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Use of the *piggyBac* transposon to create HIV-1 *gag* transgenic insect cell lines for continuous VLP production

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Abstract

Background: Insect baculovirus-produced Human immunodeficiency virus type 1 (HIV-1) Gag virus-like-particles (VLPs) stimulate good humoral and cell-mediated immune responses in animals and are thought to be suitable as a vaccine candidate. Drawbacks to this production system include contamination of VLP preparations with baculovirus and the necessity for routine maintenance of infectious baculovirus stock. We used *piggyBac* transposition as a novel method to create transgenic insect cell lines for continuous VLP production as an alternative to the baculovirus system.

Results: Transgenic cell lines maintained stable *gag* transgene integration and expression up to 100 cell passages, and although the level of VLPs produced was low compared to baculovirus-produced VLPs, they appeared similar in size and morphology to baculovirus-expressed VLPs. In a murine immunogenicity study, whereas baculovirus-produced VLPs elicited good CD4 immune responses in mice when used to boost a prime with a DNA vaccine, no boost response was elicited by transgenically produced VLPs.

Conclusion: Transgenic insect cells are stable and can produce HIV Pr55 Gag VLPs for over 100 passages: this novel result may simplify strategies aimed at making protein subunit vaccines for HIV. Immunogenicity of the Gag VLPs in mice was less than that of baculovirus-produced VLPs, which may be due to lack of baculovirus glycoprotein incorporation in the transgenic cell VLPs. Improved yield and immunogenicity of transgenic cell-produced VLPs may be achieved with the addition of further genetic elements into the *piggyBac* integron.

Background

Human immunodeficiency virus type 1 (HIV-1) is responsible for the current infection of over 20 million people and the death of over 2 million living in sub-Saharan Africa [1]. Subtype C infections predominate in southern Africa and represent a large portion of the world wide infections [1], highlighting the need to develop a safe and effective vaccine based on Subtype C.

The HIV-1 precursor structural protein Pr55 Gag has been targeted as a potential candidate in vaccine studies as it is able to self-assemble and bud from a variety of cell systems to form non-replicating and non-infectious virus-like particles (VLPs) with good humoral and cell-mediated

immune responses in animals. To date, Gag VLPs have been generated using various eukaryotic expression systems, but most often via the baculovirus-based transient protein expression system in insect cell cultures [2,3]. We have shown that baculovirus-derived HIV-1 Pr55 subtype C VLPs are able to elicit strong cellular immune responses in mice and baboons when administered as a boost to a HIV-1 *gag* DNA vaccine prime [4,5].

However, there are significant potential drawbacks to use of the baculovirus expression system: these include the necessity for constant maintenance of baculovirus stocks, the need for fresh batch infections to be made each time the product is required, and co-purification of recombinant baculovirus or baculovirus proteins with VLPs. The creation of a transgenic cell line for continuous culture and protein production may provide a way to bypass production issues arising with the use of

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baculovirus and to overcome potential safety issues with baculovirus particle contamination of VLP preparations.

The only reported attempts to transform *Spodoptera frugiperda* insect cells in culture include the random integration of an entire expression plasmid into the insect genome through recombination under antibiotic selection [6-11]. Transposon mutagenesis is an ideal alternative as it is based on a naturally occurring system in insect cells and has been extensively used to transpose many insect species [12,13]. The *piggyBac* transposable element has been widely studied and is favoured as a useful tool in insect transgenesis due to its simplicity of movement and often high frequency of transformation [14]. This class II element is derived from the cabbage looper moth *Trichoplusia ni* and is a member of the TTAA-target site-specific class of transposable elements [15]. It exclusively targets TTAA sites and duplicates this site upon insertion. The element is 2476 bp in length and encodes a single open reading frame (1.8 kbp) and terminates with 13 bp inverted terminal repeats (ITR). The ORF encodes a putative transposase (molecular weight of 64 kDa) which is responsible for the movement of the element [13]. Transgene integration into an insect genome is made possible by replacing the transposase ORF in the *piggyBac* vector with the transgene, while supplying the transposase *in trans* [13,15].

A number of whole insects from species spanning three orders [13,16] as well as non-insect species ranging from planarian to mammalian cells [17-20] have been transformed using the *piggyBac* vector system. This wide range of utility for this element makes it an attractive genetic tool. *piggyBac* transposons are favored over other elements as they are able to transpose large DNA fragments (9.6-14 kb) [20] making them suitable for applications in dual expression vectors designed to include selection markers, transcriptional activators or immune enhancer elements. Many pest species do not have transposons closely related to this element and the chance of re-transposition has been shown to be very low in several insect species studied [18,21-23].

No studies involving *piggyBac* transposon-mediated mutagenesis of cultured *Spodoptera* insect cell lines have been reported to date. Here we report the creation of transgenic *Spodoptera frugiperda* cell lines using the *piggyBac* system to express HIV-1 Gag protein, with the aim of developing a system for continuous production of HIV-1 Gag VLPs for vaccine studies. The immunogenicity of these transgenically expressed VLPs is compared to that of baculovirus produced VLPs in BALB/c mice.

Methods

Cell lines

The *Spodoptera frugiperda*-derived Sf21 cell line (Invitrogen) was maintained as a monolayer at 27°C in TC-

100 insect medium (Sigma) supplemented with 10% (v/v) foetal bovine serum (FBS), 50 µg/ml neomycin, 69.2 µg/ml penicillin G and 100 µg/ml streptomycin. The *Spodoptera frugiperda*-derived Sf9 cell line (Invitrogen) was maintained as a shaking culture at 27°C, 140 rpm in SF900 II SFM media (Gibco) supplemented with 10 µg/ml gentamicin. The High 5™ cell line is derived from *Trichoplusia ni* (Invitrogen) and was maintained as a shaking culture in Express Five® SFM media (Gibco) supplemented with 10 µg/ml gentamicin and 18 mM glutamine.

Baculovirus produced Gag VLPs

The human codon-optimised HIV-1 subtype C *gag* DNA sequence used in this study was derived from the DNA vaccine plasmid pTHgagC [24,25]. The *gag* sequence used encodes a myristoylation signal responsible for directing the myristoylated Pr55 Gag protein to the host cell membrane where it embeds, aggregates and buds off as VLPs which are “coated” in host cell outer membrane [26,27]. Baculovirus produced Gag VLPs (BV) were generated in Sf9 cells using the Bacto-Bac® Baculovirus Expression Vector System [4,27,28]. Briefly, 1×10^6 Sf9 cells/ml were infected with baculovirus encoding human codon optimised HIV-1 subtype C *gag* under control of the *polh* promoter at a multiplicity of infection (MOI) of 2-10, and VLPs were harvested from the culture supernatant 72 h post infection.

piggyBac plasmid constructs

The *piggyBac* minimal construct, pXLBacII has been previously described [14,15]. pXLBacII contains a multiple cloning site for insertion of a transgene between the 235 bp 3' ITR (Inverse Terminal Repeat) and 310 bp 5' ITR, but lacks the open reading frame encoding the *piggyBac* transposase which is necessary for transposition to occur. Instead, the *piggyBac* transposase open reading frame is encoded on a separate helper plasmid. We used as helper plasmids; pCasper-hs-orf (pCasper), in which the transposase gene is controlled by the *hsp70* promoter, and pBSII-IE1-orf (pBSII-IE1), in which the transposase gene is controlled by the baculovirus immediate early (*ie1*) promoter. These plasmids have been described previously [15]. A transactivator plasmid (hr3IE1) to be included as a third plasmid in transfection, was supplied by K.J. Maragathavally [29,30]. Constructs harboring different regulatory elements were obtained from H. Zieler (see table 1) [31,32]. Constructs containing a 133 bp intron placed upstream or downstream to the *gag* gene (409-FOR, 410-FOR, 411-FOR, 412-FOR), were included for comparison of Gag expression levels with and without the inclusion of an intron [32].

