Embryos from "0PN" Zygotes May Be Most Likely 2PN Embryo in L-R-ICSI

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Abstract

Although some zero pronuclei (0PN) zygotes cleave and develop into good embryos, they are usually discarded because they lack two distinct pronuclei (2PN). In this case report, we followed four couples to determine whether 0PN embryo could be used in late rescue intracytoplasmic sperm injection (L-R-ICSI). Here we report five healthy infants, including one set of twins, derived from "0PN" embryos from four frozen–warmed embryo transfer (FET) cycles in L-R-ICSI between 2015 and 2017. Of nine infants born from L-R-ICSI cycles in our center, five were "0PN" embryos, and all remain healthy. Embryos from "0PNs" such as 2PN can develop into healthy babies using L-R-ICSI. This finding suggests that embryos from "0PNs" in L-R-ICSI may be different from traditional 0PN gametes and more likely to originate from 2PN gametes and they may be used in those infertile couples who lack 2PN embryos, instead of discarding them. The use of "0PN" embryos increases the cycles reaching embryo transfer, allowing some infertile couples to have healthy children and to avoid mental anguish and wasted time and money.

Introduction

Assisted reproductive technologies (ART) procedures include conventional in vitro fertilization (IVF) and intracytoplasmic sperm injection (ICSI). In a conventional IVF cycle, approximately 3–5% of fertilization failures occurred with normal sperm parameters examined [1]. This result is consistent with data from our center, but some scholars have reported that the rate of complete fertilization failure was 10%-25% [2]. This discrepancy may be related to different statistical ranges, such as obtaining a low number of eggs, having abnormal semen parameters and so on. Although the incidence of complete fertilization failure decreases as the second pole for fertilization increases at 4–6 hours after insemination, the failed process carries a high emotional and financial burden for the infertile couple. Accordingly, to avoid the catastrophe of fertilization failure, some embryologists advise ICSI on day one of unfertilized mature oocytes, applying the so-called “late rescue ICSI” (L-R-ICSI).

Until now, morphological evaluation has been used widely a relatively simple and harmless method by which to predict embryo development, including prokaryotic observation, embryo and blastocyst grading. Here, we mainly introduce prokaryotic observation. Normally, fertilized zygotes from IVF and ICSI show two pronuclei (2PN) at 16–20 hours after insemination [1, 3, 4], and embryos from 0PN, monopronuclear (1PN) and three or more PN are considered to provide evidence of failed or abnormal fertilization. Guidelines do not recommend transferring embryos that do not have two pronuclei at the time of visual assessment [5]. However, the findings of Yao et al. reported that no significant differences were shown in euploid rates between 0PN and 2PN in blastocysts cycles of PGT-M (preimplantation genetic testing for a monogenic disorder) [6]. So, it remains controversial whether or not 0PN embryos should be used.

In L-R-ICSI, that is, ICSI performed the day after conventional IVF because no or little fertilization has been achieved, the time-course of 0PN formation is complicated and still unclear. It has been reported that birth of healthy twins resulting from the transfer of embryos found to be without 2PN in the cytoplasm of the oocytes 16–18 hours post-insemination, when L-R-ICSI had been performed on day one in a fresh cycle
Here, we describe the results achieved—four successful full-term pregnancies and five infants—after the transfer of OPN embryos in L-R-ICSI in frozen–warmed embryo transfer (FET) cycles.

## Methods

### Ethics approval

This study was approved by the Ethical Committee of Guangdong Provincial People's Hospital, Guangdong Academy of Medical Sciences. All methods were performed in accordance with the relevant guidelines and regulations and the approved protocol was followed throughout the study period.

### Informed consent

For each couples entering the treatment cycle, we already signed an informed consent form to state the success rate and possible complications of IVF/ICSI, and informed them that their data might be used for scientific research but their personal information is completely confidential.

### Clinical data

In L-R-ICSI of our center, 8 patients were successfully conceived and 9 babies were born, among which 5 were from OPN, between 9 January 2015 and 23 August 2017. This paper mainly describes the data of 4 OPN cases, including a pair of twins.

