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Comet assay on thawed embryos: An optimized technique to evaluate DNA damage in mouse embryos

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ABSTRACT

Our objective was to optimize the CA technique on mammal embryos. Materials and methods: 1000 frozen 2-cell embryos from B6CBA mice were used. Based on a literature review, and after checking post-thaw embryo viability, the main outcome measures included: 1) comparison of the embryo recovery rate between 2 CA protocols (2 agarose layers and 3 agarose layers); 2) comparison of DNA damage by the CA on embryos with (ZP+) and without (ZP−) zona pellucida; and 3) comparison of DNA damage in embryos exposed to 2 genotoxic agents (H2O2 and simulated sunlight irradiation (SSI)). DNA damage was quantified by the % tail DNA.

Results: 1) The recovery rate was 3.3% (n = 5/150) with the 2 agarose layers protocol and 71.3% (n = 266/371) with the 3 agarose layers protocol. 2) DNA damage did not differ statistically significantly between ZP− and ZP+ embryos (12.60 ± 2.53% Tail DNA vs 11.04 ± 1.50 (p = 0.583) for the control group and 49.23 ± 4.16 vs 41.13 ± 4.31 (p = 0.182) for the H2O2 group); 3) H2O2 and SSI induced a statistically significant increase in DNA damage compared with the control group (41.13 ± 4.31% Tail DNA, 36.33 ± 3.02 and 11.04 ± 1.50 (p < 0.0001)). The CA on mammal embryos was optimized by using thawed embryos, by avoiding ZP removal and by the adjunction of a third agarose layer.

1. Introduction

The exposure of parents to environmental toxicants, such as polycyclic aromatic hydrocarbons (Einaudi et al., 2014; Perrin et al., 2011), Bisphenol A (Goldstone et al., 2015), solvents (Kolstad et al., 1999), metals (Thompson and Bannigan, 2008; Zhou et al., 2016) or various therapies (Bujan et al., 2014; Esquerré-Lamare et al., 2015; Pecou et al., 2009; Roti Roti et al., 2012) may induce DNA damage in male and female germ cells. DNA damage in parental germ cells can lead to reproductive issues, such as reduced fertilization, impaired early embryonic development, a decreased pregnancy rate and increased miscarriage rate (Simon et al., 2014; Zhao et al., 2014). The transmission of paternal germ cell DNA damage to preimplantation embryos has been demonstrated in humans (Zenzes et al., 1999), although DNA repair occurs in the zygote (Ménézo et al., 2010). This observation raises the question of evaluating DNA damage in preimplantation embryos because, currently, no genotoxicity test on the embryo has been validated.

The comet assay is a simple and rapid test for evaluating DNA damage in eukaryotic cells (Speit et al., 2009). It allows for the visualization of denatured DNA fragments after ex-nucleus migration by electrophoresis. After staining, the obtained shape mimics a “comet” with the head containing intact DNA and the remaining parts of the chromosome and the tail containing relaxed DNA loops or broken DNA fragments (Hovrátová et al., 2004). This test is validated by toxicological regulatory agencies for the assessment of DNA damage in somatic cells (OECD, 2014). The comet assay is also used by researchers...
for the assessment of DNA damage in sperm and male germ cells (Baumgartner et al., 2009; Perrin et al., 2007; Preaubert et al., 2016) and in oocytes (Berthelot-Ricou et al., 2013, 2011a, 2011b; Courbiere et al., 2013; Einaudi et al., 2014). Studies using the comet assay on animal embryos are scarce and use heterogeneous protocols and species (Blerkom et al., 2001; Fabian et al., 2003; Harrouk et al., 2000; Hwang et al., 2013; Ju et al., 2010; Kitagawa et al., 2004; Müller et al., 1996; Natarajan et al., 2010; Rajesh et al., 2010; Sturmey et al., 2009; Takahashi et al., 1999, 2000; Thiyagarajan and Valivittan, 2009a, 2009b; Tranguch et al., 2003; Tsuda et al., 1998; Webster et al., 2000). As comet assay is a very sensitive test, the analysis of fresh embryos by comet assay requires the proximity between animal facilities and laboratory. Moreover, the analysis of fresh embryos does not allow control or additional analyses if required, notably after in vivo exposure or for an hypothetical use for regulatory issues.

