Re-visiting the endogenous capacity for recombinant
glycoprotein sialylation by baculovirus-infected Tn-4h and
DpN1 cells

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It was previously reported that Tn-4h and DpN1 cells have the endogenous capacity to efficiently sialylate secreted alkaline phosphatase (SEAP) when infected with a baculovirus expression vector. In contrast, it has been found that lepidopteran insect cell lines that are more widely used as hosts for baculovirus vectors typically fail to sialylate SEAP and other recombinant glycoproteins. Thus, the N-glycan processing capabilities of Tn-4h and DpN1 cells are of potential interest to investigators using the baculovirus expression system for recombinant glycoprotein production. In this study, we experimentally re-assessed the ability of Tn-4h and DpN1 cells to sialylate SEAP with Sf9 and glyco-engineered Sf9 cells (SfSWT-1) as negative and positive controls, respectively. Our results showed that the SEAP purified from SfSWT-1 cells was strongly sialylated and initially indicated that the SEAP purified from Tn-4h cells was weakly sialylated. However, further analyses suggested that the SEAP produced by Tn-4h cells only appeared to be sialylated because it was contaminated with an electrophoretically indistinguishable sialoglycoprotein derived from fetal bovine serum. We subsequently expressed, purified, and analyzed a second recombinant glycoprotein (GST-SfManI) from all four cell lines and found that only the SfSWT-1 cells were able to detectably sialylate this product. Together, these results showed that neither Tn-4h nor DpN1 cells efficiently sialylated SEAP or GST-SfManI when infected by baculovirus expression vectors. Furthermore, they suggested that previous reports of efficient SEAP sialylation by Tn-4h and DpN1 cells probably reflect contamination with a sialylated, co-migrating glycoprotein, perhaps bovine fetuin, derived from the serum used in the insect cell growth medium.

Keywords: baculovirus/insect/N-glycosylation/recombinant glycoproteins

Introduction

The production of therapeutic glycoproteins requires an expression system with the capacity for appropriate post-translational processing in order to ensure that the recombinant and native products will be structurally and functionally equivalent. The insect cell/baculovirus expression vector system (BEVS; Summers and Smith 1987; O’Reilly et al. 1992; Jarvis 2009) is commonly used to produce recombinant proteins requiring post-translational processing, including N-glycosylation. However, the BEVS falls short of the basic requirement given above because insect cell lines typically produce recombinant glycoproteins with N-glycans that are quite different from those found on the native products. More specifically, the processed N-glycans produced by insect cell lines usually have trimannosyl core (“paucimannosidic”) structures, whereas those on the native mammalian products have complex or hybrid structures, which have been elaborated beyond the trimannosyl core to include N-acetylgalactosamine, galactose, and often, terminal sialic acid residues (reviewed by Marz et al. 1995; Altmann et al. 1999; Marchal et al. 2001; Tomiya et al. 2004; Harrison and Jarvis 2006; Shi and Jarvis 2007; Geisler and Jarvis 2009). This difference in N-glycan processing is a serious limitation of the BEVS because terminal sugars, particularly sialic acids, can dramatically influence the biological functions and clinical efficacies of therapeutic glycoprotein products.

In order to critically assess this important biotechnological limitation of the BEVS, considerable effort has been expended to study the protein N-glycosylation pathways of insect cell systems (reviewed by Marz et al. 1995; Altmann et al. 1999; Marchal et al. 2001; Tomiya et al. 2004; Harrison and Jarvis 2006; Shi and Jarvis 2007; Geisler and Jarvis 2009). The majority of these studies have generally supported the view that lepidopteran insect cells process N-glycans less extensively than mammalian cells, as detailed above. In stark contrast, a few reports have claimed that some lepidopteran insect cell lines can produce recombinant glycoproteins with high proportions of complex, even terminally sialylated N-glycans (Davidson et al. 1990; Davis and Wood 1995; Joshi et al. 2001; Joosten and Shuler 2003a, 2003b; Palomares et al. 2003).

