High Cell Density Cultivation of A Recombinant Bacillus Subtilis for Nattokinase Production

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Research

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Abstract

**Background:** Nattokinase (NK), a fibrinolytic enzyme, can be produced by culturing recombinant *Bacillus subtilis* in Luria-Bertani broth in a shaking flask. For use as a nutraceutical, however, a large-scale preparation and a simple purification process are required.

**Results:** The present study utilized a fed-batch process to cultivate a *B. subtilis* strain carrying a pHT01 plasmid with an NK-encoding gene (*B. subtilis*/pHT01-*aprN1*). For batch A (FB A), with a pH-stat two-stage fermentation strategy, we achieved an activity of 2910.5 ± 21.6 U mL\(^{-1}\) and a specific activity of 30.32 U mL\(^{-1}\) OD\(_{600}\)\(^{-1}\). Then, we changed the strategy with a later induction and lower feeding rate to pursue higher cell density and thus higher enzyme activity, a 11.9-fold activity of 4521.8 ± 23.8 U mL\(^{-1}\) was acquired, however, the specific activity was lower than FB A. For the third batch, low-glycerol-level-maintain feeding strategy was followed, and finally, a NK activity of 7778 ±17.28 U mL\(^{-1}\) was obtained, according to our knowledge, it was the highest activity assayed by the fibrin plate method ever reported. Furthermore, fermentation supernatant was successively purified by ammonium sulfate precipitation and nickel column affinity chromatography with a total NK recovery rate of 65.2%.

**Conclusions:** Our results indicate that there is a balance between the cell growth rate and NK expression when recombinant *Bacillus subtilis* is cultured with a fed-batch process. The equilibrium state can be attained by optimizing the induction and feeding strategy, and thus a high cell density and enzyme activity can be achieved.

Introduction

Thrombosis, which is responsible for high morbidity and mortality in humans [1], can be effectively treated by thrombolytic drugs, but these are associated with adverse effects [2, 3]. Therefore, it is necessary to develop new biological substances, especially prophylactic food-source thrombolytic agents with low immunogenicity and preventative, long-term effects that are convenient for oral administration and stable in the gastrointestinal tract.

Nattokinase (NK), which decreases the ability of blood to clot, is traditionally taken from natto, a Japanese solid-state fermented soybean food [4, 5]. Currently, NK is used as a dietary supplement as well as a prophylactic or a curative thrombosis medicine [5, 6]; however, the process used to purify it from natto is complicated and accompanied by significant loss of bioactivity. Accordingly, fed-batch fermentation by adopting genetically modified bacteria may improve enzyme yield [7–10].

Several reports have shown that fed-batch culture led to increases in NK activity of 2.1–25-fold relative to batch culture and that the addition of glycerol during the cell growth phase increased NK production significantly [10–16]. Moreover, various protein purification methods have been utilized for NK purification [6, 16]. Taken together, these findings suggest that the expression of NK by genetically...
modified bacteria may reach a much higher level if efficient protocols for high cell density fermentation and subsequent purification are obtained.

We previously constructed a *B. subtilis* 168 strain containing a pH101-aprN plasmid [17]. In this study, we further investigated the expression of NK by culturing this strain under different induction and feeding strategies. In addition, we investigated the efficiency of purification with ammonium sulfate precipitation and Ni–NTA affinity chromatography.

**Results And Discussion**

**NK expression by flask culture and fed-batch experiment**

At least five different NK activity assay methods have been reported to date [6, 11, 12, 18, 19]. Among these, the fibrin plate method and chromogenic method are the most widely used, but the casein-degradation and JBSL (Japan Bio Science Laboratory Co., Ltd) methods may show much higher values of enzyme activity [11, 12, 14].

The NK activity obtained at 24 h in flask culture was 380.14 ± 5.71 U mL⁻¹, while the OD₆₀₀ reached 13.45 ± 0.45.

