Artificial insemination (AI) is the most commonly used biotechnology for the genetic enhancement of cryopreserved sperm (Philpott, 1993). Cryopreservation is a technique that uses liquid nitrogen as a cold source to pre-

Assessment of cryopreserved sperm functions of Korean native brindled cattle (Chikso) from different region research centers of Korea

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ABSTRACT Sperm cryopreservation is an important method of assisted reproductive techniques and storing genetic resources. It plays a vital role in genetic improvement, livestock industrial preservation of endangered species, and clinical practice. Consequently, the cryopreservation technique is well organized through various studies, especially on Korean native cattle (Hanwoo). However, the cryopreservation technique of Korean native brindled cattle, which is one of the native cattle species in Korea, is not well organized. Therefore, it is necessary to develop a Supplementary Table technique for the cryopreservation of Korean native brindled cattle. For this purpose, it is important to first evaluate the quality of the currently produced cryopreserved sperm of Korean native brindled cattle. In this study, we randomly selected 72 individual Korean native brindled cattle semen samples collected from 8 different region research centers and used them to evaluate sperm functions. We focused on the quality evaluation of cryopreserved Korean native brindled cattle semen following the measurement of motion kinematics, capacitation status, intracellular ATP level, sperm motility, and cell viability. Then, the values of each of the eight groups were derived from various sperm parameters of nine individual samples, including sperm motility, kinematics, cellular motility, and intracellular ATP levels, which were used to compare and evaluate sperm function. Overall, differences in various sperm parameters were observed between most of the research centers. Particularly, the deviations of motility and motion kinematics were high according to the sample. Therefore, we suggest that it is necessary to develop a standard method for the cryopreservation of Korean native brindled cattle semen. We also suggest the need for sperm quality evaluation of the cryopreserved semen of Korean native brindled cattle before using artificial insemination to attain a high fertility rate.

Keywords: cryopreservation, Korean native brindled cattle, sperm functions, spermatozoa

INTRODUCTION

Artificial insemination (AI) is the most commonly used biotechnology for the genetic enhancement of cryopreserved sperm (Philpott, 1993). Cryopreservation is a technique that uses liquid nitrogen as a cold source to pre-

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serve cells for long periods of time (Karlsson and Toner, 1996). Sperm cryopreservation has aided the advancement of reproductive technologies, such as AI and in vitro fertilization (IVF). It is also used in a variety of clinical and research settings, including assisted reproductive technology, biological storage, and genetic improvement (Zavos, 1992; Curry, 1995). Cryopreservation technology solves the challenge of long-term preservation of semen and renders semen utilization unrestricted by time, geography, and the life of breeding animals. However, the genetic influence of AI is limited by the quality of the sperm (Yoon et al., 2015). A rapid, accurate, and objective evaluation of the functional state of frozen-thawed sperm is beneficial in the process of frozen semen research to facilitate the conservation of genetic resources of good breeding bulls and the establishment of gene pools, allowing complete utilization of the excellent genetic resources (Sansone et al., 2000). In contrast, it can accelerate the progression of breed improvement, improve breeding efficiency, and then maximize its optimal production performance and improve the economic benefits of animal husbandry (Andrabi and Maxwell, 2007). Cryopreservation of sperm from agriculturally and genetically important species would protect them from extinction.

Research conducted till date on semen cryopreservation in Hanwoo has been more extensive, and commercial frozen semen production is also more common. The technique of cryopreservation is well developed, the recovery rate is also good, and several studies also have been performed. Cryopreserved Hanwoo semen samples are well controlled and managed by a specific organization. Due to the limitation of genetic resources, freezing sperm obtained from Korean native brindled cattle would be a better method to conserve genetic resources. However, there is only limited experimental information on semen evaluation from Korean native brindled cattle in previous studies. Several research centers from each province of Korea are producing Korean native brindled cattle cryopreserved semen, and the methods of cryopreservation are different between each research center. A comprehensive management of the Korean native brindled cattle cryopreserved semen has not yet been implemented. Currently, a specific cryopreservation method has not been established. Therefore, quality evaluation of cryopreserved semen of the Korean native brindled cattle is necessary.

In this study, we randomly selected 72 individual Korean native brindled cattle samples from 8 different region research centers of Korea and used them for evaluating sperm functions. We focused on the quality evaluation of the cryopreserved Korean native brindled cattle semen following the measurement of capacitation status, motion kinematics, intracellular ATP level, sperm motility, and cell viability. Then, the values of each of the eight groups were derived from various sperm parameters of nine individual samples, including sperm motility, kinematics, cellular motility, and intracellular ATP levels, which were used to compare and evaluate the sperm functions.

