rRNA Genes Are Not Fully Activated in Mouse Somatic Cell Nuclear Transfer Embryos*1,2

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Background: The nucleolus always shows delayed development and malfunction in somatic cell nuclear transfer (NT) embryos. NT embryos showed low rDNA activity and developmental competence when donor cells with low rDNA activity were used. rDNA reprogramming efficiency in NT embryos was determined by the rDNA activity in donor cells from which they derived. Developmental potential of NT embryos with rDNA activity in the donor cells was correlated.

Eukaryotic ribosomal genes (rDNA) are arranged in clusters of tandem repeats known as nucleolar organizer regions (NORs),2 which locate on the short arm of acrocentric chromosomes (1). Active NORs are associated with ribosomal RNA (rRNA) transcription machinery including upstream binding factor (UBF) and RNA polymerase I (RPI). These regions often stain positive for silver and are, therefore, called Ag-NORs. NT embryos showed low rDNA activity and developmental competence when donor cells with low rDNA activity were used. rDNA reprogramming efficiency in NT embryos was determined by the rDNA activity in donor cells from which they derived.

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The well known and most important function of nucleoli is ribosome biogenesis. However, the nucleolus showed delayed development and malfunction in somatic cell nuclear transfer (NT) embryos. Previous studies indicated that nearly half rRNA genes (rDNA) in somatic cells were inactive and not transcribed. We compared the rDNA methylation level, active nucleolar organizer region (NORs) numbers, nucleolar proteins (UBF, nucleophosmin (B23)) distribution, and nucleolar–related gene expression in three different donor cells and NT embryos. The results showed embryonic stem cells (ESCs) had the most active NORs and lowest rDNA methylation level, whereas mouse embryonic fibroblasts (MEFs) were the opposite. After the donor cells were injected into enucleated MII oocytes, admixed cell nuclei lost B23 and UBF signals in 20 min, whereas in ESC-NT embryos, B23 and UBF signals could still be detected at 60 min post-NT. The embryos derived from ESCs, collums, cumulus cells, and MEFs showed the same trend in active NORs numbers and rDNA methylation levels. As an overall pattern, functional nucleoli are established in the oocyte upon activation of oocyte growth, which provided enough ribosome content for RNA and protein synthesis at a high level to support early development (5). By the end of oocyte growth, rRNA synthesis ceases. The nucleoli disassemble and change into non-functional nucleolar remnants (6, 7). It is intriguing that mouse oocytes destroy a large amount of rRNA, ribosomal protein, and mRNAs during maturation (8).

Significance: Developmental potential of NT embryos with rDNA activity in the donor cells was correlated.

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So the maternal ribosome storage is evidently not sufficient to support embryonic development beyond the zygotic genome activation. Then in late 2-cell stage, the zygotic genome activation becomes apparent (9), and genes involved in ribosome biogenesis and assembly are included in the first gene activation burst (10). However, fibrillo-granular nucleoli with full rRNA synthesis, processing, and ribosome subunit assembly functions are only established at the 4-cell stage in mouse (11), which then provides a large amount of ribosomes for early development. Therefore, the reactivation of rDNA during zygotic genome activation and establishment of full functional nucleolus at the 4-cell stage are very important to preimplantation development of mouse embryos.

Actually, in mammalian cells only ~50% rDNA is transcribed (1), with the remaining genes inactivated through a combination of different epigenetic mechanisms including methylation (12). With the new cytological technique, researchers indicated that rDNA is regulated allelically in a regional manner in human, with one parental copy of each NOR repressed in any individual cell, which is similar to X-chromosome inactivation in females (13). Whether the same mechanism exists in other species is unknown. When the somatic cell was fused into bovine enucleated oocyte, the nucleolus progressively transformed into nucleolar bodies and lost vacuoles structure (14). The NT embryo needs to rebuild full functional nucleolus, which is important to preimplantation development (15). However, the inactive rDNA copies in donor cells may become a barrier for full activation of rDNA in NT embryos. Recently, several studies reported that the nucleolus biogenesis in NT embryos was abnormal. In bovine granulosa cell nuclear transfer embryos, the reformation of fibrillo-granular nucleoli was initiated one cell cycle earlier than that in in vivo derived embryos. But the nucleolus compartment was one cell cycle later, which indicated the NT embryos were lacking in development potentials (16). In pig, only half late 4-cell fibroblast NT embryos had transcriptionally active nucleoli, whereas in in vivo embryos the portion was 92% (17). Moreover, in mouse embryonic fibroblast or stem cell-cloned embryos, the activation of functional nucleolus was also one cell cycle-delayed (18).