Many studies have demonstrated the significant contribution of media ingredients [20, 21] and nutrient feeding strategy to NK production [6, 15]. Berenjian confirmed that glycerol was a noteworthy carbon source influencing cell density during the fermentation of *B. subtilis* natto, and that the highest activity of NK was obtained by adding 3% glycerol as a carbon source [15]. It has also been reported that 2,6-pyridine dicarboxylic acid (PDCA) and metal ions such as Ca²⁺ and Mg²⁺ could improve osmotic pressure and help to maintain enzyme conformation, thereby improving the NK activity [17, 22]. Moreover, Wang demonstrated that glucose, K₂HPO₄·3H₂O and MgSO₄·7H₂O played key roles in the production of NK, and they obtained an activity of 12.34 FU mL⁻¹ [23]. Taken together, the results of these studies showed that NK activity could be improved dozens of times by media optimization.

In addition to media, feeding solutions are critical factors that influence NK activity, which should support cell growth and recombinant protein production while avoiding substrate inhibition and other related problems [12, 15].

Based on this information, we selected a mixture of glycerol, yeast extract, PDCA and a concentrated inorganic mixture solution as the fermentation broth, and a mixture of glycerol and yeast extract as the feed broth.

The strategy of induction, including cell density at the time of induction, inducer concentration, pre-induction growth and post-induction incubation time, can also affect the efficiency of protein expression. The aim of this study was to investigate the effects of using pH-stat and low-glycerol-level-maintaining strategies on NK expression by *B. subtilis* 168/pHT01-aprN1.
Three experiments were performed to examine the effects of induction time, feeding time and feeding rate on the NK activity of fermentation broth. The results are presented in Figure 1 and Table 1.

Two-stage fermentation strategy has been used for production of recombinant protein and other metabolites [24-27]. For FB A, we proposed a two-stage fermentation strategy and expected to get a high production. In the first stage (cell growth stage), we fed 600 mL of media into the fermenter with a high flow rate at the eighth hour. Because the nutrition was not sufficient for cell growth, the glycerol concentration decreasing rapidly to about 70 mmol L$^{-1}$ at 12 h and the OD$_{600}$ not varying markedly after 13 h. However, the NK activity still increased significantly until 18 h. The final NK activity was 2910.5 ± 21.6 U mL$^{-1}$ and the specific activity was 30.32 U ml$^{-1}$ OD$_{600}^{-1}$.

Although we achieved a 7.7-fold activity compare to flask culture, the cell density and the total activity were not as high as reported previously [6, 12]. It suggested that later induction and lower feeding rate maybe to the benefit of higher cell density and thus higher enzyme activity [28, 29]. According to this hypothesis, for FB B, the expression was induced at the fifth hour, which was 1 h later than FB A, and the induced OD$_{600}$ was up to 27.3 ± 1.0, which was higher than that of FB A (17.6 ± 0.4). A three-stage feeding strategy was used, and the feeding of FB B started at 14 h, when the glycerol concentration was as low as 119.8 ± 1.3 mmol L$^{-1}$, which might have favored cell growth by reducing substrate inhibition. The glycerol content was sufficient to support the cell growth for 24 h. Although the OD$_{600}$ values from 15 h to 19 h were not varied significantly due to dilution and a new medium environment, it finally reached a high value of 208.8 ± 1.9 at 24 h, with a 11.9-fold activity of 4521.8 ± 23.8 U mL$^{-1}$. However, the specific activity was 21.66 U ml$^{-1}$ OD$_{600}^{-1}$ at 24 h, which was lower than that of FB A, and the NK activity did not synchronously increase following cell growth during the late fermentation stage, which implied that there should be a balance between the cell growth rate and enzyme expression.