The aim of our study was to optimize the comet assay for application on frozen mouse embryos, in order to simplify the handling and allow subsequent analysis.

2. Material and methods

All of the chemicals were purchased from Sigma-Aldrich (Saint-Quentin Fallavier, France) unless otherwise stated.

2.1. Institutional Review Board

Institutional Review Board approval (C2EA-14) was obtained after submission to the National Ethics Committee on Animal Experimentation. All of the experimental protocols and animal handling procedures were approved by The National Ethics Committee on Animal Experimentation.

2.2. Source of embryos

In total, 1000 frozen 2-cell embryos were purchased from the ImmunoPHEnomique Center (CIPHE, Luminy, Marseille, France). Briefly, the embryos were obtained by natural mating on superovulated B6CBA females aged 5 weeks. 48 h after mating, the oviduct was flushed and the two-cells embryo obtained were cryopreserved. We used a slow freezing protocol with a controlled rate freezing machine and 1.5 M propanediol as a cryoprotectant, according to the procedure described by the European Mouse Mutant Archive (EMMA) (EMMA—The European mouse mutant archive, 2013a; Hagn et al., 2007).

2.3. Embryo thawing

The embryos were thawed according to the procedure described by the European Mouse Mutant Archive (EMMA) (EMMA—The European mouse mutant archive, 2013b; Hagn et al., 2007). Briefly, the straw was thawed at room temperature (RT), and then the embryos were rinsed in 4 successive M2 medium drops and placed in KSOM medium (EmbyoMax®, Merck Millipore, Darmstadt, Germany).

The number of thawed embryos used for each experiment performed in this study is presented in Table 1.

2.4. Embryo viability and culture conditions

To validate the use of frozen embryos instead of fresh embryos for comet assay, we assessed viability and blastulation rates in thawed embryos. After 1 h of recovery in KSOM, the 2-cells embryos were examined under a microscope and were classified as lysed or intact. The survival rate was defined as the ratio of the intact embryos to the total thawed embryos.

The embryos were then cultured for 24 h in a humidified chamber at 37 °C (95% air/5% CO2) to determine viability. The viability rate was defined by the number of 2-cell embryos moving to the 4–8 cell stage to the total intact embryos. The embryos were cultured for 24 h more to determine the evolution rate to the blastocyst stage, assessing the number of blastocysts compared to the total intact embryos. For each separate experiment, 20 embryos were used (5 experiments = 100 embryos) (Table 1).

Table 1: Experiments performed in the study and number of thawed embryos used for each one.

<table>
<thead>
<tr>
<th>Experiments performed</th>
<th>Embryo stage used</th>
<th>Protocol used after embryo thawing</th>
<th>Nb of separate experiments</th>
<th>Total nb of 2-cells embryos used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Assessment of embryo viability and blastulation rates</td>
<td>2-Cell</td>
<td>2-Cells embryo in vitro culture to blastocyst stage</td>
<td>5</td>
<td>20 × 5 = 100</td>
</tr>
<tr>
<td>Impact of the nb of agarose layers on embryo recovery rate</td>
<td>2-Cell</td>
<td>CA Protocol 1</td>
<td>5</td>
<td>30 × 5 = 150</td>
</tr>
<tr>
<td>Impact of ZP on CA results</td>
<td>2-Cell</td>
<td>CA Protocol 2</td>
<td>5</td>
<td>30 × 5 = 150</td>
</tr>
<tr>
<td>Impact of genotoxic agents exposure on DNA damage</td>
<td>2-Cell</td>
<td>CA Protocol 2</td>
<td>5</td>
<td>60 × 5 = 300</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td></td>
<td></td>
<td>1000</td>
</tr>
</tbody>
</table>

CA: comet assay (Protocol 1 uses 2 agarose layers; Protocol 2 uses 3 agarose layers), ZP: zona pellucida.
Table 2
Published protocols of comet assay on preimplantation embryos.

