Identification of Biotinylated Molecules Using a Baculovirus-Expressed Luciferase-Streptavidin Fusion Protein

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INTRODUCTION

Streptavidin produced by the soil bacterium, Streptomyces avidinii, is a tetrameric protein where each subunit is capable of binding one molecule of biotin. The gene encoding streptavidin has been cloned and sequenced (2), and some chimeric proteins, such as streptavidin-protein A of Staphylococcus aureus, have been constructed and expressed in Escherichia coli (19). In addition, avidin that is both structurally and functionally homologous to streptavidin has been recently produced in E. coli (1).

Genes coding for various light-producing enzymes, luminesases, of bacterial as well as of insect origin have been previously expressed in both prokaryotes and eukaryotic organisms (4-6,8). We have used these genes as genetic reporters and for FACS-applications (9) as well as for different diagnostic applications such as production of reagents useful for enzyme-linked immunosorbent assays (ELISA) (15). The latter was conducted by constructing a genetic fusion of the genes encoding luciferase (LucGR) of the insect Pyrophorus plagiophthalmus, a Jamaican click beetle, and protein A of S. aureus.

The baculovirus expression vector system (BEVS) utilizes the strong polyhedrin gene promoter of the Autographa californica nuclear polyhedrosis virus (AcNPV). The genes of interest are inserted under the transcriptional regulation of the corresponding gene promoter, and the recombinant proteins are produced in lepidopteran insect cell cultures or larvae during viral infection (11,12,20). In most cases, the protein products have turned out to be processed similarly if not identically when compared to their authentic counterparts, thereby retaining their biological activity. Here, the abundant production of a LucGR-streptavidin fusion protein in Spodoptera frugiperda insect cells using the baculovirus system is reported.

MATERIALS AND METHODS

Cells and Virus

The lepidopteran insect S. frugiperda cell line (Sf9) was maintained as previously described (16,20). The cells were grown at 27°C in TNM-FH culture medium (Sigma Chemical, St. Louis, MO, USA) supplemented with 10% fetal bovine serum, 100 U/mL penicillin, 40 μg/mL streptomycin, and 2.5 μg/mL Fungizone® (Life Technologies, Gaithersburg, MD, USA) either in suspension or as monolayer cultures.

A recombinant baculovirus containing the streptavidin-luciferase fusion construct under the transcriptional regulation of the polyhedrin gene promoter (PHP) was produced by co-transfection of the plasmid pVL1393-LucGR-StreptAv, revealed that the fusion protein migrated with an apparent molecular weight of 75 kDa. Light emission measurements showed that the infected cells produced about 255 mg of the chimeric protein per liter of cell culture (127.5 μg/mL × 1×10^6 cells). Precipitation of the LucGR-StreptAv fusion protein with biotinylated acrylic beads as well as immunoblot analyses using biotinylated immunoglobulins indicated that both fusion moieties of the chimeric protein product were functional with respect to their physical and enzymatic activities.

