Quantitative analysis of mitochondrial DNAs in macaque embryos reprogrammed by rabbit oocytes

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Abstract

In cloned animals where somatic cell nuclei and oocytes are from the same or closely related species, the mitochondrial DNA (mtDNA) of the oocyte is dominantly inherited. However, in nuclear transfer (NT) embryos where nuclear donor and oocyte are from two distantly related species, the distribution of the mtDNA species is not known. Here we determined the levels of macaque and rabbit mtDNAs in macaque embryos reprogrammed by rabbit oocytes. Quantification using a real-time PCR method showed that both macaque and rabbit mtDNAs coexist in NT embryos at all preimplantation stages, with maternal mtDNA being dominant. Single NT embryos at the 1-cell stage immediately after fusion contained 2.6 × 10⁴ copies of macaque mtDNA and 1.3 × 10⁶ copies of rabbit mtDNA. Copy numbers of both mtDNA species did not change significantly from the 1-cell to the morula stages. In the single blastocyst, however, the number of rabbit mtDNA increased dramatically while macaque mtDNA decreased. The ratio of nuclear donor mtDNA to oocyte mtDNA dropped sharply from 2% at the 1-cell stage to 0.011% at the blastocyst stage. These results suggest that maternal mtDNA replicates after the morula stage.


Introduction

Mammalian cells contain two distinct genomes; one is located in the nucleus, and the other in mitochondria (mtDNA). mtDNA is a double-stranded DNA circle of about 16–17 kb, which encodes only 13 proteins involved in the process of oxidative phosphorylation (Smith & Alcivar 1993). The rest of the proteins in the mitochondrial compartment are encoded by the nuclear genome. Interactions between nuclear and mitochondrial genomes are critical for the subsequent development of the embryo.

In normal mammalian fertilization, the sperm mitochondria are destroyed during preimplantation stages and the oocyte-derived mitochondria are transmitted to the offspring, so mtDNAs are inherited maternally or homoplasmically (Giles et al. 1980).

Thus far, the fate of foreign mtDNA following nuclear transfer (NT) is still controversial. In intraspecies cloned animals, mtDNA primarily arises from the oocytes (Evans et al. 1999, Takeda et al. 1999, Do et al. 2001), whereas in the others mtDNA appears to be heteroplastic (Steinborn et al. 2000, Hiendleder et al. 1999, Do et al. 2002). In interspecies cloning where nuclear donor and oocyte are from closely related species, mtDNAs are primarily oocyte derived (Loi et al. 2001, Meirelles et al. 2001, Lanza et al. 2002). However, our previous study (Chen et al. 1999) showed that the mitochondria from donor panda cells and recipient rabbit oocytes coexist in preimplanted embryos. Similarly, heteroplasmy of mtDNA has been identified in cloned macaque–rabbit embryos by direct PCR product sequencing (Yang et al. 2003). These results indicate that mtDNA of nuclear donor origin may persist in NT embryos in interspecies NT if the nuclear donor and oocyte are from distantly related species.

In the present study, the fate of mtDNAs in preimplantation stage macaque embryos derived from interspecies cloning was quantitatively analyzed. We examined copy numbers of both macaque and rabbit mtDNAs in cloned macaque–rabbit embryos by a real-time PCR assay.
Materials and Methods

Reconstruction of macaque–rabbit cloned embryos

Ear skin was collected from a 7-year-old female macaque (Macaca mulatta tcheliensis). Primary cell culture was performed with the same method as that described previously (Han et al. 2001). Fibroblasts at passages 4–10 were used as nuclear donors. Recipients were matured metaphase II stage (MII) rabbit oocytes flushed from oviducts with M2 medium (Sigma). Using the NT procedure described previously (Chen et al. 1999, 2002, Yang et al. 2003), MII oocytes were incubated for 10 min in M2 medium containing 7.5 µg/ml cytochalasin B (Sigma), 7.5 µg/ml Hoechst 33342 (Sigma) and 10% fetal bovine serum, and then manipulated under an inverted microscope (Nikon’s ECLIPSE TE300, Nikon Corporation, Japan). The first polar body with approximately one quarter to one third of the adjacent cytoplasm containing metaphase II spindle was removed. The aspirated karyoplast was exposed to u.v. light to confirm the removal of chromosomes. A single cell was then placed in the perivitelline space. The couplets were transferred into a fusion chamber overlaid with the electrical fusion solution comprising 0.25M sorbitol, 0.5 mM Hepes, 0.1 mM Ca(CH3COO)2, 0.5 mM Mg(CH3 COO)2 and 1 mg/ml bovine serum albumin. Cell fusion was induced with two direct current (DC) pulses (1.2 kV/cm, 20 s, and then 40 cycles at 95°C for 10 s, 68°C for 10 s, 72°C for 10 s. The rabbit mtDNA was amplified using the following parameters: 95°C for 10 min, followed by 40 cycles at 95°C for 10 s, 60°C for 10 s, 72°C for 10 s. The rabbit mtDNA was amplified using the following parameters: 95°C for 10 min, and then 40 cycles at 95°C for 10 s, 68°C for 10 s. Each experiment included standards for macaque/rabbit mtDNAs, NT embryos were transferred into G2.2 medium (Vitrolife) for subsequent development.

