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Impact of Bovine Oocyte Maturation Media on Oocyte Transcript Levels, Blastocyst Development, Cell Number, and Apoptosis

Andrew J. Watson,2,4,5 Paul De Sousa,3,4,5 Anita Caveney,4,5 Lisa C. Barcroft,4,5 David Natale,4,5 Jennifer Urquhart,4,5 and Mark E. Westhusin6

INTRODUCTION

Research investigating oocyte maturation, fertilization, and embryonic development is necessary for improved assisted-reproduction technologies in animals and humans and to reveal the causes of abnormal embryonic development [1–4]. Broad variation exists in development frequency in vitro for early embryos from different mammalian species. Effective defined media (most notably CZB and KSOMaa) were first developed for the murine early embryo culture atmosphere [15–18]. These studies have reported influences of amino acid supplementation and culture atmosphere on developmental frequencies and embryo metabolism [18–25]. Studies have not, however, examined influences of serum-free oocyte maturation on oocyte transcript levels or blastocyst cell number and apoptosis. Cell number and apoptosis levels are important parameters that are emerging as useful indicators of embryo development and health [26–28]. The objectives of the present study were 1) to investigate the effects of oocyte maturation in serum-free and amino acid-supplemented defined media on oocyte transcript levels, blastocyst cell number, and apoptosis; 2) to investigate the influence of epidermal growth factor (EGF) during oocyte maturation on blastocyst cell number and apoptosis; and 3) to examine the influence of epidermal growth factor (EGF) during oocyte maturation on blastocyst cell number and apoptosis. The results demonstrate that blastocysts derived from in vitro maturation, fertilization, and embryo culture protocols undergo apoptosis but that apoptotic levels are not greatly influenced by the oocyte maturation environment. Amino acid supplementation of oocyte maturation media was associated with enhanced developmental frequencies, increased blastocyst cell number, and elevated oocyte maternal mRNA levels. Oocyte maturation with supplemented synthetic oviduct fluid medium (cSOFMaa) resulted in blastocyst cell numbers comparable to those observed with Tissue Culture Medium 199 + newborn calf serum. Blastocyst development was reduced following oocyte maturation under a 5% CO₂, 7% O₂, 88% N₂ culture atmosphere. EGF supplementation of oocyte maturation medium resulted in a concentration-dependent increase in blastocyst development but did not influence blastocyst total cell number or apoptosis. Our findings indicate that cSOFMaa medium is an effective base medium for bovine oocyte maturation.

MATERIALS AND METHODS

Oocyte Collection, Insemination, and Embryo Culture

COCs were collected by razor blade slashing of slaughterhouse ovaries within 4 h of removal from the animal [13,29]. The COCs were collected in oocyte collection medium (Hepes-buffered TCM199 + 2% v:v newborn calf serum [NCS]; Gibco BRL, Burlington, ON, Canada) and then were washed 4 times in serum-free medium prior to placement in oocyte maturation medium. Only denuded oocytes were discarded, and a COC selection strategy was not employed in this study. After oocyte maturation (see below for specific experimental conditions), oocytes were insem-
inated in vitro with frozen-thawed bovine semen (Semex Canada, Guelph, ON, Canada) prepared using a “swim-up” method in sperm TL medium (Hepes-buffered modified Tyrode’s solution as described previously [30]). Matured COCs were washed in sperm TL and placed in equilibrated fertilization drops (50 COCs/300-μl drop) composed of bicarbonate-buffered modified Tyrode’s solution under light paraffin oil ([30]; BDH, Toronto, ON, Canada). COCs and sperm (2.25 × 10⁵ motile spermatozoa per drop) were incubated for 18 h at 39°C under 5% CO₂-air atmosphere before removal, with a fine-bore glass pipette, of the cumulus investment including all corona cells. Inseminated oocytes (40–50) were placed into embryo culture consisting initially of 20-μl microdrops of citrate (0.5 mM)-supplemented and polyvinyl alcohol (PVA; 3 mg/ml)-supplemented synthetic oviduct fluid medium (cSOFMaa) [15,16] + single-strength nonessential amino acids (NEA; Sigma-Aldrich Canada Ltd, Oakville, ON, Canada) and single-strength essential amino acids (MEM; Gibco BRL) under paraffin oil in a humidified 5% CO₂, 7% O₂, 88% N₂ culture atmosphere. Two days after initiation of culture, the microdrops were increased in volume by addition of 20 μl of cSOFMaa medium. On Days 5 and 7 of culture, 20 μl of medium was removed from each and replaced with 20 μl of fresh medium; this kept the microdrop volume constant for the remainder of the 8-day culture interval. Cleavage and blastocyst frequencies were assessed on Days 3 and 8 postinsemination, respectively.

**Experimental Design**

Experiments employed a randomized design that allocated equivalent numbers of nonselected COCs to each oocyte maturation treatment group.

