ABSTRACT

In the current study, a simple, sensitive, and specific ELISA assay using a high-affinity anti-bovine β-casein monoclonal antibody was developed for the rapid detection of cow milk in adulterated yak milk. The developed ELISA was highly specific and could be applied to detect bovine β-casein (10–8,000 μg/mL) and cow milk (1:1,300 to 1:2 dilution) in yak milk. Cross-reactivity was <1% when tested against yak milk. The linear range of adulterant concentration was 1 to 80% (vol/vol) and the minimum detection limit was 1% (vol/vol) cow milk in yak milk. Different treatments, including heating, acidification, and rennet addition, did not interfere with the assay. Moreover, the results were highly reproducible (coefficient of variation <10%) and we detected no significant differences between known and estimated values. Therefore, this assay is appropriate for the routine analysis of yak milk adulterated with cow milk.

Key words: yak milk, adulteration, monoclonal antibody, indirect competitive ELISA

INTRODUCTION

Yak milk is an important income source for rural residents of Qinghai-Tibetan Plateau, China. As a result of its high levels of proteins, lactose, conjugated linoleic acids, and calcium, yak milk is considered to be a naturally concentrated milk (Sheng et al., 2008; Li et al., 2011; Livingstone et al., 2012). Yak milk and yak milk-derived products (e.g., yogurt and cheese) have become increasingly popular in recent years (Bai et al., 2011; Nikkhah, 2011). Yak is a species that lives in mountainous areas of Central Asia, at an altitude of 2,500 to 5,500 m (Zi et al., 2008). Yak lactation is seasonal, thus, yak milk production is quite limited (Wiener et al., 2003). Therefore, for legal, consumer protection, and consumer confidence reasons, it is important to be able to rapidly detect the presence of cow milk in yak milk to ensure yak milk quality.

To date, several analytical approaches have been developed to detect the adulteration of milk of different species in dairy products (Moatsou and Anifantakis, 2003; Hurley et al., 2004b). Infrared spectroscopy was applied to detect the adulteration of goat or sheep milk with cow milk (Rodriguez-Otero et al., 1997; Nicolaou et al., 2010). High-performance liquid chromatography coupled with electrospray ionization mass spectrometry was used to detect cow milk in goat milk (Chen et al., 2004). Brescia et al. (2004) used a nuclear magnetic resonance method to differentiate buffalo and cow milk samples according to species. Lee et al. (2004) detected the adulteration of cow milk in goat milk by PAGE. Two-dimensional electrophoresis was applied to detect bovine milk in buffalo Mozzarella cheese (Chianese et al., 1990). Enzyme-linked immunosorbent assay is a rapid, sensitive, and specific method that is widely used in the food industry (Giovannacci et al., 2004). Adulterated goat, sheep, or buffalo milk and milk products have been successfully detected by ELISA (Garcia et al., 1990; Rodriguez et al., 1991; Haza et al., 1997, 1999; Hurley et al., 2004a, 2006; Song et al., 2011). However, no studies have applied ELISA to the detection of yak milk adulterated with cow milk.

To differentiate among different milk species, it is important to select species-specific antigens. Compared with whey protein, casein has higher heat stability, and β-CN has the highest specific antigenicity among caseins (Anguita et al., 1996). Therefore, we selected bovine β-CN as the target antigen in this study. The objectives of this study were to produce a high-affinity monoclonal antibody (mAb 1–9B), which is specific against bovine β-CN, and to develop a rapid and sensitive indirect competitive ELISA for the detection of cow milk in yak milk. Furthermore, we assessed the effect of different treatments (heat, acidification, and rennet addition) on the performance of the ELISA.
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MATERIALS AND METHODS

