A mouse model for improving cell survival of bisected cattle embryos

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Morula and blastocyst stage embryos recovered from B6D2F1 mice were bisected with a metal microblade in M2 medium with or without sucrose and/or cytochalasin B supplementation. Cell lysis was determined by staining the embryos with Hoechst 33258 and propidium iodide. Lysed cells take up both stains but non-lysed cells only the Hoechst 33258 stain, resulting in pink fluorescence for lysed cells and blue fluorescence for non-lysed cells under UV excitation. During bisection of morulae, the presence of cytochalasin B decreased the proportion of lysed cells in both the absence (P=0.0001) and presence of sucrose (P=0.001). During bisection of blastocysts the average proportion of lysed cells was slightly lower in the presence of cytochalasin than that in the control medium, but the effect was not statistically significant (P=0.34). No effect of sucrose was observed in either demi-morulae or demi-blastocysts. These results are essentially similar to those obtained in simultaneous experiments with cattle embryos, suggesting that the simpler mouse model might be useful for developing less traumatic bisection protocols for cattle embryos.

Key words: demi-embryo, splitting, cell lysis, propidium iodide, Hoechst 33258, cytochalasin, sucrose

Introduction

Embryo bisection can be used in embryo transfer programmes either to increase the number of transferable (demi)embryos or to produce identical twin calves for research or other purposes. The first successful transfers of bisected bovine embryos were reported in the early 1980s (Willadsen et al. 1981, Ozil et al. 1982).

Although embryo bisection yields more offspring per original embryo under appropriate conditions, subsequent splitting of an embryo does not seem to give any further benefit, as the viability of quarter embryos is roughly half that of demi-embryos (Willadsen et al. 1981, Bredbacka et al. 1992). Demi-embryos apparently are particularly sensitive to further invasive treatments. Substantial cell loss in bisected embryos will either compromise further development before implantation.

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Manuscript received September 1996
or result in an insufficient embryonic signal for maternal recognition, or both.

The success of producing live foetuses after splitting varies, depending upon factors such as bisection technique (Mertes and Bondioli 1985), embryo age and stage (Williams et al. 1984, McEvoy and Sreenan 1990), method of transfer (Takeda et al. 1986), and type and status of recipient (Arave et al. 1987, Bredbacka et al. 1992). However, data on the survival of cells during bisection are sparse, as are the published data on attempts to increase viability of blastomeres during the splitting process. A recent report (Bredbacka 1995) demonstrates that a considerable proportion (about 20% to 30%) of blastomeres lyses in embryos bisected using a microblade-assisted technique. The effectiveness of the technique is nevertheless confirmed by a 60% pregnancy rate of demi-embryos transferred singly (Bredbacka et al. 1996). Survival after serial splitting might be significantly improved, if cell lysis could be prevented more efficiently.

The major factor holding up improvements to bovine embryo bisection protocols is the cost of embryos. Although in vitro produced (IVP) embryos represent a relatively inexpensive alternative to in vivo produced embryos, the IVP technology is more elaborate and requires embryo culture facilities. Furthermore, regardless of the method of embryo production, bovine embryos exhibit a large variation in morphology, which sometimes complicates interpretation of results.

The aim of my study was to demonstrate the feasibility of a mouse model for predicting cell survival in bovine embryos. The effect of sucrose and cytochalasin B on cell survival was investigated with differential staining of lysed and non-lysed cells. The results were compared with those of a parallel study in the bovine (Bredbacka 1995).

Material and methods

Female B6D2F1 (C57BL6 x DBA F1) mice were first superovulated by injecting 5 to 7.5 IU PMSG and 5 IU hCG 48 h apart and then mated with B6D2F1, males. Compacted morulae were recovered 73 to 75 hours after the hCG injection and blastocysts were recovered 90 to 94 h after the hCG injection using the method devised by Hogan et al. (1986).

The nuclei of blastomeres were stained by exposing the embryos to 20 µg/ml Hoechst 33258 stain in 4-well Nunc dishes (Nunc, Cat No. 176740, Roskilde, Denmark) for 3 to 5 h at 37°C in air. The stain was diluted in 500 µl of M2 medium (Quinn et al. 1982) containing 4 mg/ml bovine serum albumin (M2+BSA). The M2 was made up of analysis grade chemicals and the BSA was from Sigma (Cat. No. A-9647, St. Louis, MO, USA). Each embryo was evaluated immediately before splitting using a Leitz Fluovert inverted microscope; embryos with any indication of abnormal morphology were excluded from the experiments.

