Serum-free culture of the suspension cell line QB-Tn9-4s of the cabbage looper, *Trichoplusia ni*, is highly productive for virus replication and recombinant protein expression

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**Abstract**

Serum-free cultures of insect cells play an important role in the fields of protein engineering, medicine, and biology. In this paper, the suspension cell line QB-Tn9-4s of *Trichoplusia ni* (Hübner) (Lepidoptera: Noctuidae) was successfully adapted to serum-free Sf-900 III medium and passaged for 52 generations. The adapted QB-Tn9-4s cells grew faster. Their population doubling time shortened from 27.4 hr in serum-containing medium to 24.1 hr, and their maximal density increased by 1.83-fold, reaching 3.50 × 10^6 cells/mL in serum-free culture in T-flasks. The cells readily adapted to spinner culture, with maximum cell density of 4.40 × 10^6 cells/mL in a spinner flask. Although the infection rate of *Autographa californica* multiple nucleopolyhedrovirus and production of occlusion bodies (OBs) of the adapted QB-Tn9-4s cells were 91.0% and 85.4 OBs/cell, respectively, similar to those of QB-Tn9-4s cells cultured in serum-containing medium and control BTI-Tn5B1-4 cells, their budded virus titer was 4.97 × 10^7 TCID\(_{50}\)/mL, significantly higher than those of the latter two. In addition, the expression levels of β-galactosidase at six days post-infection and secreted alkaline phosphatase at seven days post-infection in the adapted QB-Tn9-4s cells reached 2.98 ± 0.15×10^4 IU/mL and 3.34 ± 0.13 IU/mL, respectively, significantly higher than those of QB-Tn9-4s and control BTI-Tn5B1-4 cultured in serum-containing media. The above findings establish a foundation for industrial production of virus and recombinant proteins in QB-Tn9-4s serum-free culture.

**Abbreviations:** AcMNPV, *Autographa californica* multiple nucleopolyhedrovirus; BV, budded virus; OB, occlusion body; SEAP, secreted alkaline phosphatase

**Keywords:** insect cell lines, population doubling time, virus production

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Introduction

Insect cell lines are of great importance in the production of baculovirus and recombinant proteins. They are generally cultured in media containing a certain percentage of serum to support cell growth and proliferation. However, serum is expensive and contains complex components detrimental to separation, purification, and detection of culture products, limiting the application of insect cells. Thus, developing serum-free cultures of insect cell lines is desirable in cell, genetic, and protein engineering, medical biology, biotechnology, and the production of baculovirus and recombinant proteins (Agathos 2007; Hashimoto et al. 2010). A variety of insect cell lines have been successfully cultured in serum-free media (Ikonomou et al. 2002; Lua et al. 2003; Imanishi et al. 2012). Among them, SF-21 and its clonal isolate SF-9 of the fall armyworm, Spodoptera frugiperda (Smith) (Lepidoptera: Noctuidae), and BTI-Tn5B1-4 (High Five) of the cabbage looper, Trichoplusia ni (Hübner) (Lepidoptera: Noctuidae), have been widely applied to virus production and recombinant protein expression and cultured in serum-free media (Granados et al. 2007). Inlow et al. (1989) showed that SF-9 had a shorter population doubling time in a serum-free suspension culture than in a serum-containing culture. Kwon et al. (2003) compared the growth and protein expression of SF-9, SF-21, and BTI-Tn5B1-4 in four different serum-free media and found that both SF-9 and BTI-Tn5B1-4 cells possessed advantages and disadvantages in actual application. BTI-Tn5B1-4 cells are highly susceptible to baculovirus and could provide superior production of occlusion bodies (OBs) and recombinant proteins when compared to other insect cell lines. On a per milliliter basis, BTI-Tn5B1-4 cells produce five- to seven-fold of heterologous proteins compared with SF-9 cells (Wickham et al. 1992; Davis et al. 1993). However, an alphaneovirus named Tn5 cell line virus was identified during production of hepatitis E virus-like particles in BTI-Tn5B1-4 cells infected with a recombinant baculovirus vector (Li et al. 2007), thus there is a serious risk of contamination when using virus-like particles to produce vaccines or recombinant proteins for therapeutic purposes in BTI-Tn5B1-4 cells (Merten 2007). Although SF-9 cells could yield more budded virus (BV), but they produce less OBs and recombinant proteins. In addition, both SF-9 and BTI-Tn5B1-4 are adherent cells. QB-Tn9-4s is a suspension T. ni cell line established in our laboratory. It has comparable production levels of OBs and recombinant proteins to BTI-Tn5B1-4 cells and does not agglomerate at high density in culture (Meng et al. 2008). In addition, QB-Tn9-4s cell line does not contain Tn5 cell line virus, thus it has application potentials in large-scale industrialized cultures (Shan et al. 2011). Therefore, in this study, the QB-Tn9-4s cell line was adapted to a serum-free medium and tested for its biological characteristics. The results showed that in serum-free medium, QB-Tn9-4s cells could grow well and produce high levels of OBs and recombinant proteins, showing broad application potentials.