In experiment 1, a total of 2950 COCs (40–50 COCs per treatment per replicate) were placed into individual wells of 4-well culture plates containing 0.5 ml of either 1) TCM199 and 10% NCS (T199+NCS); 2) cSOFM +10% NCS (cSOFM+NCS); 3) cSOFM+ single-strength essential (EAA) and nonessential (NEA) amino acids (cSOFMaa); and 4) cSOFM minus amino acids (cSOFM). All media were supplemented with 1 μg/ml FSH (Follitropin; Vetrapharma, London, ON, Canada), 5 μg/ml LH (Vetrapharm), and 1 μg/ml estradiol-17β (Sigma-Aldrich). Fifteen replicate experiments were conducted employing a 5% CO₂-in-air atmosphere at 39°C. After maturation, one tenth (i.e., 4–5 oocytes from each treatment) of the matured oocytes from each replicate were frozen in lysis buffer for RNA analysis (see below). The remaining oocytes in each treatment were inseminated, and zygotes were placed into culture for assessment of developmental frequencies, blastocyst cell number, and apoptosis.

In experiment 2, a total of 1518 COCs were allocated to oocyte maturation treatment groups. COCs (35–50 COCs per treatment per replicate) were placed into individual wells of 4-well culture plates containing 0.5 ml of either 1) cSOFMaa or 2) cSOFM. All treatments were supplemented with 1 μg/ml FSH, 5 μg/ml LH, and 1 μg/ml estradiol-17β. Fourteen replicate experiments were conducted in which half of the COCs (including both + and – amino acid groups) were matured under a 5% CO₂, 7% O₂, 88% N₂ culture atmosphere and the other half under a 5% CO₂-air culture atmosphere. After maturation, oocyte pools were inseminated, and cleavage and frequency of development to the blastocyst stage, blastocyst cell number, and apoptosis were measured.

In experiment 3, a total of 471 COCs were allocated to oocyte maturation treatment groups. COCs (35–40 COCs per treatment per replicate) were placed into individual wells of 4-well culture plates containing 0.5 ml of either 1) cSOFMaa alone or cSOFMaa supplemented with either 1, 10, or 100 ng/ml of murine EGF (Sigma-Aldrich). None of these four treatments were supplemented with gonadotropins or estradiol-17β. Three replicate experiments were conducted under a 5% CO₂-in-air culture atmosphere. For this experiment, influences on cumulus cell expansion were determined along with cleavage, frequency of development to the blastocyst stage, blastocyst cell number, and apoptosis.

**Terminal Deoxynucleotidyl Transferase-Mediated dUTP Nick End-Labeling (TUNEL) and Propidium Iodide Labeling**

Zona pellucida-intact Day 8 (postinsemination) blastocysts were fixed in 1% paraformaldehyde in PBS, pH 7.4, for 1 h at room temperature. The blastocysts were washed twice in 60 mM PIPES/25 mM Hepes buffer, pH 7.4, and stored in the second wash in a sealed 4-well plate at 4°C until ready for labeling. An in situ cell death detection kit using fluorescein-conjugated dUTP and TUNEL (Roche Molecular Biochemicals, Laval, PQ, Canada) was used for labeling apoptotic cells. Blastocysts were washed with PBS and then permeabilized with 0.1% Triton X-100 in 0.1% sodium citrate for 15 min; they were then washed twice with PBS before labeling. Positive control embryos were treated with 50 U/ml RQ1 RNase-free DNase for 20 min at 37°C and then washed twice with PBS prior to labeling. The TUNEL reagent was prepared immediately before use and kept on ice. Blastocysts were placed in 50-μl drops of TUNEL reagent, covered with filtered paraffin oil, in 35-mm Petri dishes and incubated in the dark at 37°C for 60 min in a humidified chamber. The blastocysts were washed three times with 1% BSA in PBS and incubated with 50 μg/ml RNase A in the dark at room temperature for 60 min. Total cell nuclei were labeled with 40 μg/ml propidium iodide (Sigma-Aldrich) in sterile H₂O in the dark at room temperature for 60 min. The embryos were washed twice with 1% BSA in PBS and mounted in fluoroguard antifade mounting reagent (Bio-Rad, Mississauga, ON, Canada).

**Imaging Microscopy and Analysis**

Initially, the blastocysts were viewed with confocal microscopy; however, this proved to be inefficient for accurate cell counting of total nuclei in Day 8 bovine blastocysts. We instead adapted standard epifluorescence microscopy methods applied to flattened blastocysts to generate digitized images that were analyzed with Northern Exposure image analysis software (Empix Imaging, Mississauga, ON, Canada) to quantify cell counts. Digital still images were captured through an integrating Sony (Park Ridge, NJ) XC-75 CCD video camera module. The camera was controlled through a personal computer (P1-400, 128 MB RAM, 8.4 GB HD, Hewlett-Packard 800 series CD-writer; Palo Alto, CA) running Empix Imaging software. Images of the fluorescein-labeled apoptotic cells and the propidium iodide-labeled nuclei were recorded using a Leica Orthoplan (Milton Keynes, Bucks, UK) microscope equipped with a Ploempak epifluorescence module and a full turret of high N.A. Plan-APo fluorite lenses.
Detection of Transcript Levels in Bovine Oocytes

The objective was to investigate the differences in transcript levels, if any, in oocytes matured under four different culture conditions: 1) T99+NCS, 2) cSOFM+NCS, 3) cSOFMaa, or 4) cSOFM. All the media were supplemented with gonadotropins (FSH, LH) and estradiol-17β as described above.