Because the reprogramming competence of oocyte to somatic cell nuclear is limited, we wonder whether all those inactive rDNA/NORs could be fully activated at the 4-cell stage in mouse NT embryos when compared with normal ones. If not, will it impair ribosome synthesis and intracellular metabolism of early embryonic development? Besides, donor cells with a different differentiation status would yield different outcomes in somatic cell cloning experiments (19–21). We also want to know if the rDNA epigenetic status in donor cells will result in different RNA synthesis and processing activities in NT embryos and, furthermore, affect preimplantation developmental competence.

In this study we chose mouse embryonic cells (ESCs), cumulus cells (CCs), and embryonic fibroblast cells (MEFs) as donor cells to reconstruct different NT embryos. Intracytoplasmic sperm injection (ICSI) embryos were used as control. The rDNA methylation level, active NORs numbers, and nucleolar-related gene expression were compared in donor cells and in corresponding NT embryos at different preimplantation development stages. The distribution of nucleolar protein (B23 and UBF) was also compared before and after NT.

EXPERIMENTAL PROCEDURES

Animal—B6D2F1 (C57BL/6 × DBA/2) female/male mice were obtained at 8–10 weeks of age from Vital River (Beijing, China). Animals were conformed to the Guide for the Care and Use of Laboratory Animals. All animal experiments were performed under the Code of Practice Harbin Medicine University Ethics Committees.

Cell Culture and Treatment—Derivation and culture of mouse ESCs were according to a previous protocol (22). Cells were cultured in DMEM containing 15% FBS, 50 μg/ml penicillin/streptomycin (Invitrogen, 15140-148), 100 μM nonessential amino acids (Invitrogen, 11140-050), 100 μM β-mercaptoethanol (Sigma, M7522), and 1000 units/ml leukemia inhibitory factor (Chemicon, ESG1107). The medium were changed every day, and the cells were passed every 2 days. The day before nuclear transfer, 3 μg/ml nocodazole (Sigma, M1404) was added to culture medium overnight to synchronize the cells to metaphase (23, 24). Then the cells were harvested and used as donor cells. All the ESCs used in this study were within 10 passages.

CCs were obtained during oocyte collection, then washed in HEPES-buffered CZB medium (HEPES-CZB) several times and cultured in DMEM containing 50 ng/ml FSH (Sigma, F2297) and 20 ng/ml EGF (Sigma, E4127). Partial cells were resuspended in HEPES-CZB containing 3% PVP (Polyvinylpyrrolidone, Sigma, PVP360) and used as donor cells (G0/G1) for NT directly.

MEFs were isolated from 13.5 post-coitum B6D2F1 mouse fetus as previous reported (25). Cells were cultured in DMEM containing 10% FBS under 5% CO2 in humidified air at 37 °C within three passages. Partial MEFs were treated with 10 μg/ml mitomycin C (Sigma, M4287) for 2.5 h then used as feed layers for ES cells. MEFs at the third passage were kept in DMEM containing 0.5% FBS for 72 h as serum starvation treatment and then used as NT donor cells (G0/G1).

Metaphase Chromosome Spreads Preparation, Karyotyping, and Silver-staining of Donor Cells—The ESCs, CCs, and MEFs were kept in culture medium containing 3 μg/ml nocodazole for 1, 12, and 12 h, respectively, to synchronize the cells to metaphase. Then the cells were made into metaphase chromosome spreads as previous reported (26). For karyotype observation, several slides of each group were stained with 4% Giemsa in PBS for 10 min. For silver-staining of active NORs, slides were first incubated in a humidity box at 37 °C for at least 30 min and then stained with fresh mixed silver-stain medium (2% agar medium with 1% formic acid, 50% AgNO3 medium (1:2)) in the dark at 40–50 °C for 8 min. The slides were washed, dried, mounted, and finally observed with a 100× oil-immersion objective.