<table>
<thead>
<tr>
<th>Reference</th>
<th>Species</th>
<th>Embryo production</th>
<th>Zona pellucida</th>
<th>% agar in NMP/LMP layers</th>
<th>Vol of LMP agar in which embryos are placed/vol of mix deposited to form the 2nd agar layer</th>
<th>Number of embryos per slide/embryo stage</th>
<th>Third layer agarose</th>
<th>Lysis solution composition</th>
<th>Lysis duration (min)</th>
<th>Lysis temperature (°C)</th>
<th>Incubation/electrophoreseduration (min)</th>
<th>Quantification of DNA damages</th>
</tr>
</thead>
<tbody>
<tr>
<td>Müller et al. (1996)</td>
<td>Mouse</td>
<td>Mating</td>
<td>Removed</td>
<td>0.1/0.8</td>
<td>500 μL</td>
<td>?/C</td>
<td>No</td>
<td>LS + SLS + SDS</td>
<td>15</td>
<td>?</td>
<td>7/5</td>
<td>Tail/head ratio</td>
</tr>
<tr>
<td>Takahashi et al. (1999)</td>
<td>Bovine</td>
<td>IVF</td>
<td>Intact</td>
<td>1/1</td>
<td>200 μL/10 μL</td>
<td>10 to 20/C</td>
<td>No</td>
<td>LS + SLS + PK</td>
<td>180</td>
<td>RT</td>
<td>20/20</td>
<td>Tail length</td>
</tr>
<tr>
<td>Takahashi et al. (1999)</td>
<td>Hamster</td>
<td>Mating</td>
<td>Intact</td>
<td>1/1</td>
<td>200 μL/10 μL</td>
<td>4 to 5/Z and C</td>
<td>No</td>
<td>LS + SLS</td>
<td>120</td>
<td>RT</td>
<td>20/20</td>
<td>Tail length</td>
</tr>
<tr>
<td>Harrouk et al. (2000)</td>
<td>Rat</td>
<td>Mating</td>
<td>Removed</td>
<td>7/0.25</td>
<td>Mixed with 5 μL of PBP recovered with LMP</td>
<td>?/Z</td>
<td>No</td>
<td>LS + SLS + SDS + DMSO</td>
<td>120</td>
<td>4 °C</td>
<td>40/5</td>
<td>Tail/head ratio</td>
</tr>
<tr>
<td>Fabian et al. (2003)</td>
<td>Mouse</td>
<td>Mating</td>
<td>Intact</td>
<td>1/1</td>
<td>10 μL</td>
<td>2 to 6/B</td>
<td>No</td>
<td>LS + SDS + PK</td>
<td>180</td>
<td>RT</td>
<td>20/20</td>
<td>Undamaged, necrotic, apoptic</td>
</tr>
<tr>
<td>Tranguch et al. (2003)</td>
<td>Mouse</td>
<td>Mating</td>
<td>Intact</td>
<td>?</td>
<td>75 μL</td>
<td>10/C</td>
<td>No</td>
<td>LS + SLS</td>
<td>60</td>
<td>4 °C</td>
<td>60/10</td>
<td>Tail: yes/no</td>
</tr>
<tr>
<td>Kingawa et al. (2004)</td>
<td>Porcine</td>
<td>IVF</td>
<td>Intact</td>
<td>1/1</td>
<td>50 μL</td>
<td>10/C</td>
<td>No</td>
<td>LS</td>
<td>120</td>
<td>?</td>
<td>20/20</td>
<td>Tail length</td>
</tr>
<tr>
<td>Thiyagarajan and Vallivitran (2009a)</td>
<td>Buffalo</td>
<td>IVF</td>
<td>Intact</td>
<td>1/1</td>
<td>200 μL/10 μL</td>
<td>10 to 20/C</td>
<td>No</td>
<td>LS + SLS + PK</td>
<td>180</td>
<td>RT</td>
<td>20/20</td>
<td>Tail length</td>
</tr>
<tr>
<td>Thiyagarajan and Vallivitran (2009a)</td>
<td>Buffalo</td>
<td>IVF</td>
<td>Intact</td>
<td>1/1</td>
<td>200 μL/10 μL</td>
<td>10 to 20/C</td>
<td>No</td>
<td>LS + SLS + PK</td>
<td>180</td>
<td>RT</td>
<td>20/20</td>
<td>Tail length</td>
</tr>
<tr>
<td>Sturmey et al. (2009)</td>
<td>Porcine, Human, Bovine</td>
<td>IVF</td>
<td>Removed</td>
<td>1/0.8</td>
<td>4 μL</td>
<td>1/B</td>
<td>No</td>
<td>LS + DMSO</td>
<td>Overnight</td>
<td>4 °C</td>
<td>40/20</td>
<td>% DNA migration</td>
</tr>
<tr>
<td>Ju et al. (2010)</td>
<td>Porcine</td>
<td>IVF</td>
<td>Removed</td>
<td>1/1</td>
<td>50 μL</td>
<td>10 to 15/C and M</td>
<td>NMP</td>
<td>LS + SDS</td>
<td>60</td>
<td>4 °C</td>
<td>15/15</td>
<td>Undamaged, apoptic</td>
</tr>
<tr>
<td>Natarajan et al. (2010)</td>
<td>Sheep</td>
<td>IVF</td>
<td>Intact</td>
<td>1/1</td>
<td>200 μL/10 μL</td>
<td>10/C</td>
<td>No</td>
<td>LS + SLS + PK</td>
<td>180</td>
<td>RT</td>
<td>20/20</td>
<td>Tail length</td>
</tr>
<tr>
<td>Rajesh et al. (2010)</td>
<td>Sheep</td>
<td>IVF</td>
<td>Intact</td>
<td>1/1</td>
<td>200 μL/10 μL</td>
<td>10 to 20/C</td>
<td>No</td>
<td>LS + SLS + PK</td>
<td>180</td>
<td>RT</td>
<td>20/20</td>
<td>Tail length</td>
</tr>
<tr>
<td>Hwang et al. (2013)</td>
<td>Bovine</td>
<td>IVF</td>
<td>Removed</td>
<td>?</td>
<td>75 μL</td>
<td>10/Z</td>
<td>No</td>
<td>LS + DMSO</td>
<td>180</td>
<td>4 °C</td>
<td>30/20</td>
<td>Tail length</td>
</tr>
</tbody>
</table>