RNA Isolation and Reverse Transcription (RT)

Three replicate polymerase chain reaction (PCR) experiments were conducted for all genes of interest using oocytes collected from all 15 experimental replicates as described for experiment 1. Lysed pools of matured oocytes from each treatment were stored at −70°C until all replicates were ready for simultaneous RNA isolation and RT. Likewise, the PCRs for all three replicates were performed simultaneously in order to minimize interexperimental variation. RNA was isolated from pooled oocytes as described previously [31,32]. Briefly, cumulus-denuded oocytes washed with serum-free handling medium were transferred in a minimum volume (< 1 µl) to a 0.5-mL tube to which 10 µl of lysis buffer (4 M guanidine thiocyanate, 0.1 M Tris, pH 7.4, 1 mM β-mercaptoethanol) was added; they were then frozen at −70°C. In this state, lysed embryos are stable for at least 1 yr. At the time of thawing, 0.1 pg of rabbit globin mRNA (Gibco BRL), in a 1- to 2-µl volume, was added per embryo and mixed by pipetting. For each sample, a 2 × 2-mm square of Hybond-messenger affinity paper (mAP; Amersham International, Buckinghamshire, UK) was cut and pre-wet with 0.5 M NaCl before being soaked in the lysed sample for 2–3 h at room temperature to allow for binding of poly(A)+ RNA. Each unabsorbed lysate was pipetted onto its respective mAP square supported on Whatman 1 filter paper (Whatman International Ltd, Springfield Mill, UK) on parafilm. The mAP squares were then individually transferred into separate 0.5-ml tubes and washed by gentle inversion with 200-µl volumes of 0.5 M NaCl-0.1 M Tris (3 times), 0.5 M NaCl (3 times), and 70% ethanol (2 times). Poly(A)+ RNA was eluted from each mAP square in fresh tubes in 11 µl of sterile H2O containing 0.5 µg oligo(dT)12–18 (Gibco BRL) by incubation at 70°C for 10 min, followed by cooling on ice for 5 min. RT reactions were in a final volume of 20 µl consisting of 50 mM Tris-Cl (pH 8.3), 75 mM KCl, 3 mM MgCl2, 10 mM dithiothreitol, 500 µM dNTPs, and 300 units of Superscript RNase H+ (Gibco BRL) for 90 min at 43°C. Reactions were terminated by 5 min at 95°C and then placed on ice. Reverse-transcribed cDNA was either used directly for PCR or stored at −20°C. As a negative control for RNA isolation and RT, a blank mAP square was carried along with samples during the procedure.

PCR

Experiments were conducted to contrast relative levels of transcripts encoding five marker genes including Na+/K+ ATPase α-1 isoform, copper/zine superoxide dismutase (Cu/Zn SOD), basic fibroblast growth factor (bFGF), and cyclins A and B. Oligodeoxynucleotide primers for Na+/K+ ATPase α-1 were (5’ primer) 5’-ACCTGGTTTGGCGA- TCCGAGACGAC-3’ and (3’ primer) 5’-AGGGAAAGGCA- CAGAACCACCA-3’. These primers amplify a 336-base pair (bp) product from bovine cDNA [33]. Primers for bFGF were (5’ primer) 5’-TACAACCTCAGGGAGGAGG-3’ and (3’ primer) 5’-CAGGCTTACAAGCAGGACATGG-3’.

These primers amplify a 282-bp product from bovine cDNA [33]. Primers for Cu/Zn SOD were (5’ primer) 5’-AAGGCCGTGTCCGTGCTGAA-3’ and (3’ primer) 5’-CAGTGCTCCAACTAGCCTCT-3’. These primers amplify a 246-bp product from bovine cDNA [33]. Primers for cyclin A were (5’ primer) 5’-GTACGAGACTTACATCATGAG-GAC-3’ and (3’ primer) 5’-TCCGTGACGTGTA- GAGTCG-3’. These primers amplify a 585-bp product from bovine cDNA. Primers for cyclin B were (5’ primer) 5’-ATGTTGAGATCAGAACGAGGAG-3’ and (3’ primer) 5’-CCGTGACAGCAGTGAC-3’. These primers amplify a 576-bp product from bovine cDNA. Primers for α-globin were (5’ primer) 5’-GACGCCAAGTGCCGAG-3’ and (3’ primer) 5’-GGAGGACGGAGCTT- GAAAT-3’. These primers amplify a 257-bp product [31].