Collection of NT embryos

NT embryos were collected directly into PCR tubes containing 10 µl lysis solution, ReadyAmp genomic DNA purification resin (Promega Corporation, Madison, WI, USA) and 200 µg/ml added proteinase K (Sigma). The mixtures were incubated for 30 min at 55°C, boiled for 10 min, centrifuged for 1 min, and then used as PCR templates (Wan et al. 2003).

Species-specific mtDNA primers

Sequences of macaque (GeneBank accession number U38273) and rabbit (GeneBank accession number NC-001913) mtDNAs were compared with design species-specific primers. The macaque-specific primers were forward, 5'-CCT TCT CCT CAA TCG CAC-3' and reverse, 5'-GGA GAT ATG AGC CGT AG-3'. The rabbit mtDNA specific primers were forward, 5'-CCC ATA CGA CTA TCC CTC TCC C-3' and reverse, 5'-CGT GTG GCC GAT CTT AGG TTC-3'.

Standard curves

PCR products were purified from agarose gel and inserted into pGEM-T-easy PCR cloning vector (Promega Corporation). The identities of the inserts were confirmed by sequencing. Standard curves covering 10^6 to 10^9 copies (with 10-fold dilutions) for macaque mtDNA and 10^2 to 10^9 copies for rabbit mtDNA were constructed (Fig. 1).

Real-time PCR

Embryos at seven preimplantation stages, including 1-cell (immediately after fusion), 2-cell, 4-cell, 8-cell, 16-cell, morula and blastocyst stages, were collected. Five embryos were analyzed for each of the seven stages. To perform assays, lysates from five embryos were mixed and used as templates. Each mixed sample was repeatedly measured three times using macaque-specific primer pairs and rabbit-specific primer pairs respectively. Unknown concentrations of samples were extrapolated by comparing with standards amplified under the same conditions using Lightcycler software 3.1 (Roche Diagnostics, Germany). The means and standard errors of triplicates were calculated and are presented in Table 1.

PCR reactions were prepared as follows: 6.4 µl distilled water, 1.2 µl MgCl2 (25 mM), 0.2 µl forward primer (20 µM), 0.2 µl reverse primer (20 µM), 1 µl NT embryo DNA template, 1 µl reagent mixture including dNTP, Taq polymerase and SYBR Green dye (Roche diagnostics, Germany). Real-time PCR assays were performed on a lightcycler (Roche diagnostics, Germany). The macaque mtDNA was amplified using the following parameters: 95°C for 10 min, followed by 40 cycles at 95°C for 10 s, 60°C for 10 s, 72°C for 10 s. The rabbit mtDNA was amplified using the following parameters: 95°C for 10 min, and then 40 cycles at 95°C for 10 s, 68°C for 10 s. Each experiment included standards for macaque/rabbit mtDNAs, NT embryos were transferred into G2.2 medium (Vitrolife) for subsequent development.

Figure 1

Regression lines of the standards from 10^6 copies/µl to 10 copies/µl, with log copy numbers plotted on the X axis and the threshold cycle on the Y axis. (A) Macaque mtDNA standards and (B) rabbit mtDNA standards.
embryo, rabbit/macaque DNA controls, lysis buffer control and water control. Identities of PCR products were confirmed by their sizes in agarose gel and by sequencing.

Results

Specificity of the assay

In a cross-species test, the macaque mtDNA-specific primers detected PCR product in only macaque mtDNA but not in rabbit and human mtDNA; vice versa, the rabbit mtDNA-specific primers detected PCR products only in rabbit mtDNA but not in macaque and human mtDNA. The two primer pairs were species specific and therefore used to construct standard curves and to detect mtDNAs derived from donor cell and recipient oocyte in NT embryos at different stages.

