific primers (SRV Mf-Mn 5737-LTR: 216U20, 727U20, 1284U20 and 1770U20).

Spodoptera frugiferda (Sf9) Insect Cell Preparation. Sf9 cells were seeded into a six-well plate (8x10⁵ cells/well) in 2 mL of complete Grace’s insect medium (GIBCO) supplemented with 10% FBS (fetal bovine serum), 100U/mL Penicillin, and 100 µg mL⁻¹ Streptomycin. The cells were incubated at 27°C for one hour to allow the cells fully attach to the bottom of the plate and then were verified by inspecting them under an inverted microscope.

LR Recombination. The LR recombination reaction was prepared by adding pENTR/D/TOPO plasmid as entry clone (100-300 ng), BaculoDirect linear DNA (300 ng), 5x LR clonase reaction buffer, TE buffer and LR clonase enzyme mix. The mixture was incubated at 25 °C for one hour, then added with

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2 mL of protease K solution and incubated for 10 min at 37 °C. LR recombination reaction and cellfectin (Invitrogen) reagent were combined in 800 mL Grace’s insect medium unsupplemented to make the transfection mix. The medium in the cells was removed and added to the entire transfection mix dropwise onto the cells, then incubated at 27 °C for 5 h. The transfection mixture was later on removed and added with 2 mL of complete growth medium with antibiotics and 100 μM ganciclovir to each well. The plate was incubated at 27 °C for 96 h and visual inspection of the cells was daily conducted to observe signs of the infections using inverted microscope. Once the transfected cells demonstrated signs of infection, the medium was collected from each well and transferred to a sterile snap cap tube. This was used as the P1 viral stock, kept at 4 °C and protected from light. The recombination reaction was performed according to standard procedure by Invitrogen.

High-Titer Viral Stock Preparation. The Sf9 cells at density 8x10⁵ cells per well was seeded in 2 mL of complete growth medium with 100 mM ganciclovir in a six-well tissue culture plate. The cells were incubated at 27 °C for 72 h, then the medium was collected and centrifuged at 1000 x g (Beckman GS-6R) for 5 minutes. The supernatant is P2 viral stock and kept at 4 °C. To generate the high titer viral stock P3, the Sf9 cells was seeded at density 8x10⁵ cells mL⁻¹ in 20 mL growth medium in T75 flask. Cells was incubated in 27 °C for 48 h then infected with 0.5 mL P2 viral stock and incubation was continued for 72 h. The supernatant was collected as P3 viral stock.

PCR and Sequencing Analysis of P3 Viral Stock DNA Recombinant. DNA was isolated both of supernatant and cells P3 viral stock using QiaAmp DNA Minikit procedure (QIAGEN). DNA recombination between BaculoDirect N-Term and SRV-2 Env was amplified using polyhedral forward primer and SRV-2 Env 5974L and 6243L. PCR product was purified using Qiagen Gel Extraction kit (Qiagen, USA) and cloned into pCR 2.1 TOPO 10 (Invitrogen, USA). DNA was isolated both of supernatant and cells P3 viral stock using QiaAmp DNA Minikit procedure (QIAGEN). DNA recombination between BaculoDirect N-Term and SRV-2 Env was amplified using polyhedral forward primer and SRV-2 Env 5974L and 6243L. PCR product was purified using Qiagen Gel Extraction kit (Qiagen, USA) and cloned into pCR 2.1 TOPO 10 (Invitrogen, USA). Nucleotide sequence was carried out on automatic DNA sequencing (ABI, USA) and alignment of the obtained sequences was performed with computer software BLAST program (NCBI).

Recombinant Protein Purification. Sf9 cells were grown in T225 flask to a density of 8 x 10⁵ cells mL⁻¹ and infected with P3 viral stock. Media were harvested at 72 h post-infection and centrifuged at 528 x g for 10 min at 4 °C (Beckman GS-6R). The collected supernatant were concentrated by tangential flow filtration with Pellicon XL device (Millipore) then centrifuged at 191 000 x g for 3 h at 4 °C (Beckman XL90 Optima, fixed angel Ti90). The pellet were diluted in 1 mL PBS and loaded onto a 2 mL Probond chelating column (Invitrogen) charged with nicel-nitriloacetic acid (Ni²⁺-NTA) and equilibrated with native binding buffer. After washing the column with 8 mL native washing buffer supplemented with 20 mM imidazole, recombinant protein were eluted with elution buffer containing 250 mM imidazole. Fractions were collected in 2 mL and the protein concentrations were analyzed by bicinchronic assay kit (Pierce). The Ni²⁺-NTA purification procedure was referred to Probond purification system from Invitrogen.