Materials and Methods

Materials and reagents

T. ni embryonic cell line BTI-Tn5B1-4 (High Five) (Granados et al. 1994) and S. frugiperda ovarian cell line SF-9 (Pasumarthy and Murhammer 1994) were provided by Dr. Blissard, Boyce Thompson Institute of Cornell University. T. ni embryonic suspension cell line QB-Tn9-4s was
established and preserved in our laboratory (Meng et al. 2008).

*Autographa californica* multiple nucleopolyhedrovirus (AcMNPV-1A) (Wood 1980) and its β-galactosidase expressing recombinant strain AcMNPV-β-gal (Wickham et al. 1992) and secreted alkaline phosphatase (SEAP) expressing recombinant strain AcMNPV-SEAP (Davis et al. 1992) were kindly provided by Dr. Granados of Cornell University. All of the viruses were amplified and titrated following the plague assay method described by Wood (1977) using Sf-9 cells.

TNM-FH insect medium was prepared by supplementing Grace medium (Invitrogen, www.lifetechnologies.com) with 0.3% lactalbumin hydrolyzate (BD, www.bd.com), 0.3% yeast extract (BD) and 10% fetal calf serum (FBS) (Thermo Scientific, www.thermoscientific.com). Serum-free Sf-900 III medium was from Invitrogen and serum-free EX-CELL 420 medium was from SAFC Biosciences (Sigma-Aldrich, www.sigmaaldrich.com). O-nitrophenyl-β-D-galactopyranoside (ONPG) and p-nitrophenyl phosphate (PNPP) were from Sigma-Aldrich.

**Cell culture**

Cells at logarithmic growth phase were prepared as a suspension by gently pipetting up and down in a sterile culture hood. After being diluted five-fold with fresh medium, 5 mL of suspension was inoculated into each 25 cm² flask (Corning, www.corning.com) and cultured at 27°C in an incubator. Cells were observed with an IX71 inverted phase contrast microscope (Olympus, www.olympus-global.com) and passaged every four days.

**Adaptation and culture in serum-free medium**

QB-Tn9-4s cells stably-grown in TNM-FH medium were passaged and consecutively cultured in Sf-900 or EX-CELL 420 medium containing 6%, 4%, 2%, and 1% FBS for three passages each and then in serum-free Sf-900 III or EX-CELL 420 medium. After 10 passages in serum-free medium, they were used to measure their biological characteristics and observe their morphology. One hundred randomly selected cells were observed and photographed under a microscope and their size was measured using a microscopic scale.

