Gene Expression of *Autographa californica* multiple Nucleopolyhedrovirus (*AcMNPV*) in Mammalian Cells and Upregulation of Host $\beta$-actin Gene

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Abstract

The gene expression of Autographa californica multiple nucleopolyhedrovirus (AcMNPV) was examined in two types of mammalian cells, human HeLa14 and hamster BHK cells. DNA microarray analysis followed by RT-PCR identified at least 12 viral genes transcribed in both HeLa14 cells and BHK cells inoculated with AcMNPV. 5’ RACE was carried out to examine the transcriptional fidelity of these genes in HeLa14 cells. The transcription of ie-1, ie-0 and gp64 was initiated at a baculovirus early gene motif, CAGT, accompanied by a TATA motif. In addition, the same splicing observed for ie-0 mRNA in Sf9 cells occurred in HeLa14 cells. While the transcription initiation sites for pe38 and p6.9 were not located in the CAGT motif, most of them were in a typical eukaryotic RNA polymerase II promoter structure (a conventional TATA motif and/or an INR). Interestingly, the expression of β-actin was upregulated in the mammalian cells inoculated with AcMNPV. Subsequent experiments using UV-inactivated virus confirmed the upregulation, suggesting that de novo synthesis of viral products is not required for the event. These results indicated that the AcMNPV genome acts as a template for transcription in mammalian cells through the usual infection pathway, though there is no evidence for the functional expression of viral genes at present.
Introduction

*Autographa californica* multiple nucleopolyhedrovirus (*AcMNPV*), a type member of the insect-specific virus family *Baculoviridae* (genus *Nucleopolyhedrovirus*, NPV), infects several important pest insects and has been used as a safer biopesticide in Integrated Pest Management (IPM) program. The nucleopolyhedroviruses are also used as a gene-transfer vector in insect cells, providing one of the most efficient systems for the expression of foreign genes in eukaryotic cells (19). Furthermore, recent studies demonstrated that *AcMNPV* can enter the nucleus of mammalian cells independently of the cell cycle without multiplication (28, 29) and effectively express foreign genes (15) without expressing its own genes (28), suggesting that the baculovirus can be used as a low risk viral vector in the field of gene therapy (10). Recently, *AcMNPV* was shown to be capable of stimulating production of interferon (IFN) in mammalian cell lines and when injected into mice conferred protection from lethal infections of encephalomyocarditis virus (13). It was also reported that the intranasal inoculation of a wild-type *AcMNPV* induced a strong innate immune response, which protected mice from a lethal challenge of influenza virus, through the Toll-like receptor 9 signaling pathway (1, 3). These reports indicate that baculoviruses affects the physiology of mammalian cells though they are not pathogenic to and have no ability to replicate in mammalian cells.
NPV is a large rod-shape virus with a double-stranded closed circular DNA genome (130~160 kbp) containing approximately 150 genes (4). NPV genes are usually categorized into 4 groups; immediate-early, delayed-early, late, and very-late (8). After infection, the immediate-early and delayed-early genes are transcribed by host RNA polymerase II and their promoters are characterized by the transcription initiation site consensus motif CAGT (25). Most of them are thought to code for the transcriptional transactivator of viral genes (33). Late and very-late genes are transcribed by viral RNA polymerase from the late promoter motif TAAG (22, 24) whose expression is regulated by the early genes. Thus, the senseful expression of early genes is a key to NPV infection. The mechanism of transcription by RNA polymerase II is highly conserved among eukaryotes (14). Indeed, the promoter for immediate-early gene 1 (ie-1) of Bombyx mori NPV (BmNPV) showed the ability to promote transcription in mammalian cells in transfection experiments using reporter plasmids (21). Furthermore, the viral essential regulatory protein encoded by ie-1 (IE-1) is functional in mammalian cells (21, 23). More recently, the acidic activation domains (AADs) of AcMNPV IE-1 were demonstrated to be functional for transcriptional activation in mammalian cells (12). These observations suggest that NPV could express at least part of the viral genomic information in mammalian cells and may affect the physiology of the host cells. Progress in the application of baculoviruses requires more
information about the functions of AcMNPV in mammalian cells. In this study, we reexamined the gene expression of AcMNPV in mammalian cells using a DNA microarray, which suggested transcription from several viral genes and increased transcription from cellular $\beta$-actin.