SDS-PAGE and Western Blot Analysis. The purified SRV-2 Env recombinant protein was detected by electrophoresis on a ready gel 4-15% gradient Tris-HCl SDS-PAGE (Biorad) and then either stained in a gel with Coomassie blue or transferred onto a nitrocellulose membrane. The membrane was rinsed with PBST 0.1%, blocked with BLOTTO (5% skim milk in PBST 0.1%) for 2 h at room temperature and cut into strips. Individual strips were incubated with M. fascicularis plasma/serum containing primary antibody anti-SRV-2 diluted in BLOTTO for overnight. After being washed three times with PBST 0.1%, the membrane was incubated with anti-human IgG alkaline phosphatase conjugate (1:5000 dilution in BLOTTO). The membrane was washed and the bands were developed by reaction with 5-bromo-4-chloro-3-indolyl phosphate and nitro blue tetrazolium (BCIP-NBT) as substrate.

RESULT

The SRV-2 Env provirus was isolated from PBMCs of Indonesian M. fascicularis using specific primer SRV-2 BacNTerm 5823U (upper primer) and SRV-2 LTR90L (lower primer) with Pfu polymerase enzyme, resulting in blunt PCR product about 2000 bp (Fig 1).

In order to facilitate the insertion of the gene of interest SRV-2 Env to baculovirus vector, we used pENTR/D/TOPO cloning kit (Invitrogen) as entry clone. This entry clone utilized a highly efficient method to clone a blunt end PCR product directly into a baculovirus vector with no additional ligation or restriction enzyme required. In this system, PCR products are directionally cloned by adding four bases to the forward primer (CACC). The overhang in the cloning vector (GTGG) invades the 5’end of the PCR product, anneals to the added bases and stabilized the PCR product in the correct orientation (Fig 2).

Sequencing to the pENTR/D/TOPO plasmid was done using M13 forward primer and some specific SRV-2 Env primers (216U20, 727U20, 1284U20, 1770U20). Analysis of the sequencing result with nucleotide alignment using BLAST program indicated the presence of the gene of interest SRV-2 Env in pENTR/D/TOPO plasmid with proper orientation (data not shown). This plasmid was then recombined to

![Fig 1 Amplification of SRV-2 Env provirus isolated from Indonesian Macaca fascicularis using specific primer BacNTerm 5823U-LTR90L. 1, 1 kbp DNA ladder (Invitrogen); 2-3, PBMCs cell; 4, Reagent control.](image-url)
baculovirus genome using LR recombination reaction with LR clonase enzyme mix. In this case, we used linear BaculoDirect Baculovirus Expression System (Fig 3) containing the entry clone to transfect onto the Sf9 cells using cellfectin cationic-lipid, N,N,I,NII,NIII-tetramethyl-N,NII,NIII-tetrapalmitylspermine (TM-TPS) with dioleoyl phosphatidylethanolamine (DOPE).

The SRV-2 Env gene recombinant was expressed using polyhedrin promoter and proliferated in Sf9 cells. Negative selection of the recombinant with correct expression was done using ganciclovir. Infection of the Sf9 cells typically displayed the specific characteristic of cell morphology as observed from visual inspection using inverted phase microscope. Characteristics of infected cells were shown with increased cell diameter, cessation of cell growth, detachment and cell lysis.

In order to verify the recombination between SRV-2 Env with baculovirus genome in Sf9 cells, we amplified the passage-3 (P3) viral stock using PCR technique with specific primer of polyhedrin forward to SRV-2 5974L and SRV-2 6243L. The specific bands about 400 bp and 600 bp was indicated the presence of recombination between SRV-2 Env and baculovirus genome (Fig 4). Additionally, to ensure the recombination of SRV-2 Env with baculovirus genome, the PCR product then cloned to pCR 2.1 TA TOPO cloning kits (Invitrogen), and the plasmid was then sequenced using M13 forward primer. Analysis of the sequencing result indicated the presence of recombination between SRV-2 Env and baculovirus genome in correct direction (data not shown).

The SRV-2 Env recombinant protein expression was demonstrated by specific bands for envelope glycoprotein (gp70) and gp20 on SDS-PAGE analysis, although there were some non specific bands (Fig 5). The purification to the viral stock using affinity chromatography purification system containing metal chelating resin Ni$^{2+}$-NTA was minimizing the presence of non-specific protein, specifically designed to purify 6xHis-tagged protein. It was shown by SDS-PAGE analysis with the specific band of SRV-2 gp70 Env (Fig 6). Western blot analysis of this purified protein indicated a specific reaction with anti-SRV-2 antibody positive of M. fascicularis serum shown by SRV-2 gp70 Env band (Fig 7).

**DISCUSSION**

The gene of interest to be expressed using baculovirus vector is SRV-2 Env isolated from Indonesian M. fascicularis. This gene will be expressed to surface glycoprotein and transmembrane that will first be recognized by host receptor in the initial infection (Brody et al. 1994; Rasko et al. 1999). This SRV-2 Env protein is very stable suggesting high degree of adaptation of SRV-2 to its host (Staheli et al. 2006).