While these reports have been greeted with skepticism, there is growing evidence that insects have at least a limited capacity...
for glycoprotein sialylation. For example, low levels of sialic acids have been detected in various tissues of various insects, including Lepidoptera (Roth et al. 1992; Karacali et al. 1997; Karacali et al. 1999; Malykh et al. 2000). In addition, extremely low levels of complex, sialylated N-glycans have been found in Drosophila melanogaster (Aoki et al. 2007; Koles et al. 2007). Finally, it is now well established that D. melanogaster encodes key enzymes involved in sialic acid and CMP-sialic acid biosynthesis and glycoprotein sialylation (Kim et al. 2002; Koles et al. 2004; Viswanathan et al. 2006). In light of these findings, it is inappropriate to simply dismiss the claim that native, non-glyco-engineered lepidopteran insect cell lines can produce sialylated recombinant glycoproteins. Instead, this claim should be assessed more objectively by further experimentation. Thus, the purpose of this study was to experimentally assess the claim that Tn-4h and DpN1 cells can efficiently sialylate recombinant secreted alkaline phosphatase (SEAP) when infected with a baculovirus expression vector (Joshi et al. 2001; Palomares et al. 2003).

Tn-4h and DpN1 are native, non-glyco-engineered lepidopteran insect cell lines derived from Trichoplusia ni (cabbage looper) and Danaus plexippus (monarch butterfly), respectively. We expressed and purified SEAP with these cell lines using the same conditions and methods described in previous reports (Joshi et al. 2001; Palomares et al. 2003). We also included Spodoptera frugiperda (Sf9; fall armyworm; Summers and Smith 1987) and glyco-engineered Sf9 cells (SfSWT-1; Hollister et al. 2002) as negative and positive controls, respectively. The N-glycan structures of the recombinant SEAP preparations produced by these four cell lines were then analyzed by lectin blotting with rigorous specificity controls. Finally, these analyses were extended to include another model glycoprotein, glutathione-S-transferase-tagged S. frugiperda mannosidase I (GST-SfManI), which was used previously to assess the N-glycan processing capabilities of native and glyco-engineered lepidopteran insect cell lines (Seo et al. 2001; Hollister et al. 2002; Aumiller et al. 2003). Ultimately, we found that neither SEAP nor GST-SfManI were detectably sialylated by Tn-4h or DpN1 cells. Furthermore, we discovered an explanation for the claims of SEAP sialylation in previous reports, which was that those preparations were probably contaminated with a co-purifying serum sialoglycoprotein.

Results

Purification and preliminary characterization of SEAP produced by Tn-4h, DpN1, SfSWT-1, and Sf9 cells

To re-examine SEAP sialylation by Tn-4h and DpN1 cells, we were obliged to precisely replicate the cell culture, baculovirus infection, and SEAP purification methods used in the relevant previous studies (Joshi et al. 2001; Palomares et al. 2003). Tn-4h cells were cultured in T. ni medium—formulation Hink (TNM-FH) medium supplemented with 10% fetal bovine serum and 5 mM N-acetylmannosamine. DpN1 cells were cultured in TNM-FH medium supplemented with 10% fetal bovine serum alone. Both cell lines were infected with AcSEAP, which is the same recombinant baculovirus, and cell-free media were harvested at the same times post-infection and used to purify SEAP by phosphate affinity chromatography, as described in the previous reports. In the present study, we also introduced Sf9 cells (Summers and Smith 1987) as a negative control and SfSWT-1 cells, a subclone of Sf9 cells glyco-engineered to produce sialylated recombinant glycoproteins (Hollister et al. 2002), as a positive control for recombinant glycoprotein sialylation.

The relative purity of the SEAP preparations isolated from each of the four cell lines was examined by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) with Coomassie Brilliant Blue R250 staining or SDS-PAGE and immunoblotting with a SEAP-specific antibody, as described in Materials and methods. The arrows and numbers on the left-hand side of the figure indicate the positions of protein standards and their sizes in kilodaltons.