**Measurement of cell growth and cell viabiliity**

Cells at the logarithmic growth phase were counted using a hemacytometer and diluted with fresh medium to $2 \times 10^5$ cells/mL. For the T-flask culture test of QB-Tn9-4s and BTI-Tn5B1-4 in different media, 5 mL of cells were inoculated in each 25 cm² culture flask and cultured at 27°C. Every 24 hr, cells in three randomly selected flasks were harvested and counted. For the spinner flask suspension culture test of QB-Tn9-4s in serum-free medium Sf-900 III, 80 mL of cells ($2 \times 10^5$ cells/mL) were seeded into a 125 mL spinner flask (Corning). Three cultures were prepared. The spinner flasks were placed on a magnetic stir plate and cultured at 27°C under stirring at 100 r/min. Each 1 mL of the suspended cells were removed every 24 hr. The cell density was determined using a hemacytometer (Freshney 2005). The cell growth curves were generated, and the population doubling times were determined as previously reported (Hayflick 1973). The viability of cells was measured by trypan blue staining method (Freshney 2005). In brief, an equal volume of cell suspension and 0.4% trypan blue solution was
mixed and transferred to the chambers of the hemacytometer, and all cells (stained and unstained) in the 1 mm² were counted. At least 500 cells were observed in the different squares for each culture. The cell viability was calculated using the following equation: cell viability (%) = viable cells (unstained) / total cells (stained and unstained) × 100.

**Determination of virus infection rate and OB production**

1 × 10⁵ of QB-Tn9-4s and BTI-Tn5B1-4 cells in TNM-FH medium and QB-Tn9-4s cells in Sf-900III medium were separately seeded in each well of 24-well plates (Corning) in triplicate, infected with AcMNPV-1A BV at a multiplicity of infection of 10 as previously reported (Meng et al. 2008), and incubated at 27°C. The cultures were collected at four days post-infection, and infected and uninfected cells were counted using a hemacytometer to determine the ratio of infection. Cells were then disrupted by sonication to release OBs. OB concentration was measured using a hemacytometer, and the average production of OBs per infected cell was calculated as the ratio of the number of total OBs produced to the number of cells infected, as reported previously (Wang and Granados 1992).

**Determination of BV production**

1 × 10⁵ of QB-Tn9-4s and BTI-Tn5B1-4 cells in TNM-FH medium and QB-Tn9-4s cells in Sf-900III medium were separately seeded in triplicate into a well of 24-well plates, and infected with AcMNPV-1A BV at a multiplicity of infection of 10, as previously reported (Meng et al. 2008). Culture medium was collected at four days post-infection by centrifugation at 5000 r/min for 3 min and was used as the virus source to measure BV titer, i.e. TCID₅₀ per mL, using Sf-9 cells and the limiting dilution method (Reed and Muench 1938).

**Measurement of recombinant protein production**

1 × 10⁵ of QB-Tn9-4s and BTI-Tn5B1-4 cells in TNM-FH medium and QB-Tn9-4s cells in Sf-900III medium were separately seeded in triplicate into a well of 24-well plates and infected with recombinant virus AcMNPV-β-gal or AcMNPV-SEAP at a multiplicity of infection of 10, as described by Meng et al. (2008). Samples were collected at various days post-infection and kept at -20°C.

The production of recombinant β-galactosidase and SEAP was measured as reported previously by Davis et al. (1992) and Zheng et al. (2005). In brief, for examining the level of SEAP expression, samples were sonicated and then heated at 65°C for 5 min. 2 µL of each sample was mixed with 200 µL of SEAP assay buffer (1.0 mol/L diethanolamine, 0.5 mmol/L MgCl₂, 10 mmol/L homoarginine, pH 9.8) in a 96-well microtiter plate (Corning) and incubated at 37°C for 10 min. Following the addition of 20 µL of 120 mmol/L p-nitrophenyl phosphate, absorbance at 405 nm was recorded at 1 min intervals in an MRX microplate reader (DYNEX Technologies, www.dynextechnologies.com). SEAP activity was calculated according to the formula IU/mL = (ΔOD₄₀₅/min) × (0.222 mL) / (18.8 mL/µmol cm) × (0.002 mL sample) × (0.56 cm path length).

