The Impact of High Temperature on the Development and Quality of In Vitro Matured Beef Cattle Oocytes and Their Potential to Subsequent Embryonic Development

NKADIMENG MARBLE¹,², VAN MARLE-KÖSTER², LEKOLA KGOMOTSO⁴, MPHAPHATHI MASINDI LOTUS¹, LEHLOENYA KHOBOSO CHRISTINA⁴, PILANE CYRIL MPHO¹, NEDAMBALE TSHIMANGADZO LUCKY¹,³

¹Agricultural Research Council, Animal Production Institute, Germplasm Conservation and Reproductive Biotechnologies, Private bag X2, Irene 0062, South Africa; ²Department of Animal and Wildlife Sciences, University of Pretoria, Private bag X20, Hatfield 0028, South Africa; ³Tshwane University of Technology, Faculty of Science, Department of Animal Sciences, Private bag X680; ⁴Department of Agriculture, University of Zululand, Private Bag X1001, KwaDlangezwa 3886, South Africa.

Abstract | The study aimed to evaluate the effect of high temperature on cattle oocytes matured in vitro at 39, 41, 42, 43°C and to evaluate apoptosis through Caspase-activity and DNA fragmentation on produced embryos. Cumulus oocyte complexes (COC) were matured at 39, 41, 42 and 43°C. Two best maturation temperatures were selected for maturation throughout embryo production. Post maturation, matured oocytes were exposed to standard embryonic conditions. Embryos were evaluated at 3 stages: ≥ 2-4 cell, ≥ 8cell and blastocyst to determine Caspase-3 activity and DNA fragmentation. Maturation of oocytes at 41 and 39°C demonstrated no significant difference (p>0.05) however, higher maturation rate compared to 42 and 43°C. Matured oocytes at 39 and 41°C showed no difference (p>0.05) in embryo development. Blastocysts at 39°C maturation had higher (p<0.05) nuclei cells compared to those at 41°C. Caspase-3 activity was comparable at ≥ 2-4 cell and ≥ 8cell for both 39 and 41°C maturation temperatures, however, 41°C showed high DNA fragmentation compared to 39°C maturation group. Blastocyst from oocytes matured at 41°C had higher Caspase-3 activity and DNA fragmentation than those at 39°C. Embryos derived from 41°C maturation temperature resulted in evidence of apoptosis compared to 39°C.

Keywords | Oocyte maturation, High temperature, Embryos, Cell apoptosis, Blastocyst

INTRODUCTION

To date in vitro production and developmental rate of cattle embryos attained a great deal of attention (Sakatini, 2017; Kidie, 2019). The technique is widely implemented by animal breeders around the world to accelerate animal genetics for improved animal production and livestock sustainability (Hafez, 2015; Blondin, 2015). However, in South Africa (SA) the viability of cattle embryos produced in vitro over the past decade has been reported to be less viable than the in vivo counterparts (Arlotto et al., 2001; Mapeka et al., 2019). Additionally, this technology has not yet been widely used in most African countries with the global in vitro embryo production (IVEP) statistics reporting a 3.51% adoption of the technique as compared to 98% implementation in North
America and 71.7% in European countries (IETS, 2015; Perry 2018; Kidie, 2019).

Amongst other advantages, the IVEP technology is one of the embryo biotechnologies that plays a crucial role in evaluating heat stress mechanisms (Bó et al., 2019). The artificial culture environment of IVEP allows investigation of the quality of oocytes and embryos at a preferred temperature such as the maternal temperature during hot summer months (Sakatani, 2017). Heat stress has been reported to compromise the competence of embryos in different developmental stages from maturation to subsequent development (Hanson, 2009; Payton et al., 2018). It alters Luteinizing (LH) and Follicular stimulating hormone (FSH) secretion required for follicular function (Paula-lopes and Hansen, 2002; Wolfenson and Roth, 2019). Alterations of these hormones can result in the delay of ovulation and aged oocyte that has low fertilization rate and poor quality embryos (Al-Katanani et al., 2002). The report by van Wettere et al. (2021) demonstrated that heat stress affects follicular development and growth, furthermore impair nuclear maturation on in vitro cattle oocytes.