PCRs were performed in 50 µl of single-strength GeneAmp PCR buffer II (10 mM Tris-HCl, pH 8.3, 50 mM KCl; Perkin-Elmer, Canada Ltd, Mississauga, ON, Canada) containing 200 µM dNTPs (Gibco BRL); 1 unit of AmpliTaq Gold DNA polymerase (Perkin-Elmer); 1 µM (for Na+/K+ ATPase α-1 and α-globin) or 2 µM (for Cu/Zn SOD, bFGF, and cyclins A and B) of each of the appropriate 3’ and 5’ gene-specific primers; 1 mM (for Na+/K+ ATPase α-1 and Cu/ZnSOD), 1.25 mM (α-globin), 2 mM (bFGF), or 2.5 mM (cyclins A and B) MgCl2; and a volume of the RT reaction equivalent to 2 oocytes from the pools of reverse-transcribed oocytes. Reactions were cycled in either a Perkin-Elmer GeneAmp 2400 thermal cycler (bFGF; reactions not overlaid with oil), or a Perkin-Elmer Cetus 480 thermal cycler (Na+/K+ ATPase α-1 and cyclins A and B; reactions overlaid with light liquid paraffin oil [BDH]), or a Thermolyne Amplitron (VWR Canlab, Mississauga, ON, Canada) thermal cycler (Cu/ZnSOD and α-globin; reactions overlaid with light liquid paraffin oil). The basic program for amplification of gene transcripts consisted of a 95°C soak for 10 min followed by a cycle program of 95°C for 1 min, a transcript-specific annealing temperature (60°C, 55°C, 58°C, 49°C, and 55°C for Cu/ZnSOD, bFGF Na+/K+ ATPase α-1, cyclins A and B, and α-globin, respectively) for 30 sec, and 72°C for 1 min. Extension was conducted at 72°C for 10 min. Each transcript was amplified for a number of cycles at which the amplicon was accumulating exponentially (34 for α-globin, 41 for bFGF, 42 for Na+/K+ ATPase α-1, and 45 for Cu/ZnSOD and cyclins A and B). The optimal cycle number for each transcript was established by running a linear cycle series with 2-cycle increments. The identity of each PCR product was confirmed by sequence analysis.

RT-PCR products were visualized by separation on 2% agarose gels in single-strength TAE buffer (40 mM Tris acetate, 1 mM EDTA) run at 100 V for 50 min, with all three replicates run on the same gel. Gels were stained for 30 min at room temperature with 0.5 µg/ml ethidium bromide in single-strength TAE buffer, followed by destaining in deionized water for 5–10 min. The products were imaged using an ImageMaster VDS (Pharmacia Biotech, Baie d’Urfe, PQ, Canada). The fluorescence of ethidium bromide-stained DNA, as determined by quantitation image analysis, is proportional to the amount of DNA in each band and is independent of size and sequence of the DNA. The relative band intensities were determined from images imported into a personal computer using the Image Capture Kit and ImageMaster VDS software (Pharmacia Biotech).

Data Calculations

The absolute integrated optical density (IOD), i.e., the volume of the gel band, was determined for all the ampli-
FIG. 1. Influence of serum and amino acid supplementation of oocyte maturation media on blastocyst development: frequency of cleavage, 6- to 8-cell-stage embryos (over inseminated oocytes [6–8/insem] and cleaved embryos [6–8/clvd]), and blastocysts (over inseminated oocytes [blast/insem], cleaved embryos [blast/clvd], and 6- to 8-cell embryos [blast/6–8]) displayed by oocytes matured in 1) T199 + NCS; 2) c-SOFM + NCS; 3) c-SOFMaa; and 4) c-SOFM supplemented with gonadotropins and estradiol-17β. Cleavage did not vary among treatments. Oocyte maturation in T199 + NCS resulted in a significant enhancement of development to the 6- to 8-cell and blastocyst stages. However, the proportion of 6- to 8-cell embryos that progressed to the blastocyst stage did not vary between the T199 + NCS and c-SOFMaa treatments. Values with different superscript letters are significantly different (P < 0.05).

FIG. 2. Influence of culture atmosphere and amino acids during oocyte maturation on blastocyst development displayed by oocytes matured in c-SOFMaa or c-SOFM supplemented with gonadotropins and estradiol-17β. Half of the COCs (including both + and − amino acid groups) were matured under a 5% CO₂, 7% O₂, 88% N₂ culture atmosphere and half under a 5% CO₂-in-air culture atmosphere. See Figure 1 legend for a full description of the groups. Cleavage was significantly greater in the 5% CO₂-in-air culture atmosphere and was not influenced by amino acid supplementation. Development to the 6- to 8-cell and blastocyst stages was significantly enhanced in the amino acid-supplemented 5% CO₂-in-air culture atmosphere. The proportion of 6- to 8-cell-stage embryos progressing to the blastocyst stage did not vary significantly among treatments. Values with different superscript letters are significantly different (P < 0.05).

cons including the 400-bp band of a 100-bp DNA ladder (5 μg; Gibco BRL) run on each gel. The IOD value for the 400-bp band on the gel for each specific transcript was equated with the IOD value for the 400-bp band on the α-globin gel to establish a conversion factor. This was used to calculate the IOD values for each culture group in the three replicates relative to the exogenous α-globin standard. Abundance ratios of the IODs for the specific gene transcripts relative to the comparable IODs for the α-globin were then determined.

Statistical Analysis

Data for all experiments were analyzed using the SigmaStat (Jandel Scientific, San Rafael, CA) software package. One-way ANOVA, followed by pair-wise multiple comparisons (Bonferroni’s method), was used for analysis of differences in the means for two or more populations. Differences of P ≤ 0.05 were considered significant.