MATERIALS AND METHODS

Sample collection
Semen samples were obtained from 72 individual Korean native brindled cattle from 8 different region research centers, stored on liquid nitrogen, and transferred to the laboratory.

Sample preparation
Sperm preparation was performed using the routine Percoll washing method as described previously (Lee et al., 2009; Brugnon et al., 2013). Frozen sperm samples were thawed for 20 s in a water bath at 39°C. To remove extender debris, seminal plasma, cryoprotectant and diluent added during sperm cryopreservation, the samples were centrifuged for 20 minutes at 400 × g with a discontinuous Percoll density gradient of 1 mL 90% to 1 mL 45% Percoll. The top layer of the suspension was discarded after centrifugation. The sperm pellet was washed by centrifugation at 400 × g with 1 mL Tyrode’s albumin lactate pyruvate (TALP) medium for 5 min. The supernatant was extracted, after which 500 L of TALP medium was added. The TALP medium contained 1.0 mM sodium pyruvate, 100 mM NaCl, 0.4 mM MgCl·6H₂O, 3.1 mM KCl, 0.3 mM Na₂HPO₄·12H₂O, 2.0 mM CaCl·2H₂O, 25 mM NaHCO₃, 21.6 mM sodium lactate, and 0.6% bovine serum albumin.

Computer-assisted sperm analysis
Sperm motility and motion kinematics were evaluated using a computer-assisted sperm analysis (CASA) program (FSA2016, Medical supply, Seoul, Korea) with a CMOS image sensor and a 2048 × 1536 (300 M pixels), 60-frame camera (Medical supply, Seoul, Korea), and an OLYMPUS BX43 phase-contrast microscope (Olympus, Tokyo, Ja-
pan) with a 10 × objective phase-contrast mode. A 10-μL smear of the sample was evaluated in a preheated (37°C) Makler counting chamber (Sefi-Medical Instruments, Haifa, Israel). The movement of at least 250 sperm cells was recorded for each sample from random fields (> 5). The obtained images were analyzed to determine sperm motility (%), progressive sperm motility (%), rapid sperm motility (%), medium sperm motility (%), slow sperm motility (%), curvilinear velocity (VCL, μm/s), straight line velocity (VSL, μm/s), average path velocity (VAP, μm/s), beat cross frequency (BCF, Hz), and amplitude of lateral head displacement (ALH, μm).

**Hoechst 33258 (H33258)/chlortetracycline fluorescence**

The H33258/CTC dual-staining method (combined Hoechst 33258/chlortetracycline fluorescence (CTC) assessment) was used to evaluate capacitation status. Samples were centrifuged at 400 × g for 5 min. After removing the majority of the supernatant, 135 μL of DPBS and 15 μL of H33258 solution (10 g H33258/mL DPBS) were added to the remaining sample. After incubation for 3 min at room temperature (RT), 250 μL of 2% (w/v) polyvinylpyrrolidone in DPBS was added to the mixture to remove excess dye by layering. The sample was then washed by centrifugation at 400 × g for 5 min. After the complete extraction of the supernatant, the pellet was resuspended in 50 μL of CTC solution (750 mM CTC in 5 μL buffer: 20 mM Tris, 5 mM cysteine, 130 mM NaCl, pH 7.4) and in 50 μL DPBS. The samples were then smeared onto slides in 10-μL increments. For each sample, at least 400 spermatozoa were counted on each slide. For H33258 and CTC assessment, an OLYMPUS BX43 with epifluorescence illumination and ultraviolet BP 340-380/LP 425 and BP 450-490/LP 515 excitation/emission filters were used (Olympus, Tokyo, Japan). Finally, the following four capacitation patterns were observed: live acrosome-reacted sperm (AR pattern, no fluorescence over the head, or green fluorescence only in the postacrosomal region), live capacitated sperm (B pattern, the acrosomal region showed bright green fluorescence and the region showed a dark post-acrosomal), sperm that is not incapacitated and is alive (F pattern, bright green fluorescence that is evenly spread across the sperm head), and dead sperm (D pattern, blue fluorescence distributed uniformly over the sperm head).