Table 1 List of different regulatory elements used to design a set of *gag piggyBac* vector constructs

Construct	Enhancer	Promoter	Intron upstream	Gene	Intron downstream	PolyA	Final pXLBacII construct
pHSP70		<i>Drosophila hsp70</i>		<i>gag</i>		SV40sti/polyA	pXLHSP70Gag
pIE1	<i>hr5</i>	Baculovirus AcMNPV <i>ie1</i>		<i>gag</i>		(<i>Heliothis</i>) hel2polyA	pXLEI1Gag
pIE1	<i>hr5</i>	Baculovirus AcMNPV <i>ie1</i>		Neo		hel2polyA	pXLNeo
pIE1-SV	<i>hr5</i>	Baculovirus AcMNPV <i>ie1</i>		<i>gag</i>		SV40sti/polyA	pXLEI-SVGag
pActin		<i>Drosophila actin 5C</i>		<i>gag</i>		hel2-polyA	pXLActinGag
pActin-SV		<i>Drosophila actin 5C</i>		<i>gag</i>		SV40sti/polyA	pXLActin-SVGag
409-FOR	<i>hr5</i>	Baculovirus AcMNPV <i>ie1</i>	+	<i>gag</i>		hel2polyA	pXL409Gag
410-FOR	<i>hr5</i>	Baculovirus AcMNPV <i>ie1</i>		<i>gag</i>	+	hel2polyA	pXL410Gag
411-FOR		<i>Drosophila actin 5C</i>	+	<i>gag</i>		hel2-polyA	pXL411Gag
412-FOR		<i>Drosophila actin 5C</i>		<i>gag</i>	+	hel2-polyA	pXL412Gag
Hr3ieLuc	<i>hr3</i>	Baculovirus AcMNPV <i>ie1</i>		<i>gag</i>			pXIHr3ieGag
pBSII-IE1-orf		Baculovirus AcMNPV <i>ie1</i>		<i>gag</i>		pBSII-IE1-orf polyA	pXLSII-IE1Gag

The promoters tested included *Drosophila hsp70*, AcMNPV *ie1* and *Drosophila actin 5C* [30,31] (column 3). The promoter enhancers used were *hr5* [31] and *hr3* [30] (column 2). Poly-adenylation (polyA) sequences included SV40sti/polyA, hel2polyA [31] and a polyA tail isolated from the pBSII-IE1-orf helper plasmid [14,15] (column 7). 409-FOR and 410-FOR are pIE1 vectors containing a 133 bp intron [32] (column 4 and 6) upstream or downstream of the *gag* gene, respectively. 411-FOR and 412-FOR are pActin vectors containing the intron upstream or downstream of the *gag* gene, respectively. Briefly, different promoters, enhancers, polyA sequences or the intron were removed from the construct described in column 1 and coupled to *gag* in the transposable pXLBacII vector [14,15] (column 8).

***gag piggyBac* construct design**

The human codon-optimised, myristoylated HIV-1 subtype C *gag* gene was coupled to different regulatory elements and cloned into the *gag piggyBac* minimal construct pXLBacII, to be used in transfections (see table 1).

A digenic construct was designed to harbor the neomycin gene coupled to the *gag* gene (*NeoGag*) to enable antibiotic selection of transgenic cells. This construct was created by cloning the *ie1-gag-polyA* fragment from the pXLSII-IE1Gag construct, downstream of the *neo* gene in pXLNeo (see table 1).

Transfection of *Spodoptera frugiperda* cells

Each of the pXLBacII constructs containing the *gag* or *NeoGag* gene, together with one of the helper plasmids, was co-transfected into Sf21 cells with Cellfectin (Invitrogen) for 5 hours. DNA amounts of 1 to 5 µg were used to transfect 1.5×10^6 Sf21 cells per 2 ml, and the ratios of the transgene to helper plasmids were varied from 1:1, 7:1 and 1:2 [33,34]. Two different helper plasmids, pCasper and pBSII-IE1, were tested for differences in transgene integration as assessed by subsequent Gag expression levels. The IE1 transactivator (*hr3IE1*) was included during transfection at a tenth of the total DNA amount to determine its potential to improve promoter activity or transposition frequency [29,30]. A comparison of transformability and transgenic Gag expression levels was conducted between three different insect cell lines, Sf21, Sf9 and High 5[™].

PCR detection of the transgene and determination of the integration event

DNA was extracted from $\pm 5 \times 10^6$ transgenic insect cells using the Dneasy Tissue Extraction kit (Qiagen)

and subjected to PCR screening. Primers were designed to amplify the transgenes (*gag* or neomycin) to assess their presence and stability over several passages (see table 2). To determine whether the entire *gag piggyBac* vector or only the transgene integrated into the insect cell genome, primers were designed to amplify different sites on the pXLBacII *NeoGag* vector. See table 2 for the primer sequences and Figure 1 for their respective positions on the vector. PCR screening was carried out on a number of transgenic insect cell lines expressing Gag under the control of various regulatory elements (table 3).

Determination of whether Gag mRNA is transcribed from an integrated transgene or from a persisting non-integrated *gag piggyBac* vector plasmid

To confirm that the pXLBacII vector is absent from the transgenic insect cell culture, and hence not contributing to Gag expression, rolling circle amplification (RCA, GE Healthcare, Amersham) was carried out on DNA isolated from harvested cells. RCA is based on a bacteriophage φ29 DNA polymerase that exponentially amplifies single- or double-stranded circular DNA templates by rolling circle amplification. To confirm the absence of inhibitory elements, the isolated DNA sample was spiked with 0.7 ng ($\pm 7 \times 10^7$ copies) to 0.007 ng ($\pm 7 \times 10^5$ copies) of the pXLBacII*NeoGag* vector and amplified according to the manufacturer's protocol (GE Healthcare, Amersham). Amplified products were subsequently digested with MluI (which cuts twice within the *NeoGag* region) and analysed on a 0.8% agarose gel with ethidium bromide staining.