Oocyte Recovery and Spermatozoa Preparation—Female B6D2F1 mice were superovulated by intraperitoneal injections of 5 IU pregnant mare serum gonadotropin (NSH, China) followed 48 h later by 5 IU of human chorionic gonadotropin (NSH, China). Oocytes were collected from the oviducts 14 h
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after human chorionic gonadotropin injection. After collection, cumulus cells were removed from oocytes with 300 μg/ml hyaluronidase (Sigma, H4272) in droplets of HEPES-CZB by gentle pipetting. Denuded oocytes with homogeneous ooplasm were selected and kept in new droplets of CZB medium containing 5.6 μM glucose (CZBG), covered with sterile mineral oil (Fisher, O121-20), then cultured at 37 °C in a 5% CO2 atmosphere until use. Spermatozoa were collected from the cauda epididymis of 8–12 weeks age of B6D2F1 males, then kept in CZB-HEPES medium and prepared for injection.

Generation of ICSI and NT Embryos—ICSI was carried out by a piezo-driven unit using the methods as described elsewhere (27, 28), except that our experiment was performed in HEPES-CZB containing 5 μg/ml cytochalasin B (Sigma, C6762) at room temperature (29). Only the sperm head was injected into oocyte. After 10–20 min of recovery, the ICSI-generated embryos were washed several times and cultured in K-modified simplex optimized medium at 37 °C in a 5% CO2 atmosphere. To eliminate any possible effects brought by NT methods to the following investigation, we adopted the same one-step micro-manipulation technique to reconstruct ESCs nuclear transfer (ESNT), CCs nuclear transfer (CCNT), and MEFs nuclear transfer (MEFNT) embryos as described previously (30) with modifications. The outer diameters of injection pipettes used for ESCs, CCs, and MEFs were 13, 9, and 12 μm, respectively. Briefly, the donor cell membrane was broken with several Piezo pulses, then 5–10 cells were sucked into the injection pipette. The oocyte MII spindle was adjusted to 8–10 o’clock, then one donor cell was injected into the nearby plasma. The spindle was sucked into the injection pipette immediately and taken out of oocyte. One hour after NT, the reconstructed CCNT and MEFNT embryos were activated with 5 mM SrCl2 (Sigma, WH00007343M1, 1:100) overnight at 4 °C followed by 105 copies of XenoTm RNA from SYBR Green Cell-to-Cell™ Control kit (Invitrogen, 4402959) during cell lysis as external control. The total RNA of cells and embryos was extracted by RNeasy Mini kit (Qiagen, 74104) according to the manufacturer’s instructions. cDNA was synthesized from total RNA with a High Capacity cDNA Reverse Transcription kit (ABI, 4368814). qPCR reactions were performed based on 1.5 μl of cDNA sample, 10 μl of TransStart™ Top Green qPCR SuperMix (TransGen, AQL131), and gene-specific primers (supplemental Table 1) in a 20-μl reaction system on CFX96 Real-time System (Bio-Rad). H2afz, Hprt1, and ExnoTm were used as candidate references for embryos (31, 32). HP1γ, cyclophilin-A, and cyclophilin-B were used as candidate references for cells (13).

Detection of rDNA Methylation in Donor Cells and 4-Cell Embryos—The 5’-ETS sequence (81–507, GenBank™ BK000964.1) was used as the target sequence for detection of rDNA methylation levels (13). The experiment was conducted with the EZ DNA Methylation-Direct™ kit (ZYMO Research, Irvine, CA) according to the manufacturer’s instruction. Around 10,000 cells for each cell group and 70–80 embryos for each 4-cell embryo group were used as samples. The samples were digested with proteinase K, then the unmethylated cytosines were converted to uracil by bisulfite. The target sequence was amplified by PCR and subjected to sequence analysis. The primers sequences used were TATTTATTTTTTTTTTGGTGT and TAAAGAACACTTTAACACACCAAC (reverse).