Reagents and Materials

Bovine β-CN (purity ≥98%), Freund’s complete and incomplete adjuvant, hypoxanthine-thymidine, hypoxanthine-aminopterin-thymidine selective medium, 3,3′,5,5′-tetramethyl benzidine (TMB) liquid substrate, and Tween-20 were purchased from Sigma Chemical Co. (St. Louis, MO). Polyethylene glycol (PEG) 4000 was obtained from Merck (Darmstadt, Germany), and Dulbecco’s modified Eagle’s medium (DMEM) culture medium was purchased from Invitrogen (Grand Island, NY). Peroxidase-conjugated Affinipure goat anti-mouse IgG (H+L) was obtained from ZSGB-BIO (Beijing, China), and 96-well ELISA and cell culture plates were purchased from Costar (Cambridge, MA). The SP 2/0 myeloma cells were produced in our laboratory. Rennet (Stamix 1150, 1,070 international milk-clotting units/g) was purchased from Chr. Hansen (Hørsholm, Denmark). All other reagents were of analytical pure grade (Beijing Chemical Works, Beijing, China).

Fresh yak milk was obtained from Wushaoling town of the Tianzhu grassland, on the Qinghai-Tibetan Plateau. Fresh cow milk was obtained from Beijing San Yuan Foods Co. Ltd. (Beijing, China). Before analyses, all raw milk samples were first stored at 4°C and then frozen at −20°C within 1 h. Before the experiments, milk samples were skimmed by centrifugation at 3,000 × g for 20 min at 4°C and filtered through glass wool to remove lipids.

Immunization

The antigen solution was prepared by resuspending bovine β-CN in PBS (0.5 mg/mL) and stored at −20°C. The antigen solution was emulsified with an equal volume of Freund’s complete adjuvant and injected (50 μL) both subcutaneously and intraperitoneally into female BALB/c mice (6–8 wk old). Booster injections with Freund’s incomplete adjuvant were performed 3 times with the same dosage at 2-wk intervals. Blood samples were obtained from the tail 7 to 10 d after each booster injection. Antiserum was obtained by centrifugation at 10,000 × g for 3 min at 4°C.

Monitoring Antibody Titers by Indirect ELISA

The collected antiserum was assayed by indirect ELISA for antibody titer: each ELISA well was coated with 100 μL of bovine β-CN diluted in coating buffer (15 mM Na2CO3 and 35 mM NaHCO3, pH 9.6) at 37°C for 2 h. Following the 2-h incubation, the wells were washed 3 times with PBS plus 0.05% Tween 20 (150 μL/well; PBST) for 2 min on a plate shaker and coated with 100 μL of 5% defatted milk for 1 h at 37°C. Following 3 successive washes with PBST, 100 μL/well of antiserum diluted in antibody diluents (PBST with 0.1% gelatin) was added to the wells and the plates incubated for 1 h at 37°C. The plates were washed as described above. Following the addition of 100 μL of peroxidase-conjugated Affinipure goat anti-mouse IgG diluted 1:5,000 in antibody diluent, the plates were incubated for 30 min at 37°C and washed with PBST. Then, TMB was added to each well (100 μL/well) and the plates were incubated for 15 min at 37°C. The reaction was stopped with 2 M H2SO4 (50 μL/well). Absorbance was measured by using a microplate reader (model 680, Bio-Rad, Hercules, CA) at 450 nm.

Production of the Monoclonal Antibody

The mouse with the highest antibody immunization titer was injected intraperitoneally with 100 μL of antigen solution without adjuvant 3 d before fusion. The spleen was removed for hybridoma production (Devi et al., 1999). After fusion, hybridoma cells were screened by indirect ELISA for the presence of antibodies against bovine β-CN. Positive hybridoma cells were also tested against casein extracted from yak milk. Then, hybridomas selected to produce specific antibody were subcloned by the limiting dilution method (Köhler and Milstein, 1975). Cells with the highest antibody titer and specificity were selected during each subcloning until a single positive hybridoma was obtained. Subcloning was repeated 5 times to obtain positive monoclonal antibody (mAb)-producing cells. Monoclonal antibody 1–9B was produced using the mouse ascites method (Cho et al., 2005). Animals were killed by cervical dislocation under the anesthesia. All animal maintenance and experimental procedures were conducted in accordance with the principles and specific guidelines presented in Guidelines for the Care and Use of Agricultural Animals in Research and Teaching (FASS, 2010) and approved by the Animal Ethics Committee of China Agricultural University. The resulting antibodies were purified by ammonium sulfate precipitation (Walker, 1996).