The results of FB A and FB B suggested that maybe we should not unilaterally pursue high cell density, and instead the expression should be induced at an earlier time and kept a low glycerol content during the feeding period. Consequently, for FB C, continuous feeding was adapted, starting at 10 h when the OD$_{600}$ had reached 105.3 ± 1.1 and almost two-thirds of the initial glycerol had been consumed. The glycerol concentration was controlled to around 50 mmol L$^{-1}$ by adjusting the feeding rate.

As expected, an activity of 7778.0 ± 17.3 U mL$^{-1}$ and a specific activity of 44.86 U ml$^{-1}$ OD$_{600}^{-1}$ was achieved, and these values were 1.7-, 2.6-, and 26-fold higher than those of batch B, batch A and the flask culture, respectively. According to our knowledge, this was the highest ever reported activity value by the fibrin plate method [9, 10, 12, 15].

**Recovery rate of NK by purification**

The recovery rate of NK obtained by ammonium sulfate precipitation was 89.1% (Table 2), which was consistent with a study by Garg [16]. The high recovery rate and simple operation showed that this method could be used to purify NK on a large scale.
The purification was followed by Ni–NTA affinity chromatography, and the total recovery rate was 65.2% (Table 2). The high imidazole concentration in wash buffer B led to a high loss of NK when samples were purified using Ni–NTA affinity chromatography. Accordingly, the imidazole concentrations were adjusted to zero in wash buffer A and incubated supernatant, and to 10 mmol L\(^{-1}\) in wash buffer B; thus, a recovery rate of 88% was obtained in this step. Different imidazole concentrations in elution buffer C were also investigated, but there were no significant differences in the range of 100 mmol L\(^{-1}\) to 500 mmol L\(^{-1}\). According to the NK expression level and the recovery rate, the process developed here may be applied for large scale production of NK.

**SDS-PAGE and Western blotting analysis of NK**

SDS-PAGE analysis demonstrated that a 28 kDa protein was a crucial component in the supernatant from induced *B. subtilis* 168/pHT01-\(aprN1\), but that it was not present in the supernatant from *B. subtilis* 168/pHT01 and non-induced *B. subtilis* 168/pHT01-\(aprN1\) fermentation broth (Fig. 2 A/B), suggesting that recombinant NK could be expressed in a soluble form. Western blotting with a His-tag-specific monoclonal antibody also showed a specific signal at 28 kDa, whereas no cross-reaction occurred in the total soluble proteins from non-induced *B. subtilis* 168/pHT01-\(aprN1\) broth, which confirmed that the 28 kDa protein was the recombinant NK, as expected (Fig. 2 C).

In view of these reports, NK was produced by fed-batch cultures of recombinant *B. subtilis*, and its production was improved to 7,778 U mL\(^{-1}\) from 380 U mL\(^{-1}\) of flask culture using pH-stat and low-glycerol-level strategies. Future studies will design a process and set a kinetic model for fermentation optimization.

**Materials And Methods**

**Microorganism and inoculum preparation**

A gene-modified strain (*B. subtilis* 168/pHT01-\(aprN1\)) was used to express NK. The medium for flask and fed-batch culture was as follows: yeast extract, 60 g L\(^{-1}\); glucose, 40 g L\(^{-1}\); glycerol, 20 mL L\(^{-1}\); NH\(_4\)Cl, 3 g L\(^{-1}\); K\(_2\)HPO\(_4\)·3H\(_2\)O, 1 g L\(^{-1}\); MgSO\(_4\)·7H\(_2\)O, 2 g L\(^{-1}\); MnSO\(_4\)·H\(_2\)O, 0.02 g L\(^{-1}\); FeSO\(_4\)·7H\(_2\)O, 0.05 g L\(^{-1}\); CoCl\(_2\)·6H\(_2\)O, 0.01 g L\(^{-1}\); ZnCl\(_2\), 0.01 g L\(^{-1}\); 2,6-pyridinedicarboxylic acid, 0.167 g L\(^{-1}\). Glucose and MgSO\(_4\)·7H\(_2\)O were autoclaved separately and aseptically added to the medium. Nutrient solution contained 100 g L\(^{-1}\) yeast extract and 250 mL L\(^{-1}\) glycerol was used as feed medium. To avoid plasmid loss and prevent microbial contamination, chloramphenicol (5 \(\mu\)g mL\(^{-1}\)) was added to all media after being dissolved in anhydrous ethanol and filtered through a 0.22 \(\mu\)m membrane.