During oocyte maturation, heat stress further disrupts chromosome alignment, decreases oocytes number reaching metaphase II and induces oocyte apoptosis (Hansen, 2009). In the early stages of embryo development, heat stress results in organellar damage as a result of microfilament and microtubule disruption, and mitochondrial swelling (Payton et al., 2018; Sakatani, 2017). Studies have shown that heat stress activates apoptosis in oocytes and embryos by releasing oxidative stress agents in the mitochondria and disrupts the mitochondrial function (Sakatani, 2017). Apoptosis occurs due to the release of the DNase enzyme that cuts dsDNA and generates 3-hydroxy ends through activation of caspases of the cysteine proteases family (Somal et al., 2015).

High temperature due to climate change has affected livestock production worldwide (Rojas-Downing et al., 2017). In SA, extremely high temperatures reaching 43°C with 12 heat waves in 71 days were evident primarily in the Northern, Central and the East coastal regions during December-February summer months of 2015/2016 (Scholtz et al., 2016; Mbokodo et al., 2020). Moreover, estimations of 1.5 to 3°C in temperature increase is expected in 2050 as a result of climate changes (Department of Environmental Affairs, 2010). This may affect the fertility of beef cattle enormously.

South African indigenous breeds such as Nguni, Bonsmara and Brahman are known to be adaptable to harsh conditions such as extreme temperatures (Mapiye et al., 2008; Katiyatiya et al., 2017). However, the 2015/2016 drought resulted in a decline in beef cattle productivity and high mortality in the affected provinces (Mthembu and Zwane, 2017). It is therefore vital to understand “how heat stress affects the competence and development of beef cattle oocytes and embryos in SA breeds”. Thus, in SA no information is available on the evaluation of heat stress on cattle embryos in beef cattle breeds. Furthermore, there are limited studies on the physiological changes of oocytes competency at higher temperatures equivalent to most of the South African summer temperatures such as 43 °C. Therefore, the IVEP study evaluates the high temperature on embryo development in all the embryo production stages and the link between heat stress and cell apoptosis on the produced embryos.

MATERIALS AND METHODS

OOCYTES COLLECTION

Beef cattle ovaries (Nguni, Bonsmara and Brahman) were received from a local abattoir (Morgan). Ovaries were transported within an hour to the laboratory at 38 °C in buffer saline (Sigma Aldrich®, Germany). At arrival, fresh pre-warmed buffer saline was used to wash ovaries for removal of blood contamination and further sprayed with 70% alcohol for any preclusion of contamination. Aspiration method was used to retrieve oocytes according to Bohlooli et al. (2015). A total of 777 oocytes were utilized for in vitro maturation procedure to find the best suitable maturation temperatures throughout the study and 982 oocytes were used for subsequent embryo production experiments Figure 1.

Figure 1: Flow of data collected in the study period.
In vitro maturation of oocytes

Post oocytes retrieval, cumulus oocyte complexes (COCs) with ≥ 3 layers of cumulus cells (Șen and Kuran, 2018) were selected under the Olympus CX 23 microscope at 80x magnification (New York microscope Co, USA). Three ml of modified Dulbecco phosphate buffer saline (DPBSS) (lifetechnologies®, South Africa) and tissue culture medium [M-199 + 10% Fetal bovine serum (FBS)] (Sigma Aldrich®, Germany) were used to wash selected COCs. In each medium COCs were washed two times. The COCs were allocated randomly into four temperature groups: 39, 41, 42 and 43 °C (50 oocytes per 4 maturation wells for each incubation temperature). The universal M-199 medium was used for maturation of COCs. The medium was supplemented with 40 μl FSH, 20 μl LH and 20 μl Estradiol (EST) (Sigma Aldrich®, USA) and covered with 250 μl of mineral oil (Sigma Aldrich®, USA). COCs were matured at 5% CO₂, 5% O₂ and 90% N₂ at 100% humidified air for 24 hours. Post 24-hour (h) incubation, polar body extrusion was examined. Polar body extrusion was used as a maturation parameter according to Şen and Kuran (2018). In each temperature, oocytes were examined for polar body extrusion. Briefly, oocytes were submerged in 200 μl M-199 medium supplemented with FBS (Sigma Aldrich®, Germany) (M-199 + 10% FBS, Sigma Aldrich®, Germany) and vortexed for a minute and 30 seconds. Olympus 1x 71 microscope (New York microscope Co, USA) was used to assess polar bodies at 20x/0.45 Na2 magnification. The selection of two temperatures (39 and 41 °C) for subsequent embryonic development were due to polar bodies examination outcomes. Embryonic development proceeded to normal conditions (39 °C incubation temperature, 5% CO₂, 5% O₂, and 90% N₂ at 100% humidified air).