To measure β-galactosidase levels, samples were sonicated for 10 sec and centrifuged for 2 min at 12,000 × g to remove debris. 2 µL of the supernatants was mixed with 0.8 mL Z-buffer (60 mmol/L Na₂HPO₄, 40 mmol/L NaH₂PO₄, 10 mmol/L KCl, 1 mmol/L NaCl, 10 mmol/L magnesium chloride, 0.002% sodium dodecyl sulfate (SDS), 0.05% sodium deoxycholate, 0.5% NP-40, 100 µg/mL benzamidine, 100 µg/mL leupeptin, 100 µg/mL aprotinin, 0.01% phenylmethylsulfonyl fluoride), and the absorbance at 420 nm was recorded at 1 min intervals in a microplate reader (BioRad, USA).
MgSO$_4$, 50 mmol/L β-mercaptoethanol, pH 7.4), and incubated at 28°C for 10 min. The reaction was initiated by addition of 0.2 mL of pre-warmed ONPG (4 mg/mL ONPG in Z-buffer). After 2 min of incubation at 28°C, the reaction was stopped by addition of 0.5 mL of 1 mol/L Na$_2$CO$_3$. β-galactosidase activity was measured by readings at OD 420 nm and calculated according to the formula IU/mL = (OD$_{420}$ × 1.5 mL) / (0.0045 × 2 min incubation × 0.002 mL sample).

**Statistical analyses**

The data were expressed as mean ± standard deviation. Differences in AcMNPV infection rate, BV titer, OB production, and protein expression were analyzed by one-way ANOVA and Duncan’s pairwise multiple comparison test using DPS data processing system (Tang and Zhang 2013).

**Results**

**Adaptation of QB-Tn9-4s cells to serum-free medium**

QB-Tn9-4s cells cultured in TNM-FH medium and EX-CELL 420 medium containing 6%, 4%, and 2% FBS had a similar growth rate and morphology. However, when cultured in EX-CELL 420 medium containing 1% FBS, their morphologies changed significantly. Some cells clustered together and particles appeared nearby. When cultured in serum-free EX-CELL 420 medium, many cells underwent deformation. Their boundaries blurred and the number of dead cells increased. To improve cell adaptability, cells were re-cultured in EX-CELL 420 medium with 1% FBS. After being passaged for four generations, cells were in a better growth status. However, when cultured again in serum-free medium, cells began to deteriorate; they were broken and eventually died. After repeated tries, cells failed to grow in serum-free EX-CELL 420 medium.

QB-Tn9-4s cells cultured in TNM-FH medium (Fig. 1A) were well adapted to serum-free Sf-900 III medium compared to serum-free EX-CELL 420 medium. When serially passaged to Sf-900 III medium containing 6%, 4%, 2%, and 1% FBS, after three passages they were able to successfully passage at a 1:4 ratio (cell: medium). After passage to complete serum-free Sf-900 III medium, their growth became slower. Some cells underwent aggregation and deformation (Fig. 1B) and could not be normally passaged. After refreshing with 3 mL medium every four days, cells gradually grew normally (Fig. 1C) and passed at a 1.5:3.5 ratio at confluency. About 50 days later, QB-Tn9-4s were able to grow stably in serum-free Sf-900 III medium and subcultured normally (Fig. 1D). Afterwards, cells grew significantly faster and passed at a 0.8:4.2 ratio, indicating they were successfully adapted to serum-free medium. After passage 10, cells were used to measure their biological characteristics. So far, QB-Tn9-4s cells have been passaged to 52 generations in serum-free Sf-900 III medium and were able to successfully recover after three months of storage in liquid nitrogen.

**Cell morphology**

The morphological characteristics of QB-Tn9-4s cells changed in the adaption process to the serum-free culture (Fig. 1). The proportion of cells with a spindle shape increased from 70.1% in TNM-FH medium to 92.0% in serum-free medium, while that of cells with a round shape was reduced from 29.9% in TNM-FH medium to 8.0% in serum-free medium. In addition, the size of cells with a spindle shape increased from 57.7 ± 10.5 × 18.2 ± 2.1 µm in TNM-FH
medium to 97.0 ± 17.0 × 19.1 ± 2.6 µm in serum-free medium, and the size of cells with a round shape decreased slightly from 22.2 ± 3.0 µm in TNM-FH medium to 21.3 ± 3.1 µm in serum-free medium.

**The spinner culture of QB-Tn9-4s cells in serum-free medium**

QB-Tn9-4s cells were readily adapted to spinner culture, and cell density increased generally during the early stage of suspension culture. In addition, cells were singles and not aggregated, and the cell mortality was less than 10% (Fig. 2). The maximum cell density of QB-Tn9-4s was 4.40 × 10^6 cells/mL in a spinner flask at 144 hr of culture (Fig. 3). As cells grew beyond that time, some aggregation of cells was observed.