**Shake flask experiments**

To prepare a stock culture, one colony of *B. subtilis* 168/pHT01-\(aprN1\) was inoculated in 1 mL Luria-Bertani (LB) medium and then incubated overnight at 37°C in a shaking incubator at 200 revolutions per
minute (rpm). A flask with 25 mL fed-batch culture media was then inoculated with 7.5% (v/v) of stock culture and incubated at 37°C until reaching mid-log phase (OD_{600} of 0.8–1.0). Next, the temperature was adjusted to 28°C and held for 20 min, after which a 25-μL aliquot of 0.5 mmol L^{-1} isopropyl β-D-1-thiogalactopyranoside (IPTG) was added and the culture was further incubated for 45 h at 28°C in a shaking incubator (200 rpm) for NK expression.

**Fed-batch experiments**

An in-situ sterilizable 15 L bioreactor (BIOSTAT®B, Germany) equipped with pH and pO₂ probes was used for fed-batch experiments. One milliliter of glycerol stock bacteria was inoculated into 100 mL LB medium and cultured for 12 h at 37°C in a shaking incubator (200 rpm), after which the pre-culture solution was aseptically inoculated into 1 L LB medium and incubated under the same conditions for 12 h. Next, 700 ml of seed cultures were inoculated for 7-L fed-batch fermentation in the bioreactor. The value of oxygen dissolved was set at 20%, which was cascade controlled by agitation speed (300–1000 rpm) and air flow rate (3–20 L min^{-1}). The pH was set as 7.0 and automatically controlled by adding 2 mol L^{-1} HCl or 2 mol L^{-1} NaOH.

In the first fed-batch experiment (FB A), the temperature was adjusted to 28°C at 3.5 h, while IPTG was added 30 min later as the OD_{600} value reached about 18. A total of 600 mL feed media was supplemented at 8 h with a flow rate of 30 mL min^{-1}. In the second fed-batch experiment (FB B), the expression was induced at 5 h when the OD_{600} value had reached 27, and the feed media was constantly fed after 14 h of cultivation with a flow rate of 6 mL min^{-1} for 1 h, then 3.6 mL min^{-1} for 2 h, and finally 2.4 mL min^{-1} for another hour. In the third experiment (FB C), the expression was induced at 3 h when the OD_{600} reached about 9 and the feed media was fed at 10 h. The feeding rate depended on the content of glycerol, which was controlled at a concentration of 50 mmol L^{-1}. Cell growth was monitored at various times by measuring the OD_{600} values.

**NK activity assay**

Quantitative analysis of NK activity was conducted by the fibrin plate method, with slight modification [18]. Briefly, bovine fibrinogen (Sigma, St. Louis, MO, USA) and thrombin (Sigma, St. Louis, MO, USA) were dissolved in 0.1 mol L^{-1} sodium phosphate buffer (PBS) at pH 7.4. An equi-voluminal mixture of 5 g L^{-1} bovine fibrinogen solution and 12 g L^{-1} agarose solution was then warmed in a 45°C bath, after which 10 μL of 500 U thrombin was added to 15 mL of the mixture solution in a 90 mm petri dish and kept at room temperature for 1 h to form fibrin. Holes with a 2 mm diameter were made on the fibrin plate and 10 μL of each diluted supernatant of fermentation broth was then added. The plates were subsequently incubated at 25°C for 16 h, after which the areas of the lysis zones on the fibrin plates were measured and the fibrinolytic activities were determined according to the standard curve of urokinase.