In vitro fertilization and culture

Preparation of matured oocytes for in vitro fertilization

In vitro fertilization commenced with oocyte wash in 5 drops of 100 μl pre-warmed Bracket and Oliphant (BIO-IVF) medium covered with 250-μl mineral oil (Sigma Aldrich®, USA). Oocytes were transferred to seven drops of 50 μl pre-warmed Bio-IVF medium (20-25 oocytes) covered with 250 μl of mineral oil (Sigma Aldrich®, USA) for fertilization.

Semen thawing

Oocytes were fertilized by frozen-thawed Nguni semen in 0.25ml straw size. The thawing procedure proceeded with semen in air for 30 seconds and in warm water of 37 °C for one minute. Thawed semen was washed through centrifugation two times in 4 ml of pre-warmed sperm wash medium (Sigma Aldrich®, USA) at 1500 RPM for 8 minutes per centrifugation. During second centrifugation, previous sperm wash medium was removed and new 4 ml sperm wash medium was added to proceed with the centrifugation. Post second centrifugation, 50 μl sperm wash X number of fertilization drops was added to the sperm pellet. A total of 20-25 matured oocytes in each drop of Bio-IVF medium (Sigma Aldrich®, USA) was then fertilized with 50 μl sperm pellet at a concentration of 0.98×10⁶ sperm cells/ml and incubated for 18 hours (Seshoka et al., 2016).

In vitro culture

Post 18 hours of insemination, cumulus cells on presumptive zygotes were removed through vortex for one minute and 30 seconds in 200 μl of pre-warmed [M199+10% FBS (Sigma Aldrich®, USA)] medium. Five drops of 100 ml pre-warmed synthetic oviductal fluid medium supplemented with bovine serum albumen [SOF-BSA (Sigma Aldrich®, USA)] covered with 250 μl mineral oil was used to wash zygote prior in vitro culture. Zygotes (20-25) were cultured in a modular chamber that contained 5 % oxygen and CO₂ mixed gas added for 48 hours at 39 °C incubation temperature and 100 % humidity. Cleavage was examined at 48 hours of incubation. Post 48 hours, embryos were transferred to pre-warmed synthetic oviductal fluid supplemented with FBS [SOF-FBS (Sigma Aldrich®, USA)] and further incubated for 72 hours. At 72 h, a minimum of 20 μl of [(SOF-FBS (Sigma Aldrich®, USA)] old medium was removed and replaced with pre-warmed fresh medium of the same amount in each drop. Blastocyst were evaluated at day seven of embryo production (Bryla and Trzcińska, 2011).

Nuclei cell analysis of produced blastocyst

Expanded and hatched blastocyst were examined for nuclei cell count per maturation group (39 and 41°C). Blastocyst were removed and washed twice in 3 ml of pre-warmed DPBS medium (lifetechnologies®, South Africa). Blastocyst were stained using Hoechst (33342) (Sigma Aldrich®, Isreal) solution and transferred to a sterile microscope glass slide (Labcon, Germany). Four square points of Vaseline was added surrounding the blastocyst on the glasslide and the coverslip was placed on top of the four Vaseline pointers. An amount of 5 μl Hoechst solution (33342) was gently added on each side of the coverslip (Labcon, Germany) to cover the whole area under the slip. The coverslip was then moderately pressed until it touches the embryo drop and sealed with a colourless nail polish (Essence, New York). The staining and mounting procedure was performed with dimmed lights (Bohlooli et al., 2015). Computer sperm analyser® v 3.4.0 english version (Microptic, Spain) and Olympus BX51 epifluorescence was used to evaluate Nuclei cell at 60x magnification.
Caspase-3 activity evaluation of produced embryos.

