ABSTRACT

Nisin is an antimicrobial peptide, an important biopreservative, and it is produced by certain strains of Lactococcus lactis ssp. lactis. In this paper, a foam separation technique was used for the separation of nisin from its culture broth, and the effects of temperature and trehalose on the performance of foam separation of nisin were studied to increase the enrichment ratio and recovery percentage of nisin and decrease the inactivity percentage of nisin. The results showed that temperature and trehalose significantly affected the performance of foam separation of nisin. Under the optimum conditions of 50°C temperature, 150-mL/min air flow rate, 400-mL initial loading liquid volume, and 1-g/L trehalose concentration, the maximum enrichment ratio, recovery percentage, and the minimum inactivity percentage of nisin reached 23.7, 84.1%, and 5.9%, respectively, which were, respectively, 5.04, 0.93, and 1.03 times more than those under the conditions of 20°C temperature, 150-mL/min air flow rate, 400-mL initial loading liquid volume, and no trehalose addition. These results indicated that the change of temperature and the addition of trehalose could improve the performance of foam separation of nisin.

Key words: nisin, foam separation, temperature, trehalose

INTRODUCTION

Nisin is an antimicrobial peptide, and it is produced by certain strains of Lactococcus lactis ssp. lactis (Delves-Broughton et al., 1996; Delves-Broughton, 2005; Cao et al., 2007; Jozala et al., 2008; Trmčić et al., 2011). Nisin can intensely inhibit the growth of a wide range of gram-positive microorganisms, which are seriously harmful to foods and it can also inhibit the growth of Salmonella or some other gram-negative microorganisms by combing EDTA. In addition, nisin is nontoxic to humans. Therefore, nisin is widely used in dairy products, plant protein foods, canned foods, and meat products as a natural preservative in many countries and regions around the world (Cheeseman and Berridge, 1957; Liu and Hansen, 1990; Roberts et al., 1992; de Vos et al., 1993; Cheigh et al., 2004; Trmčić et al., 2011).

Nisin is produced by liquid culture using strains of Lactococcus lactis ssp. lactis coupled with downstream purification (Cheigh et al., 2004). The main separation methods of nisin reported in the literature are organic solvent extraction (de Arauz et al., 2009), cell adsorption (Yang et al., 1992; Van’t Hul and Gibbons, 1996), membrane separation (Daoudi et al., 2001), salting (Daoudi et al., 2001), and other adsorption methods (Coventry et al., 1996; Cheigh et al., 2004). These separation methods have the drawbacks of high cost, complicated operation steps, and serious pollution (Parente and Ricciardi, 1999), which have restricted the widespread application of nisin. Therefore, it is necessary to develop a new separation technology with low cost, simple operation, and no pollution.

Foam separation technology has the advantages of simple equipment, low investment, low energy consumption, and environmental compatibility (Pilon et al., 2002). Early in the 20th century, it had been applied in the metallurgy industry and environmental engineering. During recent years, more attention has been paid to the application of biochemical engineering, particularly the separation of proteins. Foam separation technology uses bubbles as the media and concentrates surfactants on the basis of their adsorption properties on the gas-liquid interface due to their hydrophobicity. Sarachat et al. (2010) studied the concentration of a rhamnolipid biosurfactant produced by Pseudomonas aeruginosa SP4 using foam separation and they obtained good separation efficiency of recovery percentage (97%) of the biosurfactant and enrichment ratio (4) of the biosurfactant. Vanhoute et al. (2008) studied the advancement of foam separation of bioactive peptides and found that a peptidic fraction (3,500–7,000 Da) having
antimicrobial activity with a 5% extraction yield could be separated from the 3% degree of hydrolysis. Hirsch (1950) suggested that foam separation could be used for the separation of nisin, which is also a biosurfactant. Wu et al. (2009) have been studying the separation of nisin from its broth by foam separation for many years and the results have been applied to industrial production (Guo et al., 2006; Liu et al., 2010; Zhang et al., 2011). Enrichment ratio and recovery percentage are used to describe the performance of foam separation. At present, enrichment ratio and recovery percentage in the industrial production process of nisin separation from its broth by foam separation are 5 and 85%, respectively. However, the performance of foam separation for concentrating nisin from its broth should be further improved, especially enrichment ratio, to reduce the separation cost. So, it is necessary to study how to increase the efficiency, especially enrichment ratio, of foam separation for concentrating nisin from its broth.