RESULTS

Experiment 1

The first experiment was an examination of the influences of serum, amino acids, and culture media (TCM199 vs. c-SOFM), employed for bovine oocyte maturation, on development to the blastocyst stage. No significant differences in cleavage were observed among the four oocyte maturation treatment groups (P < 0.05; Fig. 1). Development to the 6- to 8-cell stage was significantly greater (P < 0.05) in the T199 + NCS treatment, with no difference in this parameter among the remaining three treatments (Fig. 1). Blastocyst development was significantly greater (P < 0.05) in the T199 + NCS oocyte maturation treatment group than for the other treatments (Fig. 1). Blastocyst formation expressed over the proportion of cleaved embryos (blast/clvd), however, was significantly greater (P < 0.05) in the c-SOFMaa oocyte maturation treatment than for the c-SOFM maturation treatment (Fig. 1). The T199 + NCS treatment also displayed a significant increase (P < 0.05) in the proportion of 6- to 8-cell embryos that developed to the blastocyst stage over that with the c-SOFM + NCS and c-SOFM treatment groups (Fig. 1). Interestingly, however, the proportion of 6- to 8-cell-stage embryos progressing to the blastocyst stage did not vary significantly between the T199 + NCS and c-SOFMaa treatment groups (Fig. 1).

Experiment 2

The next investigation concerned the influence of culture atmosphere during bovine oocyte maturation in vitro on development to the blastocyst stage. Oocyte maturation under a 5% CO₂-in-air atmosphere supported significantly greater (P < 0.05) cleavage frequencies than for oocytes matured under a 5% CO₂, 7% O₂, 88% N₂ (5/7/88) culture atmosphere (Fig. 2). Amino acid supplementation of oocyte maturation media did not significantly influence cleavage. Development to the 6- to 8-cell stage did not vary significantly between the two 5% CO₂-in-air treatments but was significantly greater (P < 0.05) in both of these groups than in the c-SOFM 5/7/88 group (Fig. 2). Development to the blastocyst stage was significantly higher (P < 0.05) in the c-SOFMaa 5% CO₂-in-air oocyte maturation group than in the other treatments (Fig. 2). No significant difference in blastocyst formation frequency was observed in the c-SOFM 5% CO₂-in-air and the two 5/7/88 oocyte maturation treat-
Influence of oocyte maturation on blastocyst development displayed by oocytes matured in 0, 1, 10, and 100 ng/ml of murine EGF-supplemented cSOFMaa. See Figure 1 legend for a full description of the groups. Cleavage and development to the 6- to 8-cell and blastocyst stages varied significantly among the treatment groups in a concentration-dependent manner. The proportion of 6- to 8-cell embryos that progressed to the blastocyst stage did not vary among the four treatments. Values with different superscript letters are significantly different (P < 0.05).

Experiment 3

The influence of EGF-supplemented cSOFMaa oocyte maturation medium on development to the blastocyst stage was investigated next. We first examined influences on cumulus cell expansion. EGF stimulated cumulus cell expansion in a concentration-dependent manner with the greatest amount of expansion occurring in the 100 ng/ml EGF treatment (data not shown). However, the proportion of fully expanded COCs observed even at this EGF concentration was not as extensive as we normally observe with FSH-supplemented oocyte maturation media. EGF stimulated a concentration-dependent increase in cleavage and 6- to 8-cell-stage embryos, as both the 10 ng/ml and 100 ng/ml concentration-dependent increase in cleavage and 6- to 8-cell-stage embryos was not as extensive as we normally observe with FSH-supplemented oocyte maturation media (data not shown). However, the proportion of fully expanded COCs observed even at this EGF concentration was not as extensive as we normally observe with FSH-supplemented oocyte maturation media. EGF stimulated a concentration-dependent increase in cleavage and 6- to 8-cell-stage embryos, as both the 10 ng/ml and 100 ng/ml treatments displayed a significant increase (P < 0.05) in these parameters over that for the control and 1 ng/ml treatment (Fig. 3). Interestingly, the proportion of 6- to 8-cell-stage embryos over cleaved embryos (6- to 8-cell/clvd) did not vary significantly among the four treatments (Fig. 3). There was a trend for a dose-dependent increase in blastocyst formation, but only in the 100 ng/ml EGF treatment, which displayed a significant (P < 0.05) increase in blastocyst formation (Fig. 3). The proportion of 6- to 8-cell-stage embryos that progressed to the blastocyst stage did not vary significantly among the four treatments (Fig. 3).

Blastocyst Total Cell Numbers and Apoptosis

In experiment 1, blastocyst total cell number was significantly greater (P < 0.05) in the T199+NCS treatment than in the cSOFM+NCS and cSOFM treatment groups (Table 1). However, the total blastocyst cell number did not vary significantly between the T199+NCS and cSOFMaa treatments (Table 1). The incidence of apoptotic cells did not vary significantly among these four treatment groups (Table 1). No significant differences in blastocyst total cell number or number of apoptotic cells were observed in blastocysts derived from the four oocyte maturation treatment groups in the second experiment (Table 2). Likewise, EGF supplementation of oocyte maturation media did not significantly influence blastocyst cell number or apoptosis (Table 3). Day 8 blastocyst total cell numbers ranged from 53 to 317 cells from all three experiments. Representative images of labeled blastocysts for total cell counts and measurement of apoptosis are displayed in Figure 4.

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### Table 1. Blastocyst total cell number and apoptosis following oocyte maturation in TCM-199+NCS, cSOFM+NCS, cSOFM, or cSOFM (means ± SEM).