ABSTRACT

A genetic fusion between streptavidin of Streptomyces avidinii and luciferase of Pyrophorus plagiophthalmus was constructed. The fusion protein was produced in the Sf9 insect cell line using the baculovirus expression vector system (BEVS). Sodium dodecyl sulfate polyacrylamide gel electrophoresis of the proteins from cells infected with the recombinant virus, VL1393-LucGR-StreptAv, revealed that the fusion protein migrated with an apparent molecular weight of 75 kDa. Light emission measurements showed that the infected cells produced about 255 mg of the chimeric protein per liter of cell culture (127.5 μg/mL × 1×10^6 cells). Precipitation of the LucGR-StreptAv fusion protein with biotinylated acrylic beads as well as immunoblot analyses using biotinylated immunoglobulins indicated that both fusion moieties of the chimeric protein product were functional with respect to their physical and enzymatic activities.
pLucGR(BS) (24) using the polymerase chain reaction (PCR) (17) and the following primers: forward primer: 5'-TTT TAA GGA TCC ATG ATG AAG AGA GAG AAA AAT GTT G-3' and the reverse primer: 5'-AA TTA ATC TAG AAG TTT AGA ACT CTG CTC AGC-3', where the underlined sequences denote the recognition sites for BamHI and XbaI, respectively. The streptavidin-coding fragment was amplified from the vector pUC8-SZ (2) using the following PCR primers: forward primer: 5'-T TAT ATT CTG GAC TGC-3' and the reverse primer: 5'-T TTT ATT GTA CTA GAT CTA CTG TGT AAC GGC GTG-3', where the underlined sequences represent the recognition sites XbaI and BglII, respectively. The PCR products were carried out using Vent polymerase (New England Biolabs, Beverly, MA, USA), and the fragments were purified on a low-gelling temperature agarose gel (FMC BioProducts, Rockland, ME, USA). The PCR fragment containing the luciferase gene was digested with BamHI and XbaI and ligated to pVL1393 (10), which had been cut with the same enzymes and treated with calf intestinal phosphatase (CIP), resulting in transfer vector pVL1393-LucGR-StreptAv (Figure 1). The recombinant DNA techniques and cultivation of the cloning host, E. coli MC1061, were performed essentially according to Sambrook et al. (18). Transformation of the bacterial cells was carried out using electroporation (3).

**Light Emission Measurements**

At each step of the cloning procedures, bacterial cells were measured for their luminescence by taking 100 µL of the E. coli MC1061 miniprep culture and adding 50 µL of substrate solution (1 mM d-luciferin [BioTools Oy, Turku, Finland], 100 mM Na-citrate, pH 5.0). Light emission was detected with a BioOrbit Oy 1250 manual luminometer (Turku, Finland). The light-emitting bacterial clones (5–10 mV at OD600 of 1) were collected and used for restriction analyses.

Similarly, light emission measurements were used for generating the recombinant baculoviruses as recently described (14). The SF9 cells infected with wild-type baculovirus (strain E2), or the recombinant viruses VL1392-LucGR (6), VL1392-ProtA-LucGR (15) or VL1393-LucGR-StreptAv, were grown as monolayer cultures in 24-well plates (Nunc, Roskilde, Denmark) at a density of 3 × 10^6 cells per well. The infected cells were harvested at 12, 24, 36, 48 and 72 h post infection (p.i.). Aliquots (1–10 µL) of the cell suspensions (measurements in vivo) or cell lysates (measurements in vitro) were mixed with 200 µL substrate solution (in vivo: 1 mM d-luciferin in 100 mM Na-citrate buffer, pH 5.0; in vitro: luciferase monitoring reagent [BioTools Oy] containing d-luciferin and ATP) and measured as described previously (6,15). Purified firefly luciferase (Sigma Chemical) was used as a standard in all experiments.

**Sample Preparation**

The samples exposed for analyses in vitro were prepared essentially as described previously (15). Briefly, at indicated time points p.i., cells were washed once with phosphate-buffered saline (PBS) (pH 7.4) and cytoplasmic extracts prepared using ice-cold disruption buffer (50 mM Tris-HCl [pH 7.6], 1 mM EDTA, 1% Nonidet® P40, 15% glycerol). Lysates were kept on ice for 15 min, sonicated 3 times for 5 s (MSE 100; Measurement Scientific Equipment, London, England, UK) and clarified by centrifugation at 12 000× g for 10 min. Light emission measurements were performed both from the soluble as well as the insoluble fractions. In the latter case, the precipitates were carefully overlaid with the same buffer once to eliminate the residual soluble supernatant fraction, after which they were resuspended several times prior to the light emission measurements.