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Experiment Design and Statistical Analysis—Each experiment was repeated at least three times. More than 5 slides and 50 metaphase plates were checked for each group in karyotype and silver-stain experiment. More than five embryos in each group were selected for nucleolar protein observation by confocal microscope. More than 20 4-cell embryos blocked at metaphase in each group were observed for detection of UBF spots. The data were compared using the Least Significant Difference test after analysis of variance in SPSS 13.0. In qPCR experiments, 20 2-cell embryos, 15 4-cell embryos, 10 morulas, and 10 blastocysts of each group were used as one repeat. The raw data were first imported to qBasePLUS 2.2 software (Bio-gazelle) for reference gene selection and data calculation. Then the relative gene expression in ICSI embryos was set to 1. Finally, the relative amount of gene expression was compared using the Least Significant Difference test after analysis of variance in SPSS. A value of p < 0.05 was considered statistically significant.
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RESULTS

rDNA Activity in ESCs, CCs, and MEFs—We first examined the karyotypes of ESCs, CCs, and MEFs. All the 3 cell lines showed normal karyotypes that contained 40 acrocentric chromosomes (Fig. 1, A–C). The metaphase slides were stained with AgNO₃ to show the active NORs. At interphase, the silver-stain signals showed as big brown clusters in the nuclei in all the three cell lines (A”, B”, C”). At metaphase, the silver-stain signals showed as brown spot pairs on certain chromosomes (arrows). ESCs had more spot pair numbers than CCs and MEFs at metaphase (A’, B’, C’). After injection, the B23 and UBF signals in reconstructed embryos distributed in the whole cytoplasm (Fig. 3, A–D) as mentioned previously (H). We then studied the distribution of nucleolar protein UBF in reconstructed embryos 1 h after NT—Because several differences were observed among the donor cells, we then checked the B23 and UBF distribution after NT to study whether they have different responses after being injected into ooplasm. Our pre-experiments showed the B23 and UBF signals of cumulus cell nucleoli disappeared within 1 h by 1 μg/ml actinomycin D (an RPI inhibitor) treatment (data not shown). We then fixed the embryos at 0, 20, 40, and 60 min post-NT/sperm injection. The results indicated that B23 and UBF signals co-localized within donor cell nuclear, except for B23 in the ESNT group (Fig. 5, E, F, G, and H). Immediately after injection, the B23 and UBF signals in reconstructed embryos displayed the same distributions as seen in cells. The B23 and UBF signals vanished 20 min after NT in CCNT and MEFNT embryos (Figs. 5, I, J, M, and N, and 6 I, J, M, and N), and indicated the rRNA transcription and processing machineries were immediately disassembled in ooplasm. In ESNT embryos, both signals could still be detected at 60 min post-NT (Figs. 5H and 6H), although the signals were getting weaker over time. As a comparison, no B23 or UBF signal could be detected at the injected sperm head or maternal chromosomes in ICSI embryos (Figs. 5, A–D, and 6, A–D).

B23 and UBF Distribution in Reconstructed Embryos during Preimplantation Development—We also studied the distribution of B23 and UBF in ICSI, ESNT, CCNT, and MEFNT embryos at pronuclear, 2-cell, 4-cell, and morula stages. B23 recruited in pronuclei at 6 hpa, and the signal became stronger.
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FIGURE 2. Active NORs in donor cells and in 4-cell embryos. Metaphase spreads of ESCs, CCs, and MEFs were stained with AgNO₃, then the active NORs numbers were counted (A). ICSI, ESNT, CCNT, and MEFNT embryos at the 4-cell stage were blocked to metaphase by nocodazole treatment, then the UBF signal spots were labeled and counted (B). Columns with different letters indicate significant difference (p < 0.05). ESCs had significantly more active NORs than CCs and MEFs (7.66 versus 6.45 versus 4.70, p < 0.05), and MEFs had the least. ICSI and ESNT embryos had no significant difference in active NORs numbers at the 4-cell stage, but their active NORs numbers were significantly more than CCNT and MEFNT embryos (7.44 versus 7.19 versus 6.68 versus 5.77).

FIGURE 3. Distribution of B23 and UBF in ESCs, CCs, and MEFs. B23 and UBF were labeled by monoclonal anti-B23 antibody/monoclonal anti-UBF antibody, respectively, and fluorescent conjunct second antibody (green). The nuclei/chromosomes were stained with Hoechst 33342 (blue). Typical areas (dotted line rectangle) were magnified to show the details (solid line rectangle). B23 showed as three or four condensed clusters within the nucleus in CCs (F’). In MEFs, B23 signals showed as 1–3 big rings (J’). ESCs showed much a stronger B23 signal with an irregular shape (A). At metaphase, the B23 signal distributed to the whole cytoplasm (B’). The UBF signal showed as several clusters in the nuclei of all the three groups of cells (C, G, and K). During mitosis, the UBF signal showed as pairs of small spots in metaphase plate (C’) or spindle polar (K’).