Table 2 Primer sequences used in the screening of transgenic insect cell lines expressing Gag protein

Primer pair	Product
sense CTA AATTGTAAGCGTTAATATTTG antisense GGTAGCGACCCCTCAGCCAATAAGAATTC	Product 1 (3' ITR and 3' end of <i>gag</i>)
sense: GGGTTAATCTAGCTGCATCAGGATC antisense: GAGCGCACGAGGGAGCTTC	Product 2 (5' ITR)
sense: CTGGGATGGGTGGGTTGCTGG antisense:CTTCTGAGAGCTCGATCGATAC	Product 3 (Junction between neomycin and <i>gag</i>)
sense ATGGGTGCTCGGCATCTATCC antisense ATTCTTGCTGAGGGGGTCCG	Gag
sense GCGGTACCATGATTGAACAAG antisense CCGAGCTCTCAGAAGAACTCG	Neomycin
sense CAGTTACCAATGCTTAATCAG antisense GTATGAGTATTCAACATTCCG	Ampicillin

Screening for optimal protein expression using ELISA

To determine which method of transfection and which regulatory variable generated a cell line capable of secreting optimal Gag yields, cell supernatants were screened for the p24 component of Gag using a p24 antigen ELISA kit (Vironostika, BiomeriueX) according to the manufacturer’s instructions.

Selection assay

The *NeoGag* construct, together with the pCasper helper, was transfected into Sf21 cells. Selection using 1 mg/ml geneticin (G418, Sigma) [7,6] began two days

post transfection. Transgenic cells were exposed to geneticin after 40 passages (20 weeks) and examined for viability using an inverted light microscope (Carl Zeiss “Axioskop”). Non-transgenic cells were exposed to the antibiotic and used as a negative control.

VLP isolation

piggyBac (PB)-produced VLPs were harvested from 1.5×10^9 transgenic *NeoGag* insect cells and baculovirus produced (BV) VLPs were harvested from 24×10^7 baculovirus infected cells. Cell culture supernatants were clarified by low speed centrifugation at 1000 g for 10

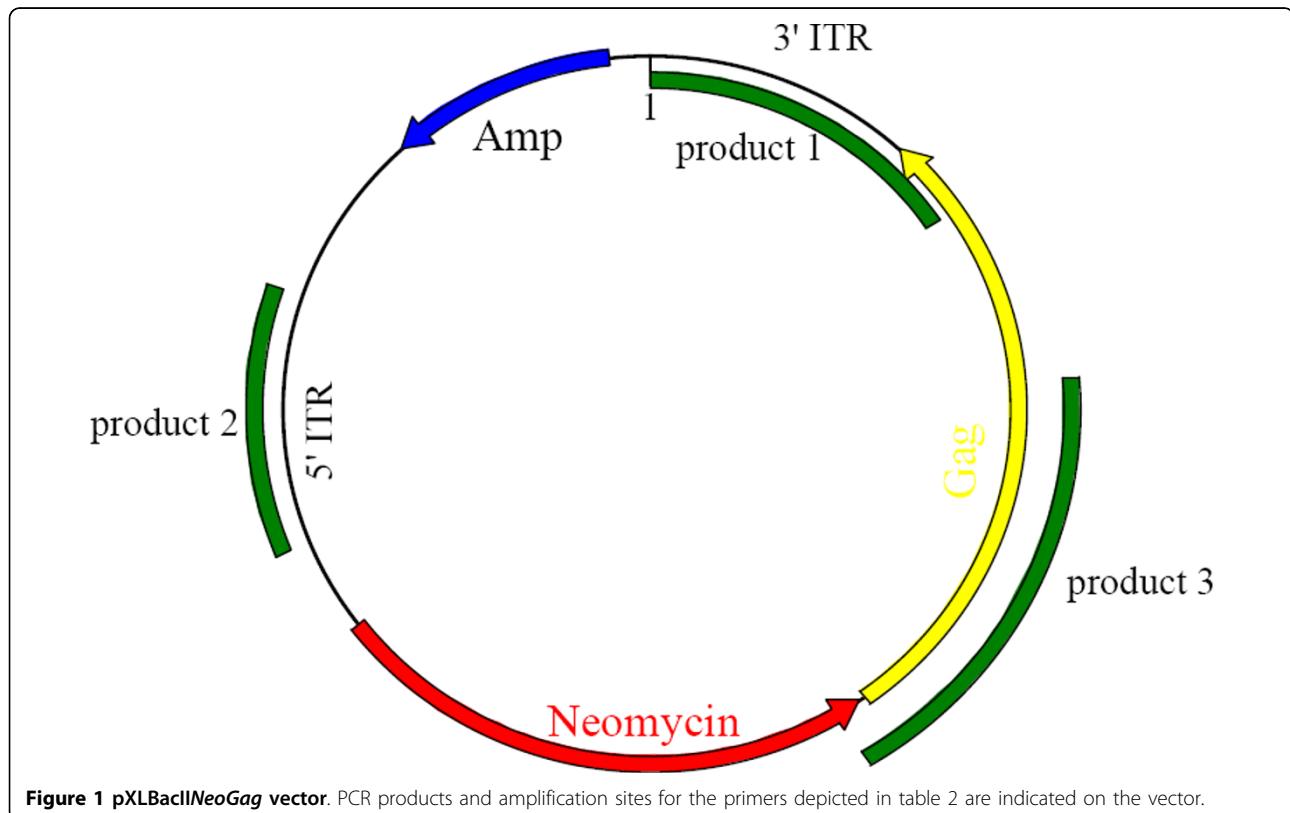


Figure 1 pXLBacII/NeoGag vector. PCR products and amplification sites for the primers depicted in table 2 are indicated on the vector.

Table 3 PCR screening results of transgenic insect cell lines

Construct	Product 1	Product 2	Product 3	<i>gag</i>	Neo	Amp
pXLNeoGag passage 15 in Sf21 cells	+	+	+	+	+	+
pXLNeoGag passage 26 in Sf21 cells	+	+	+	+	+	+
pXLNeoGag passage 23 in Sf9 cells	X	+	+	+	+	+
passage 7				+		
passage 13				+		
passage 93				+		
pXLI-E-SVGag passage 16 in Sf21 cells	X	+	X	+	X	+
pXIHr3ieGag passage 9 in Sf21 cells	X	+	X	+	X	+
pXLHSP70Gag passage 6 in Sf21 cells	X	+	X	+	X	+

Screening was carried out using primers complementary to different sections of the *pXLBacII NeoGag* vector. The presence of a PCR product is indicated with a plus (+) and absence with a cross (X). Confirmation of stable integration up to 93 passages was indicated by performing PCR amplification on several passages of pXLNeoGag in Sf9 cells (see bold font).

minutes. VLPs in these supernatants were pelleted at 4°C by ultracentrifugation in a Beckman SW32ti rotor at 26 000 rpm for 90 minutes. The pellets were then resuspended in phosphate buffered saline (1 × PBS, pH 7.4) and fractionated by centrifugation using a 10-50% Optiprep® (Sigma) step gradient at 26 000 rpm in a Beckman SW32ti rotor for 4 hrs at 4°C. Three light-scattering bands observed at the 10/20%, 20/30% and 30/40% Optiprep® interphases (identical for both *PB* and *BV* VLP preparations) were collected and pooled. VLPs were pelleted, after dilution of the Optiprep® with PBS, at 26 000 rpm for 90 minutes in a SW32ti rotor then resuspended in 1 × PBS. Both *PB* and *BV* VLP preparations were confirmed negative for the presence of endotoxins (QCL-1000® Chromogenic LAL Kit, Cambrex).

Electron microscopy

Gradient purified samples were adsorbed onto carbon coated copper grids, stained with 2% uranyl acetate and visualized with a LEO 912 transmission electron microscope.