**Cell growth curve and population doubling time**

The growth curves of QB-Tn9-4s cells in serum-containing and serum-free media in a T-flask and in a spinner flask were plotted using culture time as abscissa and cell density and cell viability as ordinate, respectively, and compared to that of control BTI-Tn5B1-4 cells. All cultures showed greater than 90% cell viability except BTI-Tn5B1-4 cells in TNM-FH at 24 hr after inoculation, which had only 88.7% cell viability (Fig. 2). As shown in Fig. 3, cell density increased significantly 24 hr after inoculation, indicating that the inoculated cells were less affected by mechanical injury and able to quickly adapt to the new culture environment. 144 hr after inoculation, the density of QB-Tn9-4s cells in serum-free Sf-900 III medium reached its peak of 4.40 × 10^6 cells/mL in spinner flasks and 3.50 × 10^6/mL in the T-flask, respectively. The cell density (3.50 × 10^6/mL) was 1.83-fold of that in TNM-FH medium (1.91 × 10^6/mL) and 1.87-fold of that of control BTI-Tn5B1-4 cells in T-flask culture (1.88 × 10^6/mL). With the culture time increasing, nutrients in the medium were consumed and cellular metabolites were increased, resulting in slowly decreased cell growth rate, declined cell density, and an increased amount of aggregated cells.

The population doubling times of QB-Tn9-4s cells in serum-free medium in the spinner suspension and in the T-flask were 22.9 hr and 24.1 hr, respectively, both shorter than that of QB-Tn9-4s (27.4 hr) and of the control BTI-Tn5B1-4 cells (28.0 hr) in TNM-FH medium, indicating that QB-Tn9-4s grew faster in serum-free medium than in serum-containing medium and faster than the control BTI-Tn5B1-4 cells.

**Virus infection rate and yield**

AcMNPV infection similarly changed the morphology of QB-Tn9-4s cells in TNM-FH medium and serum-free Sf-900 III medium. The cells grew slower and had condensed, aggregated chromatins; at three days post-infection, some cells appeared to have an enlarged nucleus and a small amount of OBs with strong refractivity in their nucleus; at four days post-infection, cells produced large amounts of OBs; at five days post-infection, large amounts of OBs were released into medium. Table 1 shows AcMNPV infection rate, OB yield, and BV titer in cells at different culture conditions. The infection rates of AcMNPV in QB-Tn9-4s cells cultured in both serum-free and serum-containing medium and control BTI-

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**Table 1.** The infection rate, occlusion body (OB) yield, and budded virus titer of AcMNPV in different cell lines. Values in the table are mean ± SD.

| Cell line     | Medium         | Infection rate (%) | OB yield (OBs/cell) | TCID₅₀/mL (×10^7) |
|--------------|----------------|--------------------|---------------------|-------------------|
| QB-Tn9-4s    | SF-900 III     | 91.0 ± 1.2         | 85.4 ± 3.0          | 4.97 ± 0.00       |
| QB-Tn9-4s    | TNM-FH         | 90.6 ± 2.8         | 84.0 ± 3.3          | 3.70 ± 0.06       |
| BTI-Tn5B1-4  | TNM-FH         | 91.1 ± 1.3         | 83.1 ± 2.7          | 3.16 ± 0.00       |
Tn5B1-4 cells in TNM-FH medium were greater than 90% with no significant difference. In addition, the average OB yields were higher than 83 OBs/cell. Although OB yields in QB-Tn9-4s cells cultured in serum-free and serum-containing media were higher than that of the control BTI-Tn5B1-4 cells, there were no significant differences.

The titer of BV was determined in triplicate in sf9 cells using culture supernatants collected at four days post-infection. As shown in Table 1, the titer of BV produced by QB-Tn9-4s cells in serum-free SF-900 III medium was the highest, followed by QB-Tn9-4s cells in TNM-FH medium, while that of BV produced by the control BTI-Tn5B1-4 cells in TNM-FH medium was the lowest. ANOVA showed extremely significant differences among the titer of BV produced at the three different conditions (P < 0.01).