IVF: in vitro fertilization; LMP: low melting point; NMP: normal melting point; LS: lysis solution (2.5 M NaCl; 100 mM Na2EDTA; 10 mM Tris-HCl, pH 10; 1% Triton X-100); PK: proteinase K; DMSO: dimethyl sulfoxide; SLS: sodium lauroyl sarcosinate; SDS: sodium dodecyl sulfate; RT: room temperature; OTM: Olive Tail Moment; Z: zygote (one cell); C: cleavage stage embryo; M: morula stage embryo; B: blastocyst stage embryo.
2.5. Impact of the number of agarose layers on the embryo recovery rate

All of the published comet assay protocols on mice embryos (Fabian et al., 2003; Harrouk et al., 2000; Hwang et al., 2013; Kitagawa et al., 2004; Müller et al., 1996; Natarajan et al., 2010; Rajesh et al., 2010; Sturme et al., 2009; Takahashi et al., 1999, 2000; Thiyagarajan and Valivittan, 2009a, 2009b; Tranguch et al., 2003; Webster et al., 2000) used 2 layers of agarose except one (Ju et al., 2010). A glass microscope slide is initially coated with 1% high or normal-melting point (HMP, NMP) agarose (the first agarose layer), and then, the embryos are mixed with a drop of low-melting point (LMP) agarose and placed on the first layer (the second agarose layer). We compared the impact of two protocols on the rate of cell loss. In Protocol 1, as described by Berthelot-Ricou et al. (2011a) we used 2 layers. In protocol 2, we used 3 layers (as described in the “Alkaline comet assay” section). For each protocol, the recovery rate was defined by the number of embryos found and analyzed on a slide at the end of the protocol compared to the total embryos initially deposited on the slide. For each of the 5 separate experiments, 30 embryos were used with protocol 1 and 30 embryos with protocol 2 (300 embryos) (Table 1).