We used pENTR Directional TOPO cloning kit (Invitrogen) that facilitated the recombination of SRV-2 Env gene to
baculovirus genome. This is a universal cloning method using the site-specific recombination properties of bacteriophage lambda (Landy 1989). BaculoDirect has evolved from gateway technology (Invitrogen, USA) platform and enables the direct transfer of the gene into a baculovirus genome without the need for the propagation of the recombinant bacmid DNA. Briefly, attR1 and attR2 gateway sites have been introduced into the viral genome to enable the recombinatorial cloning of gene interest from the gateway entry clone directly into baculovirus DNA. This feature make BaculoDirect system ideal for protein expression system, since it immediately removes one of the more-time consuming stages of the entire process (Hunt 2005).

The specially engineered BaculoDirect linear DNA (Fig 3) contains a strong polyhedrin promoter (P_H) for high level protein expression. Att recombination sites for efficient recombination with any attL-flanked gateway entry vector and herpes simplex virus thymidine kinase (HSV-1 TK) and LacZ genes located between attR sites for the selection of recombinant baculovirus. During the LR reaction with gateway entry clone, the LacZ and TK genes are recombined out as by-products. Sf9 cells are then placed under selection with ganciclovir [9-(1,3-dihydroxy-2-propoxymethyl) guanine], which is enzymatically phosphorylated by HSV-1 TK. Once phosphorylating, the active analog incorporates into DNA and inhibits DNA replication. Ganciclovir selection has been used in Sf9 cell to purify the recombinant viruses, therefore eliminating any remaining parental non-recombinant virus (Hunt 2005).

We observed the SRV-2 Env expression both in pellet and in supernatant cells indicated by specific bands of gp70 and gp20 after SDS-PAGE analysis (Fig 5). It means that the recombinant protein has already been released to the cell supernatant. In this study, we used the full-length gene of SRV-2 Env, and the presence of gp20 band indicated that the SRV-2 Env protein has been cleaved by cellular protease enzyme, although the band was thinner compared to whole SRV-2 antigen (Brody et al. 1994). It was assumed that to produce both gp70 and gp20 recombinant protein the use of full-length SRV-2 Env protein was not efficient. The expression level of full-length Env protein was lower compared to truncated Env protein (Yao et al. 2000).

**Fig 5** Expression of SRV-2 Env recombinant protein using SDS-PAGE technique. 1, broad low marker protein standar; 2, whole SRV-2 Antigen; 3, supernatant cell P3 viral stock; 4, pellet cell P3 viral stock.

**Fig 6** Ni$_2^+$-NTA purification analysis of SRV-2 Env recombinant protein in P3 viral stock by SDS-PAGE staining with Coomassie brilliant blue. 1, protein standard broad low marker; 2, whole SRV-2 antigen virus; 3, elution buffer fraction; 4, washing buffer fraction; 5, binding buffer fraction; 6, recombinant protein pre-purification.

**Fig 7** Western blot analysis of Ni$_2^+$-NTA purified SRV-2 Env recombinant protein probed with positive antibody anti-SRV-2 of _M. fascicularis_ serum (a) and negative antibody anti-SRV-2 of _M. fascicularis_ serum (b). 1, 10, kaleidoscope protein standard; 2, 9, SRV-2 antigen virus; 3, 8, binding buffer fraction; 4, 7, washing buffer fraction; 5, 6, elution buffer fraction.
The recombinant protein was purified using immobilized metal ion affinity chromatography (IMAC) containing metal chelating resin Ni\(^{2+}\)-NTA to bind the 6x His-tagged protein. This His-tagged protein will compete with imidazole and will be removed from the column when the concentration was increased to 250 mM in elution buffer. This purification system has already removed some non-specific proteins, shown by elution buffer fraction in SDS-PAGE analysis (Fig 6). We could not observe the gp20 Env protein, probably it was also removed during the early elution process due to its low level protein expression.

In this study, we observed the antigenicity of this purified SRV-2 Env recombinant protein on western blot analysis using antibody anti-SRV-2 positive of M. fascicularis serum. It was shown by a specific band of SRV-2 gp70 Env (Fig 7). Thereby, this gp70 Env recombinant protein retained its antigenicity and could be used as alternative antigen source for serological test in ELA (enzyme immunosorbant assay) or western blot analysis against antibody SRV-2 and could be applied for routine diagnostic purposes in supporting the Macaca Specific Pathogen Free (SPF) colony program. In the future, the development of SRV-2 recombinant protein will substitute the use of SRV-2 active viral protein as antigen source due to its pathogenicity and biosafety level requirement.

REFERENCES