**Materials and Methods**

**Cells and viral inoculation**

*Spodoptera frugiperda* (Sf) 9 cells were grown in TC-100 medium (Sigma) containing 10% FBS at 26°C. Mammalian cells (HeLa and BHK) were maintained in MEM (Nissui) containing 10% FBS in a CO$_2$ incubator. Treatment (inoculation) of the mammalian cells with AcMNPV was carried out by incubating the cells in DMEM (FBS-free) containing AcMNPV for 2 hrs at 37°C followed by culture in DMEM containing 10% FBS at 37°C.

**Viruses**

AcMNPV was grown in Sf9 cells in TC-100 medium containing 10% FBS at 26°C. We constructed the recombinant AcMNPV, Ac-dhspp-EGFP, using the Bac-to-Bac system (Invitrogen). This virus has an EGFP-coding sequence under the control of the *Drosophila*
heat shock protein 70 promoter (nucleotides (nts) 1 to 1927, GB No., AY032740). Viral titers were determined in Sf9 cells by plaque assay according to Maeda’s method (19). Inactivation of viruses was carried out by exposing the viral suspension (3×10^7 pfu/ml DMEM) in a sterile tissue culture dish to shortwave UV (UVC, 254nm) at a distance of 30 cm for 30 min in a laminar flow hood. The inactivation of infectivity was verified by a plaque assay with Sf9 cells. The recombinant virus carrying luciferase under the control of the ie-1 promoter (rBACie1-luc) (2) was also used to examine the inactivation of viruses and demonstrated that UV-irradiation for 30 min under such conditions resulted in the loss of 99.7% of the activity to express luciferase and a lack of plaque formation (data not shown). The approach (irradiation for 30 min) was then used to prepare UV-inactivated virus.

DNA microarray

A microarray analysis using a baculovirus DNA chip consisted of 192 spots including 140 AcMNPV genes was carried out as described elsewhere (32). In brief, total RNA was isolated using Trizol reagent (Invitrogen) and 400 pg of synthesized RNA, λpolyA+RNA-A (TaKaRa), was added in each series for normalization. The fluorescent-labeled cDNA prepared for hybridization was generated with an RNA Fluorescent Labeling Core Kit
(Takara) using 30 µg of total RNA and oligo(dT) according to the manufacturer’s instructions. DNA microarrays were hybridized for 10 h under coverslips with Cy5 or Cy3-dCTP (Amersham)-labeled cDNA probes, washed, dried and scanned immediately in an Affymetrix 428 Array Scanner (Affymetrix Instruments). Data were analyzed using ImaGene software (Biodiscovery Inc.).

5’ RACE and Northern blot analysis

Total RNA was isolated from Sf9 or HeLa14 cells at 48 hpi as above. 5’ RACE was performed using the GeneRacer core Kit (Invitrogen) according to the manufacturer’s instructions. Briefly, mRNAs were dephosphorylated, decapped and ligated with an adaptor oligo RNA supplied in the kit. A 1st PCR was then carried out with an adaptor primer from the kit and gene specific 1st PCR primers (ie-1 and ie-0: 5’-GTCTGTTCAGGGTGCCACGAC-3’, pe38: 5’-GGCTGGCGCACTGTCGTCAC-3’, gp64: 5’-AGACTGTTGCCACGCCC-3’, p6.9: 5’-GCGTGTTCTGTAACCTCGACC-3’). A 2nd PCR was performed using an adaptor-specific nested primer and gene-specific nested primers (ie-1 and ie-0: 5’-AACTGTTCCACACCCGTG3’, pe38: 5’-CCGTAATGCCCACGTTGCGAC-3’, gp64: 5’-CGACCAGCCGCTGGCATCTTTC-3’, p6.9: 5’-CGACCAGCCGGTGATCTTTC-3’).
5'-GGGGTCTACCCGGCGGCGT-3'). PCR products were cloned using an AT cloning pGEM-T Easy vector system (Invitrogen) and at least four cDNA were sequenced using the ABI PRISM 310 Genetic Analyzer for each gene.

Total RNA (10µg) isolated at the time presented in the figure was denatured and separated by electrophoresis in a 1% agarose gel, transferred to a nylon membrane (Hybond-N+; Amersham) and hybridized with oligonucleotide probes (human β-actin: 5’-ACGTCACACTTCATGATGGAGTTGAAGGTAGTTTCGTGGATGCCACAGGACTCC ATGCT-3’, human 18SrRNA: 5’-CTGGACC GGCCCTGCGTACTTAGACATGTATGGCTTAATTTGAGACAAGCATATGGTT-3’) labeled with [γ³²-P]ATP (Amersham) using MEGALABEL (Takara). Quantitative analysis was performed with an Image Reader BAS1000 (Fuji film).