The main studied parameters affecting foam separation are air flow rate, liquid loading volume, foam height, surfactant concentration, pH, and bubble size. However, temperature is usually neglected. Kumpa-booth et al. (1999) and Qu et al. (2008) researched the effect of temperature on the recovery of SDS by foam separation, but the temperature used only ranged from 10 to 35°C and from 20 to 25°C, respectively. Grieves and Bhattacharyya (1965) studied the effect of temperature on foam separation efficiencies using the system of cationic surfactant EHDA-Br, and they did not study the effect of temperature on foam properties. In addition, only a few researchers have investigated the effect of temperature on foam properties (foaming ability and foam stability; Pradhan et al., 1990; Cóceca et al., 2003; Zhang et al., 2008). The effects of temperature on the protein separation from whey wastewater and tea saponin separation using 2-stage foam separation were studied (Jiang et al., 2011; Yan et al., 2011). The results showed that the increase in temperature could significantly affect the increase in enrichment ratio. However, no reports exist about the effects of temperature on foam separation of nisin from its culture broth. Therefore, it is very important to study the effects of temperature on foam separation of nisin from its culture broth.

It is possible for the proteins to be denatured during the process of foam separation (Burapatana et al., 2005). Because nisin is a biologically active peptide, it can be denatured using foam separation. To decrease nisin inactivity caused by foam separation, Zhang et al. (2011) researched the effects of ionic strength on nisin inactivity from its culture broth with the addition of sodium chloride and other salts and the results showed that ionic strength was a parameter affecting nisin inactivity caused by foam separation. So, it should also be noted that inactivity percentage of nisin cannot be ignored in foam separation at high temperature because protein thermal denaturation is a more important parameter that affects the functional properties of the proteins (Bellavia et al., 2011). Guo et al. (2006) had studied nisin thermal stability using the different protective agents and the results showed that trehalose could effectively decrease inactivity of the nisin product, which was heated to 115°C for 15 min. Trehalose is an effective protective agent that can preserve the structure and functions of the proteins (Yoshii et al., 2008). Furthermore, trehalose is also an important food additive and it does not affect the quality of the nisin product.

In this paper, nisin culture broth, produced by Lactococcus lactis ssp. lactis W28, was used as a system. The effects of temperature on the properties of the system, including viscosity, surface tension, foaming ability, and foam stability, and the efficiency of foam separation were studied for increasing the enrichment ratio of nisin. Then, the effects of trehalose on the efficiency of foam separation were studied for decreasing the inactivity percentage of nisin. So, a new technology could be developed for not only increasing the enrichment ratio of nisin but also for decreasing the inactivity percentage of nisin and increasing the recovery percentage of nisin using foam separation. Therefore, the application of foam separation technology can be promoted in biological and chemical industries.

**MATERIALS AND METHODS**

**Bacterial Strains and Media**

The nisin-producing strain (Lactococcus lactis ssp. lactis W28) was provided by Tianjin Kangyi Bioengineering Co. Ltd. (Tianjin, China). The culture medium consisted of 50 g of corn syrup (liquid), 40 g of sucrose, 20 g of KH₂PO₄, 10 g of peptone, 10 g of yeast extract, 2 g of NaCl, and 0.2 g of MgSO₄·7H₂O per liter of distilled water and its initial pH was adjusted to 7.6 by 1 mol/L of NaOH. Prior to culture, the inoculums were propagated twice at 30°C for 8 h in the seed medium (initial pH 6.9), which was composed of 20 g of KH₂PO₄, 15 g of sucrose, 15 g of peptone, 15 g of yeast extract, 2 g of NaCl, and 0.2 g of MgSO₄·7H₂O per liter of distilled water. The culture technology for the production of nisin was the fed-batch culture, which was detailed by Wu et al. (2009). Micrococcus flavus NCIB 8166, purchased from the China General Microbiological Culture Collection Center (Beijing, China), was detailed by Wu et al. (2009).
was used as the indicator strain in the nisin bioactivity assay. It was grown in SI medium (initial pH 7.2; Wu et al., 2009; 10 g of agar, 10 g of Tween-20, 8 g of tryptone, 5 g of glucose, 5 g of NaCl, 3 g of yeast extract, and 2 g of Na₂HPO₄ per L of distilled water). This medium was used in the bioassay of nisin. All media were autoclaved at 121°C for 20 min and stored at 4°C.

Nisin culture broth was obtained from Tianjin Kangyi Bioengineering Co. Ltd. The titer and pH of nisin culture broth were 4,200 IU/mL and 3.0, respectively. All above chemical reagents of analytical grade were purchased from Tianjin Yingtaxigui Chemical Reagent Co. (Tianjin, China).