Lectin blotting analysis of SEAP sialylation

Subsequently, lectin blots with Sambucus nigra agglutinin (SNA) were performed to probe the SEAP preparations from all four cell lines for the presence of terminal α2,6-linked sialic acids (Figure 2B). The specificity of the lectin blotting method...
was monitored by pre-treating the SEAP preparations with either Flavobacterium meningosepticum peptide-N-glycosidase (PNGase-F; lanes marked P in Figure 2) or Arthrobacter ureafaciens neuraminidase (lanes marked N in Figure 2). The integrity and electrophoretic mobility of each enzymatically digested, purified protein was monitored by immunoblotting with a SEAP-specific antibody (Figure 2A). The results showed that PNGase-F increased the electrophoretic mobilities of all four SEAP preparations relative to the untreated controls, indicating that each had been successfully deglycosylated by this enzyme. In contrast, neuraminidase had no detectable effect on the mobilities of the SEAP preparations, including the one from SfSWT-1 cells. This result was inconclusive because desialylation was not necessarily expected to detectably alter the electrophoretic mobility of these proteins.

The lectin blotting results showed that only the untreated SEAP preparation from SfSWT-1 cells reacted strongly with SNA (Figure 2B). Pre-treatment with PNGase-F or neuraminidase abolished the SNA reactivity, confirming that the lectin blotting assay was specific for terminally sialylated N-glycans. The lectin blotting results also revealed that the untreated SEAP preparation from Tn-4h cells reacted very weakly with SNA, although the band was very faint and does not show up very well in the image (Figure 2B). Again, pre-treatment with either PNGase-F or neuraminidase abolished the SNA reactivity, demonstrating the specificity of the lectin blotting analysis. From these results, we initially concluded that the SEAP preparation from SfSWT-1 cells was highly sialylated, the SEAP preparation from Tn-4h cells was marginally sialylated, and the SEAP preparations from DpN1 and Sf9 cells were not sialylated, at least not at levels that could be detected in these assays. However, the SNA reactivity of the SEAP preparation from Tn-4h cells was inconsistent with the fact that it had a faster electrophoretic mobility than the control, sialylated SEAP preparation from SfSWT-1 cells (Figure 1). Thus, we began to consider alternative explanations for the weak SNA reactivity of the SEAP preparation from Tn-4h cells.

A serum protein contaminant co-purifies with SEAP via phosphate affinity chromatography

One potential alternative explanation was that the SEAP preparation from Tn-4h cells was contaminated with a sialoglycoprotein from the bovine serum in the growth medium, which had been co-purified with SEAP in the phosphoaffinity method and co-migrated with SEAP in SDS-PAGE. To examine this possibility, we performed sham phosphoaffinity purifications using the cell-free supernatant from mock-infected Tn-4h cells or fetal bovine serum alone, as the starting material. The eluted fractions from the sham purifications were then examined by immunoblotting and SNA blotting, with the eluted SEAP fractions from AcSEAP-infected Tn-4h and SfSWT-1 cells as controls. Neuraminidase treatments were used to monitor the specificity of the SNA blotting analysis. The immunoblotting results confirmed that the control SEAP preparations from SfSWT-1 and Tn-4h cells contained immunoreactive bands with slightly different electrophoretic mobilities that were not altered by neuraminidase treatment (Figure 3A). These results also showed that there was no detectable immunoreactive SEAP in the elution fractions from the sham purifications performed with the cell-free supernatant from mock-infected Tn-4h cells alone (Figure 3B). These results demonstrated the specificity of the lectin blotting analysis.

Fig. 2. N-Glycosylation of SEAP purified from various insect cell lines. Tn-4h, DpN1, SfSWT-1, and Sf9 cells were infected with AcSEAP, and recombinant SEAP was purified as described in Materials and methods. Equal aliquots of each SEAP preparation were then treated with buffer alone (−), PNGase-F (+ P), or neuraminidase (+ N) and analyzed by (A) SDS-PAGE and immunoblotting with a SEAP-specific antibody or (B) SDS-PAGE and SNA lectin blotting. The arrows and numbers on the left-hand side of the figure indicate the positions of protein standards and their sizes in kilodaltons.