FIGURE 4. Nucleolar-related gene expression in ESCs, CCs, and MEFs. The expression level of ESCs was set as 1. Different letters indicate significant differences at p < 0.05.

at 8 hpa because this protein was inherited and reused in early cleavage (33). There were no obvious differences among the four groups. Meanwhile, UBF could not be detected at pronuclear stage (supplemental Figs. 2 and 3). At metaphase of the first mitosis (17 hpa), B23 signals distributed to the whole cytoplasm (Fig. 7, A, E, I, and M). No UBF signals could be detected on metaphase chromosomes of each group (Fig. 8, A, E, I, and M). The B23 protein re-accumulated in nuclear at 2-cell stage (24 hpa). As blastomere started cleavage, B23 distributed to the whole plasma again (Fig. 7, B, F, J, and N). Meanwhile, no UBF signal could be detected in ESNT, CCNT, and MEFNT embryos at this stage (Fig. 8, F, I, and N). However, several ICSI embryos (3 in 5) showed weak UBF spots around the nucleoli (Fig. 8B’), which indicated they may start to synthesis rRNA earlier than NT counterparts. At 4-cell and morula stage, all groups showed ring-like B23 signals around nucleoli without any obvious differences (Fig. 7, C, D, G, H, K, L, O, and P). UBF signals showed as discontinuous small clusters around nucleoli at the 4-cell stage (Fig. 8, C, G, K, and O). At the morula stage, many blastomeres were under mitosis, and the UBF signals distributed in the whole cytoplasm with chromosomes or at the spindle poles (Fig. 8, D, H, L, and P). No obvious differences were observed among the four groups at the 4-cell or morula stages.

Active NORs Numbers and rDNA Methylation Level in 4-Cell Stage Embryos—Although silver stain is an effective method to distinguish the transcription active NORs from inactive ones, for 4-cell embryos it was difficult to prepare metaphase plates and conduct silver-stain procedures. Because UBF remain associated with active NORs throughout mitosis (5, 34), we then blocked the 4-cell embryos to metaphase by nocodazole and detected the UBF signals as an alternative way to count the active NORs. As shown in Fig. 9, in every blastomere at metaphase, the UBF signals strictly adhered to certain chromosomes and were revealed as bright signal spots in pairs. After comparison of the pair numbers of UBF spots in each group, we found ICSI and ESNT embryos owned significant more pair numbers than CCNT and MEFNT embryos (7.44 and 7.19 versus 6.68 versus 5.77).
FIGURE 5. Changing of B23 signal in NT and ICSI embryos during a 1-h post-NT/sperm injection. B23s were labeled by monoclonal anti-UBF antibody and fluorescent conjugate second antibody (green). The donor cell chromatin was stained with Hoechst 33342 (blue). Arrowheads indicate the donor cell chromatin/sperm head. The B23 signals in CCNT and MEFNT were showed as several obvious clusters (I and M). But after 20 min, the signals got much weaker (J and N). No B23 signal could be detected after 40 min (K, L, O, and P). The B23 in ESNT group lasted for 60 min, although the signal got weaker over time, but the signal was not companied with chromatins (E–H). No B23 signal could be detected at sperm head or maternal chromatins (A–D).

FIGURE 6. Changing of UBF signal in NT and ICSI embryos during a 1-h post-NT/sperm injection. UBFs were labeled by monoclonal anti-UBF antibody and fluorescent conjugate second antibody (green). The donor cell chromatin was stained with Hoechst 33342 (blue). Arrowheads indicate the donor cell chromatin/sperm head. The UBF signals in CCNT and MEFNT were showed as several obvious clusters (I and M). After 20 min, their signals got much weaker (J and N). Almost no B23 signal could be detected at and after 40 min (K, L, O, and P). The UBF in ESNT group showed as small bright spots in pairs and lasted for 60 min (E′–H′). No UBF signal could be detected at sperm head or maternal chromatins (A–D).
FIGURE 7. Distribution of B23 in NT and ICSI embryos at different preimplantation developmental stages. B23s were labeled by monoclonal anti-B23 antibody and fluorescent conjugated secondary antibody (green). The nuclei/chromosomes were stained with Hoechst 33342 (blue). B23 distributed to the whole cytoplasm at metaphase of the first mitosis (A, E, I, and M) and then recruited in the nuclei at the 2-cell stage (B, F, J, and N). When the blastomere started to cleavage, B23 distributed to the cytoplasm again (N). All the four groups of embryos showed ring-like B23 signals around the nucleoli at 4-cell and morula stages (C, D, G, H, K, L, O, and P).