Immunoblotting and quantification of *PB* and *BV* VLPs

The banding patterns of purified *PB* and *BV* VLP samples were analysed on Coomassie stained gels and anti-p24 western blots. Aliquots of purified *PB* and *BV* VLPs were fractionated by electrophoresis through a 10% denaturing SDS polyacrylamide gel and either stained with 0.1% Coomassie Blue stain to assess the relative purity of the preparation, or blotted to a nitrocellulose membrane (Nitrobond, Osmonics Inc.) using a semi-dry electroblotter (Hoefler) for 1.5 h at 15 V. Pre-stained molecular weight standards (PageRuler, Fermentas) and a serial dilution of a HIV-1 Pr41 Gag positive control (41 kDa; Quality Biological Inc., USA) was included on the gel. Membranes were probed with 1:10 000 dilution of anti-p24 rabbit antiserum (ARP432, NISBC Centralised Facility for AIDS reagents, MRC, UK) followed by

a 1:5000 dilution of anti-rabbit alkaline phosphatase-conjugated secondary antibody (Sigma). Membranes were developed with Nitro blue tetrazolium chloride/5-bromo-4-chloro-3-indolyl phosphate (NBT/BCIP, Roche).

Gag concentrations were determined by comparing calculated densities of the Pr55 bands in experimental and control samples using gel imaging software (Syngene). VLP concentrations were determined by calculating densities of the Pr55 band on western blots rather than using p24 ELISA quantification, as the Pr55 content is a more reliable indicator of the actual VLP concentration in the sample than the p24 content. VLP preparations were formulated with 15% trehalose to 100 ng Gag/100 µl PBS then stored at -70°C.

Baculovirus gp64 envelope glycoprotein content of *PB* and *BV* VLPs was assessed by western blot analysis as described above using a 1:10 000 dilution of anti-baculovirus envelope gp64 antibody (Clone: AcV5, eBioscience), followed by a 1:5000 dilution of anti-mouse alkaline phosphatase-conjugated secondary antibody (Sigma). Membranes were developed with NBT, as above.

Immunization of mice and detection of cellular immune responses

We compared the immunogenicity in BALB/c mice of *PB* VLPs and *BV* VLPs when given alone, and their ability to boost a response to a prime with the matched HIV-1 subtype C DNA vaccine, pTHgagC [24,25,35]. VLPs (100 ng Gag protein in 100 µl PBS) and DNA (100 µg DNA in 100 µl PBS) were given to groups of female BALB/c mice (5 mice per group) via the intramuscular route with 50 µl injected into each quadriceps muscle. To test responses to VLPs alone, mice were inoculated with a single dose of either *PB* Gag VLPs or *BV* Gag VLPs on day 0, and spleens from VLP-inoculated mice were harvested on day 12. To assess VLP

boosting of a DNA prime, mice were given a single dose of pTHGagC on day 0 and boosted with a single dose of either PB Gag VLPs or BV Gag VLPs given on day 28 and spleens were harvested on day 40. Mice vaccinated with pTHGagC on day 0 without any boosting on day 28 were used as a DNA primed, unboosted control group and their spleens were likewise harvested on day 40. All procedures were carried out according to guidelines and with approval of the UCT Animal Research Ethics Committee. Cell mediated immune responses were determined using splenocytes in Interferon gamma (IFN- γ) and interleukin 2 (IL-2) ELISPOT assays (BD Pharmingen) with Gag CD8 peptide (AMQMLKDTI), and Gag CD4 (13) and Gag CD4 (17) peptides (NPPIPVGRYKRWIILGLNK and FRDYVDRFFKTLRAEQATQE, respectively) as previously described [27].

Results

Transgene expression

p24 ELISA was used to screen the culture supernatants from different transgenic cell lines maintained in culture. All preliminary experiments were carried out in Sf21, Sf9 and High 5[™] cell lines in order to confirm that these lines were amenable to *piggyBac* transformation and subsequent Gag VLP expression. For reasons of practical handling, all further experiments were carried out using only Sf21 cell lines, for small scale amplification, and Sf9 cell lines, for large scale amplification (Sf9 cells being more suitable to this because they were maintained as a shaking culture versus Sf21 cells that were maintained as a monolayer culture). Sf21 cells and Sf9 cells expressed the Gag protein at similar levels for each construct tested. The highest p24 expression level obtained was from cell lines transgenic for the *gag* gene coupled to the *ie1* promoter and the pBSII-IE1 polyA tail (derived from pXLBacSII-IE1Gag). Expression from these cell lines ranged between 100-1000 pg per 1.5×10^6 cells. Previous work suggested that inclusion of an intron adjacent to a gene can stimulate its transcription and possibly enhance protein expression levels [32]. No significant expression variations were observed when the same intron was used in this study. No p24 expression was

evident when the *gag* gene was placed under the *hsp70* or actin 5C promoters. Further investigation into transgenic *hsp70-gag* cell lines confirmed the integration of the construct (see table 3) ruling out the possibility that no transgene integration had occurred.

Inclusion of a third plasmid expressing a transactivator is known to improve protein expression levels by enhancing promoter activity [30] or by increasing transpositional frequency of the element [29]. No significant variation in Gag expression was observed when the IE1 transactivator was included at transfection. Use of different helper plasmids and transfection ratios did not affect the expression either.

Persistence of expression

Transgenic Sf21 and Sf9 cell lines continued to express Gag protein at constant levels (as determined using p24 ELISA) for at least 100 cell passages. Table 3 and 4 indicate the PCR amplification results determined for several transgenic insect cell lines. PCR amplification of the *gag* transgene from insect cell genomic DNA remained positive after 93 cell passages. PCR amplification of the neomycin transgene from insect cell genomic DNA remained positive for 30 cell passages and was not tested further. Exposure of the Sf21 *NeoGag* cells to geneticin at passage 40 had no effect on their survival, which was in contrast to the non-transgenic control cells that lost viability within three to four days. The geneticin was not administered prior to passage 40 indicating that it is not a requirement for stable integration. These results strongly indicate stable integration and expression of the transgene over many cell generations.

Integration of the *piggyBac* vector

To examine the nature of the integration event that had occurred, PCR screening of the *piggyBac* vector and the transgenes was carried out on several transgenic insect cell lines. Table 3 indicates the absence or presence of the PCR products in selected cell lines (refer to table 2). Product 1 is an amplification of the 3' end of the *gag* gene and the 3' *piggyBac* border (3' ITR), product 2 is an amplification of the 5' *piggyBac* border (5' ITR). Product 1 and 2 would be absent if canonical *piggyBac* transposon

Table 4 Continuous expression of p24 protein from several transgenic insect cell lines as determined by ELISA

Cell line and passage number	<i>gag</i> transgene presence (PCR)	Gag expression: pg p24/ml culture supernatant (1.5×10^6 cells)
pXLNeoGag passage 7 in Sf9 cells	+	129.87 pg/ml
pXLNeoGag passage 13 in Sf9 cells	+	129 pg/ml
pXLNeoGag passage 23 in Sf9 cells	+	75.47 pg/ml
pXLNeoGag passage 93 in Sf9 cells	+	606.6 pg/ml

The presence and active transcription of the *gag* transgene in the Sf9 cell line pXLNeoGag, was confirmed over several passages using PCR and p24 ELISA screening, respectively.

mutagenesis had occurred. Product 3 is the 3' end of the neomycin gene and the 5' end of the *gag* gene, providing evidence of transgene integrity. The majority of results indicate that the plasmid was linearised at the 3' ITR prior to integration of the entire intact *piggyBac* plasmid, possibly through a recombination mechanism unrelated to transposition. A study in *Aedes aegypti* using *piggyBac* transformation revealed similar non-canonical integration of sequences from both donor and helper plasmids [22,36]. No evidence of helper plasmid integration was evident in our study as confirmed by PCR screening for the *hsp70* promoter in transgenic cell lines transfected with the pCasper helper plasmid (data not shown). However, these results are inconclusive in predicting whether recombination or transposon insertion occurred, as single insect cell clones were not isolated post transfection and mixed populations may exist.