Fig. 3. Co-purification of SEAP and a co-migrating sialoglycoprotein. Phosphoaffinity purifications were performed with the cell-free medium from AcSEAP-infected SfSWT-1 cells, AcSEAP-infected Tn-4h cells, or mock-infected Tn-4h cells or with fetal bovine serum alone, as the starting materials as described in Materials and methods. Equal aliquots of the purified column eluants were then treated with buffer alone (−) or neuraminidase (+) and analyzed by (A) SDS-PAGE and immunoblotting with a SEAP-specific antibody or (B) SDS-PAGE and SNA lectin blotting, as described in Materials and methods. The arrows on the left-hand side of the figure mark the position of the 66.4-kDa standard.
cells or serum, alone, as expected. The lectin blotting results confirmed that the SEAP preparation from SfSWT-1 cells contained a strongly SNA-reactive protein and the SEAP preparation from Tn-4h cells contained a weakly SNA-reactive protein (Figure 3B), as observed previously (Figure 2B). However, the lectin blots also showed that both of the sham preparations also contained weakly SNA-reactive proteins that co-migrated with bona-fide SEAP (Figure 3B). In every case, the SNA reactivity was eliminated by pre-treatment with neuraminidase, indicating that the reactivity was sialic acid-specific (Figure 3B). These results strongly suggested that the weakly SNA-reactive protein observed in purified SEAP preparations from Tn-4h cells (Figures 2B and 3B) was not SEAP. Rather, the SNA-reactive protein appeared to be a contaminating sialoglycoprotein with an indistinguishable electrophoretic mobility from Tn-4h cells (Figures 2B and 3B). These results strongly suggested that the weakly SNA-reactive protein appeared to be a contaminating sialoglycoprotein(s) in our purified SEAP preparations to small peptides that would run off the bottom of the gel, while leaving the ∼55 kDa protease-resistant SEAP fragment intact for analysis by our lectin blotting method.

Thus, SEAP was purified from AcSEAP-infected Tn-4h, DpN1, SfSWT-1, and Sf9 cells, incubated with buffer alone or buffer plus trypsin, resolved by SDS-PAGE, and analyzed by immunoblotting or SNA blotting, as described in Materials and methods. The immunoblotting results showed that trypsin produced immunoreactive SEAP fragments that were ∼10 kDa smaller than the buffer-treated controls, as expected (Figure 4A). The SNA blotting results showed that there was a strongly SNA-reactive protein in the untreated SEAP preparation from SfSWT-1 cells and very weakly SNA-reactive proteins in the untreated SEAP preparations from the other three cell lines (Figure 4B, lanes marked with negative symbol). However, the only SNA-reactive protein observed after trypsin treatment was the ∼55 kDa SEAP fragment from the control SfSWT-1 cells (Figure 4B, lanes marked with positive symbol). None of the SEAP fragments from any of the other insect cell lines, all of which were clearly visible in the immunoblot (Figure 2A), had any detectable SNA reactivity. These results strongly supported the idea that the SEAP produced by Tn-4h and DpN1 cells was not detectably sialylated but only appeared to be sialylated due to the presence of one or more contaminating, trypsin-sensitive sialoglycoproteins.

**Lectin blotting analysis of GST-SfManI sialylation**

Subsequently, we used a second model glycoprotein and protein purification method to circumvent the complication of a contaminating, co-migrating sialoglycoprotein(s) and extend our analysis of the endogenous capacity of Tn-4h and DpN1 cells for recombinant glycoprotein sialylation. We chose GST-SfManI as the second model for these studies because it is larger (∼96 kDa) than SEAP, it can be effectively purified by a different affinity method, and we had used it for several previous studies on the protein glycosylation patterns of native and engineered lepidopteran insect cell lines (Hollister and Jarvis 2001; Hollister et al. 2002; Aumiller et al. 2003; Hollister et al. 2003).