**Instruments**

The instruments used in the experiments were a pH meter (research model pHS-25; Shanghai Leici Instrument Factory, Shanghai, China), electronic balance (model FA1204B; Shanghai Precision Scientific Instruments Co. Ltd., China), a Ross-Miles latherometer (model 2151; Shanghai Jianqiang Glass Instrument Co. Ltd., Shanghai, China), an automatic tensiometer (model JYW-200B; Chengde Experimental Machine Co. Ltd., Chengde, China), an air compressor (model AC0-318; Guangdong Hailea Group Co. Ltd., Guangdong, China), a glass rotameter (model LZB-3WB, 30~300 mL/min; Tianjin Hedongwuhan Factory, Tianjin China), an ultra thermostat (model 501; Shanghai Experimental Instrument Factory Co. Ltd., Shanghai, China), and an Ubbelodhe viscometer (I.D.3–0.49 mm; Shanghai Sendi Scientific Instrument Co. Ltd., Shanghai, China).

**Measurement Methods of Broth Properties** *(Surface Tension and Viscosity)*

Nisin culture broths of different temperatures were prepared from 20 to 80°C. The broth properties were characterized by surface tension and viscosity. The surface tension and viscosity of the broths with different temperature were measured by the automatic tensiometer and the Ubbelodhe viscometer, respectively.

**Measurement Methods of Foam Properties** *(Foaming Ability and Foam Stability)*

Nisin culture broths of different temperature were prepared from 20 to 80°C. The foam properties of the broths were characterized by the foaming ability and the foam stability, which were measured by the Ross-Miles latherometer (Carey and Stubenrauch, 2010; Yulin et al., 2010). The foaming ability and the foam stability were described by foam height (mm) and half collapse time (t₁/₂; min) of the foam phase.

**Experimental Process**

Figure 1 shows the experimental apparatus for foam separation. The foam separation column with marked scales was 1,200-mm tall with an i.d. of 40 mm and it was manufactured from transparent polymethyl methacrylate. The air compressor and the glass rotameter were used to control the flow of the compressed air, which passed through a sintered glass gas distributor of apertures about 40 to 60 μm at the bottom of the column. Bubbles formed in the bulk liquid phase when the air passed through the distributor. The column was tightly wound by a silica gel tube. The silica gel tube was connected to the ultra thermostat by which the temperature of the column was adjusted. Foam from the top of the column overflowed into a collection vessel. The process was operated until no foam overflowed from the column. The volume and the titer of the foam concentrate liquid and the residual broth were measured to calculate the enrichment ratio (E), recovery percentage (R), residual percentage (r), and inactivity percentage (I) using Equations 1 to 4:

\[
E = \frac{T_f}{T_i} \times 100%; \quad [1]
\]

\[
R = \frac{T_f}{T_i} \cdot \frac{V_f}{V_i} \times 100%; \quad [2]
\]

\[
r = \frac{T_r}{T_i} \cdot \frac{V_r}{V_i} \times 100%; \quad [3]
\]

\[
I = 100% - R - r, \quad [4]
\]

where \(T_f, T_i, \) and \(T_r\) are the titers of nisin in the foam concentrate broth, the initial broth, and the residual broth, respectively; and \(V_f, V_i, \) and \(V_r\) are the volumes of the foam concentrate liquid, the initial broth, and the residual broth, respectively.

The titer of nisin culture broth (4,200 IU/mL) could be maintained when the pH of the broth was 3.0. However, the titer of the broth decreased easily with the increase in pH (Jozala et al., 2008). Therefore, the pH of the broth was at 3.0 in the experiments and the effects of pH on the performances of foam separation of the broth were not studied. All of the above experiments were performed in triplicate.

**Measurement Method of Nisin Titer**

The agar diffusion method was used to measure the titer of nisin, and it is a bioassay used for the quantification of nisin by measuring the response of nisin
RESULTS AND DISCUSSION

Effect of Temperature on the Broth Properties and the Foam Properties of Nisin Culture Broth

Effect of Temperature on the Broth Properties of Nisin Culture Broth. The effect of temperature on viscosity and surface tension of nisin culture broth is shown in Figure 2a. The viscosity of the nisin culture broth decreased significantly with the increase in temperature from 20 to 80°C. The increase in temperature led to the expansion of the liquid volume and the increase of the distance between molecules. Therefore, intermolecular forces were weakened. Furthermore, the increase in temperature intensified the molecular motion of nisin and some of the micelle formed by nisin molecules were destroyed. So, the viscosity of nisin culture broth decreased significantly with the increase of temperature.