Results

DNA microarray analysis in mammalian cells

We adopted a DNA microarray to analyze the transcripts of AcMNPV in the mammalian cells. Cy3 or Cy5-dCTP labeled cDNA probes were simultaneously hybridized to probe sequences on the microarray, and the amount of fluorescence seen with the individual dyes
was determined by microarray scanner. The positivity of the signals of the gene spots on
the DNA array chips was judged under the criteria “significantly positive: Signal mean –
(Background mean + 2×Background Standard deviation) > 0” (Fig. 1A). Forty-three viral
genes were detected in the HeLa14 cells and 14 genes in the BHK cells (Table 1). The
experiment was duplicated and the similar results were obtained. Twelve of these genes
overlapped in the two cell types, as confirmed by nested RT-PCR using gene-specific
primers (data not shown).

Fidelity in transcription

The fidelity of transcription initiation for the AcMNPV genes in HeLa14 cells was then
analyzed by 5’RACE. First of all, transcription initiation sites of ie-0, ie-1, gp64 and pe38
in Sf9 cells were determined at 48 hpi (Fig. 2). The transcription start site of ie-1 in Sf9
cells was located in a CAGT motif (early gene motif) located 77 nts upstream of the
translation start codon, consistent with that reported previously (25). In HeLa14 cells, ie-1
was also transcribed from this site (Fig. 2). ie-0 and gp64 of AcMNPV have both a CAGT
early motif (16, 9) and a TAAG late motif (4). 5’ RACE using mRNA purified from Sf9
cells at 48 hpi demonstrated that both genes were transcribed from the TAAG motif as
previously reported (16, 7, 34) while the transcription start sites of these genes in HeLa14
cells were located in the CAGT early motif (Fig. 2). Furthermore, sequencing of the cDNA synthesized against the 5’ terminal region of \( \text{ie-0} \) transcripts, which is known as the only baculoviral transcript produced by splicing (11), revealed that the intron sequence (nts 122946 to 127149, GB No., L22858) was precisely spliced out in HeLa14 cells. On the other hand, the transcription of \( \text{pe38} \) was rather complicated. At 48 hpi, two transcription initiation sites for \( \text{AcMNPV pe38} \) were elucidated in Sf9 cells, one in a TAAG motif located 4 nts upstream of the early CAGT motif and another located 452 nts downstream of the early motif (Fig. 2A) (20). This feature of the transcription, ie an internal transcription initiation site, corresponded with that observed for the \( \text{Orgyia pseudotsugata NPV (OpNPV) p34} \) gene, a homologue of \( \text{AcMNPV pe38} \) (31). The internal transcription initiation site generates a small mRNA with the capacity to code for a protein with a calculated molecular mass of 20 k both in \( \text{OpNPV} \) (31) and in \( \text{AcMNPV} \). The transcription of \( \text{pe38} \) in HeLa14 cells started at several sites which were not located in the CAGT motif (Fig. 2). The transcription start site of the late gene \( \text{p6.9} \) in HeLa14 cells was located 99 nts downstream of the late TAAG motif (Fig. 2).

**Upregulation of \( \beta \)-actin expression in mammalian cells inoculated with \( \text{AcMNPV} \)**

As described elsewhere (32), the NPV DNA microarray contains human \( \beta \)-actin and \( \lambda \).
phage sequences as internal controls for normalization between chips. Interestingly, the relative intensity of the signal for β-actin on the DNA array was clearly increased in BHK cells (Fig. 1) and moderately increased in HeLa14 cells (data not shown) inoculated with AcMNPV. We then carried out Northern blot hybridization using total RNA isolated from HeLa14 cells to analyze the kinetics of the transcription of β-actin (Fig. 3). The amount of transcript from β-actin in HeLa14 inoculated with AcMNPV increased until 6 hpi, reached a level about 2.5-fold that of the control and then rapidly decreased, while the amount of transcript in the control cells gradually decreased until 24 hpi. A similar profile for the transcription of β-actin was obtained from BHK cells (data not shown). UV-inactivated AcMNPV also induced upregulation of β-actin expression at a level comparable to that induced by non-irradiated AcMNPV at 6 hpi (Fig. 3B lanes 1-3).