Embryos produced from oocytes at 39 and 41°C maturation temperature were removed at day two (≥ 2-4 cell), day five (≥ 8 cell) and day seven (blastocyst) of embryo production for evaluation of Caspase-3 activity. Drops of 50 μl pre-warmed phosphate buffer saline-polyvinylpyrrolidone (PBS-PVP) (Sigma Aldrich®, Germany) was used to wash all embryos prior the Caspase-3 activity procedure. The procedure was performed using Caspase-3 Colorimetric Activity Assay Kit (EMD Millipore, USA). Prior the procedure positive control samples were incubated in 50 μl drops of hydrogen peroxide (H₂O₂) (Sigma Aldrich®, Germany) for two hours. The reactions of enzyme were monitored with 96 microtiter Thermo scientific plate reader (Power Wave XS; BioTek, USA) at 405 nm and 450 nm reference filter through the OD readings (Optimal density) values (Hwang et al., 2012).

TUNEL assay procedure for DNA fragmentation on produced embryos from oocytes matured at 39 and 41°C

Produced embryos were removed and washed using a similar procedure as in the Caspase-3 activity experiment. Embryos were permeabilized in 50 μl drop of 0.5 % (v/v) Triton X-100 (Sigma Aldrich®, USA), 0.1 % (w/v) and sodium citrate (Sigma Aldrich®, USA) solution incubated for 30 minutes. Embryos were fixed in 50 μl drops of 4 % paraformaldehyde (Sigma Aldrich®, USA) solution for one hour at room temperature prior permeabilization. After fixation, embryos were washed 3 times in 50 μl drops of PBS/ PVP (Sigma Aldrich®, Germany) and incubated for 30 minutes to permeabilize. Embryos were stored for 30 min at room temperature in a humidified box post permeabilization prior TUNEL assay procedure (Silva et al., 2013).

Post preparation of all embryos, samples continued to the Alexa Fluor® 488 Click-iT® TUNEL assay kit (Eugen, USA) according to the manufacturer’s instruction for the examination of fragmented DNA. Post completion of the TUNEL assay, all samples were mounted on a slide and covered with a coverslip in a parallel procedure performed on blastocyst nuclei cell analysis experiment for imaging. TUNEL index was also performed on produced blastocysts from both maturation groups (39 and 41°C) to evaluate the percentage of fragmented nuclei. Click-iT® TUNEL assay (Eugen, USA) was used for determining the apoptotic nuclei. The total nuclei number was determined by Hoechst (33342) stain and nuclei cells were recorded as positive for TUNEL assay procedure when they had visible green fluorescence from Alexa 488-stain. The nuclei fragmented percentage was done by TUNEL index = (no. TUNEL-positive nuclei, fragmented and condensed)/

RESULTS

In vitro maturation of oocytes from four different maturation temperatures

Extrusion of polar body demonstrated no difference (p>0.05) between oocyte maturation at 41 and 39°C incubation temperature. However, 42 and 43°C had lower extrusion of polar bodies (p<0.05) than 39 and 41°C (Table 1). Figure 2 illustrates images of oocytes matured at four temperatures (39, 41, 42 and 43°C).

Table 1: Polar body extrusion of cattle oocytes matured at 39, 41, 42 and 43°C incubation temperatures (Mean ± SD).

| Treatment | Oocytes (no) | % of Polar body | % of Non-polar body |
|-----------|--------------|-----------------|---------------------|
| 39°C      | 192          | 57.3 ± 1.0b     | 42.1 ± 1.5b         |
| 41°C      | 201          | 60.0 ± 1.2a     | 40.1 ± 1.9h         |
| 42°C      | 188          | 20.5 ± 1.0b     | 80.3 ± 4.8a         |
| 43°C      | 196          | 10.3 ± 1.0b     | 89.8 ± 1.0h         |

* Mean within columns with different superscripts differ at (p<0.05). Non-polar body is the number of oocytes presented without extrusion of polar body post 24h maturation.