The control medium for bisection was M2+BSA. In the treatment groups the sucrose was added at a concentration of 200 mM, and cytochalasin B (Sigma, Cat. No. C6762, St. Louis, MO, USA) was used at 7.5 µg/ml. The combination of cytochalasin B and sucrose was also tested in the experiments with morulae. Before being split in sucrose-containing medium, the embryos were allowed to equilibrate for 3 to 5 min in the medium at room temperature. When bisected in the presence of cytochalasin B, the embryos had an equilibration time of 5 to 15 min at room temperature.

The embryos were bisected with a Leitz micromanipulator using a microblade prepared from a razor blade (Williams and Moore 1988) and the “scratched bottom technique” (Bredbacka 1991). With this technique, scratches are produced parallel to the orientation of the microblade on the bottom of the lid of a plastic petri dish (Nunc, Cat. No. 153066, Roskilde, Denmark) to prevent the embryo from slipping during bisection. The embryo is then bisected by a vertical movement of the microblade in a drop of 100–200 µl of medium. Blastocysts are bisected so that both the trophoderm and inner cell mass are halved. To reduce the stickiness of
blastomeres the stage was chilled to about 10–15°C by placing precooled aluminium blocks on the Leitz Fluovert microscope stage. The temperature of the stage was monitored by a Linkam CO60 stage warmer with the heating off.

After the splitting, each demi-embryo pair was placed in a 50 μl-drop of M2+BSA. Propidium iodide (Sigma, Cat. No. P-4170, St. Louis, MO, USA) was added to a final concentration of 10 μg/ml to stain nuclei of lysed cells. After 45 to 60 min of staining at room temperature, each demi-embryo pair was washed and placed into 50 μl of medium in a well of a 96-well plastic dish. The nuclei were counted by fluorescence microscopy using a Leitz Fluovert microscope with filter block A.

Cells of each embryo were classified as lysed (pink fluorescence) or viable (blue fluorescence), with the results being expressed as the proportion of the total number of cells. Data were analysed by variance analysis using the Statistical Analysis System (SAS Institute, Cary, NC, USA).

Results

The average number of nuclei (±SEM) in morulae was 18.5 (±0.6) and in blastocysts 47.3

Table 1. Cell viability of mouse morulae after splitting in different media.

| Treatment          | n  | Proportion (±SEM) of viable cells | P value* |
|--------------------|----|----------------------------------|----------|
| Control            | 24 | 0.41 (±0.03)                     | –        |
| Cytochalasin B     | 24 | 0.56 (±0.02)                     | 0.0001   |
| Sucrose            | 24 | 0.43 (±0.02)                     | 0.52     |
| Cytochalasin B + sucrose | 24 | 0.55 (±0.03)                     | 0.0001   |

* In comparison with control medium

Table 2. Cell viability of mouse blastocysts after splitting in the presence of cytochalasin B.

| Treatment      | n  | Proportion (±SEM) of viable cells | P value* |
|----------------|----|----------------------------------|----------|
| Control        | 19 | 0.47 (±0.02)                     | –        |
| Cytochalasin B | 20 | 0.49 (±0.02)                     | 0.34     |

* In comparison with control medium

Table 3. Cell viability of mouse blastocysts after splitting in the presence of sucrose.

| Treatment | n  | Proportion (±SEM) of viable cells | P value* |
|-----------|----|----------------------------------|----------|
| Control   | 20 | 0.45 (±0.02)                     | –        |
| Sucrose   | 20 | 0.45 (±0.02)                     | 0.96     |

* In comparison with control medium.
Cytochalasins inhibit reversibly the polymerization of actin, thus enhancing the flexibility of cell membranes. This property has been used to decrease cell damage in nucleus transfer experiments (for review, see Loskutoff 1990). Cytochalasin B has been shown to decrease cell lysis during bisection of Day 7 bovine embryos (Bredbacka 1995). In the present study, treatment of mouse morulae with cytochalasin B significantly decreased the proportion of lysed cells in bisected morulae. Note that a highly significant difference (P=0.0001) from the control medium could be shown with only a small number of embryos (24 per treatment).