Gag expression is not due to persisting non-integrated plasmid

Rolling circle amplification (RCA) was carried out on DNA extracted from transgenic Sf21 *NeoGag* cells to eliminate the possibility of transient Gag expression from a persistent non-integrated circular *piggyBac* vector. As seen in Figure 2, lane 3, no circular DNA that resembled the pXLBacII*NeoGag* plasmid (lane 2) was detected in the transgenic DNA sample. The minimum amount of pUC19 DNA that can be amplified using RCA is 1 pg ($\pm 3 \times 10^5$ copies) (RCA instruction manual, GE Healthcare, Amersham). To discount the possibility that non-amplification could be attributed to a low copy of the circularised DNA or the presence of inhibitors in the isolated DNA sample, transgenic insect DNA extract was spiked with 10-fold dilutions of the pXLBacII*NeoGag* plasmid prior to amplification. A minimum

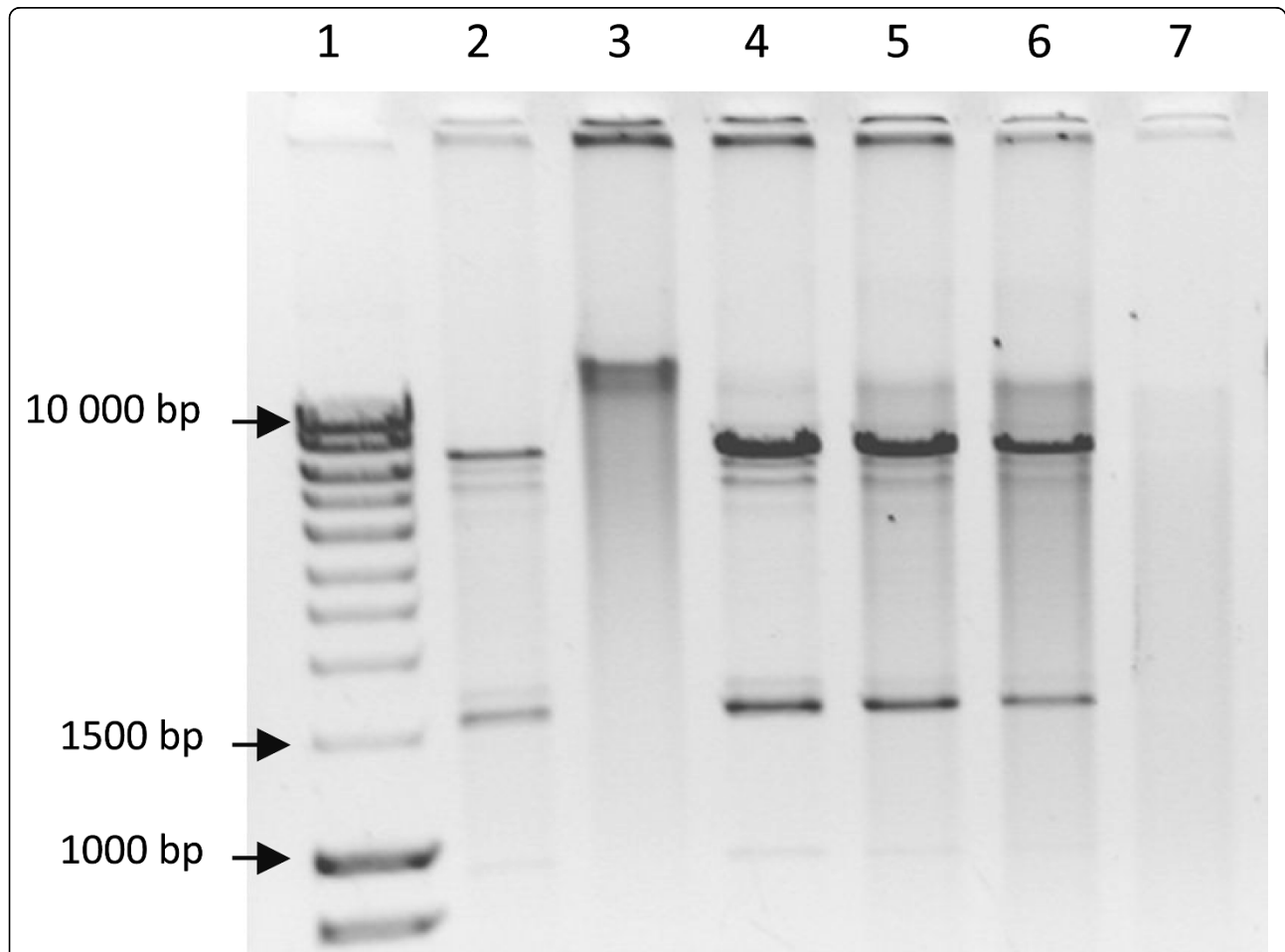


Figure 2 RCA amplification carried out on DNA extracted from *NeoGag* expressing transgenic cells. The RCA product was digested with MluI and visualised on an 0.8% agarose gel. Lane 1- Fermentas Bioline hyperladder 1 with sizes indicated in basepairs. Lane 2- positive control (pXLBacII*NeoGag*). Lane 3- Sf21 *NeoGag* genomic DNA. Lane 4- Sf21 *NeoGag* genomic DNA spiked with 7 ng of the positive control. Lane 5- Sf21 *NeoGag* genomic DNA spiked with 700 pg of positive control. Lane 6- Sf21 *NeoGag* genomic DNA spiked with 70 pg of positive control. Lane 7- Sf21 *NeoGag* genomic DNA spiked with 7 pg of positive control.

of 70 pg ($\pm 7 \times 10^6$ copies) of spiked construct was detectable. No amplification of 7 pg of spiked plasmid ($\pm 7 \times 10^5$ copies) occurred. This copy number approaches the limits of detectability for the RCA assay. Together these results suggest that absence of detectable RCA amplification in transgenic Sf21 *NeoGag* cells is not due to inhibitor presence but is rather due to absence of non-integrated plasmid in the samples.

Size and morphology of *PB* and *BV* VLPs

EM analysis of the three protein bands isolated from the *PB* and *BV* gradients revealed particles of size 120-150 nm in the 10/20% and 20/30% Optiprep® inter-phases. VLP size range and morphology resembled that of VLPs previously reported [3,4,27]. *PB* VLPs and *BV* VLPs were similar in size and shape although *BV* VLPs appeared more defined and compact than the *PB* VLPs (see Figure 3).