Dynamic changes of mtDNA in NT embryos

The copies of mtDNA from donor cells and recipient oocytes in cloned embryos at 1-cell, 2-cell, 4-cell, 8-cell, 16-cell, morula and blastocyst stages were detected separately using the rabbit-specific and macaque-specific primer pairs. Results in triplicate showed that rabbit/macaque DNA controls, lysis buffer and water control were negative and the macaque/rabbit DNA standards were well proportioned (Fig. 2), and so the detected mtDNA copies were authentic. In the 1-cell stage NT embryo, the number of mtDNA copies from donor macaque somatic cells was \(2.6 \times 10^4\), while there were \(1.3 \times 10^6\) mtDNA copies from recipient rabbit oocytes. In subsequent in vitro development of NT embryos, from the 2-cell stage to the morula stage, there were no significant changes in copy numbers of both mtDNA species. At the blastocyst stage, however, the copy number of macaque mtDNA was reduced to \(5.4 \times 10^3\) while the copy number of rabbit mtDNA was increased to \(4.7 \times 10^7\), bringing the ratio of macaque/rabbit mtDNAs from 2% at the 1-cell stage to 0.011% at the blastocyst stage (Table 1). These results demonstrated that the mitochondria from both macaque and rabbit coexisted in these early stage NT embryos, which is consistent with our previous qualitative PCR analysis of mitochondrial fate in cloned macaque–rabbit embryos (Yang et al. 2003).

Table 1 The mtDNA copy number and ratios of macaque mtDNA to rabbit mtDNA in NT embryos at different stages. Values are means of triplicates.

<table>
<thead>
<tr>
<th>Stages</th>
<th>Macaque mtDNAs</th>
<th>Rabbit mtDNAs</th>
<th>Macaque mtDNA/rabbit mtDNA (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-cell</td>
<td>(2.6 \times 10^4) ± 1.7 \times 10^4)</td>
<td>(1.3 \times 10^6) ± 0.7 \times 10^6)</td>
<td>2.00 ± 0.83</td>
</tr>
<tr>
<td>2-cell</td>
<td>(2.8 \times 10^4) ± 1.8 \times 10^4)</td>
<td>(2.1 \times 10^6) ± 1.3 \times 10^6)</td>
<td>1.33 ± 0.57</td>
</tr>
<tr>
<td>4-cell</td>
<td>(2.5 \times 10^4) ± 2.0 \times 10^4)</td>
<td>(1.5 \times 10^6) ± 1.0 \times 10^6)</td>
<td>1.67 ± 1.27</td>
</tr>
<tr>
<td>8-cell</td>
<td>(1.0 \times 10^4) ± 0.8 \times 10^4)</td>
<td>(1.6 \times 10^6) ± 0.9 \times 10^6)</td>
<td>0.63 ± 0.34</td>
</tr>
<tr>
<td>16-cell</td>
<td>(2.1 \times 10^5) ± 1.3 \times 10^5)</td>
<td>(2.5 \times 10^6) ± 1.3 \times 10^6)</td>
<td>0.84 ± 0.16</td>
</tr>
<tr>
<td>Morula</td>
<td>(2.8 \times 10^4) ± 1.8 \times 10^4)</td>
<td>(2.2 \times 10^6) ± 1.0 \times 10^6)</td>
<td>1.27 ± 0.50</td>
</tr>
<tr>
<td>Blastocyst</td>
<td>(5.4 \times 10^3) ± 3.5 \times 10^3)</td>
<td>(4.7 \times 10^7) ± 3.5 \times 10^7)</td>
<td>0.011 ± 0.007</td>
</tr>
</tbody>
</table>

Confirmation of PCR products

PCR products from both mtDNA species were analyzed by electrophoresis and showed the expected sizes (143 bp for macaque mtDNA and 196 bp for rabbit mtDNA). The identities of these bands were further confirmed by sequencing (data not shown).