**Binding to Biotinylated Acrylic Beads**

Acrylic beads labeled with biotin (Sigma Chemical) (binding capacity 1 mg biotin/25 mg beads) were incubated in buffer A (50 mM Tris-HCl pH 7.4, 1 mM EDTA, 15% glycerol) and washed 3 times with the same buffer (1 mL/25 mg beads) prior to the experiments. For each sample, 12.5 mg of beads in 200 µL buffer A were mixed with lysates corresponding to 60 000 cells. Samples were agitated for 1 h at room temperature (RT). Beads were collected by centrifugation at 280× g for 2 min, washed 3 times with 1 mL of buffer A and resuspended in 400 µL buffer A for light emission measurements. Supernatants were centrifuged at 280× g for an additional 2 min to ensure the absence of beads. Amounts of 20–200 µL of sample and 100–500 µL of substrate were used for the light emission measurements performed as described above.

**Sodium Dodecyl Sulfate Polyacrylamide Gel Electrophoresis (SDS-PAGE)**

Cytoplasmic extracts of cells infected with the recombinant viruses and the wild-type baculovirus were prepared as described above and mixed with an equal volume of 2× Laemmli sample buffer (7). Samples (20 µL) were boiled for 5 min and loaded on 10% SDS polyacrylamide slab gels along with molecular weight markers (Bio-Rad, Hercules, CA, USA). Proteins

![Figure 1. The transfer-plasmid construct pVL1393-LucGR-StreptAv used for generation of the recombinant virus.](image-url)
were separated by electrophoresis and visualized by Coomassie Brilliant Blue staining.

**Immunoblotting**

Immunoblot procedures were performed essentially as described by Towbin et al. (22). Proteins (9.3 μg of protein A purified human Ig per lane) were separated on 10% polyacrylamide gels (SDS-PAGE) according to Laemmli (7) and electrophoretically transferred to nitrocellulose membranes in transfer buffer (25 mM Tris, 192 mM glycine, 20% methanol) using a Bio-Rad Trans-Blot® System. The nitrocellulose sheets were then soaked in blocking buffer (PBS + Tween 20 [0.05%] + 2% skimmed milk [Valio Oy, Helsinki, Finland]) to prevent nonspecific antibody binding (1 h, RT). After two washes (PBS + Tween 20 [0.05%]), filters were immersed into a biotinylated mouse anti-human Ig antibody solution (Sigma Chemical) (1:500 dilution in PBS + Tween 20 [0.05%]). After a 36-h incubation at 4°C and two washes (PBS + Tween 20 [0.05%]), indicated dilutions of the recombinant streptavidin-LucGR lysates were added and incubated for 1.5 h at RT. The filters were washed twice in PBS containing 5% glycerol followed by the addition of substrate (luciferase monitoring reagent; 1 mL per filter per lane). Finally, the filters were exposed on Hyperfilm™-MP (Amersham International, Little Chalfont, Bucks, England, UK) for 1 h.

**RESULTS**

**Expression of the LucGR-Streptavidin Fusion Construct**

Sf9 cells infected with VL1393-LucGR-StreptAv, AcMNPV, VL1392-LucGR and Ac701-ProtA-LucGR were collected at 48 h p.i. The protein contents were analyzed by SDS-PAGE and Coomassie blue staining (Figure 2A). A major protein of about 75 kDa, corresponding to that of the fusion construct, was clearly identified in cells infected with VL1393-LucGR-StreptAv. A 32-kDa polyhedrin expressed by wild-type, baculovirus-infected cells and a 61-kDa LucGR are seen as somewhat weaker bands in this figure. The chimeric protein, ProtA-LucGR of 80–85 kDa, was not clearly identified in the Coomassie blue-stained gel (15). As can be seen from Figure 2B, the overall expression of the LucGR-StreptAv fusion protein as a function of time correlated with that of polyhedrin and luciferase (6). Some minor proteins also increasing in intensity are related to viral infection and thus are also seen

Figure 2. Synthesis of the chimeric protein, LucGR-StreptAv, in Sf9 cells during viral infection. (A) Total cell extracts were prepared at 48 h, p.i. and separated on 10% vertical SDS-PAGE slab gels followed by Coomassie blue staining. Cells infected with VL1392-LucGR and Ac701-ProtA-LucGR were treated similarly and used as controls. The molecular weight markers in thousands are shown on the left. (B) SDS-PAGE analysis and Coomassie blue staining of Sf9 cells infected with the recombinant virus, VL1393-LucGR-StreptAv. Cells were collected at 12, 24, 36, 48 and 72 h p.i. and treated similarly as in Figure 2A. The molecular weight markers in thousands are shown on the left. For details, see the Materials and Methods section.
in the wild-type, virus-infected cells. None of the indicated proteins were seen in the mock-infected Sf9 cells (data not shown).