Western blot immunodetection and quantification of *PB* and *BV* VLPs

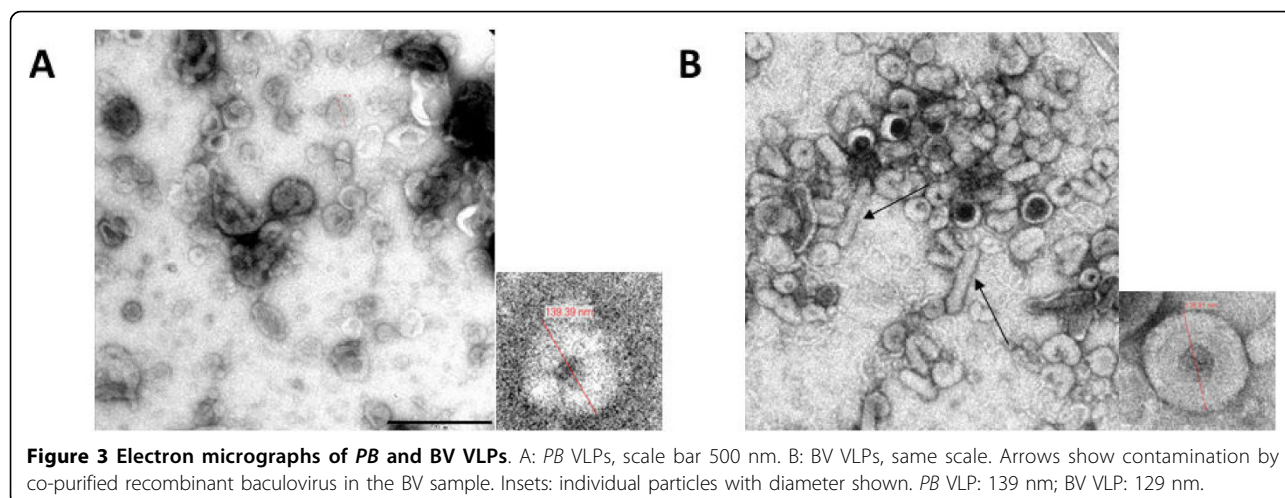
Purified Gag VLPs extracted from the transgenic cell culture and from recombinant baculovirus infected cells were analysed and quantified on an anti-p24 western blot using densitometry. An identical Pr55 Gag band was detected in both *PB* and *BV* VLP preparations (Figure 4A). Quantification of purified *PB* VLPs on western blots against a serial dilution of a HIV-1 Pr41 positive control indicated yields of 1-2 ng per 1×10^6 Sf9 cells, at least 1000 times lower than yields obtained for *BV* expressed VLPs. Anti-gp64 western blot results show the absence of gp64 envelope protein in *PB* VLP samples and its presence in *BV* VLP samples (Figure 4B).

As seen in Figure 5, the Pr55 Gag protein of the *BV* VLP sample was readily detected on a Coomassie stained gel as well as on an anti-p24 western blot. The

Pr55 Gag protein of the *PB* VLP sample was detected at the same protein size location on the anti-p24 western blot, but was not detectable against the background on the Coomassie stained gel, as was expected given the low p24 concentration of the loaded sample).

Immunogenicity of VLPs

It has previously been shown that *BV* Gag VLPs enhance a cellular immune response in mice when used as a boost to a pTHgagC DNA prime [4,27]. To assess whether the *PB* VLPs elicited a similar response in the absence of baculovirus elements, mice were inoculated with equal quantities of either *BV* or *PB* VLPs. Because the study reported here was designed in proof of concept, we conducted only a standard heterologous prime-boost immunogenicity experiment, in which we compared the boost potential of *PB* VLPs versus *BV* VLPs, using the same dosage which our group had previously found optimal when using *BV* VLPs to elicit a boosted immunological response in DNA inoculated mice: that is, 100 ng of HIV-1C VLPs administered as 2×50 ng inoculated into each quadriceps muscle alone or as a boost to DNA primed mouse at day 28 (ms. submitted; S Pillay, A Meyers, EG Shephard, A-L Williamson, EP Rybicki). IFN- γ and IL-2 ELISPOT analysis (Figure 6) showed that *PB* VLPs did not induce an immune response nor did they boost an immune response to a DNA vaccine prime. In contrast the ELISPOT assays detected a low magnitude of GagCD4(13)-specific IFN γ and IL-2 producing cells but no GagCD8-specific cells in response to *BV* VLP vaccination. These *BV* VLPs boosted pTHGagC responses. A 1.8 fold increase in the response to the GagCD8 peptide was detected in the IFN- γ ELISPOT assay. The low cumulative response to the GagCD4(13) and GagCD4(17) peptide of 50 IFN- γ sfu/ 10^6 splenocytes induced by pTHGagC increased 12 fold to 602 sfu/ 10^6 splenocytes when pTHgagC primed



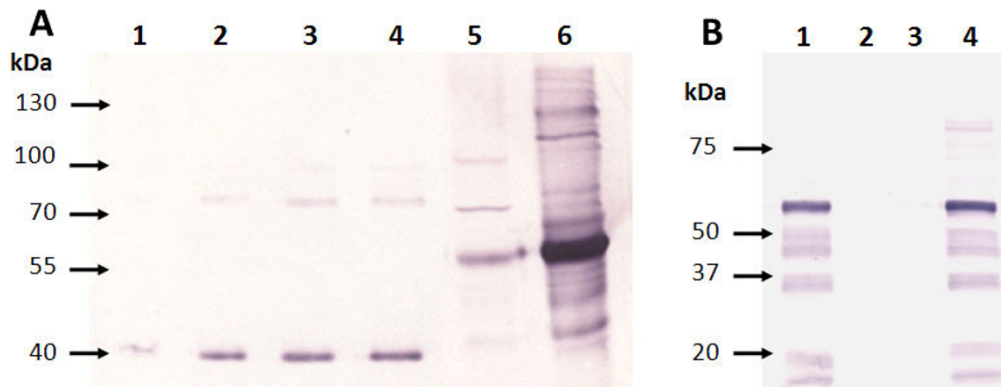


Figure 4 Western blot immunodetection of PB and BV VLPs. A: Anti-p24 western blot. Lane 1-1.2 ng Pr41 std. Lane 2-2.4 ng Pr41 std. Lane 3-4.8 ng Pr41 std. Lane 4-7.2 ng Pr41 std. Lane 5- purified PB VLPs from the 20/30% Optiprep® interphase (22 ug of total soluble protein loaded). Lane 6- purified BV VLPs from the 20/30% Optiprep® interphase (0.68 ug of total soluble protein loaded). B: Anti-gp64 western blot. Lane 1- Supernatant from cells infected with negative baculovirus (no gag insert). Lane 2- purified PB VLPs from the 10/20% Optiprep® interphase. Lane 3- purified PB VLPs from the 20/30% Optiprep® interphase. Lane 4- purified BV VLPs from the 20/30% Optiprep® interphase. The respective sizes from markers run on the same gels are indicated in kDa on the left of both blots.

mice were boosted with these BV VLPs. No boost of the DNA vaccine induced GagCD8 response was detected in IL-2 assays. Although pTHgagC did not induce Gag-specific CD4+ T cells that produced IL-2, a high cumulative magnitude of 390 sfu/10⁶ splenocytes Gag-specific CD4(13) and GagCD4(17) T cells producing IL-2 was detected in response to a DNA prime BV VLP boost.