<table>
<thead>
<tr>
<th>Treatment*</th>
<th>n</th>
<th>Total cell no. (mean ± SEM)</th>
<th>Apoptosis (cell no.) (mean ± SEM)</th>
<th>Apoptosis (% of total cells) (mean ± SEM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TCM-199+NCS</td>
<td>34</td>
<td>168.2 ± 6.1</td>
<td>16.3 ± 1.7</td>
<td>9.5 ± 0.8</td>
</tr>
<tr>
<td>CSOFM+NCS</td>
<td>34</td>
<td>138.2 ± 7.0</td>
<td>11.8 ± 0.9</td>
<td>9.1 ± 0.8</td>
</tr>
<tr>
<td>CSOFMaa</td>
<td>34</td>
<td>157.6 ± 7.1</td>
<td>13.4 ± 1.5</td>
<td>8.5 ± 0.9</td>
</tr>
<tr>
<td>cSOFM</td>
<td>34</td>
<td>132.8 ± 7.7</td>
<td>16.3 ± 2.1</td>
<td>13.0 ± 1.6</td>
</tr>
</tbody>
</table>

* All cumulus-oocyte-complexes were matured in one of the four treatment groups under a 5% CO₂ in air at 39°C culture atmosphere; inseminated oocytes from each oocyte maturation treatment were cultured to the blastocyst stage in cSOFMaa medium under a 5% CO₂/7% O₂/88% N₂ culture atmosphere at 39°C.

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### Table 2. Blastocyst total cell number and apoptosis following oocyte maturation in cSOFMaa or cSOFM under 5% CO₂ in air and 5% CO₂/7% O₂/88% N₂ culture atmospheres (means ± SEM).

<table>
<thead>
<tr>
<th>Treatment*</th>
<th>n</th>
<th>Total cell no. (mean ± SEM)</th>
<th>Apoptosis (cell no.) (mean ± SEM)</th>
<th>Apoptosis (% of total cells) (mean ± SEM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>cSOFMaa</td>
<td>38</td>
<td>133.8 ± 8.0</td>
<td>13.2 ± 1.1</td>
<td>10.4 ± 0.8</td>
</tr>
<tr>
<td>5% CO₂</td>
<td>38</td>
<td>129.0 ± 6.0</td>
<td>12.1 ± 1.1</td>
<td>9.4 ± 0.8</td>
</tr>
<tr>
<td>cSOFM</td>
<td>38</td>
<td>125.6 ± 7.5</td>
<td>10.9 ± 1.0</td>
<td>9.1 ± 0.7</td>
</tr>
<tr>
<td>5% CO₂, 7% O₂, 88% N₂</td>
<td>38</td>
<td>131.5 ± 7.9</td>
<td>9.6 ± 0.9</td>
<td>7.7 ± 0.6</td>
</tr>
</tbody>
</table>

* All inseminated oocytes from each oocyte maturation treatment were cultured to the blastocyst stage in cSOFMaa medium under a 5% CO₂/7% O₂/88% N₂ culture atmosphere at 39°C.

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### Table 3. Blastocyst total cell number and apoptosis following oocyte maturation in cSOFMaa supplemented with either 1, 10, or 100 ng/ml of EGF (means ± SEM).

<table>
<thead>
<tr>
<th>Treatment*</th>
<th>n</th>
<th>Total cell no. (mean ± SEM)</th>
<th>Apoptosis (cell no.) (mean ± SEM)</th>
<th>Apoptosis (% of total cells) (mean ± SEM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>11</td>
<td>141.3 ± 17.3</td>
<td>54.3 ± 14.2</td>
<td>34.5 ± 6.2</td>
</tr>
<tr>
<td>1 ng/ml EGF</td>
<td>22</td>
<td>147.3 ± 12.6</td>
<td>54.5 ± 11.2</td>
<td>39.5 ± 6.8</td>
</tr>
<tr>
<td>10 ng/ml EGF</td>
<td>23</td>
<td>149.3 ± 10.1</td>
<td>36.7 ± 3.7</td>
<td>25.5 ± 2.3</td>
</tr>
<tr>
<td>100 ng/ml EGF</td>
<td>29</td>
<td>158.3 ± 11.8</td>
<td>40.3 ± 7.6</td>
<td>24.6 ± 3.3</td>
</tr>
</tbody>
</table>

* All cumulus-oocyte-complexes were matured in one of the four treatment groups under a 5% CO₂ in air at 39°C culture atmosphere; inseminated oocytes from each oocyte maturation treatment were cultured to the blastocyst stage in cSOFMaa medium under a 5% CO₂/7% O₂/88% N₂ culture atmosphere at 39°C.
cell number | apoptosis
---|---
| | |

**FIG. 4.** Representative images of blastocyst total cell number and apoptosis. Shown are blastocysts consisting of **A** 161 total cells and 21 apoptotic cells produced from the T199 + NCS oocyte maturation treatment; **B** 198 total cells and 16 apoptotic cells produced from the cSOFM + NCS oocyte maturation treatment; **C** 177 cells and 16 apoptotic cells produced from the cSOFMaa oocyte maturation treatment; **D** 148 cells and 25 apoptotic cells produced from the cSOFM treatment. Bars = 50 μM.