2.6. Zona pellucida impact on comet assay.

To determine the impact of the ZP on the comet assay, we tested normal embryos (ZP+) and zona-free embryos (ZP−). In the ZP− embryos, the ZP was digested by an acidic Tyrode’s solution at RT and was neutralized 3 times with 0.4 M Tris-HCl (pH 7.5), rinsed with ultrapure water, dipped into 100% methanol (HPLC grade purity solvent), and then kept 20 min at 4 °C for solidification. DNA unwinding was performed with an alkaline solution (1 mMol Na2EDTA and 300 mMol NaOH) for 20 min at 20 °C in a 30.5 × 22 cm horizontal electrophoresis unit. Next, electrophoresis was conducted for 30 min (25 V, 300 mA) in the same buffer solution at RT. After the electrophoretic run, the slides were neutralized 3 times with 0.4 M Tris-HCl (pH 7.5), rinsed with ultrapure water, dipped into 100% methanol (HPLC grade purity solvent), and dried overnight at RT. Staining was performed with a propidium iodide solution (0.1 μg/mL), and the slides were examined with a fluorescence microscope at a 400 × final magnification (Carl Zeiss Axios Imager A2, Bayern, Germany). The digital pictures were analyzed, and the determination of the % tail DNA was performed with a CCD Nikon camera (DS-Fi2, Nikon, Champigny-sur-Marne, France) and Komet software (version 7.0, Nottingham, UK).

2.7. Exposure conditions

The subsequent experiments were performed on embryos with an intact zona pellucida (ZP+). To validate the sensitivity of the assay, we used two positive controls of well-known chemical and physical genotoxic agents (Berthelot-Ricou et al., 2011a; Kumaravel and Jha, 2006; Miranda-Vilela et al., 2010): H2O2 and Sun Simulated Irradiation (SSI). For the H2O2 group, the embryos were placed in a 220 μM H2O2 solution for 6 min at 4 °C in the dark. SSI induces DNA damage by irradiation to light. For the SSI group, light irradiation was carried out with a Suntest CPS + solar simulator (Atlas Material Testing Technology BV, Moussy le Neuf, France) equipped with a xenon arc lamp (1100 W) and special glass filters restricting transmission of light below 290 nm and near IR-blocking filter. The addition of glass filters limited the spectral emission to the 320–800 nm band (UVA–Visible light). The irradiation dose was 120 kJ/m². The temperature of the samples was kept at 4 °C by using a regulated cooling system during the irradiation experiments. For each set of experiments, the control group consisted of embryos incubated for 1 h in KSOM medium. For each of the 5 separate experiments, 20 embryos were analyzed for each condition (control, H2O2 or SSI) (300 embryos) (Table 1).

2.8. Alkaline comet assay

Protocol 2 for the alkaline comet assay was based on the protocol described by Berthelot-Ricou et al. (2011a) for oocytes with some modifications to adapt the method for embryos. We first covered glass microscope slides with HMP agarose (1.6% in PBS) and then dried these slides overnight at 20 °C. A 50 μL drop of 1% LMP agarose in PBS at 37 °C was placed on the first agarose layer, and the embryos were immediately transferred from the culture medium into this drop (under a stereo microscope to visualize the embryos). The resulting suspension was then gently mixed with the 1% LMP agarose. After a few seconds of solidification, the agarose drop with the embryos was surrounded by an 80 μL ring of 1% LMP agarose, covered with a coverslip (24 × 32 mm) and then kept 20 min at 4 °C for solidification (the second layer). After removing the coverslip, a third layer of 80 μL of 1% LMP agarose was deposited and was covered with a coverslip (24 × 32 mm) before solidification at RT. After removing the coverslips, slides were lyed in a freshly prepared lysis solution (LS) (2.5 M NaCl, 100 mMol Na2EDTA, 10 mMol Tris-HCl, pH 10, 1% Triton X-100, and 10% DMSO in ultrapure water) for 90 min at 4 °C. DNA unwinding was performed with an alkaline solution (1 mMol Na2EDTA and 300 mMol NaOH) for 20 min at 20 °C in a 30.5 × 22 cm horizontal electrophoresis unit. Next, electrophoresis was conducted for 30 min (25 V, 300 mA) in the same buffer solution at RT. After the electrophoretic run, the slides were neutralized 3 times with 0.4 M Tris-HCl (pH 7.5), rinsed with ultrapure water, dipped into 100% methanol (HPLC grade purity solvent), and dried overnight at RT. Staining was performed with a propidium iodide solution (0.1 μg/mL), and the slides were examined with a fluorescence microscope at a 400 × final magnification (Carl Zeiss Axios Imager A2, Bayern, Germany). The digital pictures were analyzed, and the determination of the % tail DNA was performed with a CCD Nikon camera (DS-Fi2, Nikon, Champigny-sur-Marne, France) and Komet software (version 7.0, Nottingham, UK).