The surface tension of the nisin culture broth decreased significantly with the increase in temperature. The increase in temperature led to the decrease in viscosity of the nisin culture broth. As a result, nisin molecules were more easily adsorbed onto the gas-liquid interface. Therefore, the surface tension of the nisin culture broth decreased significantly with the increase in temperature.

Figure 2b shows the effect of temperature on foaming ability and foam stability of the nisin culture broth. The foaming ability of the nisin culture broth decreased with the increase in temperature. This is because the decrease in viscosity of the nisin culture broth made it easier for nisin molecules to diffuse. Consequently, nisin molecules were more easily adsorbed onto the gas-liquid interface, resulting in a decrease in foaming ability.

Effect of Temperature on the Foam Properties of Nisin Culture Broth. The effect of temperature on foaming ability and foam stability of the nisin culture broth is shown in Figure 2b. The foaming ability of the nisin culture broth decreased with the increase in temperature. The increase in temperature led to the decrease in viscosity of the nisin culture broth. As a result, nisin molecules were more easily adsorbed onto the gas-liquid interface, resulting in a decrease in foaming ability.

The experimental data are presented in terms of arithmetic averages of at least 3 replicates, and the standard deviations are indicated by error bars.
result, it was hard to form enough thick liquid film, resulting in an increase in bubble coalescence. Therefore, the foam height was smaller when the temperature was higher.

The foam stability of the nisin culture broth decreased with the increase in temperature. The resistance of foam drainage decreased due to the decrease in viscosity of the nisin culture broth with the increase in temperature and the liquid film became thinner. Furthermore, the liquid film of the upper foam always protruded upward, which made the film sensitive to evaporation and the increase in temperature led to a decrease in the relative humidity, so the evaporation of the liquid film accelerated (Li et al., 2010). As a result, the thickness of the liquid film became thinner and the foam stability of the nisin culture broth decreased.

**Effect of Temperature on the Foam Separation Efficiency of Nisin**

**Effect of Temperature on the Enrichment Ratio and Recovery Percentage of Nisin.** The effects of temperature on the enrichment ratio and recovery percentage of nisin were carried out under the conditions of 150-mL/min air flow rate and 400-mL initial loading liquid volume. The temperature ranged from 20 to 80°C. The effects of temperature on the enrichment ratio and recovery percentage of nisin are shown in Figure 3a.

The recovery percentage of nisin decreased with the increase in temperature. The enrichment ratio of nisin increased and then decreased slightly with the increase in temperature from 20 to 50°C and from 50 to 80°C, respectively. The viscosity of nisin culture broth decreased significantly with the increase in temperature from 20 to 50°C and the foam drainage was improved. So, the coalescence of the bubbles in the foam phase increased and the foam concentrate liquid volume decreased. Therefore, the enrichment ratio of nisin increased and the recovery percentage of nisin decreased significantly. However, inactivity of nisin occurred in the foam separation process. The effect of inactivity of nisin at the higher temperature was more dominant than the effect of the foam drainage. Therefore, the enrichment ratio of nisin decreased when the temperature was above 50°C.

**Effect of Temperature on the Inactivity Percentage of Nisin.** The effect of temperature on the inactivity percentage of nisin was carried out under the same conditions as above. The effect of temperature on inactivity percentage of nisin is shown in Figure 3b.

The inactivity percentage of nisin increased with the increase in temperature. In the foam separation process, many factors, especially the formation, coalescence, and rupture of the bubbles and oxidation, led to the inactivity of biosurfactants, such as enzymes (Liu et al., 1998; Clarkson et al., 1999a,b; Burapatana et al., 2004). The coalescence and rupture of the bubbles in the foam phase and the oxidation of the biosurfactants increased with the increase in temperature and so the inactivity percentage of nisin increased during foam separation. Therefore, the optimal temperature was 50°C.

**Effect of Air Flow Rate on the Enrichment Ratio and Recovery Percentage of Nisin at 50°C**

The effects of air flow rate on the enrichment ratio and recovery percentage of nisin were carried out under
the conditions of initial 400-mL loading liquid volume and 50°C temperature. The air flow rate ranged from 100 to 300 mL/min. The effects of air flow rate on the enrichment ratio and recovery percentage of nisin are shown in Figure 4.

An increase in air flow rate increased both the transportation of the bulk liquid into the foam and the biosurfactant adsorption onto the increasing surface area of the bubbles. Therefore, the foam became wetter and the foam concentrate liquid volume was greater, leading to a decrease in the enrichment ratio of nisin. In contrast, at a lower air flow rate, a longer residence time of the bubbles in the rising foam increased the foam drainage, resulting in a dryer foam with a higher nisin concentration in the foam concentrate liquid. So, the larger enrichment ratio of nisin was obtained at the lower gas velocity rate. Therefore, the optimal air flow rate was 150 mL/min.