FIGURE 8. Distribution of UBF in NT and ICSI embryos at different preimplantation developmental stages. UBFs were labeled by monoclonal anti-UBF antibody and fluorescent conjugated secondary antibody (green). The nuclei/chromosomes were stained with Hoechst 33342 (blue). No UBF signal could be detected at metaphase of the first mitosis (A, E, I, and M). At the 2-cell stage, no UBF signal could be detected in ESNT, CCNT, and MEFNT embryos (B, F, J, and N). But several ICSI embryos showed weak UBF spots around the nucleoli (B'). At the 4-cell stage, the UBF signals showed as discontinuous small clusters around nucleoli (C, G, K, and O). At the morula stage, many blastomeres were under mitosis, and the UBF signals distributed in the whole cytoplasm with chromosomes or at the spindle pole (D, H, L, and P).
versus 5.77, p < 0.05). However, there was no significant difference between ICSI and ESNT groups (Fig. 2B).

rDNA methylation status of 4-cell embryos were also evaluated. The rDNA methylated level at 5’-ETS sequence of ICSI, ESNT, CCNT, and MEFNT embryos were 9.38, 6.36, 9.67, and 15.52 %, respectively (see Fig. 10). MEFNT embryos still had the highest methylation level, indicating those methylated rDNA copies in donor cells were not fully activated at 4-cell stage.

Preimplantation Development and Nucleolar-related Gene Expression of ICSI and NT Embryos—We then compared preimplantation development rates of the four groups of embryos. ICSI embryos had the highest development rate at every preimplantation developmental stage, reaching 97% in blastocyst rate. Among the NT groups, ESNT embryos had significantly higher morula and blastocyst rates (80.7 and 69.3%) than that of CCNT (72.2 and 62.0%) and MEFNT (60.2 and 35.6%) embryos (supplemental Table 2). MEFNT had the lowest development potential after the 2-cell stage and, moreover, showed an obvious drop of development rate from morula to blastocyst stage (Fig. 11).

Finally, we compared the nucleolar-related gene expression of these embryos at different preimplantation developmental stages. Hrpt1 and Exno™ were used as internal and external reference genes, respectively, according to geNormPLUS analysis. At the 2-cell stage, ICSI embryos had significantly higher UBF and RPI expression levels than NT groups, which was consistent with earlier immunofluorescent results that UBF could only be detected in ICSI embryos at the 2-cell stage. The ESNT group showed significant higher 47 S rRNA expression than CCNT embryos and significantly higher RPI levels than CCNT and MEFNT counterparts. There were no significant differences in 18 S rRNA, UBF, FBL, and B23 expression levels among the NT groups (supplemental Fig. 4). At the 4-cell stage, ICSI embryos had significant higher 47 S and FBL levels than the
three NT groups. There were still no significant differences among ESNT, CCNT, and MEFNT groups at the expression levels of most genes except RPI (Fig. 12). When it came to the morula stage, we found MEFNT embryos had significant lower expression of FBL and 18 S rRNA compared with other groups. Moreover, its UBF and 47 S rRNA expression levels were significant lower than that of ICSI embryos (Fig. 13). At the blastocyst stage, the three NT groups had significant higher 47 S rRNA levels than that of the ICSI group. MEFNT blastocysts had significantly lower B23 levels than ICSI embryos, and the ESNT embryos had significantly lower RPI expression than the CCNT counterpart. No other differences existed between the four groups (Supplemental Fig. 5).

DISCUSSION

The successful reprogramming of the donor cell genome to the state of totipotency requires both the continued silencing of the somatic-specific genes and the activation of essential embryonic genes, enabling further embryonic development (35–37). Moreover, the developing preimplantation embryo has a profound need for synthesis of proteins supporting both housekeeping and cell differentiation. So it is not surprising that the rRNA genes are among the earliest genes being activated. A proper activation of these genes is crucial for continued embryonic development (38, 39).