**Intracellular ATP level**

Intracellular ATP generation was quantified using an ATP assay kit (ab83355, Abcam, Cambridge, UK) as instructed by the manufacturer. Briefly, Samples (1 × 10⁶ cells) were centrifuged at 10,000 × g for 5 min, and removed the majority of the supernatants. The sample was then washed by 1 × PBS (4°C). The washed samples was added to 96-well plates. Before the experiment, the ATP reaction mixture was primed and equilibrated to RT. The reaction mixture was then added in equal volumes to each well and incubated at RT for 30 min shielded from light. Finally, luminescence was measured using a microplate reader at OD 570 nm (Gemini Em: Molecular Devices Corporation, Sunnyvale, CA, USA) and analyzed using the SoftMax Pro 7 software (Molecular Devices Corporation, Sunnyvale, CA, USA).

**Sperm viability**

The Cell Assay Kit was used to evaluate cell viability (ab112118, Abcam, Cambridge, UK). This kit uses a patented water-soluble dye that adjusts its absorption spectra in response to cellular reduction. The variation in absorption ratio is proportional to the number of live cells. The assay fluid was thawed and preheated to 37°C before the tests. Then, 100 μL of spermatozoa and 20 μL of assay solution were added to a 96-well plate. The sample was then incubated for 2 h at 37°C in 5% CO₂ in a light-protected incubator. Cell viability behavior, as well as absorbance at OD 570 and 605 nm, was measured (Gemini Em: Molecular Devices Corporation, Sunnyvale, CA, USA). The ratio of OD 570 nm to OD 605 nm was used to evaluate cell viability (SoftMax Pro 7; Molecular Devices Corporation, Sunnyvale, CA, USA).

**Statistical analysis**

The samples were divided into eight categories according to their source. In each category, numerical values were derived from various sperm parameters (such as motility, motion kinematics, capacitation state, viability, and ATP level) in nine individual samples. Data were interpreted using SPSS (Version 25.0, IBM, Armonk, NY, USA). One-way ANOVA was used to compare the various parameters of eight groups. Data were expressed as mean ± SEM. p values of < 0.05 were considered to be statistically significant.
RESULTS

Sperm motion parameters

The CASA program was used to monitor spermatozoa motility and motion kinematics after Percoll washing. Results showed that the deviation of motility (%) and motion kinematics was high according to the sample ($p < 0.05$; Supplementary Table 1 and Fig. 1). No differences were observed between samples collected from the eight different region research centers in terms of medium motility and slow motility. The other eight parameters (sperm motility, progressive motility, rapid motility, VCL, VSL, VAP, BCF, and ALH) of samples collected from the G research center showed significantly higher values than those of samples collected from other research centers. Sperm motility and progressive sperm motility of samples from the F research center were significantly lower than those of samples from other research centers. VCL, VSL, VAP, BCF, and ALH of samples from B, E, and F research centers were significantly lower than those of samples from other research centers ($p < 0.05$; Supplementary Table 2 and Fig. 2).

Assessment of sperm capacitation status

CTC/H33258 dual staining was used to determine sperm capacitation status. Results showed that the proportions of AR, B, and F patterns were $0.72 \pm 0.59\%$, $11.25 \pm 5.13\%$, and $88.03 \pm 5.32\%$, respectively ($p < 0.05$; Fig. 3). The deviation of capacitation status was high according to the sample. The proportion of AR-pattern spermatozoa was the lowest ($0.03 \pm 0.10\%$) in samples collected from the F research center, whereas it was the highest ($1.45 \pm 0.27\%$) in samples collected from the G research center ($p < 0.05$; Fig. 4). The proportion of B-pattern sperma-

![Fig. 1. Measurement of sperm motility of 72 individual Korean native brindled cattle after Percoll washing. (A) Sperm motility (%). (B) Progressive sperm motility (%). (C) Rapid sperm motility (%). (D) Medium sperm motility (%). (E) Slow sperm motility (%). (F) Curvilinear velocity (VCL, $\mu m/s$). (G) Straight line velocity (VSL, $\mu m/s$). (H) Average path velocity (VAP, $\mu m/s$). (I) Beat cross frequency (BCF, Hz). (J) Amplitude of lateral head displacement (ALH, $\mu m$).]
Fig. 2. Measurement of sperm motility of samples collected from eight different research centers. (A) Sperm motility (%). (B) Progressive sperm motility (%). (C) Rapid sperm motility (%). (D) Medium sperm motility (%). (E) Slow sperm motility (%). (F) Curvilinear velocity (VCL, μm/s). (G) Straight line velocity (VSL, μm/s). (H) Average path velocity (VAP, μm/s). (I) Beat cross frequency (BCF, Hz). (J) Amplitude of lateral head displacement (ALH, μm). Sperm motility and motion kinematics are presented as the mean ± SE. Superscripts a, b, c, d and e indicate significant differences between sperm parameters by one-way ANOVA of variance (p < 0.05, n = 9).