In vitro embryo production and blastocyst evaluation from oocytes matured at 39 and 41°C

Table 2 represents produced embryos at different in vitro embryo production stages and nuclei cell count of produced blastocyst from oocytes matured at 39 and 41°C. Produced blastocysts at day 7 from oocytes matured at two temperatures 39 and 41°C and images of Hoechst (33342) stained blastocysts from 39 and 41°C maturation temperatures are presented in Figure 1. There was no difference (p>0.05) on all the embryonic developmental stages between 39 and 41°C. Blastocyst from oocytes matured at 39°C had higher (p<0.05) nuclei cell number...
than blastocyst of oocytes matured at 41°C.

Figure 2: Assessment of heat stress on cattle oocytes at four temperatures: 39, 41, 42 and 43 ºC. Produced blastocyst at day 7 and Hoechst (33342) staining of nuclei cells of produced blastocysts. Suitable maturation temperatures were selected according to the extrusion of polar bodies at a significant of (P < 0.05). Matured cattle oocytes from (A) 39 ºC, (B) 41 ºC, (C) 42 ºC and (D) 43 ºC. Day 7 Blastocyst from cattle oocytes matured at (E) 39 ºC and (F) 41 ºC, Hoechst 33342 (Blue) stained blastocyst from (G) 41 ºC and (H) 39 ºC matured cattle oocytes.

Table 2: In vitro embryo development from cattle oocytes matured at 39 ºC and 41 ºC incubation temperatures (Mwan ± SD).

| Temperature | No embryos | 2-4 cell (d2) | 8 cell (d2) | Morula (d5) | Blastocyst (D7) | Cell nuclei (n) |
|-------------|------------|---------------|-------------|-------------|----------------|----------------|
| 39 ºC       | 241        | 65.0 ± 6.8    | 39.2 ± 6.6  | 19.0 ± 14.6 | 19.4 ± 2.6     | 133 ± 57.9     |
| 41 ºC       | 250        | 62.4 ± 6.5    | 35.0 ± 6.4  | 15.2 ± 4.8  | 11.2 ± 6.3     | 45.8 ± 11.2    |

CASPAR-3 ACTIVITY OF PRODUCED EMBRYOS FROM OOCYTES MATURED AT 39 AND 41 ºC

Embryos at ≥ 2-4 cell and ≥ 8 cell from oocytes matured at both 39 and 41 ºC showed no differences (p>0.05) on Caspase-3 activity (Table 3). However, Caspase-3 activity was higher (p<0.05) on blastocyst produced from at 41 ºC matured oocytes compared to oocytes matured at 39 ºC.

Table 3: Caspase-3 activity on different stages of produced cattle embryos in vitro (Mean±SD).

| Temperature | Embryos (no) | Positive control | ≥ 2-4 cell embryo | ≥ 8 cell embryo | Blastocyst |
|-------------|--------------|------------------|-------------------|----------------|------------|
| 39 ºC       | 90           | 0.815 ± 0.049    | 0.015 ± 0.001    | 0.022 ± 0.007  | 0.037 ± 0.012 |
| 41 ºC       | 87           | 0.016 ± 0.002    | 0.032 ± 0.013    | 0.053 ± 0.005  | 0.057 ± 0.006 |

a,b,c mean value within the rows with different superscripts differ at (p<0.05).

TUNEL ASSAY EVALUATION ON PRODUCED EMBRYOS FROM OOCYTES AT 39 AND 41 ºC MATURATION TEMPERATURE

Significantly higher (p<0.05) DNA fragmentation was observed in ≥ 2-4 cell embryos from 41 ºC compared to 39 ºC. Embryos at ≥ 8 cell had higher TUNEL positives (p<0.05) at 41 ºC compared to embryos produced from oocytes at 39 ºC. The was no significant difference (p>0.05) on TUNEL positives between embryos at ≥ 2-4 cell and ≥ 8 cell from 39 ºC incubation temperature (Table 4). The 41 ºC maturation temperature on produced blastocyst showed higher (p<0.05) percentage of fragmented DNA compared to 39 ºC (Table 5). Figure 3, illustrates stained ≥ 2-4 cell, ≥ 8 cell embryos and images of stained blastocyst post TUNEL assay procedure from both temperatures groups (39 and 41 ºC).