**FIG. 5.** RT-PCR analysis of oocyte transcripts. Experiments were conducted to contrast relative levels of transcripts encoding five marker genes including Na+/K+ ATPase α-1 isoform, bFGF, Cu/ZnSOD, cyclins A and B, and exogenously supplied α-globin in oocytes matured in 1) T199 + NCS, 2) cSOFM + NCS, 3) cSOFMaa, and 4) cSOFM. Ethidium bromide-stained gels displaying the amplified product of each transcript in all three experimental replicates. map, a blank mAP paper control; H2O, a minus cDNA blank control. The expected sizes of each cDNA amplicon are displayed.

**Oocyte Transcript Levels**

The abundance of each specific gene transcript was expressed relative to an exogenously supplied α-globin mRNA standard as described above. Figure 5 displays the ethidium bromide-stained gels for each oocyte transcript and experimental replicate. The mean relative mRNA abundance for the five specific transcripts is displayed in Figure 6. The relative abundance of transcripts encoding the Na+/K+ ATPase α-1 isoform did not vary significantly between the T199 + NCS, cSOFM + NCS, and cSOFMaa treatments; but all three of these treatments displayed a significantly higher (P < 0.05) relative abundance of this gene product than in oocytes matured in the cSOFM treatment (Figs. 5 and 6). An identical outcome was observed for transcripts encoding cyclin A (Figs. 5 and 6). No significant differences in the relative abundance of transcripts encoding Cu/Zn SOD, cyclin B, and bFGF were observed among the oocyte maturation treatment groups. There was, however, a noteworthy trend toward a lower relative abundance of transcripts encoding bFGF within the serum-free treatments (Fig. 5).

**DISCUSSION**

This study has contrasted the influences of bovine oocyte maturation environments on blastocyst formation frequency, total cell number, apoptosis, and oocyte transcript levels. Our findings indicate not only that cSOFMaa medium is an effective base medium for embryo culture [14,15,18,29,36] but also that it can be effectively employed as a base medium for oocyte maturation. We employed for the first time an assessment of blastocyst total cell number and apoptosis to examine influences of oocyte maturation environments on bovine blastocyst development. The results demonstrate that blastocysts derived from in vitro maturation, fertilization, and embryo culture protocols undergo apoptosis. However, apoptotic levels are not greatly influenced by the oocyte maturation environment. Amino acid supplementation of oocyte maturation media was associated with enhanced developmental frequencies, increased blastocyst cell number, and elevated oocyte maternal mRNA levels. Oocyte maturation with cSOFMaa medium resulted in blastocyst cell numbers comparable to...
FIG. 6. Variation in oocyte marker gene transcripts. The relative abundance of each marker gene transcript for oocytes cultured in 1) T199 NCS, 2) cSOFM NCS, 3) cSOFMaa, and 4) cSOFM is presented. Transcripts encoding Na⁺/K⁺ ATPase α1 isoform and cyclin B were significantly reduced in the amino acid-free cSOFM. Transcripts encoding Cu/ZnSOD, bFGF, and cyclin A were not present at significantly different levels among the oocyte maturation treatments. However, there was a trend for transcripts encoding bFGF to be reduced in the serum-free culture treatments. Values with different superscript letters are significantly different (P < 0.05).
medium, it can be easily modified to investigate the effects of specific components. SOFM was designed from concentrations of salts and energy metabolites found in sheep oviductal fluid [44], and this medium is “the standard” for ovine embryo culture. SOFM medium has been used effectively to support bovine development in vitro [15,16,18]. Keskinetepe et al. [15,16] observed an important benefit from the addition of citrate to SOFM media. Citrate may act as a regulator of fatty acid synthesis, ion chelation, and solute transport, and its addition to SOFM in combination with NEA is beneficial for obtaining maximal blastocyst formation frequencies with this medium [15,16]. For several years now we have employed this base medium with phosphate removed and glucose levels reduced to support bovine embryo development in vitro. This medium consistently supports high frequencies of development of bovine embryos to the blastocyst stage in vitro. For these reasons we proposed that cSOFMaa medium would be a useful base medium for oocyte maturation studies as well. The results from this study certainly support this view.

The culture atmosphere plays a critical role during oocyte maturation and embryo culture. Tervit et al. [45] first reported a benefit of culture under reduced O₂ (5% CO₂, 5% O₂, 90% N₂) atmosphere, but this approach was not widely adopted for the culture of mammalian embryos from other species until it became apparent that reduced O₂ atmospheres eliminated a requirement for embryo coculture with somatic cells and also increased blastocyst frequencies in all mammalian species tested including the mouse [15–17,29,44–49]. This benefit may stem from a reduction in the deleterious effects of reactive oxygen species on early development. A 5% CO₂-in-air culture atmosphere is typically employed for the in vitro maturation of mammalian oocytes. Our study examined whether this culture atmosphere is optimal for bovine oocyte maturation. Blastocyst development was reduced following oocyte maturation under a 5% CO₂, 7% O₂, 88% N₂ culture atmosphere. We would speculate that cumulus cell metabolism is adversely affected by exposure to reduced O₂ culture atmospheres, and this impairs cumulus cell-oocyte interactions during oocyte maturation, thus reducing overall developmental potential. Cumulus cell expansion and attachment were markedly reduced in COCs matured under a reduced O₂ culture atmosphere, and these observations support this hypothesis.