Tn-4h, DpN1, SfSWT-1, and Sf9 cells were infected with AcGST-SfManI under the same conditions used for SEAP production, as described above, and the GST-SfManI products were purified from the cell-free supernatants by a glutathione-affinity method, as described in Materials and methods. The purified GST-SfManI preparations were then analyzed by SDS-PAGE with immunoblotting or SNA blotting, as for SEAP. The results of the immunoblots showed that the GST-SfManI fusion proteins had similar electrophoretic mobilities, although the purified GST-SfManI from SfSWT-1 cells appeared to migrate as a poorly resolved doublet in which the upper band had a slightly retarded electrophoretic mobility (Figure 5A). The immunoblotting results also showed that neuraminidase had no detectable impact on the migration of any of the GST-SfManI preparations (Figure 5A). The SNA blotting results showed that the only clearly SNA-reactive
Glycoprotein sialylation by Tn-4h and DpN1 cells

We focused our attention on previous reports of SEAP sialylation by baculovirus-infected Tn-4h (Joshi et al. 2001) and DpN1 (Palomares et al. 2003) cells because these cell lines were relatively newly described, under-utilized hosts for baculovirus expression vectors. In fact, there were no data to refute the claim that these cells could produce recombinant glycoproteins with sialylated N-glycans. Moreover, the production of sialylated SEAP by baculovirus-infected adherent Tn-4h cell cultures required the addition of N-acetylmannosamine, which was consistent with the idea that these cells might utilize this sialic acid precursor in an endogenous pathway leading to recombinant glycoprotein sialylation. This pathway appeared to be quite efficient, as it was reported that baculovirus-infected Tn-4h cells cultured in the presence of N-acetylmannosamine produced SEAP with ~20% terminally sialylated N-glycans (Joshi et al. 2001). In related studies, it was reported that a baculovirus-infected Tn-4h cell variant (Tn-4s) also produced recombinant SEAP with up to ~20% sialylated N-glycans when cultured under various conditions (Joosten et al. 2003; Joosten and Shuler 2003a, 2003b). Finally, baculovirus-infected DpN1 cells, another relatively newly described host cell line, reportedly produced recombinant SEAP with ~13% sialylated N-glycans when grown in a standard medium lacking N-acetylmannosamine (Palomares et al. 2003). Thus, these reports appeared to document examples of highly efficient recombinant glycoprotein sialylation by relatively new, potentially advantageous hosts for baculovirus expression vectors.

Our analysis of SEAP preparations isolated from Tn-4h, DpN1, SfSWT-1, and Sf9 cells revealed that they each consisted of single bands of immunoreactive protein with the expected electrophoretic mobilities. Notably, the SEAP preparation isolated from SfSWT-1 cells had a slightly slower electrophoretic mobility than the SEAP preparations from the other cell lines, reflecting the ability of SfSWT-1 cells to produce complex, terminally sialylated N-glycans (Hollister et al. 2002). This difference in their electrophoretic mobilities provided an early clue that the SEAP preparations from Tn-4h, DpN1, and Sf9 cells might not, after all, contain any complex, terminally sialylated N-glycans. However, SNA, which is a lectin specific for terminal α2,6-linked sialic acids, bound weakly, though sporadically, to SEAP preparations isolated from Tn-4h, DpN1, and Sf9 cells. While we expected the SEAP from Sf9 cells to serve as a negative control, we nevertheless considered that the SNA binding results might truly reflect the ability of all three of these cell lines to produce a sialylated form of SEAP. We also considered the possibility that the SEAP preparations isolated from the non-glyco-engineered insect cell lines were not sialylated but only appeared to be sialylated because they might be contaminated with a mammalian sialylglycoprotein that was electrophoretically indistinguishable from SEAP.