Discussion

Gag epitopes elicit robust CTL responses that effectively control HIV-1 viral load in the early phases of HIV-1

infection, and Gag is therefore thought to be highly suitable as a vaccine candidate to elicit CTL [3]. HIV-1 Gag VLPs, in particular, are widely accepted as being strongly immunogenic particulate antigens that stimulate good CTL responses in prime/boost vaccination strategies [3-5,27,37]. Baculovirus production of HIV-1 Gag VLPs is a well documented method for generating immunogenic HIV-1 particles, but contamination of VLPs with co-purified baculovirus particles (Figure 3) is not favourable for their subsequent use in vaccine studies. To address this problem and the problem of

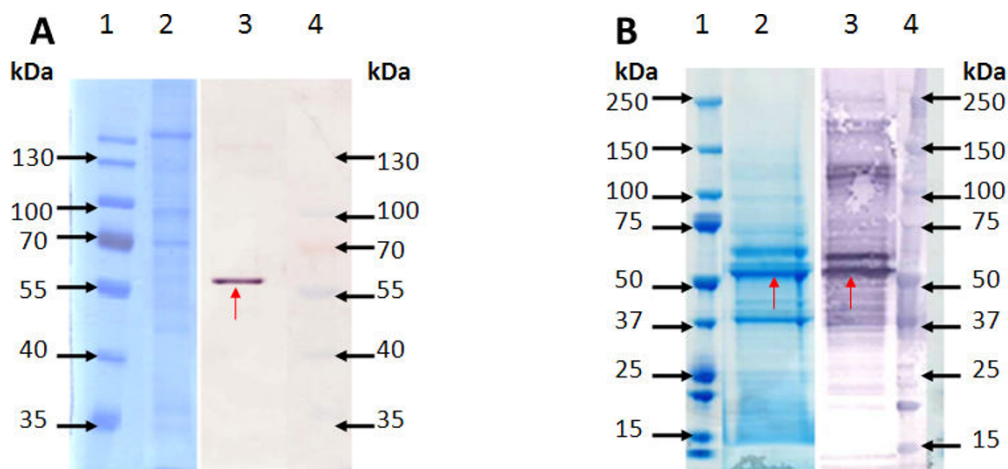
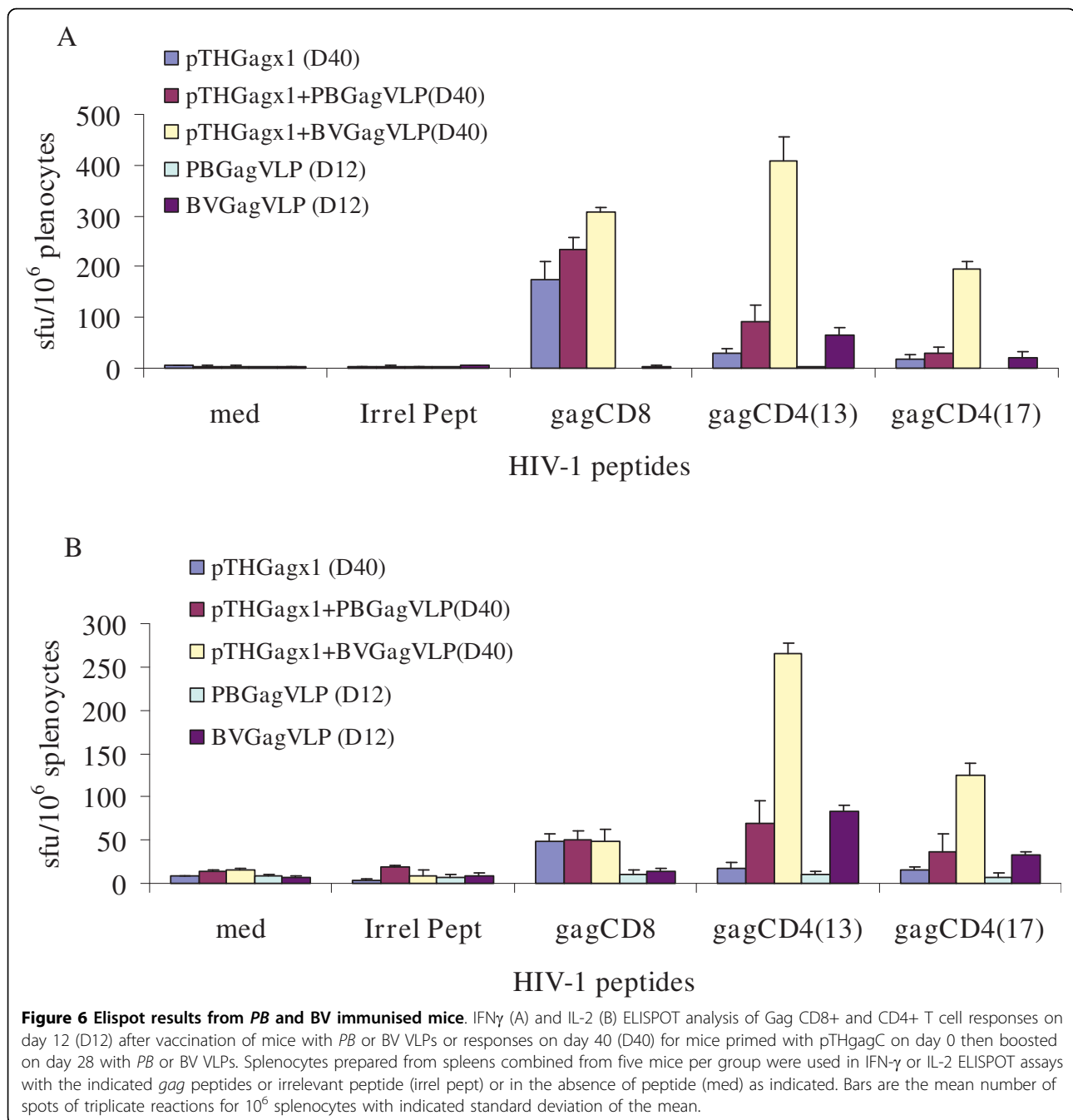


Figure 5 Comparison of PB and BV VLP banding pattern differences as visualised on a Coomassie stained gel and western blot. Coomassie Blue stained 10% denaturing polyacrylamide gels are shown on the left of 5A and 5B and anti-p24 stained western blots are shown on the right of 5A and 5B. Protein size markers on gels and blots are indicated in kDa. Red arrows indicate the Gag Pr55 band. A- Purified PB VLPs. Lanes 1 and 4 - MW marker. Lane 2- Coomassie Blue stained PB VLP preparation (30 ul of purified sample, corresponding to 6 pg p24). Lane 3- PB VLP preparation detected with anti-p24 antibody (30 ul of purified sample, corresponding to 6 pg p24). B- Purified BV VLPs. Lanes 1 and 4 - MW marker. Lane 2- Coomassie Blue stained BV VLP preparation (28 ul of purified sample, corresponding to 170 ng p24). Lane 3- BV VLP detected with anti-p24 antibody (28 ul of purified sample, corresponding to 17 ng p24).



routine maintenance of infectious stocks, we utilised *piggyBac* transposon mutagenesis as a novel method of generating gag-transgenic insect cell lines for continuous production of HIV-1 Gag VLPs. Expression of Gag VLPs from these transgenic cell lines proved stable for at least 100 cell passages: this is a novel result, which may have valuable implications in future HIV and other vaccine work.