EGF positively influences oocyte maturation in a number of species, including cattle [50–59]. Its addition to serum-free TCM199 is sufficient to support high frequencies of bovine oocyte maturation and subsequent development to the blastocyst stage [50–53]. Its role, however, in regulating bovine blastocyst cell number or apoptosis after its addition to oocyte maturation medium has not been investigated. Our results clearly demonstrated that EGF has a distinct influence on cumulus cell expansion and cleavage. Our results, however, do not support an influence of EGF during oocyte maturation on development of 6- to 8-cell-stage embryos to the blastocyst stage or on blastocyst cell number or apoptosis. This of course does not suggest that EGF does not serve as a regulator of blastocyst formation. However, EGF-mediated influences on oocyte maturation simply may not persist beyond these early cleavage stages. However, by increasing the proportion of cleaved embryos, EGF treatment during oocyte maturation indirectly promotes increased development to the blastocyst stage.

Programmed cell death, or apoptosis, has been detected at the blastocyst stage, in both the inner cell mass and trophectoderm, as a normal feature of murine development [26,27]. We have now demonstrated that bovine blastocysts also display apoptosis. Although the role of cell death in the early embryo is unknown, it may involve the removal of cells with abnormal properties or inappropriate potential. There is evidence to suggest that certain “survival” factors, such as growth factors, produced by the embryo and maternal reproductive tract play a role in controlling apoptosis levels [26,27]. EGF receptor ligands have been linked to reduced apoptosis in murine blastocysts [26,27]. Although the addition of EGF to oocyte maturation medium did not influence blastocyst apoptosis, we would expect that treatment of bovine embryos with EGF or transforming growth factor-α would reduce bovine blastocyst apoptosis in a similar fashion. This outcome, however, awaits further experimentation. We believe that the majority of the apoptotic cells observed in our bovine blastocysts were located in the inner cell mass as opposed to the trophectoderm. This can be confirmed only by differential staining techniques, which are incompatible with the double-labeling procedure employed in this study. The level of apoptosis we observed is similar to that reported for murine embryos [26,27], and therefore we would predict that it represents an expected level of apoptosis for cultured blastocysts. The blastocyst total cell number values reported in the present study are in agreement with values for Day 8 bovine blastocysts reported for oocytes matured in TCM199+serum and then cultured in serum-supplemented medium [28]. We can conclude, therefore, that serum-free oocyte maturation and embryo culture environments do not result in reduced blastocyst cell numbers. A proven semiquantitative RT-PCR method was applied to examine relative variations in the levels of transcripts encoding five marker genes [31]. We chose Na⁺/K⁺ ATPase α-1 isoform, bFGF, Cu/ZnSOD, and cyclins A and B as genes of interest for the following reasons: 1) Na⁺/K⁺ ATPase α-1 isoform is an important housekeeping gene controlling plasma membrane ionic concentration gradients, and it facilitates the trophectoderm ion transport mechanisms that control blastocyst formation [33,60]; 2) bFGF is a maternally expressed growth factor gene, as transcripts encoding bFGF are reported in bovine and ovine embryos up to the 8- to 16-cell stage and are absent in blastocysts [34]; 3) Cu/ZnSOD is an antioxidant gene, especially important in high oxygen concentration environments [35]; 4) cyclins A and B are important genes controlling cell cycle events and germinal vesicle breakdown [61]. Cyclin A controls the progression of the cell from G1 to S phase and peaks during the S phase. Cyclin B is an active mediator of meiotic progression, controlling the entry into and exit of the cell from the M phase [61]. Our results have clearly demonstrated that oocyte maturation media can influence levels of oocyte transcripts. Furthermore, a strong influence of EGF on the greatest deficit was observed using the amino acid-free cSOFM medium. This medium also supported the lowest developmental frequencies. Therefore our studies have linked oocyte mRNA levels with the capacity of the oocyte, once fertilized, to develop to the blastocyst stage in vitro. These findings are supported by other studies that have examined influences of culture media on embryonic transcript levels [39,40,62]. The decreased relative levels of Na⁺/K⁺ ATPase α-1 isoform and cyclin A mRNAs observed in oocytes matured in amino acid-free cSOFM medium may reflect the acceleration of mechanisms that direct the degradation of maternal mRNAs prior to eventual activation of the embryonic genome [63–66]. In particular, it is possible that these mRNAs are subject to early deadenylation that would result in truncated poly(A)-tails and re-
duced RT by oligo(dT) [63–66]. The precise cause of the observed reduction in transcript levels and their full functional significance await further experimentation. Our results demonstrate that oocyte culture media can influence transcript levels and that the use of serum-free amino acid-supplemented defined media for oocyte maturation maintains transcripts at levels observed in oocytes matured in serum-supplemented media.

The characterization of fully defined optimized culture conditions for preimplantation development in vitro is an obvious priority, especially when there is increasing evidence demonstrating negative consequences of culture in suboptimal environments on patterns of embryonic gene expression and, more dramatically, on fetal phenotype in animal models [2–4,8,35,39,40]. These points are of particular importance as the assisted-reproduction technology field prepares to adopt human in vitro oocyte maturation and embryo culture to the blastocyst stage as emerging advancements. Our results support the use of serum-free defined conditions for both oocyte maturation and embryo culture in vitro. These approaches will allow for studies aimed at defining the precise roles of hormonal and growth factor modulators in supporting early development and should result in the eventual characterization of optimal conditions for early mammalian embryo culture.

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