Detection of β-CN and Cow Milk and Assessment of Cross-Reactivity

Following optimization of the assay by checkerboard titration, mAb 1–9B was used to detect β-CN and cow milk by indirect competitive ELISA. The inhibitory concentration was calculated as follows: % inhibition = %B/B0, where B and B0 are the...
absorbance in the presence and absence of the competitor, respectively. Fifty percent inhibition was the competitor concentration at which the absorbance was half of B₀; 50% inhibition for each compound was determined based on its inhibition curve.

To assess the specificity of mAb 1–9B, its cross-reactivity (CR) with yak milk was evaluated by indirect competitive ELISA based on comparison of the half-maximal (50%) inhibitory concentrations of yak milk and cow milk. The CR was calculated as follows: CR (%) = (concentration of standard cow milk inhibiting 50% antibody binding)/(concentration of yak milk inhibiting 50% antibody binding) × 100.

The indirect competitive ELISA was performed as follows. Plates were coated at 37°C for 2 h with 100 μL/well of bovine β-CN in coating buffer. The plates were then emptied, washed, and blocked with 100 μL/well of coating buffer plus 5% dried nonfat milk powder for 1 h at 37°C. After the plates were washed 3 times with PBST, bovine β-CN, cow milk, or yak milk, previously diluted in antibody diluents and mixed with an equal volume of ascites antibodies at 37°C for 1 h, was added to the wells (100 μL/well) and incubated at 37°C for 1 h. The plates were then washed 3 times with PBST before being incubated with 100 μL/well of peroxidase-conjugated AffiniPure goat anti-mouse IgG diluted 1:5,000 in antibody diluent for 30 min at 37°C. The plates were washed as described above and then incubated with 100 μL/well of TMB for 15 min at 37°C. The reaction was stopped with 50 μL/well of 2 M H₂SO₄ and the absorbance was measured at 450 nm on a microplate reader.

**Detection of Cow Milk in Yak Milk**

The adulteration standards were prepared in 3 independent batches in which the reference skimmed cow and yak milk samples were mixed at varying percentages: 1, 5, 10, 20, 40, 60, 80, 90, and 100% (vol/vol). The samples were detected in triplicate by the optimized indirect competitive ELISA as described above. The standard curves were validated by quantifying the adulteration in blind samples.

**Detection of Adulterated Milk Samples with Different Treatments**

**Heat Treatment.** Cow milk and adulterated yak milk (containing 1–100% cow milk) were subjected to different heat treatments, including pasteurization (65°C × 10 min), heat treatment at 95°C for 10 min, UHT treatment (138°C × 4 s), or spray drying (150 to 160°C × 10–20 s). Fresh (untreated) and heat-treated samples were analyzed in triplicate by indirect competitive ELISA.

**Acidification Treatment.** The adulteration standards were prepared as described above. Each adulterated yak milk sample (containing 1–100% cow milk) was acidified to pH 6.3, 5.8, 5.0, or 4.6 with 0.5 M HCl. Four different pH and raw milk samples were prepared and analyzed in triplicate by indirect competitive ELISA.

**Rennet Treatment.** The adulteration standards were prepared as described above. Rennet solution, which consisted of rennet and sterile water (1:40, wt/wt), was added (60 μL) to each adulterated yak milk sample (30 g), mixed in a vortex for 30 s, and incubated in a water bath at 32°C for 1 h. The coagulum formed was cut and rested for 5 min. The expelled whey was centrifuged at 2,500 × g for 15 min at room temperature and decanted, and the remaining curd was obtained and readjusted with PBS to the natural concentration of milk (the concentration of casein was about 28 g/L) and homogenized in a mechanical blender (Haza et al., 1999). The samples were prepared in triplicate and then analyzed by indirect competitive ELISA. The coagulation test was run in triplicate. The coagulated samples were re-adjusted to the natural concentration of milk with the addition of PBS, homogenized in a mechanical blender (Haza et al., 1999), and analyzed by indirect competitive ELISA.