Expression of recombinant β-galactosidase

The expression levels of β-galactosidase at two, four, six, and eight days post-infection in cells infected with recombinant virus AcMNPV-β-gal are shown in Figure 4. As shown, the expression level of β-galactosidase increased with the gradual increase of viral infection time, except that the highest level was seen six rather than eight days post-infection in QB-Tn9-4s cells cultured in serum-free SF-900 III medium. Moreover, at the same days post-infection, β-galactosidase expression level was the highest in QB-Tn9-4s cells cultured in serum-free SF-900 III medium, followed by QB-Tn9-4s cells cultured in TNM-FH medium, and the lowest was in control BTI-Tn5B1-4 cells. For example, at six days post-infection, β-galactosidase expression level reached a peak of 2.98 ± 0.15 × 10^4 IU/mL in QB-Tn9-4s cells cultured in serum-free SF-900 III medium, followed by 2.65 ± 0.11 × 10^4 IU/mL in QB-Tn9-4s cells in TNM-FH medium and 2.33 ± 0.09 × 10^4 IU/mL in control BTI-Tn5B1-4 cells in TNM-FH medium. There were significant differences among β-galactosidase expression levels in these three conditions at six days post-infection (P < 0.05) based on Duncan’s multiple comparison test.

Expression of recombinant secreted alkaline phosphatase (SEAP)

The expression levels of SEAP in cells infected with recombinant virus AcMNPV-SEAP at three, five, seven, and nine days post-infection are shown in Figure 5. As shown, SEAP expression levels gradually increased with time in all cells except in the control BTI-Tn5B1-4 cells, which showed the highest SEAP expression at seven days post-infection rather than at nine days. At the same days post-infection, SEAP expression level in QB-Tn9-4s cells was higher when cultured in serum-free SF-900 III medium than in TNM-FH medium. In addition, SEAP expression in QB-Tn9-4s cells was higher than in the control BTI-Tn5B1-4 cells. For example, at seven days post-infection, SEAP expression reached its peak level of 3.34 ± 0.13 IU/mL in QB-Tn9-4s cells in serum-free SF-900 III medium, which was extremely different from that of 2.32 ± 0.10 IU/mL in QB-Tn9-4s cells in TNM-FH medium and that of 2.23 ± 0.09 IU/mL in the control BTI-Tn5B1-4 cells at seven days post-infection (P < 0.01) based on Duncan’s multiple comparison test. Moreover, SEAP expression in QB-Tn9-4s cells in TNM-FH medium was not significantly different from that in the control BTI-Tn5B1-4 cells at seven days post-infection (P > 0.05).
Discussion

With the rapid advancement of serum-free cell culture techniques in recent years, a variety of insect cell lines have been successfully cultured in corresponding serum-free media (Ikonomou et al. 2002; Lua et al. 2003; Imanishi et al. 2012). However, the application of each medium is restricted to cell types because of their different adaptabilities to various serum-free media. Kwon et al. (2003) showed that serum-free Sf-900 II medium was suitable for Sf-9 and Sf-21 cells, whereas serum-free Express Five medium was preferable for Tn5B1-4 cells. In this paper, QB-Tn9-4s cells were trained to survive in both serum-free EX-CELL 420 and Sf-900 III media. The results showed that they were able to grow in serum-free Sf-900 III medium, but not in EX-CELL 420 medium. Currently, QB-Tn9-4s cells have been passaged for 52 generations, and are still in good status, laying a foundation for further research and application of the cell lines.

In insect cell cultures, serum provides not only growth factors for cell growth and reproduction but also detoxifying and antioxidant proteins and protease inhibitors for cell protection (Agathos 2007). In the adaption process of QB-Tn9-4s cells to serum-free medium, with reduction of serum concentration, cells were prone to aggregate, form many particles around areas with high cell density, and grow slowly. These phenomena may be due to the decreased protective effects in low serum medium and the mechanical damages caused by pipetting during the passaging process, which eventually led to poor cell morphology and aggregation growth. Renner et al. (1993) believed that damaged cells tend to aggregate because they could release a kind of DNA to bring cells together and thereby promote cell adhesion.