Fig. 3. Assessment of sperm capacitation status of 72 individual Korean native brindled cattle. (A) Patterns of live acrosome-reacted (AR pattern). (B) Patterns of live capacitated (B pattern). (C) Patterns of live noncapacitated (F pattern).
Spermatozoa was the lowest (4.55 ± 1.44%) in samples collected from the G research center, whereas it was the highest (15.13 ± 2.82%) in samples collected from the D research center (p < 0.05; Fig. 4). Spermatozoa exhibiting the F pattern were detected in the highest proportion (95.35 ± 1.44%) at the B research center, whereas it was the lowest (83.79 ± 2.69%) at the E research center (p < 0.05; Fig. 4).

**ATP generation in spermatozoa**

Quantitative ATP measurements were conducted using an ATP assay kit. The deviation of ATP measurements was high according to the sample (p < 0.05; Fig. 5). No differences were detected between samples collected from C, D, E, and H research centers. The intracellular ATP levels of samples collected from G (0.083 ± 0.0005) and A (0.0460 ± 0.0005) research centers were significantly higher than those of samples collected from other research centers. The intracellular ATP levels of samples collected from F (0.042 ± 0.0005) and B (0.044 ± 0.0004) research centers were significantly lower than those of samples collected from other research centers (p < 0.05; Fig. 6).

**Cell viability**

Monitoring cell viability is one of the most important activities for comparing sperm functions. Results showed that the deviation of sperm cell viability was high according to the sample (p < 0.05; Fig. 7). There were no differences between samples collected from C, D, and H research centers. However, significant differences were observed between samples collected from A, B, E, F, and G research centers. The proportions of the cell viability of samples collected from G (0.92 ± 0.005) and A (0.89 ± 0.004) research centers were significantly higher than those of samples collected from other research centers. The proportions of the cell viability of samples collected from F (0.87 ± 0.005) and B (0.84 ± 0.004) research centers were significantly lower than those of samples collected from other research centers (p < 0.05; Fig. 6).
from E (0.78 ± 0.003), B (0.77 ± 0.003), and F (0.76 ± 0.003) research centers were significantly lower than those of samples collected from other research centers (p < 0.05; Fig. 8).

**DISCUSSION**

Currently, assessment of semen quality contributes substantially to the AI industry around the world (Larsson and Rodriguez-Martínez, 2000). In the bovine industry, > 70% of cows are used for reproduction using frozen-thawed sperm obtained from superior genetic bulls through AI, but only 50% of full-term pregnancy complications are confirmed to be effective (Kastelic and Thundathil, 2008). As the quality of semen contributes significantly to subsequent fertility, it is essential to evaluate the quality of semen (Bonde, 1998). The research performed the more exact evaluation on bull semen quality certifies the breeding effects and the successful calvings. However, in previous research, only the limited data on Korean Native Brindled Cattle could be found. Therefore, we performed the frozen semen evaluation to provide an effective and widely applicable system for the assessment of Korean native brindled cattle semen as a foundation for improving semen evaluation standards.