**RESULTS AND DISCUSSION**

**Optimization of ELISA**

Highly specific mAb 1–9B against bovine β-CN was successfully produced by monoclonal antibody technology. An indirect competitive ELISA of high sensitivity and specificity was designed to detect the presence of cow milk in yak milk using mAb 1–9B. The optimum plate-coating concentration and primary antibody dilution were determined by checkerboard titration. As shown in Figure 1, absorbance remained high at a concentration of 1/128,000, indicating high sensitivity of the monoclonal anti-bovine β-CN antibody. Based on the results, the optimum plate coating concentration was 5 μg/mL and the optimum primary antibody (mAb 1–9B) dilution was 1:2,000.

**Detection Limit of Bovine β-CN and Cow Milk**

The ability of mAb 1–9B to detect bovine β-CN was assessed by indirect competitive ELISA. The working concentration of bovine β-CN detection ranged from 10 to 8,000 μg/mL (Figure 2a). The minimum detection
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Limit and 50% inhibition for β-casein were 10 and 300 μg/mL, respectively. The linear correlation coefficient was −0.99.

As shown in Figure 2b, cow milk at different dilutions was detected. The linear range was from 1:1,300 to 1:2 diluted cow milk. The minimum detection limit and 50% inhibition for cow milk were 1:1,300 and 1:50 dilution, respectively. The linear correlation coefficient was −0.99. The content of β-CN is approximately 36.9% in cow milk proteins (Braga et al., 2006; De Marchi et al., 2010). The linear range of cow milk (i.e., 1:1,300 to 1:2 dilution) was equal to 9 to 6,000 μg/mL β-CN. This observation was similar to the detecting range of β-CN described above, suggesting that the detection of β-CN using mAb 1–9B was not affected by other cow milk proteins.

ELISA Specificity

The CR of mAb 1–9B with yak milk was determined by indirect competitive ELISA. As illustrated in Figure 3, the results revealed that 1/50 and 1/0.42 dilutions of cow milk and yak milk resulted in 50% inhibition, respectively. The CR of yak milk was <1.0% against cow milk. According to Yin et al. (2010), our results indicate that mAb 1–9B has relatively low CR toward yak milk, and our assay can be considered to be highly specific for bovine β-CN. Therefore, mAb 1–9B can be used to detect the adulteration of yak milk with cow milk.

Detection of Cow Milk in Yak Milk

The inhibition curve of adulterated yak milk is shown in Figure 4. The linear range of detectable adulterant concentration was 1 to 80% (vol/vol) with a linear correlation coefficient of −0.98; the minimum detection limit of the assay was 1% (vol/vol). The assay was highly reproducible with intra-and interassay coefficients of variation <10% (Supplementary Tables S1 and S2; http://dx.doi.org/10.3168/jds.2014-8127). The standard curve was applied in a single blind trial. Based on the trial, the presence of cow milk could be estimated in the range from 1 to 80%; we detected no significant differences between known and estimated values (P = 0.18). The mAb AH4 anti-bovine β-CN was produced for the detection of cow milk in ewe and goat milks with a working range of 1 to 10% (Anguita et al., 1995). Song et al. (2011), who used modified polyclonal antibodies against bovine β-CN to detect the presence of bovine milk in Shaanxi goat milk, reported that the working range was 2 to 50%. Compared with their findings, our monoclonal antibody had a wider working range and applicability for the detection of adulterated yak milk with cow milk. Furthermore, at present, the reference method can detect 1% cow milk in sheep milk (European Commission, 2001). Because adulteration of less than 1% would not be economically profitable, the method described here is adequate for quantitative testing. Therefore, this assay can be adequately applied in routine adulteration testing.