This latter possibility was strongly supported by the results of sham phosphoaffinity purifications performed with the growth medium from mock-infected cells or fetal bovine serum, alone, which obviously contained no recombinant SEAP, but nevertheless yielded an SNA-reactive protein with the electrophoretic mobility of SEAP. Further support was obtained by trypsin treatment of the SEAP preparations from all four insect cell lines, as this produced an immunoreactive...
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~55 kDa SEAP fragment in each case but an SNA-reactive SEAP fragment only with the positive control preparation from SfSWT-1 cells.

Together, the results described above indicated that Tn-4h and DpN1 cells failed to efficiently sialylate recombinant SEAP when infected by a baculovirus expression vector. These cells also failed to detectably sialylate another recombinant N-glycoprotein, GST-SfManI, which was clearly sialylated by SfSWT-1 cells. Thus, the results obtained in this study do not support the idea that Tn-4h and DpN1 cells have the endogenous ability to produce recombinant glycoproteins with terminally sialylated N-glycans.

The results of this study also suggest a possible explanation for the previous reports of efficient SEAP sialylation by Tn-4h and DpN1 cells, which is that the SEAP preparations in those studies were likely contaminated with a sialylated, co-migrating glycoprotein derived from the serum used in the insect cell growth medium. We speculate that this contaminant might be bovine fetuin because fetuin has an apparent molecular weight of ~63–64 kDa (Johnson and Heath 1986), biantennary and triantennary N-glycans with both terminal α2,3- and α2,6-linked sialic acids (Rohrer et al. 1993), and is a major component of bovine serum. Notably, the N-glycans previously identified in SEAP preparations from various insect cell lines included triantennary and tetraantennary structures, some with terminal α2,3-linked sialic acids. In contrast, the only sialylated N-glycans detected in total N-glycan preparations of D. melanogaster were monooantennary structures with terminal α2,6-linked sialic acids (Aoki et al. 2007). Even native placental alkaline phosphatase has only biantennary, complex N-glycans (Endo et al. 1988), so it seems unlikely that a recombinant form of this protein would acquire more highly branched structures. Thus, the detection of more highly branched and α2,3-sialylated N-glycans in recombinant SEAP preparations is consistent with the idea that these preparations were contaminated with bovine fetuin, which has these types of oligosaccharide side chains.

Another interesting observation pursuant to the previous claims of SEAP sialylation by Tn-4h and DpN1 cells is that we detected the co-migrating, SNA-reactive protein contaminant somewhat sporadically in our SEAP preparations and found that the presence of the contaminant was inversely related to the yields of SEAP obtained in various expression and purification experiments. Generally, Sf9 and SfSWT-1 cells produced ~10-fold higher levels of SEAP than Tn-4h and DpN1 cells, and the latter SEAP preparations were more likely to contain the co-migrating, SNA-reactive protein contaminant. Based on these observations, we conclude that a lower expression level and, therefore, lower concentration of SEAP in the starting material for the phosphoaffinity purification step was a critical factor determining whether or not a given SEAP preparation would contain the co-migrating, SNA-reactive contaminant. We speculate that relatively non-specific phosphate-binding proteins can more effectively compete for sites on the affinity matrix in the presence of lower concentrations of SEAP. This speculation is indirectly supported by the fact that at least one baculovirus-derived protein was found to interact non-specifically with the phosphate affinity matrix when the SEAP concentration in the starting material was low (Zhang et al. 2001). Finally, this speculation is more directly supported by a survey of the literature reporting sialylation of SEAP by T. ni and DpN1 cells, which revealed an inverse correlation between the concentrations of SEAP in the starting material and the proportions of sialylation reported for SEAP (Table 1).

Materials and methods

Cells and viruses

DpN1 (Palomares et al. 2003), Tn-4h (Joshi et al. 2000), Sf9 (Summers and Smith 1987), and SfSWT-1 (Hollister et al. 2002) cells were all maintained at 28°C as adherent cultures in TMN-FH supplemented with 10% fetal bovine serum (Atlas Biologicals, Fort Collins, CO) and 0.1% Pluronic® F-68 (Sigma-Aldrich, St. Louis, MO).