Discussion

The host range of baculoviruses is restricted to arthropods including insects and crustaceans. The host specificity of AcMNPV was originally thought to be restricted to cells derived from insects. However, several groups recently reported that AcMNPV can transfer and express foreign genes under the regulation of appropriate eukaryotic promoters in several
types of mammalian cells and animal models (18). This observation led us to consider its
use in gene therapy given its low cytotoxicity in mammalian cells even at a high MOI, an
inherent incapability to replicate in mammalian cells, and the absence of preexisting
antibodies against baculoviruses in animals. On the other hand, Beck et al. reported that
AcMNPV infection repressed phenobarbital-mediated gene induction and stimulated
TNF-alpha, IL-1 alpha, and IL-1 beta expression in primary cultures of rat hepatocytes (5),
indicating that mammalian cells could be physiologically influenced by the experimental
attack with AcMNPV. More recently, AcMNPV was shown to induce an innate immunity
that confers protection from a lethal challenge of influenza virus in mice through the
recognition of the baculovirus genomic DNA by a Toll-like receptor (TLR) molecule (1, 3).
These observations are thus not in contradiction to previous observations that AcMNPV
genes were not transcribed in HeLa cells inoculated with AcMNPV at a MOI of 100 PFU
per cell (MOI of 100) at 24 hpi or after several passages (28). In contrast, we clearly
detected the viral transcripts in mammalian cells inoculated with AcMNPV at a lower MOI
(MOI of 30) using a DNA microarray and 5’RACE technologies. This discrepancy might
be due to the difference in the sensitivity of the detection methods used in the two
experiments.

5’RACE analysis revealed that all genes examined in this study, except pe38 and p6.9,
were transcribed from an early transcription start site motif (CAGT) preceded by a TATA-motif 30 nts upstream in HeLa14 cells (Fig. 2). The CAGT motif is a part of the eukaryotic transcription initiation site sequence (INR; YYANT/AYY) (27) and the early promoters for these viral genes with a typical eukaryotic RNA polymerase II promoter structure (a conventional TATA motif and an INR) are functional in HeLa14 cells. Furthermore, the transcripts from ie-0 were observed to be precisely spliced (11) in HeLa14 cells, showing the compatibility of the splicing machinery between mammalian cells and insect cells. These observations suggested that there was no critical disadvantage in the transcription of these genes in mammalian cells. A late gene, p6.9 was also transcribed from 25 nts downstream of a TATA-like motif in the HeLa14 cells. However, the transcription initiation site was located in the coding sequence (Fig. 2). Four transcription initiation sites (nucleotide positions +112, +454, +552 and +575) were identified for pe38 in HeLa14 cells and none of them corresponded with the start sites in Sf9 cells. The most upstream site (+112) and the second site (+454) were located in the INR-like sequences CCGCAGA and CCATTGT, respectively, which were also accompanied by a TATA-motif 25 nts upstream (Fig. 2A). The sequences of surrounding the other two sites showed no significant similarity to INR but were preceded by sequences with some similarity with the TATA-motif or TFIIB recognition element G/CG/CG/ACGCC (BRE) (17, 26) 30 nts
upstream (Fig. 2B). However, we could not obtain evidence for the transcript from a CAGT motif located 4 nts downstream of the transcription start site +1 for pe38 in HeLa14 cells though it was also accompanied by a TATA-motif 30 nts upstream. This suggested that the typical eukaryotic RNA polymerase II promoter structure was not sufficient to initiate transcription in the AcMNPV genome.

We happened to observe the upregulation of β-actin expression in AcMNPV-inoculated mammalian cells in an analysis using a NPV DNA microarray containing a human β-actin DNA spot as a control. Actin is essential for the transport of NPV to the nucleus in insect and mammalian cells (29) and is also necessary not only for constitution of the virogenic stroma but also for budding of the viruses in insect cells (6). In insect cells, AcMNPV infection induces overexpression of cellular actin and it inhibits the polyhedron synthesis and polyhedra formation, although factors stimulate actin expression are not clear (30). In order to investigate if the upregulation is related to the viral transcripts in HeLa14 cells, effects of UV-inactivated viruses on β-actin expression were examined. UV-inactivated viruses which were expected to be impaired in the fusion capability of envelope protein by the denaturation of GP64 (1) were able to induce an upregulation of β-actin expression compared to untreated control viruses, suggesting that the upregulation was caused by events occurring before the internalization of viral components. Thus, the significance of
the AcMNPV transcript to the physiology of mammalian cells remains to be solved.

In conclusion, the results of our investigation provided evidence that AcMNPV is capable of expressing some viral genes at least at the transcription level in mammalian cells through the usual pathway of infection. These results emphasize the significance of studying the molecular details of baculovirus-mammalian cell interactions to reinforce the inability of AcMNPV to replicate in mammalian cells for facilitating the use of baculoviruses in the agro-biological, pharmaceutical and medical fields.

References


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Figure legends

Fig. 1 (A) DNA microarray analysis using Cy5-labeled cDNA probes. The cDNA probes were synthesized from total RNA extracted from mock-infected cells (left) or
AcMNPV-infected BHK cells (at a MOI of 30) (right). Daggers indicate the positions of the spots for β-actin. Asterisks indicate the internal control, λ.polyA.