Light Emission Measurements

Cells infected with the recombinant virus, VL1393-LucGR-StreptAv, were collected at 24, 48 and 72 h p.i. and exposed to in vivo light emission measurements after addition of the substrate, \( \beta \)-luciferin. As shown in Figure 3A, the amount of light increased with time up to 48 h p.i. The increase in light production as a function of time correlated with that obtained from cells infected with VL1392-LucGR and Ac701-ProtA-LucGR, respectively. The amounts of light produced by cells infected with VL1393-LucGR-StreptAv (5–10 mV per cell) were similar when compared to that from cells infected with the unfused VL1392-LucGR construct, whereas the levels of light emitted from cells infected with the Ac701-ProtA-LucGR fusion construct were lower (0.5–2 mV per cell). The amount of protein produced by Sf9 cells infected with the respective recombinant viruses was determined by in vitro light emission measurements, where purified authentic firefly luciferase (LucFF) was used as a standard. In Figure 3B, the amount of protein produced by cells infected with the corresponding viruses is shown in \( \mu \)g of protein per \( 1 \times 10^6 \) cells. At 72 h p.i., cell cultures infected with the recombinant virus, VL1393-LucGR-StreptAv, produced 127.5 \( \mu \)g/1 \times 10^6 cells of the chimeric protein. The corresponding expression levels for VL1392-LucGR and Ac701-ProtA-LucGR-infected cells were 239 \( \mu \)g/1 \times 10^6 cells and 23.5 \( \mu \)g/1 \times 10^6 cells, respectively.

Binding of the fusion protein, VL1393-LucGR-StreptAv, to immobilized biotin was also analyzed by incubating biotinylated acrylic beads with cytoplasmic extracts prepared from Sf9 cells infected with VL1393-LucGR-StreptAv. Expression of the fusion product was performed in insect cell culture medium lacking biotin. Light emission measurements revealed that about 50% of the fusion protein was precipitated with the biotinylated beads (data not shown). The biotinylated acrylic beads did not precipitate any light-emitting protein from cytoplasmic extracts prepared from cells infected with VL1392-LucGR or Ac701-ProtA-LucGR, which were used as controls.

Detection of Biotinylated Mouse Anti-Human Ig Antibodies

The specific reactivity of the baculovirus-expressed chimeric protein

Figure 3. Light emission measurements of Sf9 cells infected with the following recombinant viruses: VL1393-LucGR-StreptAv, VL1392-LucGR, Ac701-ProtA-LucGR and wild-type AcNPV. (A) Cells were collected at 24, 48 and 72 h p.i., kept intact and used for light emission studies after addition of the substrate \( \beta \)-luciferin (in vivo). Cells infected with wild-type baculovirus (AcMNPV) served as a control. (B) Cells were collected at 24, 48 and 72 h p.i., disrupted and exposed to light emission measurements after addition of the substrate (in vitro). The amount of recombinant protein was determined using authentic purified firefly luciferase (LucFF; 100 ng/mL) as a standard (for details, see Materials and Methods).
product was further evaluated by immunoblotting. Purified human Ig molecules were separated on SDS-PAGE slab gels and transferred to nitrocellulose membranes as described above. The corresponding proteins were allowed to react with biotinylated mouse anti-human Ig antibodies. These antibody conjugates were then exposed to cytoplasmic extracts containing LucGR-StreptAv and LucGR, respectively. As shown in Figure 4, a protein band with an apparent molecular weight of about 50 kDa, corresponding to the heavy-chain portion of the complete Ig molecule, was clearly visible when the proteins were incubated with the extract containing LucGR-StreptAv. This protein band was not detectable with an extract containing LucGR alone.