AcSEAP and AcGST-SfManI, which are recombinant baculoviruses encoding SEAP or GST-SfManI under the control of the polyhedrin promoter, respectively, have been described previously (Davis et al. 1992; Kawar et al. 2000). All working virus stocks were produced by infecting 50–100 mL shake flask cultures of Sf9 cells in TMN-FH medium with low passage seed stocks at low multiplicities of infection (≤0.1 plaque-forming unit/cell). The cell-free culture media were harvested at 3–5 days after infection and stored at 4°C in the dark, and baculoviral titers were determined by plaque assays on Sf9 cells (Jarvis 2009).

Recombinant protein production

Recombinant proteins were produced by infecting adherent or semi-adherent cultures of Sf9, DpN1, Tn-4h, or SfSWT-1 cells in 75 cm² flasks containing a total of 5–15 million cells/flask to conform to previous experiments (Joshi et al. 2001; Palomares et al. 2003). DpN1, Tn-4h, and Sf9/SfSWT-1 cells were infected with AcSEAP or AcGST-SfManI at multiplicities of infection of 20, 10, and 4 plaque-forming units/cell, respectively. In each case, the virus was allowed to adsorb for 2 h at 28°C, and then the cells were pelleted by centrifugation at 300 × g for 5 min in a swinging-bucket rotor and re-suspended in fresh TMN-FH

<table>
<thead>
<tr>
<th>Cell line</th>
<th>Reference</th>
<th>SEAP concentration in cell-free media (U/mL)</th>
<th>SEAP sialylation (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tn-5B1-4</td>
<td>Palomares et al. (2003)</td>
<td>6.79</td>
<td>ND</td>
</tr>
<tr>
<td>Tn-4s</td>
<td>Joosten et al. (2003)</td>
<td>6.3</td>
<td>0.7</td>
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<tr>
<td>Tn-4s</td>
<td>Joosten and Shuler (2003b)</td>
<td>3.7</td>
<td>2.6</td>
</tr>
<tr>
<td>Tn-4s</td>
<td>Joosten and Shuler (2003a, 2003b)</td>
<td>1.4, 1.5</td>
<td>9.1, 20</td>
</tr>
<tr>
<td>DpN1</td>
<td>Palomares et al. (2003)</td>
<td>0.51</td>
<td>13.3</td>
</tr>
<tr>
<td>Tn-4h</td>
<td>Joshi et al. (2001)</td>
<td>NR</td>
<td>20</td>
</tr>
</tbody>
</table>

ND, not detected; NR, not reported.
medium supplemented with 10% (v/v) fetal bovine serum and 0.1% (w/v) Pluronic® F-68. The medium used to re-suspend the Tn-4h cells after infection was additionally supplemented with 5 mM N-acetylmannosamine to conform to conditions used in previous studies (Joshi et al. 2001). At 96 h post-infection, the culture media supernatants were harvested by centrifugation at 3200 × g for 10 min in a swinging-bucket rotor and were either used immediately for purification of SEAP or GST-SfManI or stored at −20°C for future purification.

**Protein purifications**

A phosphate affinity resin (4-aminobenzophosphonic acid-histidyl-agarose) was prepared and used to purify SEAP from the cell-free media of AcSEAP-infected insect cell cultures, as described previously (Kulakosky et al. 1998). Briefly, the cell-free media were centrifuged at 72,000 × g for 30 min in a Beckman 45Ti fixed-angle rotor to pellet debris and budded virus particles. SEAP and other proteins in the media were then concentrated and partially purified by adding solid ammonium sulfate to 55–60% saturation with stirring on ice. The precipitates were harvested by centrifugation at 17,000 × g in a Beckman JA20 fixed-angle rotor for 30 min at 4°C, re-dissolved in 5–10 mL of column buffer (20 mM Tris-HCl, 1 mM MgCl₂, pH 8.0), and then dialyzed overnight against 2 L of the same buffer. The dialysates were then loaded onto phosphate affinity columns with a bed volume of ~2 mL, which had been pre-equilibrated with column buffer. The columns were then extensively washed with at least 100 bed volumes of column buffer, once with 10 mL of column buffer containing 1 μM dibasic sodium phosphate, and finally eluted with 5–10 mL of 100 μM dibasic sodium phosphate in column buffer. The purity and enzymatic activity of SEAP were measured in each fraction by SDS-PAGE with Coomassie Brilliant Blue R-250 staining and a phosphatase activity assay, respectively, as described previously (Laemmli 1970; Davis et al. 1992).