Discussion

The method for studying the mitochondrial fate in NT embryos

During cloning manipulation by either whole-cell electrosurgery or karyoplast injection, donor nucleus with all or part of the cytoplasm is transferred into an enucleated recipient oocyte. Different cytoplasm from a donor cell and an enucleated recipient oocyte are mixed. Thus, the NT embryos should harbor two kinds of mtDNA. mtDNAs are the genetic material of mitochondria, a component of cytoplasm. In the early development of NT-reconstructed embryos, the detection of mtDNA is helpful to reveal changes of donor cytoplasm. Several methods have been used to explore the fate of mitochondria in cloning, including single-strand conformation polymorphism of PCR fragments and allele-specific PCR (AS-PCR) in cloned cattle (Takeda et al. 1999, Do et al. 2002), direct DNA sequencing in interspecies cloned panda embryos (Chen et al. 2002), DNA chromatography in cloned bovine embryos (Do et al. 2002) and restriction fragment length polymorphism in cloned sheep (Evans et al. 1999). Of these methods, AS-PCR is considered to be the most sensitive and suitable method for the analysis of mtDNA heteroplasmy. However, all these identification methods can be used to qualify the homoplasmy or heteroplasmy only, but not to quantify the number of mtDNAs. In the present study, real-time quantitative PCR was used to detect the slight changes of mtDNA in NT embryos accurately and the results demonstrated that this method can amplify \(10^1\) copies, sometimes even \(10^6\) copies, of macaque or rabbit mtDNA standard (as shown in Fig. 2). Thus, this method with sensitivity to detect less than ten copies of mtDNA also can be extended to nuclear gene detection.

In order to perform real-time PCR, an effective DNA extraction protocol from a single early NT embryo should...
be developed. We previously used a very simple and rapid approach described by Wan et al. (2003) for qualitative mtDNA analysis of macaque–rabbit NT embryos by direct DNA sequencing. The method of DNA extraction, by adding 100 μg/ml proteinase K and 1% Triton X-100 into 1× PCR buffer, proved to be effective in common PCR. However, when the approach was used in real-time PCR in this study, the high concentration of Triton X-100 in a raw DNA mixture decreased the activity of Taq DNA polymerase (Roche Molecular Biochemicals) and affected the accurate quantification of mtDNA. We changed the protocol and used the ReadyAmp genomic DNA purification resin containing 200 μg/ml proteinase K as the DNA extraction mixture. Results showed that this is an efficient and simple DNA extraction technique for quantitative PCR analysis of a single embryo. This method not only ensures the complete extraction of mtDNA but also skips a phenol/chloroform extraction step.

The mtDNA copies in NT embryos

The study of mitochondrial fate in NT embryos is helpful to reveal the mechanism of epigenetic reprogramming. Several reports have shown that there are $1.59 \times 10^5$ and $3.14-7.95 \times 10^5$ mtDNAs molecules per mouse and human oocyte (Steuerwald et al. 2000, Barritt et al. 2002) respectively, while there are $2-5 \times 10^3$ mtDNAs in a sheep (Evans et al. 1999) or human (King & Attardi 1989) somatic cell. The ratio of nuclear donor cell mtDNA to recipient cytoplast mtDNA is about 2–5% in the reconstructed 1-cell embryos immediately after fusion. Steinborn et al. (2000) quantified the percentage of donor mtDNA in cloned cattle by allele-specific real-time PCR. They observed heteroplasmy in seven of ten cattle clones with the donor-to-recipient ratios ranging from 0.4% to 4% and the ratios remained the same throughout development to term, while the other three clones showed a significant reduction or absence of donor mtDNA at ratios ≤0.3%. In

Figure 2 Analysis of mtDNA heteroplasmy. (A) and (B) show log linear phases of real-time PCR reactions of macaque and rabbit mtDNAs respectively. The Lightcycler software 3.1 extrapolates unknown concentrations from the threshold cycles of the standards.
In the present study, we estimated the copies of mtDNA in early macaque–rabbit NT embryos at different stages by real-time PCR. A single embryo contains $0.54 - 2.8 \times 10^4$ macaque mtDNAs and $0.13 - 4.7 \times 10^7$ rabbit mtDNAs, values higher than those in early human embryos (King & Attardi 1989). A previous study found that there was no mtDNA replication until the blastocyst stage in the mouse (Piko & Taylor 1987). In the present study, we found that the mtDNA copy number in macaque–rabbit cloned embryos from the 1-cell to the morula stage did not change significantly; however, at the blastocyst stage, the mtDNA copies derived from macaque donor cells were reduced and those from recipient rabbit oocytes were significantly increased, suggesting that the maternal mtDNA in the NT embryo replicates after the morula stage. Our results support the conclusion that mtDNA starts replicating at the blastocyst stage. Due to the reduction of donor macaque rabbit mtDNAs and the increase of recipient mtDNAs, the ratio of nuclear donor mtDNA to recipient cytoplasm mtDNA also decreased from 2% at the 1-cell stage to 0.011% at the blastocyst stage. Our results suggest that mtDNAs from donor macaque cells and those from enucleated recipient rabbit oocytes coexist in NT embryos from the 1-cell to the blastocyst stage, which is consistent with the result obtained by direct PCR product sequencing (Yang et al. 2003).

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References


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