Despite the clearly beneficial effect of cytochalasin B in morulae, no clear effect of this compound was demonstrated in blastocysts. Since the proportion of non-lysed cells was slightly higher in the presence of cytochalasin, it is possible that the number of embryos was too low to demonstrate an actual effect. On the other hand, cytochalasin may provide less protection for cells in blastocysts. It is also possible that the inner cell mass (ICM) and the trophoderm (TE) respond differently to cytochalasin. If cytochalasin protects only one of the two cell types, variation in the ICM/TE ratio between embryos would mask the actual effects. This stresses the need to use more embryos for analysis unless cell survival can be measured separately in the ICM and TE.

It has been proposed, that adding of sucrose to the splitting medium inhibits pressure-induced cell membrane rupture during the splitting process by means of dehydration (Herr et al. 1988). In the present study the addition of sucrose at a concentration of 200 mM had no beneficial effect on mouse blastomere survival. The same conclusion was drawn in the study with bovine embryos (Bredbacka 1995). Addition of sucrose at another concentration, however, might have rendered the blastomeres more resistant to the stress induced by the bisection procedure.

The observations made here with mouse morulae are essentially identical to the results obtained with cattle embryos (Bredbacka 1995), suggesting that mouse embryos might be useful...
for developing splitting techniques in cattle. In an experiment with a limited number of equine embryos, cytochalasin B tended to protect cell lysis (Huhtinen et al. 1995), implying the possibility of using the mouse model across several species.

Bringing cell lysis down to a minimum could improve the efficiency of embryo transfer programmes not only by making bisection less traumatic, but also by improving biopsy for preimplantation diagnosis. Increased cell viability of biopsied embryos could have a substantial effect on pregnancy rates when biopsy is combined with cryopreservation.

The use of mouse embryo bisection as a model for improving the cattle embryo splitting protocol has several advantages. The cost of producing mouse embryos is reasonable especially compared with in vivo produced cattle embryos. Although the use of the latter need not be ruled out, mouse embryos show less variability in quality as assessed by morphological criteria, not least because inbred lines or hybrid F1 lines can be used. The dead cells often present in intact cattle embryos complicate splitting experiments, as lysed cells take up propidium iodide even without the cell injury induced by microsurgery and so have to be excluded from analysis. Last, but not least, with about half of the cells lysing in a standard splitting protocol with mouse embryos, improvements to the protocol are easier to measure than with cattle embryos, in which roughly 25% of the cells lyse in the standard protocol.

In conclusion, the use of mouse embryos offers a valuable tool for selecting protocols for further bisection and biopsy studies with cattle embryos.

Acknowledgements. I thank T.-M. Nieminen for management of the mice and preparation of the media used in this study and K. Bredbacka for useful suggestions with regard to the manuscript.

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Hiiren alkio mallina solukestävyyden parantamiseksi naudan alkioiden halkaisussa

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Morula- ja blastokystivaiheessa olevia hiiren alkioita halkaistiin mikroterällä M2-liuoksessa. Halkaisukokeessa tutkittiin sakkaroosin (200 mM) ja sytokalasiinin (7,5 ug/ml) vaikutusta solujen kestävyyteen kaksoisvärjäyksen avulla. Värjäyksen jälkeen solut voidaan erottaa toisistaan fluoresenssimikroskoopin avulla väarin perusteella. Hoechst 33258-väri värjää kaikki solut (sininen fluoresenssi), ja propidiumjodi vain tuhoutuneet solut (vaaleanpunainen fluoresenssi).

Morulavaiheessa olevia alkioita halkaistaessa sytokalasiini vähensi hajonneiden solujen osuutta riippumatta siitä, oliko liuoksessa myös sakkaroosia. Kun käytettiin blastokystivaiheessa olevia alkioita, eri käsittelyillä ei havaittu vaikutusta hajonneiden solujen määrin. Sakkaroosilla ei ollut vaikutusta blastokystivaiheen solujen kestävyyteen. Näitä tulokset ovat samansuuntaisia kuin vastaavasta naudan alkioilla tehdystä kokeesta saadut tulokset. Halkaisussa tuhoutuneiden solujen erottaminen on hiiren alkiolla tehdystä kokeesta saadut tulokset. Edullisena ja yksinkertaisena hiirimalli voi olla hyödyllinen kehitettäessä naudan alkion soluja paremmin suojaavia mikrokirurgisia menetelmiä.