**Effect of Initial Loading Liquid Volume on the Enrichment Ratio and Recovery Percentage of Nisin at 50°C**

The effects of initial loading liquid volume on the enrichment ratio and recovery percentage of nisin were carried out under the conditions of 150-mL/min air flow rate and 50°C temperature. The initial loading liquid volume ranged from 300 to 600 mL. The effects of initial loading liquid volume on enrichment ratio and recovery percentage of nisin are shown in Figure 5.

The enrichment ratio of nisin decreased with the increase in initial loading liquid volume, whereas the recovery percentage of nisin increased from with an increase in the initial loading liquid volume from 300 to 400 mL and then decreased with an increase in the initial loading liquid volume from 400 to 600 mL. The foam height decreased with the increase in initial loading liquid volume, leading to a shorter residence time of bubbles in the foam phase and so the enrichment ratio of nisin decreased. When the initial loading liquid volume was small, the height of the foam phase was large and the increase of the foam height led to a longer foam residence time, which improved foam drainage. But inactivity percentage of nisin increased at high temperature (50°C) and so the recovery percentage of nisin was small when the initial loading liquid volume was small. So, the largest recovery percentage of nisin was 72.5% when initial loading liquid volume was 400 mL. Therefore, the optimal initial loading liquid volume was 400 mL.

**Effect of Trehalose on Foam Separation Efficiency of Nisin at 50°C**

**Effect of Trehalose on the Inactivity Percentage of Nisin at 50°C**. The results from Figure 3b show that inactivity percentage of nisin was as high as 20.2% at 50°C temperature. To decrease the inactivity percentage of nisin, the effects of trehalose on the inactivity percentage of nisin were carried out under the conditions of 50°C temperature, 150-mL/min air flow rate, and 400-mL initial loading liquid volume. The concentration of trehalose ranged from 0 to 2.5 g/L. The results are shown in Figure 6a.
From Figure 6a, trehalose had an important effect of decreasing the inactivity percentage of nisin during the foam separation process. Trehalose is an exceptional stabilizer of proteins and helps retain the activity of enzymes in solution (Kaushik and Bhat, 2003). Yoshii et al. (2008) studied the retention of the enzymatic activity of alcohol dehydrogenase on spray-drying conditions using trehalose and the results showed that trehalose was an effective protective agent that could preserve the structure and activity of nisin during the foam separation process. Therefore, the optimal trehalose concentration was 1.0 g/L.

**Effect of Trehalose on the Enrichment Ratio and Recovery Percentage of Nisin at 50°C.** The effects of trehalose on the enrichment ratio and recovery percentage of nisin were carried out under the conditions of 50°C temperature, 150-mL/min air flow rate, and 400-mL initial loading liquid volume. The trehalose concentration ranged from 0 to 2.5 g/L. The results are shown in Figure 6b.

The results from Figure 6b showed that the enrichment ratio and recovery percentage of nisin increased significantly with the increase in trehalose concentration from 0 to 1.0 g/L and the increase in the enrichment ratio and recovery percentage of nisin became insignificant when the trehalose concentration was higher than 1.0 g/L. So, the effects of trehalose on the enrichment ratio and recovery percentage of nisin were consistent with those of trehalose on inactivity percentage. The enrichment ratio and recovery percentage of nisin were 23.7 and 84.1%, respectively, at the optimal trehalose concentration 1.0 g/L.

**CONCLUSIONS**

The present experimental studies showed that temperature and trehalose significantly affected the enrichment ratio, recovery percentage, and inactivity percentage of nisin. The enrichment ratio of nisin increased significantly with the increase in temperature from 20 to 50°C. The inactivity percentage of nisin decreased significantly with the increase in trehalose concentration from 0 to 1.0 g/L. Under the optimum conditions of 50°C temperature, 150-mL/min air flow rate, 400-mL initial loading liquid volume, and 1-g/L concentration of trehalose, the maximum enrichment ratio and recovery percentage and the minimum inactivity percentage of nisin reached 23.7, 84.1%, and 5.9%, respectively, which were, respectively, 5.04, 0.93, and 1.03 times more than those under the conditions of...
20°C temperature, 150-mL/min air flow rate, 400-mL initial loading liquid volume, and no trehalose addition. Therefore, the change in temperature and the addition of trehalose can promote the application of foam separation technology in biological and chemical industries.

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REFERENCES