**DISCUSSION**

The strong binding capacity of streptavidin to D-biotin makes it an attractive molecule for many biotechnological and biomedical applications. The gene encoding streptavidin has been expressed in *E. coli*. In a study performed by Sano and Cantor (19), a truncated version of the corresponding gene was constructed and fused to protein A. In another study by Thompson and Weber (21) where a synthetic streptavidin gene was expressed, the yield of active protein after cell disruption, inclusion body solubilization and refolding appeared to be 3 mg/L. In both studies, however, the main difficulty was to obtain soluble and biologically active protein. Streptavidin has, in addition, been expressed as well as secreted from *Bacillus subtilis* cells (13). In that study, the yield of recombinant streptavidin in minimal biotin-free medium was 20 mg/L.

Here, we report the expression of a biologically functional genetic fusion containing coleopteran insect luciferase and bacterial streptavidin. The chimeric protein was produced in lepidopteran insect cells by using the baculovirus technology, where the first amino acid (D) at the mature amino terminus of streptavidin was linked to the last amino acid (L) of the carboxyl terminus of insect luciferase. The two coding sequences were linked together using an engineered XbaI recognition site yielding only one additional amino acid (L) between the domains. According to the light emission measurements, the Sf9 cell line was able to produce 127.5 μg of functionally active luciferase-streptavidin fusion protein per one million of infected Sf9 cells. These determinations were carried out using purified authentic luciferase from the firefly, *P. pyralis*, as a standard. This is in agreement with the data obtained from the SDS-PAGE analyses where the production levels were compared to polyhedrin expressed by wild-type AcMNPV (20). In addition, the light-emitting properties of the chimeric protein were almost identical compared with recombinant luciferase (LucGR). As shown by in vitro experiments (indicating the total amount of protein produced in milligrams), the Protein A-LucGR fusion protein was produced at significantly lower levels as compared with LucGR or LucGR-StreptAv (Figure 3B). This may be partly explained by the fact that a different baculovirus transfer vector was used to generate the corresponding recombinant virus (11).

The chimeric protein product described here appeared to bind specifically to vitamin D-biotin. This was shown by precipitating the fusion protein with biotinylated acrylic beads. Close to 50% of the protein was precipitated with these beads according to the light emission measurements. Furthermore, biotinylated anti-human immunoglobulins bound to human immunoglobulins on nitrocellulose sheets could be detected using the light-emitting properties of this protein and X-ray film. Interestingly, about half of the recombinant protein was soluble at 36 h p.i. when the infected cells were grown in ordinary insect cell culture medium. Expression of the fusion protein in culture medium lacking biotin appeared to improve the solubility (data not shown).

Due to the sensitive monitoring procedures by which luciferase activity can be detected, these enzymes represent interesting alternatives to the currently used genetic reporters as well as marker enzymes, e.g., for diagnostic use. Since streptavidin has been frequently used in a number of applica-

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**Figure 4. Immunoblot analysis of human immunoglobulins using biotinylated mouse anti-human Ig antibodies and the luciferase-streptavidin fusion protein (LucGR-StreptAv).** An amount of 9.3 μg of protein A purified human Ig was separated per lane. Dilutions of the recombinant fusion protein, LucGR-StreptAv, are indicated in the figure (for details, see Materials and Methods).
tions such as isolation of biotinylated macromolecules and labeling of biologically active materials, it was considered an attractive fusion partner (23). Together the data presented here show that recombinant baculoviruses can be used to produce a functional fusion protein containing the entire coding sequences of luciferase and streptavidin in lepidopteran insect cells. The dual biological activity of the corresponding protein product may be of value within several fields of the biosciences as well as diagnostic medicine.

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