Mammalian spermatozoa cannot fertilize oocytes before functional maturation. Maturity includes exercise, over-activation, capacitation, and AR (Suarez, 2008; Rahman et al., 2014). After ejaculation, spermatozoa in mammals gain motility. As only overactivated motile sperm can fertilize oocytes, sperm motility, and overactivation are considered as important factors for a successful pregnancy (Park, 2012; Rahman et al., 2013). The ratio of motile sperm is the important parameter on the semen evaluation. Therefore, it has been widely used in commercial and laboratory settings to evaluate sperm quality. Straightness, progressive trajectories, curvilinearity, and amplitude of lateral displacement were found in the sperm motility patterns, which are Supplementary Table for penetration into the cervical mucus of the female genital tract in vivo (David et al., 1981; Serres et al., 1984). The CASA system was used to evaluate motility and other kinematic parameters in this study; the analysis have been extensively used to determine the quality of sperm in animal species. The kinematic values determined for each spermatozoon include the sperm head’s velocity of moves, the width of the sperm head’s trajectory, and the frequency of the sperm head’s directional movement (David et al., 1981). Our results showed that the sperm motility, progressive motility, rapid motility, VCL, VSL, VAP, BCF, and ALH of samples collected from the G research center were significantly higher than those of samples collected from other research centers (p < 0.05; Supplementary Table 2 and Fig. 2). Sperm motility and progressive sperm motility of samples collected from the F research center were significantly lower than those of samples collected from other research centers. VCL, VSL, VAP, BCF, and ALH of samples collected from B, E, and F research centers were significantly lower than those of samples collected from other research centers. Under both *in vitro* and *in vivo* conditions, effective fertilization requires completion of spermatozoa capacita-
tion at the appropriate time. In fact, fertilization is a complicated series of events involving several regulatory steps. According to a review of the literature, no specific and direct methods exist for determining the fertilization capacity of spermatozoa, and detecting the fertilization characteristics of most of the spermatozoa is quite difficult (Gillan et al., 2005). In this regard, determining the spermatozoa’s capacitation status is believed to be an important indicator of the quality of the sperm. The status of spermatozoa capacitation can be determined directly by monitoring calcium regulation changes with chlorotetracycline, a fluorescent antibiotic. The highly fluorescent Ca\(^{2+}\) complexes bind to the cell membrane, resulting in four different spermatozoa staining patterns (Zaneveld et al., 1991; Fraser and McDermott, 1992). The acrosome reaction occurs during fertilization, and the sperm loses the integrity of the head portion of the membrane that promotes sperm-zona binding after capacitation (Luconi et al., 1996; Kirichok et al., 2006). Our results showed that samples collected from the F research center had the lowest proportion of AR-pattern spermatozoa, whereas those collected from the G research center had the highest proportion. The proportion of B-pattern spermatozoa was the lowest in samples collected from the E research center, whereas it was the highest in samples collected from the B research center. Regarding the proportion of F-pattern spermatozoa, it was the lowest at the B research center but the highest at the E research center (p < 0.05; Fig. 4).

Mammalian sperm must gradually move in a certain period of time to fertilize the oocyte (Mortimer, 1990). As the ejaculated spermatozoa demand substantial amounts of energy to facilitate in vivo transport through the female reproductive tract and for both in vitro and in vivo penetration of the oocyte zona pellucida (Mortimer, 1990; Mukai and Okuno, 2004), it is important to note that sperm motility depends primarily on the availability of intracellular ATP. Depletion of ATP during capacitation is attributed to the loss of ATP required to maintain sperm motility and activate cAMP (Wishart and Palmer, 1986). It has been suggested that the energy required to regulate sperm motility is provided by ATP, and ATP is primarily produced by mitochondrial respiration. Therefore, if mitochondrial respiration cannot synthesize ATP, it may reduce sperm motility. Mitochondria play a central role in energy metabolism and regulate sperm motility, AR, and sperm oocyte fusion (Mukai and Okuno, 2004). Our results demonstrated that the intracellular ATP levels of samples collected from G and A research centers were significantly higher than those of samples collected from other research centers (p < 0.05; Fig. 6).

Sperm viability is the key factor for determining semen quality; low sperm viability can directly affect male fertility (Tsujii et al., 2002; Kaur and Bansal 2004). In our study, the cell viability of samples collected from G and A research centers was significantly higher than that of samples collected from other research centers. However, the cell viability of samples collected from E, B, and F research centers was significantly lower than that of samples collected from other research centers (p < 0.05; Fig. 8).

The deviations of sperm motility (%) and motion kinematics were high according to the semen sample. In addition, differences in various sperm parameters were observed between most of the research centers. It may be judged methods for cryopreservation of Korea native brindled cattle are different in each research center because a Supplementary Table standard protocol for cryopreservation of Korea native brindled cattle does not exist. Therefore, we suggest that it is necessary to develop a standard method for the cryopreservation of Korean native brindled cattle semen. We also suggest that it is important to evaluate the sperm quality of the cryopreserved semen of Korean native brindled cattle before performing AI to attain a high fertility rate. Furthermore, these findings may provide a useful model for the development of semen cryopreservation strategies in other endangered species.

CONFLICTS OF INTEREST

No potential conflict of interest relevant to this article was reported.

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SUPPLEMENTARY MATERIALS

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