Researchers have studied and compared virus yield in a variety of insect cell lines in serum-free medium. Wang et al. (1992) showed that the productions of polyhedral and extracellular viruses of Sf-21, Sf-9, and Tn368 cells in serum-free medium were not significantly different from those produced in serum-containing medium. Chen et al. (1993) showed T. ni cells had higher AcMNPV titer and OB yield in serum-free EX-CELL 400 medium than in serum-containing TC199-MK medium, and OBs produced in serum-free culture were more virulent to T. ni larvae compared to those produced in serum-containing culture. In this study, the titer of AcMNPV BV produced in QB-Tn9-4s cells in serum-free Sf-900 III medium was significantly higher than that of BV produced in QB-Tn9-4s cells in TNM-FH medium and in the control BTI-Tn5B1-4 cells, indicating that QB-Tn9-4s cells in serum-free medium have application potential in studies on in vitro viral replication and proliferation.

In recent years, the insect baculovirus expression system has been widely used in theoretical research on proteins and massive production of genetically engineered vaccines and therapeutic proteins (Granados et al. 2007; Smagghe et al. 2009). Thus, breeding insect cell lines with the ability to highly express recombinant proteins is of great importance in research and application of cell engineering and in serum-free culture systems for exogenous protein expression. Wang et al. (1992) measured the β-galactosidase expression level in Sf-21, Sf-9, and TN368 cells in serum-free medium and found that β-galactosidase expression level in serum-free EX-CELL 400 cultures of all
the three insect cell lines was lower than that in serum-containing cultures. McKenna et al. (1998) measured the protein expression level in several T. ni embryonic cell lines in serum-free EX-CELL 400 medium and found that Tn-4B31 cells showed high protein expression levels, and the expression level of alkaline phosphatase was higher than that in BTI-Tn5B1-4 cells. Tatieek et al. (2001) compared the effect of different serum-free media on the expression of β-galactosidase in Sf-21 and BTI-Tn5B1-4 cells and found that β-galactosidase expression in Sf-21 cells was higher in EX-CELL 400 medium than in EX-CELL 401 and Sf900-II media, and expression of β-galactosidase in BTI-Tn5B1-4 cells was higher in EX-CELL 401 medium than in EX-CELL 405 medium. In this study, β-galactosidase and SEAP expressions in QB-Tn9-4s cells were higher in serum-free SF-900 III medium than in TNM-FH medium, indicating that protein expression of the same cell line was different in different media. Thus, it is feasible to select an appropriate serum-free medium for serum-free cultures of different insect cell lines to achieve high protein expression levels.

In this study, a new suspension cell line, QB-Tn9-4s, was successfully cultured in serum-free SF-900 III medium. The cells grew well in T-flasks and spinner flasks and had a high virus yield, titer, and recombinant protein expression level. These advantages provided broad prospects for development and in-depth research of the cell line. Further exploration on cell suspension culture in spinner flasks or shake-flasks and rational design of cell culture devices to study the metabolic characteristics of the cell line in large-scale cultivation and to optimize growth conditions would help to achieve large-scale industrialized cell cultures to express vaccines, enzymes, diagnostic reagents, and other important medical or valuable commercial goods.

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Figure 1. QB-Tn9-4s cells grown in serum-containing and serum-free media. A, QB-Tn9-4s cells in TNM-FH medium. B, The early third generation QB-Tn9-4s cells in serum-free Sf-900 III medium. C, The late third generation QB-Tn9-4s cells in serum-free Sf-900 III medium. D, The fifteenth generation QB-Tn9-4s cells in serum-free Sf-900 III medium. Scale bar: 40 µm. High quality figures are available online.

Figure 2. Viability of cells cultured in TNM-FH medium and serum-free Sf-900 III media. Data in the figure are expressed as mean ± SD. High quality figures are available online.
Figure 3. Growth curves of cells cultured in TNM-FH medium and serum-free Sf-900 III media. Data in the figure are expressed as mean ± SD. High quality figures are available online.

Figure 4. Beta-galactosidase expression level of cells cultured in different media at two to eight days post-infection. High quality figures are available online.
Figure 5. Secreted alkaline phosphatase expression level in cells cultured in different media at three to nine days post-infection. High quality figures are available online.