2.9. Statistical analysis

The % tail DNA was used to determine the DNA damage in each embryo. The experiments were replicated 5 times for each group. A t-test or Pearson’s Chi-squared test as appropriate for statistical analysis was performed using StatView software. The % tail DNA is expressed as the means ± standard error of the mean (S.E.M.). A probability of p < 0.05 was considered significant.

3. Results

3.1. Embryo survival and culture

After thawing and recovery in KSOM, the overall survival rate of the 5 separate experiments was 85% (85/100: 85 intact embryos and 15 lysed embryos). After a 24 h culture, the viability rate was 99% (84 2-cell embryos moving to the 4–8 cell stage out of 85 intact embryos). After a 48 h culture, the blastocyst rate was 82% (70 blastocysts obtained from in vitro culture of 85 intact 2-cell embryos).

3.2. Impact of the number of agarose layers on the embryos recovery rate

The overall recovery rate in the 5 different experiments using Protocol 1 (two agarose layers) was 3.3% (n = 5/150). With Protocol 2 (three agarose layers), the overall recovery rate was significantly increased to 71.3% (n = 107/150) (p < 0.0001).

3.3. Impact of the zona pellucida on the comet assay

The impact of the ZP on the Comet Assay is illustrated in Fig. 1. No statistically significant difference was observed in DNA damage for the ZP- and ZP+ groups (12.60 ± 2.53% tail DNA vs 11.04 ± 1.50, n = 150, p = 0.5837, respectively). The ZP− and ZP+ groups after H2O2 exposure were also not statistically significant different (49.23 ± 4.16% Tail DNA vs 41.13 ± 4.31, n = 150, p = 0.1820, respectively).

3.4. Alkaline comet assay

Fig. 2 shows the aspects of the comets observed in the control groups (A) and in the groups exposed to the genotoxic agents (B, C). The control embryos showed a low basal level of DNA damage. Fig. 3 shows that H2O2 and SSI induced a statistically significant increase in
is damaged, the two nuclei overlap and appear merged. (B, C). Arrows show the polar bodies. Scale bar, 100 μm. Magnification ×400.

4. Discussion

4.1. Thawed embryos can be used for comet assay

We describe, for the first time, the use of the alkaline comet assay on thawed mouse embryos by optimizing protocols that were previously published in the literature. Control mouse thawed embryos showed a low basal level of DNA damage, comparable to the level of DNA damage observed in control sperm (Preaubert et al., 2016) and control oocytes (Berthelot-Ricou et al., 2011a) in our hands. In the literature, we selected 14 studies using the comet assay on mammal embryos matching with our criteria, and no study was performed on frozen embryos. Among the 3 studies in mice (Fabian et al., 2003; Müller et al., 1996; Tranguch et al., 2003), only Fabian et al. evaluated the evolution rate to the blastocyst stage in embryos obtained by natural fertilization, and it was estimated to be 88%. This result suggests that the freezing-thawing step in our protocol did not affect the embryo quality. The use of frozen embryos simplifies the handling by allowing the researcher to dissociate the location and the timing of the embryo collection from the embryo analysis, which facilitates collaboration between laboratories. The use of frozen embryos also allows subsequent analysis if required.