In this study, ESCs, CCs, and MEFs were chosen as donor cells as they were easy to achieve, widely used in NT experiments, and were supposed to have different differentiation status. rDNA clusters (NORs) commonly exist on 5 chromosomes (12, 15, 16, 18, and 19) in standard inbred mouse strains. These strains may have 3–5 pairs of chromosomes with silver-stained active NORs (40, 41). For outbred F1 strains, they may have half of the maternal/paternal NOR-bearing chromosomes (42). However, the reported NORs numbers for C57BL/6J and DBA/2 strains varied due to different methods and samples used (40, 41, 43, 44). Moreover, there was still no available data of the NOR numbers in B6D2F1 strain. Previous studies indicated that >90% of the NORs in ES cells were marked with both H3Ac and H3K4me, which means almost all the NORs had transcription activity (13, 45). In our system 68% of the ESCs had 8 active NORs at metaphase, and the active NORs numbers in CCs, MEFs, and all the four groups of embryos did not exceed 8. We speculated that the outbred B6D2F1 mouse used in our study had eight NOR-bearing chromosomes in total. The diploid genome of mouse contains about 400 rDNA copies whose expression depends on the demand of cell biosynthetic activity. The ratio of active and inactive rRNA genes is tissue-specific and is stably propagated through the cell cycle (46). As a kind of somatic cell, the relative more active NORs in CCs may reflect its high proliferation and secretion activity during oocyte growth and maturation. MEFs had the least active NORs, which is in accordance with its high rDNA-methylated status. Therefore, the three cell lines had different rDNA activities that were intimately linked with rDNA methylation level as previously reported (12). It then provided a good model for evaluating the influence of donor cell rDNA activity on NT embryos in the following experiments.

The nucleolar protein distributions in donor cell nucleoli before and after NT have never been demonstrated in mice before. Previous work on bovine NT embryos showed that after bovine skin fibroblast was fused into enucleated MII oocyte, the donor cell nucleoli degenerated, as indicated by B23 and FBL signals. The same phenomenon could be simulated by treating the donor cells with 5 μg/ml actinomycin D, a kind of rDNA transcription inhibitor. However, the two proteins could still be detected at 2 h post-fusion (14). In our study, UBF and B23 were used as indicators of rRNA transcription and processing machineries. They both vanished 20 min post-NT in CCNT and MEFNT embryos, which was much faster than that in bovine NT embryos. This result indicated that the donor cell nucleoli lost rRNA synthesis and processing activities quickly in ooplasm and may undergo degeneration. However, UBF and B23 signals in ESCs were more resistant to ooplasm as they were still detectable 60 min after NT. Because almost all the CCs and MEFs used in NT experiments were at G0/G1 phase but ESCs were at metaphase before NT, the difference may come from the cell cycle phase of donor cell at the time of entering ooplasm. During mitosis, the nucleolar disassembly starts with the loss of RPI subunits from the fibrillar centers, and no nascent rRNA transcripts are synthesized (34, 47). So the ESNT embryos had no rDNA transcription activity after NT, like the other two somatic cell NT counterparts. No sign of rRNA transcription activity was detected during pronucleus formation and the first mitosis (17 hpa), as previously reported...
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(48). Because UBF remaining associated with chromosomes at active NORs is a typical pattern during mitosis (49), lacking a UBF signal during the first mitosis in NT embryos indicated that donor cell chromosomes had undergone deep remodeling. Until 2-cell stage, the ICSI embryos took the lead to transcribe rRNA, and then all the groups started to transcribe and process rRNAs. Thus, we speculated that all the embryos lost rRNA transcription and processing capacity shortly after NT. The different rDNA epigenetic status and rRNA synthesis activities in donor cells would barely have any impact on NT embryos before 2-cell stage.

ESNT embryo had more RPI and 47 S rRNA expressed at the 2-cell stage that of the CCNT and MEFNT counterparts. More opened NORs in chromosomes may result in earlier or more rRNA transcription. Apart from this, no other obvious differences were observed among the NT groups at 2-cell and 4-cell stages in nucleolar proteins distribution and nucleolar-related gene expression. However, ESNT embryos had almost all the NORs activated and with no significant difference compared with ICSI embryos, which indicated their rDNA activity was near the normal embryos. The active NORs numbers in MEFNT embryos obviously increased when compared with MEFs but were still much less than being sufficiently activated as in ICSI or ESNT embryos. The rDNA methylation levels in three NT groups slightly decreased when compared with the donor cells from which they derived. The result indicated that partial methylated rDNA were unmethylated during NT reprogramming. Moreover, they repeated exactly the same trend as what was seen in the donor cells; that is, ESNT embryos had the lowest rDNA methylation level, MEFNT had the highest, and CCNT embryos were in the middle. In summary, the rDNA activity in 4-cell NT embryos directly reflected the rDNA activity in donor cells from which they derived.