**Glycerol concentration measurement**
The concentration of glycerol was determined enzymatically using a free glycerol assay kit (Sigma, St. Louis, MO, USA) according to the manufacturer's procedures.

**Purification and lyophilization of NK**

The final fermentation broth was collected and centrifuged (GL-25M, Luxiangyi, Shanghai, China) immediately at 18,300 × g for 10 min. Solid \((\text{NH}_4)_2\text{SO}_4\) was then gradually added until the supernatant was 20% saturated, after which it was stored at 4°C overnight. Next, the broth was centrifuged at 18,300 × g for 10 min and the precipitate was discarded. Additional \((\text{NH}_4)_2\text{SO}_4\) was subsequently added to make the supernatant 60% saturated, after which the mixture was stored at 4°C overnight again. Following centrifugation at 18,300 × g for 30 min, the supernatant was removed and the precipitate was dissolved in 10 mmol L\(^{-1}\) PBS (pH 7.4) as a crude enzyme. The crude enzyme was further purified by Ni–NTA affinity chromatography according to the manufacturer's (Novagen, San Diego, CA, USA) protocols. After purification, 10 µl of eluted fractions of different dilution times were used to assay their fibrinolytic activities. Residual solution was desalted by dialysis and lyophilized using a GLZY-0.5B vacuum freeze drier (Pudong Freeze Drying Equipment, Shanghai, China) to obtain NK Lyophilized powder.

**SDS-PAGE and Western blotting analysis**

SDS-PAGE protein analysis was performed in a Mini-Protean Tetra system (BioRad, Hercules, CA, USA) by loading 30 µL of 1: 1 (v: v) boiled supernatant and dye buffer onto a 4–20% precast Mini protean TGX gel (BioRad) and then running the samples in 1 × Tris-glycine-SDS running buffer (BioRad, Shanghai, China) at 100 V for 15 min followed by 200 V for 25 min. A protein standard (10-250 kDa, Precision Plus Protein Kaleidoscope, Bio-Rad) was used as a ladder.

NK was identified by Western blotting analysis as described by Towbin, with slight modification [30]. Briefly, 10 µL aliquots of different supernatants were separated by 15% SDS–PAGE. After electrophoresis, the protein bands were transferred from the gel to an Amersham nitrocellulose (NC) membrane (GE Healthcare, Piscataway, NJ, USA) using a Mini Trans Blot electrophoretic transfer cell (BioRad). NK on the membrane was reacted with a mouse monoclonal antibody (Yisheng Biological Technology, Shanghai, China) against His-tag (1:1000 v/v), then incubated with anti-mouse IgG alkaline phosphatase (Sigma) conjugate (1:2000 v/v). After washing, the NC membrane was stained using Super ECL Detection Reagent (Yisheng Biology Technology, Shanghai, China) according to the manufacturer's instructions, then covered by an X-ray film and exposed for 1 min. A Tanon v.3500 Gel Imaging System (Tanon Co., Shanghai, China) was used for the analysis of proteins.

**Declarations**

**Acknowledgment**

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Conflict of interest

The authors have no conflicts of interest to declare.

References

1. Wei X, Luo M, Xie Y, Yang L, Li H, Xu L, Liu H. Strain screening, fermentation, separation, and encapsulation for production of nattokinase functional food. Appl Biochem Biotechnol. 2012;168:1753–64.

2. Aloni PD, Nayak AR, Chaurasia SR, Deopujari JY, Chourasia C, Purohit HJ, Taori GM, Daginawala HF, Kashyap RS. Effect of Fagonia arabica on thrombin induced release of t-PA and complex of PAI-1 tPA in cultured HUVE cells. J Tradit Compl Med. 2015;6:219–23.

3. Peng Y, Yang X, Zhang Y. Microbial fibrinolytic enzymes: an overview of source, production, properties, and thrombolytic activity in vivo. Appl Microbiol Biotechnol. 2015;69:126–32.