Detection of Adulterated Milk Samples with Different Treatments

Heat Treatment. Monoclonal antibody 1–9B was used to detect different heat-treated cow milk samples. The inhibition curves of pasteurized, UHT, and spray-dried samples were similar to those of fresh (untreated) samples (Supplementary Figure S1; http://dx.doi.org/10.3168/jds.2014-8127), indicating that heat treatment did not interfere with the determination of bovine β-CN in the assay. The selection of target antigen is crucial in adulteration testing. Hurley et al. (2004a) successfully used a monoclonal antibody specific
against bovine IgG to detect the presence of fresh and pasteurized cow milk. However, their assay could not be used to detect the presence of UHT or spray-dried cow milk, indicating that whey proteins are sensitive to heat, and severe heat treatment can cause denaturation of the whey protein. Casein has higher heat stability than whey. Our results revealed that the antigenic determinant of bovine β-CN was not affected by the different heat treatments. Therefore, the antibody against bovine β-CN is suitable for detecting cow milk that has undergone different heat treatments.

Because heat treatment did not affect the detection of cow milk in this study, the assay was then used to detect the presence of cow milk in adulterated yak milk (1–100%) subjected to different heat treatments. The assay generated similar inhibition curves for the different heat treatments with a working range of 1 to 80% (Supplementary Figure S1; http://dx.doi.org/10.3168/jds.2014-8127). Thus, our assay can be used to detect

Figure 2. Standard curves of monoclonal antibody 1–9B for detecting bovine β-CN (a) and cow milk (b) by indirect competitive ELISA under optimum conditions. Data points represent mean ± standard deviation (n = 3).

Figure 3. Cross-reactivity of monoclonal antibody 1–9B with yak milk by indirect competitive ELISA under optimum conditions. Cow milk was used as a reference. Data points represent mean ± standard deviation (n = 3).
heat-treated adulterated samples. Both UHT-treated yak milk and yak milk powder have a long shelf life and thus are more popular than fresh yak milk in local markets. Therefore, our assay is adequate for the detection of adulterants in these products and will help in the detection of adulterated yak milk.

**Acidification Treatment.** To assess whether our ELISA could be used in yogurt and other acidulated dairy products, milk samples acidified to pH 6.3, 5.8, 5.0, and 4.6 were analyzed. The inhibition curves of acidified samples were similar to those of fresh (untreated) samples (Supplementary Figure S1; http://dx.doi.org/10.3168/jds.2014-8127). During acidification, as colloidal calcium phosphate is released from the casein micelles, β-CN dissociates and aggregates with further acidification (Lucey, 2002; Dalgleish and Corredig, 2012). Our results imply that this process did not denature the target epitope. Therefore, acidification did not affect the linear range of ELISA. Our developed method could be applied to detect the adulteration of yak milk yogurt with cow milk.

**Rennet Treatment.** To assess whether our developed method could be used in cheese, ELISA was used in rennet-treated adulterated milk samples (ranging from 1 to 100%). We found no significant differences between the standard curves of raw and rennet-treated milk samples (Supplementary Figure S1; http://dx.doi.org/10.3168/jds.2014-8127). Today, yak cheese has an increasing number of market prospects (Or-Rashid et al., 2008). Combined with the results obtained from the acidified samples, this assay could be used to detect the adulteration of yak cheese.

**CONCLUSIONS**

We developed an optimized indirect competitive ELISA to detect the adulteration of yak milk with cow milk using a monoclonal antibody specific for bovine β-CN (mAb 1–9B). The minimum detection limit of this assay was 10 μg/mL of bovine β-CN and the linear range for detecting adulteration was 1 to 80%, which is sufficiently broad for routine adulteration testing. The ELISA method was not affected by heat, acid, or rennet treatment of milk; therefore, the assay could be applied to detecting adulteration with cow milk in different yak dairy products.

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**REFERENCES**


Figure 4. Standard curves of indirect competitive ELISA for the detection of cow milk in yak milk under optimum conditions. Data points represent mean ± standard deviation (n = 3).