Full functional nucleolus is established at the 4-cell stage in in vitro fertilization mouse embryos, accompanied with formation of fully fibrillo-granular nucleolus in ultra-structure (11), which is the most important step of nucleolus biogenesis during preimplantation development. However, researchers demonstrated that mouse MEFNT embryos had poor nucleolus development at the 4-cell stage, as the fibrillo-granular structure only appeared at the surface of the nucleolus (18). This phenomena could be partially explained by our results that MEFNT embryos had the least active NORs and the functional nucleoli formation was based on the active NORs (4). As a kind of somatic cell NT embryo, CCNT embryos had relatively more active NORs and almost the same nucleolar-related genes expression level when compared with ESNT embryos. This may be one of the reasons why CCs always yielded better cloning outcomes than other somatic cells and were wildly used in NT experiments of different animals (50). Because the oocytes were supposed to have the same reprogramming competence against different donor cells, we believed that the rDNA methylation level and active NOR numbers in 4-cell embryos were determined by the rDNA epigenetic status of donor cells.

When it came to the morula stage, the MEFNT embryos showed obvious weakness in 47 S rRNA, 18 S rRNA, UBF, and FBL expressions. We examined the rRNA transcripts level from 2-cell to blastocyst in ICSI embryos and found they reach the peak at morula stage (supplemental Fig. 6). The immunofluorescent results also revealed that many blastomeres were undergoing mitosis at this stage. These indicated the embryos may have the highest proliferation and metabolism level around morula stage. Because the rRNA transcription, pre-rRNA processing, and ribosome assembly are well linked and coordinated (51), we could speculate that the low rDNA activity in MEFNT embryos would result in insufficient rRNA transcription, processing, and ribosome assembly, which could not fulfill the high ribosome requirement around morula stage and, finally, lead to poor developmental capability from morula to blastocyst stage (Fig. 11).

The qPCR data of blastocyst stage showed almost no significant differences among the NT groups, which indicated that those embryos that successfully reaching the blastocyst stage might have a similar gene expression profile. However, all the NT groups had significant higher 47 S rRNA expression levels than that of the ICSI group. This phenomenon has been reported previously between mouse in vitro fertilization and tail tip cell NT embryos. The author believed that, in NT blastocyst, the accumulation unprocessed 47 S rRNA and rRNA intermediates failed to be processed into mature rRNAs (31). Because no significant differences were found in FBL and B23 expressions between ICSI and NT groups in our study, the exact mechanism might be more complicated and need further investigations.

rDNA reactivation plays a basic but very important role in preimplantation development. Insufficient reactivation may lead to poor ribosomal biogenesis and impair developmental competence. Moreover, in human blastocyst, half NORs could be inactivated post-implantation (13). If the same mechanism is also adopted in mice, the embryos reconstructed from donor cells with low rDNA activity, like MEFNT embryos in our study, may face ever more problems after implantation. Thus, according to our results, choosing a cell line that has a lower rDNA methylation level and/or more active NORs as donor cell will be a good strategy for NT experiments. Artificially improving the rDNA epigenetic status before or immediately after NT is another option. Recently, several histone deacetylase inhibitors were used to treat the NT embryos to achieve better clone efficiency (52–55). The histone deacetylase inhibitor can increase the overall histone acetylation level of donor cell chromatin, then make the genes more reachable to maternal factors, therefore, promoting reprogramming. In a recent study, scriptaid (a kind of histone deacetylase inhibitor) helped NT embryos overcome the asynchronous rRNA transcription at the 2-cell stage by activation of rDNA and promotion of nucleolar protein allocation (56). This study revealed how histone deacetylase inhibitor could improve NT embryo development in rRNA activation levels. We may also expect that, for those terminally differentiated cells, if the inactive rDNA could be activated before or after entering the oocyte appropriately, better NT outcomes can be achieved thereafter.

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