**Fig. 2** Mapping of the 5’ ends of the transcripts for ie-1 and ie-0, gp64, and pe38 in Sf9 and HeLa14 cells at 48 hpi. (A) Schematic representation of the gene structure for ie-1 and ie-0, gp64, and pe38. The open box shows the open reading frame. Arrows on the diagram for each gene indicate the initiation sites in Sf9 cells at the early (E) or late (L) stage and in HeLa14 cells (M), respectively. (B) Sequences around the transcription initiation sites in Sf9 cells and HeLa14 cells. The arrow shows the transcription initiation site and the number in parenthesis indicates the nucleotide position relative to the transcription initiation site in Sf9 cells (+1) at the early stage (for ie0 (16), ie1 (25), gp64 (9), and pe38 (20)) or the late stage (for p6.9 (this paper)). The CAGT early motif and the TAAG late motif are underlined. Bold type indicates TATA-box like and BRE-like sequences.

**Fig. 3** Northern hybridization assays using ³²P-labeled β-actin and 18SrRNA cDNA probes. (A) Northern hybridization was performed using total RNA purified from AcMNPV-infected HeLa14 cells at the times shown in the panel. Kinetics of the expression of β-actin is presented as a sequential line graph below the autoradiograph and the values
are shown as the ratio of $\beta$-actin to 18SrRNA on the basis of the signal intensities obtained from the Northern hybridization. Circles and boxes in the graph indicate the kinetics for AcMNPV-infected and mock-infected cells, respectively. (B) Northern hybridization assay using total RNA from Mock-infected (1), AcMNPV-infected (MOI of 30) (2) and UV-inactivated virus-infected (3) cells. The bar graph below the autoradiograph shows the ratio of hybridization signal for $\beta$-actin to that for 18SrRNA.
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* E or L indicates the presence of the following promoter motifs. E, early promoter motif (TATA box followed by CAGT motif); L, late promoter motif ((A/T/G)TAAG). Underlines indicate the genes which detected both in HeLa14 and in BHK cells.
Fig. 2

A

exon0

4.1kb splicing intron

gp64

pe38 L1,E

pe38 L2

pe38 M1

pe38 M2

pe38 M3

pe38 M4

predicted 20kDa region

pe38 coding region

p6.9 L1, L2

p6.9

B

ie-0

ie-0 L

GCACAGAGGGCTTGAGGCCAAGCTATAAAGCCCGTTGTGCCAATCCTGAAATCTAATGATC

ie-0 M

AATGTGTCCTCGGCAACAGCAGTCGGTTCGATGGCAGTATATGCAATTGATAAAGG

ie-1

ie-1 L

TATCGTGTCAGGAGGCTAGTATAAAATGACGATGCTAGTTGGATATTGGTTT

ie-1 M

TTTACAAGAAAACTCAAGAAAGTT

gp64

gp64 L

CCGTGGTCTGATCCGGGGATGGTTTTGCTTATCGAGATAATTGATGTGCAGTAC

gp64 M

ACGTAGGCCAGATAACGGTCGGG

pe38

pe38 L1,E

GTCCCCAATCGCAGCAGCTATAAAAGCAGAGCCACTCAACACTGTAAGCACAGTTCTGTTT

pe38 M1

ACAAACAATCGCAGCCCTAGCTACGATATAATGCAAGTATCTCGCAAGACAGTT

pe38 L2,M2

ACTTGTCCTTTTTGATTATGAAATATTCCGCGATTGTGCTGTTCACTTGGTTTTAC

pe38 M3,M4

TAAGTTTTACAGAAAACTCAAGAAAGATGTTCCGCGCCGCCGTGCCAGATATAAAACATATTATGCT

p6.9

p6.9 L1,L2

TAAACATATATTTGGGAGTTCAGTCGTCGAATGCAAAGCGTAAAAAATATTAA

p6.9 M

CCGCCGCCGTTCTTCAACCGGTCAGTAGAGGCTGAAGAGGCGGCCAAGAAGCTCGGTT
Fig. 3

A

\[ \beta \text{-actin} \]

0 6 12 24 48

mock-infection

AcMNPV-infection

\[ 18S \text{ rRNA} \]

B

\[ \beta \text{-actin} \]

1 2 3

18S rRNA

Ratio of \( \beta \)-actin/18S rRNA signal

0 6 12 24 48 (hpi)

Ratio of \( \beta \)-actin/18S rRNA signal

1 2 3

(hpi)