4. Fujita M, Nomura K, Hong K, Ito Y, Asada A, Nishimuro S. Purification and Characterization of a Strong Fibrinolytic Enzyme (Nattokinase) in the Vegetable Cheese Natto, a Popular Soybean Fermented Food in Japan. Biochem Biophys Res Commun. 1993;197:1340–7.

5. Yanagisawa Y, Chatake T, Chibakamoshida K, Naito S, Ohsugi T, Sumi H, Yasuda I, Morimoto Y. Purification, crystallization and preliminary X-ray diffraction experiment of nattokinase from Bacillus subtilis natto. Acta Crystallogr Sect F: Struct Biol Cryst Commun. 2010;66:1670–3.

6. Yoo HJ, Kim M, Kim M, Lee A, Jin C, et al. The effects of nattokinase supplementation on collagen–epinephrine closure time, prothrombin time and activated partial thromboplastin time in nondiabetic and hypercholesterolemic subjects. Food Funct. 2019;10:2888–93.

7. Chiang CJ, Che HC, Chao YP, Tzen JT. Efficient system of artificial oil bodies for functional expression and purification of recombinant nattokinase in Escherichia coli. J Agric Food Chem. 2005;53:4799–804.

8. Liang X, Jia S, Sun Y, Chen M, Chen X, Zhong J, Huan L. Secretory Expression of Nattokinase from Bacillus subtilis YF38 in Escherichia coli. Molecul Biotechnol. 2007;37:187–94.

9. Ni H, Guo PC, Jiang WL, Fan XM, Luo XY, Li HH. Expression of nattokinase in Escherichia coli and renaturation of its inclusion bod. J Biotechnol. 2016;231:65–71.

10. Liu Z, Zheng W, Ge C, Cui W, Zhou L, Zhou Z. High-level extracellular production of recombinant nattokinase in Bacillus subtilis WB800 by multiple tandem promoters. BMC Microbiol. 2019;19:89.

11. Chen PT, Chiang CJ, Chao YP. Medium optimization for the production of recombinant nattokinase by Bacillus subtilis using response surface methodology. Biotechnol Prog. 2007;23:1327–32.

12. Cho YH, Song JY, Kim KM, Kim MK, Lee IY, Kim SB, Kim HS, Han NS, Lee BH, Kim BS. Production of nattokinase by batch and fed-batch culture of Bacillus subtilis. New Biotechnol. 2010;27:341–6.

13. Dabbagh F, Nagahdaripour M, Berenjian A, Behfar A, Mohammadi F, Zamani M, Irajie C, Ghasemi Y. Nattokinase: production and application. Appl Microbiol Biotechnol. 2014;98:9199–206.
14. Kwon EY, Kim KM, Kim MK, Lee IY, Kim SB. Production of nattokinase by high cell density fed-batch culture of *Bacillus subtilis*. Bioprocess Biosyst Eng. 2011;34:789–93.

15. Berenjian A, Mahanama R, Kavanagh J, Dehghani F, Ghasemi Y. Nattokinase production: Medium components and feeding strategy studies. Chem Ind Chem Eng. 2014;20:541–7.

16. Garg R, Thorat BN. Nattokinase purification by three phase partitioning and impact of t-butanol on freeze drying. Sep Purif Technol. 2014;131:19–26.

17. Cui Q, Qian BJ, Yao XM, Ji SL, Zhang JH. Construction of nattokinase-producing genetic engineering strain and its activity analysis. Food Sci China. 2017;38:1–8.

18. Kim W, Choi K, Kim Y, Park H, Choi J, Lee Y, Oh H, Kwon I, Lee S. Purification and characterization of a fibrinolytic enzyme produced from *Bacillus* sp. strain CK 11 – 4 screened from Chungkook-Jang. Appl Environ Microbiol. 1996;62:2482–8.

19. Friberge P, Knos M, Gustavsson S, Aurell L, Claeson G. Methods for determination of plasmin, antiplasmin and plasminogen by means of substrate S-2251. Haemostasis. 1978;7:138–45.