Four female patients and their domestic partners with unexplained infertility for 3.4-7 years and aged from 26 to 32 years were scheduled for fertilization (Data of fresh cycles as table 1). Three patients used the standard long stimulation GnRH agonist protocols and the other one used GnRH antagonist protocol. Human chorionic gonadotrophin at a dose of 5000–10000 IU was administrated after at least two follicles of 18 mm or greater in size were visualized by means of transvaginal ultrasound scanning. Fifty cumulus oocyte complexes (COCs) were retrieved by ultrasound-guided needle aspiration in the four infertile couples. The COCs were collected and washed fourfold and transferred into a Center-Well Organ Culture Dish (353037,60 × 15 mm; Becton Dickinson, USA) containing 1 mL of G-IVF medium (Vitrolife, Sweden) for pre-insemination culture. Oocytes were inseminated in the same medium with approximately 3 ~ 4 × 10^5 motile spermatozoa/mL after hCG injection 40–42 hours later, and incubated at 37°C in a mixed atmosphere of 6% CO₂ and 5% O₂ with high humidity. A fertilization check was performed 18–20 hours later (about 8 a.m.) on the next morning (D1) after insemination and only five oocytes were fertilized. After patients provided appropriate signed informed consent, L-R-ICSI as an alternative treatment was performed on the thirty-five metaphase II stage oocytes with sperm from D0 (day of oocyte pick-up, OPU) insemination at about 9 a.m.. Embryos were cultured in the G-1 medium drops (Vitrolife, Sweden), and covered with mineral oil in Falcon tissue culture dishes (353001, 35 × 10 mm, Becton Dickinson, USA). Around 8 a.m. on the D2 morning after OPU, only seven of these oocytes had two distinct pronuclei (2PN) in the cytoplasm, and the other 23 oocytes had no pronuclei but showed cleavage on the following culture.
All rescued embryos were observed on D3 or D5 or D6 after L-R-ICSI. Here, we introduced two systems of classifying the embryos. On day 3, the blastomeres and fragments were evaluated and classified as four types separately by means of the following scoring system. Blastomere: grade I-even sized blastomeres with regular morphology; grade II-slightly uneven sized blastomeres with regular morphology; grade III-asymmetrical blastomeres and irregular morphology; grade IV-severely asymmetrical blastomeres and irregular morphology or significant cytoplasmic particles. Embryo fragments were classified into I, II, III and IV grades according to fragmentation < 5%, 6%-20%, 21%-50% and > 50%. Number of cells 7–9, embryos and fragments were classified as I and II, but those with different levels of II were defined as top-quality embryos. Cell numbers > 4, blastomeres grade I-III and fragments classified as I-II were considered suitable embryos for transfer, excluding embryos with blastomere as III and fragment as II. The blastocyst grading system proposed by Gardner and Schoolcraft was based on blastocyst morphology parameters [7]; that is, blastocyst development was divided into 6 stages according to the size of blastocoele expansion degree, blastocyst cavity and whether it hatched or not. The inner cell mass (ICM) and trophoblast ectoderm (TE) were classified into three levels: A, B and C according to cell number and intercellular adhesion (A: a good number of cells and good intercellular adhesion; B: small cell numbers and good intercellular adhesion; C: almost no cells). The blastocysts with blastofoles were greater than or equal to 3, ICM and TE scores of A, B and C, but not both C, were considered as eligible and transferred, and those defined as good blastocysts for ICM and trophoblast ectoderm (TE) were graded A or B.

Avoiding dyssynchrony between the endometrium and embryonic development, fresh cycles of embryo transfers were cancelled. The couples were informed of the uncertainty associated with the transfer of 0PN-derived embryos in terms of the health of any resulting babies. All embryos and blastocysts that reached the standard of transfer, either from 2PN or 0PN, were frozen in vitrification. The freezing process between embryos and blastocysts differed in that the blastocysts needed to be shrunk artificially with a laser pulse prior to vitrification by the embryologist. Embryos/blastocysts were first transferred to an equilibration solution (VT101-®, KITAZATO, Japan) for 10 minutes, and subsequently transferred to a vitrification solution (VT101-®, KITAZATO, Japan) for 1 minute. Then, one or two embryos/blastocysts was/were placed into a cryotop (Cryotop®, KITAZATO, Japan), which was quickly plunged into liquid nitrogen, and then covered with a tube and stored in liquid nitrogen.