Deleting the removal of zona pellucida improves feasibility.

Many authors remove the ZP before the comet assay, in order to study individual blastomeres and to allow DNA migration (Harrouk et al., 2000; Hwang et al., 2013; Ju et al., 2010; Müller et al., 1996; Sturme et al., 2009; Webster et al., 2000); nevertheless, they did not study the impact of the ZP on the downstream results. Moreover, this step requires the use of a chemical and physical stress that could unnecessarily damage the embryo and induce DNA damage. In a previous study of our group (Berthelot-Ricou et al., 2011a), we performed the comet assay on mouse oocytes with and without ZP, and no significant difference was found in the DNA damage in both of the groups. In the present study, we did not observe significant differences in DNA damage in the ZP- and ZP+ embryos, indicating that the presence of the ZP did not affect the embryo lysis and/or DNA migration. Deleting the ZP removal step simplifies the protocol, saving time and improving feasibility.

4.2. Adding a third layer of agarose decreases the embryo loss

One of the main issues of our study was to reduce embryo loss during lysis and electrophoresis. As the main difficulty of embryos studies is the low number of available samples, this issue was very important. The significant embryo loss we experienced with Protocol 1 (two agarose layers) was surprising. Indeed, we never observed such a significant loss during the comet assay with Protocol 1 on mouse oocytes (Berthelot-Ricou et al., 2011b, 2011a, 2013; Courbiere et al., 2013; Einaudi et al., 2014; Greco et al., 2015), which sizes are comparable to the embryo size. In the studies using the comet assay on embryos, selected by our literature review, the authors used protocols with 2 agarose layers and deposited between 1 and 20 embryos per glass slide (Table 2), which is comparable with Protocol 1 (20 embryos, 2 agarose layers). None of these studies described the ratio between the number of embryos deposited and the number of recovered embryos after lysis and electrophoresis. Nevertheless, we can assume that embryo loss was not as high as in the present study. Our hypothesis is that the important embryo loss we observed with Protocol 1 could be related to the alteration of ZP due to the freezing-thawing protocol: dehydration due to cryoprotectant could induce hardening and thinning of the ZP (Cavusoglu et al., 2016; Trounson and Mohr, 1983). These alterations could modify the interaction between the ZP and the agarose, and increase the embryo loss during the steps of coverslip removal and/or lysis. The use of a third agarose layer dramatically decreased the rate of embryo loss.
4.3. Limitations

In our study, we measured DNA damage by the % Tail DNA, as recently recommended by the OECD (2014). The use of the % Tail DNA gives a clear indication of the appearance of the comets, and it is also linearly related to the DNA break frequency over a wide range of levels of damage (Ersson and Möller, 2011; Hartmann et al., 2003; OECD, 2014; Uno et al., 2015). Therefore, the amount of DNA damage we observed could not be easily compared with the DNA damage in the other studies we selected. Indeed, the tail moment and/or the tail length measurements are also reported herein (Table 2), as these studies were published before the OECD recommendations. To better evaluate the sensitivity of our optimized protocol, despite the inability to compare these factors, we used two well-known genotoxic agents as positive controls. Nevertheless, % Tail DNA should be used for DNA damage quantification. Another limitation is that we performed the Comet Assay experiments on thawed embryos after an in vitro exposure to two well-known genotoxic agents; our protocol should also be tested to detect embryo DNA damage induced after in vivo mouse exposure associated to DNA damage in gametes (Einaudi et al., 2014; Singh et al., 2015).

5. Conclusion

Our optimized protocol of the comet assay on thawed embryos allows the rapid detection of primary DNA damage in the mouse embryo. We simplified the protocol by showing that the lack of ZP removal does not alter DNA lysis and migration. In addition, we demonstrated that the adjunction of a third agarose layer decreases embryo loss and allows the rapid detection of primary DNA damage in the mouse embryo. This new optimized tool for the study of a small amount of embryos. This new optimized tool could be used to analyze DNA damage in mice embryos after in vivo exposure to environmental agents. This new optimized tool could be used to analyze DNA damage in mice embryos after in vivo exposure to environmental agents.

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