20. Liu J, Xing J, Chang T, Ma Z, Liu H. Optimization of nutritional conditions for nattokinase production by *Bacillus* natto NLSSE using statistical experimental methods. Process Biochem. 2005;40:2757–62.

21. Chen PT, Chao Y. Enhanced production of recombinant nattokinase in *Bacillus subtilis* by the elimination of limiting factors. Biotechnol Lett. 2006;28:1595–600.

22. Ikeda S, Ohsugi TT, Sumi H. Activation of fibrinolysis (Nattokinase) induced by dipicolinic acid and related compounds. Food Sci Technol Res. 2006;12:152–5.

23. Wang JK, Chiu HH, Hsieh CS. Optimization of the medium components by statistical experimental methods to enhance nattokinase activity. Foo-yin J Health Sci. 2009;1:21–7.

24. Chen SJ, Ke BS, Chiu IC. A fermentation strategy for production of recombinant protein subjected to plasmid instability. Korean J Chem Eng. 2008;25(5):1110–4.

25. Savergave LS, Gadre RV, Vaidya BK, et al. Two-stage fermentation process for enhanced mannitol production using *Candida magnoliae* mutant R9. Bioprocess Biosyst Eng. 2013;36:193–203.

26. Dusica R, Ana Lucia C, Stefan W, et al. Engineering *Corynebacterium glutamicum* for the production of 2,3-butanediol. Microb Cell Fact. 2015;14:171.

27. Mironczuk AM, Rakicka M, Biegalska A, et al. A two-stage fermentation process of erythritol production by yeast *Y. lipolytica* from molasses and glycerol. Bioresour Technol. 2015;198:445–55.

28. Hong H, Zhao X, Wu Z. A two-stage glycine supplementation strategy enhances the extracellular expression of sortase A in *Escherichia coli*. Process Biochem. 2019;76:11–7.

29. Zhu F, Deng H, He X, Song X, Chen N, Wang W. High-level expression of *Thermobifida fusca* glucose isomerase for high fructose corn syrup biosynthesis. Enzym Microb Technol. 2020;135:109494.

30. Towbin H, Staehelin T, Gordon J. Electrophoretic transfer of proteins from polyacrylamide gels to nitrocellulose sheets: procedure and some applications. Anal Biochem. 1979;76:459–71.
Tables

Table 1
Parameters for NK production process of flask fermentation and three fed-batch fermentation experiments with different induction and feeding strategies

|                | FB A  | FB B  | FB C  | LB    |
|----------------|-------|-------|-------|-------|
| Induction time | 4     | 5     | 3     | 3     |
| OD$_{600}$ at induction time | $17.64 \pm 0.44$ | $27.28 \pm 0.98$ | $9.14 \pm 0.34$ | $0.85 \pm 0.12$ |
| Feeding start time (h) | 8     | 14    | 10    | No feeding |
| Fermentation time (h) | 24    | 24    | 22    | 24    |
| Maximum value of OD$_{600}$ | $105.60 \pm 0.85$ | $208.80 \pm 1.94$ | $185.31 \pm 0.85$ | $13.45 \pm 0.45$ |
| NK activity (U mL$^{-1}$) | $2910.53 \pm 21.60$ | $4521.78 \pm 23.79$ | $7778.02 \pm 17.28$ | $380.14 \pm 5.71$ |

Table 2
Recovery rates of NK by ammonium sulfate precipitation and nickel column purification

| Sample                                      | Recovery rate (%) |
|---------------------------------------------|-------------------|
| Fermentation supernatant                    | 100               |
| 60% ammonium sulfate-precipitated protein   | $89.1 \pm 3.2$    |
| Protein adsorbed in nickel column           | $78.6 \pm 2.6$    |
| Nickel column elution buffer                | $65.2 \pm 2.4$    |