Glutathione-affinity chromatography was used in analogous fashion to purify the GST-SfManI from cell-free supernatants of AcGST-SfManI-infected insect cell cultures after ammonium sulfate concentration, as described previously (Geisler et al. 2007). At 96 h post-infection, the culture media supernatants were harvested by centrifugation at 17,000 × g for 10 min in a swinging-bucket rotor and were centrifuged at 17,000 × g for 10 min in a swinging-bucket rotor to pellet debris and budded virus particles. SEAP and other proteins in the media were then concentrated and partially purified by adding solid ammonium sulfate to 55–60% saturation with stirring on ice. The precipitates were harvested by centrifugation at 17,000 × g in a Beckman JA20 fixed-angle rotor for 30 min at 4°C, re-dissolved in 5–10 mL of column buffer (20 mM Tris-HCl, 1 mM MgCl₂, pH 8.0), and then dialyzed overnight against 2 L of the same buffer. The dialysates were then loaded onto phosphate affinity columns with a bed volume of ~2 mL, which had been pre-equilibrated with column buffer. The columns were then extensively washed with at least 100 bed volumes of column buffer, once with 10 mL of column buffer containing 1 μM dibasic sodium phosphate, and finally eluted with 5–10 mL of 100 μM dibasic sodium phosphate in column buffer. The purity and enzymatic activity of SEAP were measured in each fraction by SDS-PAGE with Coomassie Brilliant Blue R-250 staining and a phosphatase activity assay, respectively, as described previously (Laemmli 1970; Davis et al. 1992).

**SDS-PAGE, immunoblotting, and lectin blotting analyses**

SDS-PAGE assays were performed under reducing conditions with 8–10% acrylamide resolving gels (Laemmli 1970), and proteins were either stained directly with Coomassie Brilliant Blue R-250 or electrophoretically transferred to polyvinylidene difluoride (PVDF) membranes (Immobilon P, Millipore), as described previously (Towbin et al. 1979). SEAP and GST-SfManI were detected on PVDF membranes with primary rabbit polyclonal antibodies against human placental alkaline phosphatase (AbD-Seronet, Oxford, UK) or GST (Sigma-Aldrich), respectively. An anti-rabbit IgG-alkaline phosphatase conjugate (Sigma-Aldrich) was used as the secondary antibody. Glycans attached to SEAP and GST-SfManI proteins on membranes were detected by lectin blotting using biotinylated lectins (Vector Laboratories, Burlingame, CA) followed by incubation of lectin-treated membranes with alkaline phosphatase-conjugated streptavidin. Immune complexes and lectin–streptavidin complexes were visualized by the chromogenic nitroblue tetrazolium/5-bromo-4-chloro-3-indolyl phosphate reaction (Blake et al. 1984).

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**Abbreviations**

BEVS, baculovirus expression vector system; GST, glutathione-S-transferase; GST-SfManI, Spodoptera frugiperda mannosidase I-glutathione-S-transferase fusion protein; PVDF, polyvinylidene difluoride; SDS-PAGE, Sodium dodecyl sulfate-polyacrylamide gel electrophoresis; SEAP, human placental secreted alkaline phosphatase; SNA, Sambucus nigra agglutinin; TNM-FH, Trichoplusia ni medium–formulation Hink.

**References**