Using immunodetection and EM we verified that Gag VLPs were secreted from transgenic insect cell lines.

However, the yields were low (at least 1000 times lower than from the baculovirus production system) and therefore we tested the ability of various regulatory elements to improve protein expression of the gag transgene. No improvement in Gag protein expression was noted when we cloned hr5 and hr3 baculovirus-derived enhancer elements [38] or introns [32] into the *piggyBac* construct, nor when we included the transactivator during transfection [30,39]. Different molar transposase-to-transposon ratios did not affect Gag expression levels,

which also confirms previous observations that the *piggyBac* system does not demonstrate overproduction inhibition [20]. In preliminary experiments, we noted that the *hsp70* or actin 5C promoters were not active in the *Spodoptera* transgenic cell lines, and so no further work was done with these constructs. The *Bombyx mori* actin 3C promoter and *Drosophila* ubiquitin promoter have proven active in a variety of insect species [13] and should be evaluated in *Spodoptera* insect cell culture as an alternative means to possibly improve transgene expression.

Although Gag VLP yields obtained in this preliminary study were low, we are confident that there are several approaches that can be employed to improve protein expression yields. Inclusion of a Gal4 DNA binding domain as an N-terminal fusion to the transposon has been shown to increase the number of transposition events [40] which in turn can result in improved protein expression levels. It has been shown recently that the translation enhancer activity of 5'-UTR *pol* (un-translated region of the nucleopolyhedrovirus polyhedron gene) is able to improve transgene expression when placed upstream to the promoter [41]. Poor transgene expression could be attributed potentially to the integration of the gene into an unfavourable genomic site such as a silent heterochromatin region or near to unfavourable transcription enhancers. This can be overcome by directing the transposon construct to a targeted site using the *Gal4/UAS* or *FLP/FRT* system [40,42,43], or by surrounding the transgene with an insulator [44]. Transposon constructs can be designed to harbour bidirectional promoters that drive simultaneous expression of the transgene and a strong artificial transcriptional activator [45], leading to improved transgene expression levels.

However, it is possible that constitutive production of Gag protein is toxic to the cell, so only low expressers survive. In this case, inducible expression systems could be explored.

We conducted a comparative study in mice to assess the relative immunogenicity of baculovirus-produced VLPs versus *piggyBac* transgenically expressed VLPs. While BV VLPs were able to induce a good CD4 immune response in mice when administered as a boost to a DNA prime, *PB* VLPs, on the other hand, elicited no immune response, showing that Gag VLPs are not intrinsically as highly immunogenic as previously thought, and that baculovirus-derived elements probably enhance Gag VLP immunogenicity. Baculovirus-expressed Gag VLPs include trace amounts of insect cell and baculoviral contaminants (lipids, nucleic acids and proteins) that are not efficiently removed during purification, as well as incorporated baculovirus envelope proteins. Deml *et al* proposed that these

contaminating components act as “danger signals” that can activate an innate immune response [3,26]. It has also been shown that VLPs isolated from yeast [46] as well as from the baculovirus expression system [3] contain host cellular contaminants capable of stimulating human antigen presenting cells (APC) by up-regulating the maturation of cytotoxic T lymphocyte (CTL) markers and inducing cytokine secretion. Although it was beyond the ambit of the current study, it would be interesting to compare the relative ability of *PB* VLPs and BV VLPs to stimulate dendritic cells, as this could provide insight into the observed differences in immunogenicity between *PB* and BV VLPs in the study reported here.

Baculovirus has been shown to transduce mammalian cells, which could lead to adjuvanting of immune responses [47,48]. A recent study showed that intranasal inoculation of mice with a wild-type baculovirus induces a strong innate immune response, which protects mice from a lethal challenge of influenza virus [49]. Cellular uptake of baculovirus and subsequent immune response enhancement may be due largely to the presence of the IFN-stimulatory baculovirus surface envelope glycoprotein gp64, which is responsible for host cell receptor binding and membrane fusion during viral entry by endocytosis [50]. In particular, gp64 protein is known to incorporate into the outer surface of baculovirus expressed Gag VLPs [3,26,47,51]. This has been additionally demonstrated in the current study, where both BV VLPs and *PB* VLPs are “coated” in host cell outer membrane, but gp64 would be available for incorporation only into the BV VLPs (Figure 4). We observed that while *PB* VLPs were similar in size and morphology to BV VLPs, as seen by EM, they did not appear to be as compact or sharply defined in shape as BV VLPs (Figure 3). Since *PB* VLPs lacked baculovirus gp64 incorporation into the VLPs outer membrane coating, the more defined shape of the BV VLPs compared to that of the *PB* VLPs may be accounted for by the incorporation of gp64 into the BV VLP outer membrane. It is likely that incorporation of gp64 onto the surface of baculovirus expressed VLPs facilitates BV VLP uptake into APC by promoting membrane fusion between BV VLPs and host cells, thereby enhancing the resultant immune response to VLP Gag antigens conditioned by APCs. Transgenically produced VLPs lack gp64 and this may result in less efficient cellular uptake of VLPs, with resultant lower immunogenicity.

The use of molecular adjuvants or incorporation of gp64 onto VLP surfaces could be utilised to improve the immunogenicity of these particles expressed from transgenic insect cell lines. Shi *et al.* [33] demonstrated the use of *piggyBac* transposon vectors to transiently express

two gene products. In this system, the two genes were placed under the control of bidirectional promoters which in turn were enhanced by a single enhancer element. Such dual expression systems could be used to co-express immune enhancer elements [52] or immunogenic baculovirus elements together with HIV-1 Gag VLPs to improve VLP immunogenicity.

Once a cell line has been established that transgenically expresses Gag VLPs at a high level and with enhanced immunogenicity features, permanent stabilisation of the transgene in *piggyBac*-transformed insect cell lines would be carried out by transgene integration site elimination. It has been demonstrated that elimination of the *piggyBac* transposon integration sites adjacent to the integrated transgene renders the element immobile to further transposase exposure [43,53].

Conclusion

This study serves as a basis to indicate the potential of a transgenic insect cell expression system as an alternative to the baculovirus-insect cell production system. Stably transformed cells produced VLPs reliably over 100 passages; purification of VLPs was also easier than in the baculovirus system due to lack of heterologous virus particles. However, further work is needed to improve VLP expression levels and their immunogenicity.

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Authors' contributions

AL participated in the design of the study, carried out most of the experimental work and drafted the manuscript. FT conceived the study, participated in the design of the study, and helped in drafting and revising the manuscript. ES performed the immunological experiments described in the study and participated in experimental design. MF provided the *piggyBac* transposons and protocols and helped with the ms. A-LW was overall study director and obtained the funding, and helped in drafting the ms. EPR participated in study design and supervision and helped to revise the manuscript. All authors read and approved the final version.

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