Table 4: DNA fragmentation of nuclei using TUNEL assay at different embryonic stages (Mean±SD).

| Temperature | No embryos | Embryo group | Tunel positive (%) | Tunel negative (%) |
|-------------|------------|--------------|-------------------|-------------------|
| 39 ºC       | 77         | ≥ 2-4 cell   | 10.8 ± 1.2        | 89.2 ± 4.8        |
| 41 ºC       | 65         | ≥ 8 cell     | 19.6 ± 1.9        | 80.5 ± 2.7        |
| 39 ºC       | 80         | ≥ 2-4 cell   | 26.2 ± 2.9        | 73.8 ± 4.1        |
| 41 ºC       | 70         | ≥ 8 cell     | 63.3 ± 3.1        | 36.7 ± 1.6        |

Table 5: TUNEL assay indexes on in vitro cattle blastocyst (Mean ± SD)

| Temperature | Blastocysts (no) | No nuclei cell | TUNEL positive nuclei | TUNEL indexes (%) |
|-------------|------------------|----------------|-----------------------|-------------------|
| 39 ºC       | 12.0 ± 1.2       | 127.8 ± 53.4   | 10.3 ± 4.8            | 9.7 ± 6.7         |
| 41 ºC       | 10.0 ± 1.4       | 46.2 ± 10.0    | 21.8 ± 21.8           | 48.0 ± 8.2        |

a,b mean value within the rows with different superscripts differ at (p<0.05). Nuclei cell were recorded as positive for TUNEL labelling when they had visible green fluorescence.

Figure 3: Representation of images following TUNEL Assay procedure. Hoechst 33342 (Blue) stained 2-4 cell embryos from (A) 39 °C, (B) 41 °C, (C) ≥ 8 cell embryos from both 39 and 41 °C, blastocysts from cattle oocytes matured at (D) 39 °C and (E) 41 °C. Alexa 488-stained (green) 2-4 cell embryos from 41 (F), 39 °C (G), ≥ 8 cell embryos from 41 °C (H) and Alexa 488-stained (green) blastocysts from cattle oocytes matured at (I) 39 and (J) 41 °C maturation group.
Exposure of oocytes at high temperature during maturation resulted in cell apoptosis on produced embryos. The maturation experiments demonstrated no significant difference between oocytes matured at 39 and 41 °C, however, 42 and 43°C differed respectively in polar body extrusion. There was no difference in subsequent developmental stages (cleavage rate, morula and blastocyst rates) of produced embryos amongst the two maturation groups (39 and 41°C). The number of nuclei on the produced blastocyst differed significantly between the two temperature groups (39 and 41°C). The present study showed a relationship between high temperature and cell apoptosis through Caspase-3 and DNA fragmentation evaluation of embryos matured at 41°C on different embryonic stages.

According to Roth and Hansen (2004) and the report by Sakatani (2017), maturation failures are as a result of elevated temperatures within the physiological range of 40–41°C during maturation. In contrast to the current study, in vitro maturation resulted in similar polar body extrusion between heat stressed (41°C) oocytes and oocytes from 39°C. Rynkowska et al. (2011) suggested that exposure of oocytes or any other cells to stresses such as high temperature activates a set of proteins called heat shock proteins (HSPs). These proteins maintain cellular homeostasis and protection against environmental stressors. The latter might have had an influence on the survival of oocytes incubated at 41°C in the present study. The high non-polar body extrusion at 42 and 43°C maturation temperature groups in this study may be due to the reduction in the synthesis of intracellular proteins by 30 to 50 % due to exposure of oocytes to severe heat (Saeki et al., 1997). The above statement is parallel to the findings by Payton et al. (2011) that showed a reduction in the level of polyA mRNA in bovine oocytes at 12 h in vitro maturation due to heat stress. Furthermore, Campen et al. (2018) highlighted that heat stress exposure at 42 °C on oocytes results in the alterations of gap junction communications between oocytes and cumulus cells which consequently interrupt cytoplasmic maturation.

The similarities in embryo development between 39 and 41 °C maturation temperature in the current study are in line with the work of Sakatani et al. (2015) which emphasized that subsequent embryo development may not be affected by incubation of oocytes at 41 °C during maturation. Moreover, Silva et al. (2013) reported that the potential of Bos Taurus (Friesian and Brown Swiss) oocytes development during hot season becomes detrimentally reduced than in Bos indicus (Nguni and Brahman) cows. The genetic deposition of the Bos Indicus cattle allows them to have a protective mechanism against heat stress than Bos Taurus cattle (Friesian and Brown Swiss) (Silva et al., 2013). Therefore, the above statement might have an influence in the similarities observed on embryo development between oocytes matured at 41 and 39 °C.

The current study further demonstrated that embryos produced from heat stressed oocytes have the ability to develop to advanced stages of embryo development. Therefore, the produced day 7 blastocysts in the current study from oocytes matured at 41 °C were examined for their developmental quality through assessment of nuclei cell number. The findings were in line with the work of Sakatani et al. (2004) however, differed with the report by Sakatani et al. (2015). Sakatani et al. (2015) revealed no difference in the inner cell mass ratio and trophectoderm cell of blastocyst previously exposed to heat stress at the early stages of oocyte development. The low nuclei cell number of the heat stress blastocyst in this study might be due to the decrease in trophectoderm cell number. It is considered that at metaphase II of the oocyte life heat stress affects the quality of the trophectoderm cells from produced blastocysts (Ju et al., 2005; Habeeb et al., 2018). Furthermore, the report by Ascari et al. (2017) and Stamperna et al. (2020) demonstrated that the build-up of apoptosis elements and disruption of gene expression in the inner cell mass of blastocyst is as a result of exposure of oocytes to heat stress for 12 hours during in vitro maturation.

Embryos were affected differently at different stages of development during the evaluation of Caspase-3 activity in the present study. The report of Hansen (2015) emphasized that at 2-4 cell the mitochondria of the embryo becomes resistant to depolarization and inhibition of apoptosis may occur or the presence of apoptosis may become minimum. The above may be the reason for the low activity of Caspase-3 on 2-4 cell embryos in the present study. The study of Roth and Hansen (2004) and a report by Stamperna et al. (2021) demonstrated Caspase-3 activity in the cytoplasm of a single blastomere of heat-shocked 4 to 8 cell embryos, morulae and blastocyst compared to control. The present study, however, showed no significant difference at ≥ 8 cell embryos from oocytes matured at 39 and 41 °C. The similarities of the results are in line with the report by Silvia et al. (2009) which emphasizes that resistance to pro-apoptotic signals may occur during preimplantation stage of cattle embryos. Brad et al. (2007) further reported that mitochondrial membrane in cattle embryos can become artificially depolarized by carbonyl cyanide 3-chlorophenylhydrazone (CCCP). Under such circumstances, Caspase-9 and Caspase-3 activation take place however, DNA fragmentation does not transpire.

The TUNEL assay findings in the current study showed lower DNA fragmentation in 2-4 cell stage. These suggest
that the release of DNase by Caspase in 2-4 cell embryos is at minimal. According to the report by Fear and Hansen (2011) cattle embryos at lower stages of development may remain refractory to apoptotic stimuli compared to 8-16-cell developmental stage. Hence, more TUNEL positive nuclei in the present study were observed from embryos at ≥ 8 cell than ≥ 2-4 cell. This may suggest that as development progresses higher temperature in lower developmental stages affect a fraction of the blastomere at later stages of development to become TUNEL positive (Brad et al., 2007). As a result, more fragmented nuclei cells in blastocysts were evident from oocytes at 41 than in 39°C.

CONCLUSIONS AND RECOMMENDATIONS

The study showed that high temperature influences oocyte competency and compromises embryo development and survival with evidence cell apoptosis. The results expand on the importance of more implementation of assisted reproduction technologies such as IVF in South Africa to provide a controlled environment for produced embryos. This is to overcome reproductive inefficiency in response to high temperature during summer months. It is further recommended that genetic mechanisms involved in the survival of beef cattle embryos during heat stress from diverse SA genotypes should be evaluated to account for the expected future rise of temperature in SA.

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NOVELTY STATEMENT

To our knowledge, this is the first study in Southern Africa to report heat stress on beef cattle embryos produced in vitro and its effect of cell apoptosis on developmental phases of produced embryos.

AUTHOR’S CONTRIBUTION

MML, PCM and LK assisted in data collection and designing of the manuscript. VME, NTL, NM and LKC drafted, supervised and evaluated the manuscript content.

CONFLICT OF INTEREST

The authors have declared no conflict of interest.

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