About two months after oocyte pick-up, FET cycles were started, in which one or two cryopreserved embryos/blastocysts were warmed and transferred. To thaw, the embryos/blastocysts loaded into the cryotop were immersed in the first thawing solution (VT102-®, KITAZATO, Japan) at 37°C for 1 minute, and then transferred to the second and the third thawing solutions (VT102-®, KITAZATO, Japan) for 3 and 5 minutes at room temperature. Lastly, the embryos/blastocysts were washed in the fourth solution (VT102-®, KITAZATO, Japan) at 37 °C for 5 minutes. After warming, embryos/blastocysts with more than 50% intact blastomeres were regarded as viable and were transferred to the recipient uterus of all four patients.

Ten embryos/blastocysts of five FET cycles were transferred to the four patients. One patient received two transplants, the first of which had two 2-prokaryotic embryos, and was not a pregnancy. The
remaining eight fresh cycles of 0 prokaryotic embryos were transplanted four times and four clinical pregnancies resulted (Data of of FET cycles as table 2). Then, 6–8 weeks after FET, four clinical pregnancies were confirmed using transvaginal ultrasonography. Among them were two twin pregnancies with four sacs but three fetal heartbeats.

**Results**

The four recipients of the "0PN" embryos were offered pre-natal diagnosis and karyotype ploidy analysis to verify fetal normality at 16–20 weeks of pregnancy. Fortunately, the fetuses have normal ploidy and four deliveries at 36–41 weeks resulted in the births of five normal healthy babies, including a pair of live twins between 28 December 2015 and 24 May 2018. One of the four pregnant women had a natural birth and the other three women underwent lower segment cesarean sections. The data in detail of the 3 boys and 2 girls are shown in Table 2. All the children had regular physical examinations by their pediatricians and were growing normally. The oldest baby is four and a half years old and the youngest twins were more than 2 years of age by now. All children are healthy with normal developmental milestones up to today.

**Discussion**

To the best of our knowledge, this is the first report of live healthy infants of "0PN" embryos in FET cycles in L-R-ICSI. Results described in the present case report indicate that "0PN"-derived embryos may be most likely 2PN embryo in L-R-ICSI cycles and the conventional discard of embryos in c-IVF and ICSI cycles that originated from 0PN zygotes at fertilization check-up may no longer apply to L-R-ICSI. Since "0PN-derived" embryos are shown to result in live and healthy babies, these "0PN" zygotes bring good hope to infertile couples and increase the chances of having a successful pregnancy for infertile patients without 2PN embryos.

The use of 0PN zygotes in IVF has been a global topic of debate for the past 20 years because of uncertain ploidy. Some experts believe that 0PN-derived embryos should not be recommended based on the results of chromosome tests using FISH technology in which only 3%-4.5% of the 0PN embryos presented euploid karyotype [5, 8, 9]. Hence, in some IVF centers, it has become standard clinical policy to discard 0PN embryos. However, other scholars have reported different FISH results, including that 57–62% of embryos developing from 0PNs were diploid, and that 66% of diploid 0PN embryos were thrown away at the embryo stage due to erroneous classification of ploidy based on pronuclear numbers scoring[3, 10]. Lee et al. also reported that 0PN-originated embryos had a similar euploidy rate (23.1% vs. 30.0%) by array comparative genomic hybridization (aCGH) [11]. Because the results reported by different investigators and institutions are quite different, the method of chromosome detection has been questioned and whether any embryo (2PN or otherwise) can be guaranteed [12]. Therefore, in different IVF centers, transferring 0PN-derived embryos may or may not be considered in those treatment cycles in which 2PN-derived embryos are not available.
In brief, embryos derived from 0PN zygotes create a dilemma for embryo selections in clinical practice. Some studies, however, have reported 0PN-derived embryos resulted in the birth of healthy babies [3, 13, 14]. Liu et al. also reported that 13 healthy infants resulted from 0PN-originated embryos with an implantation rate of 17.0% in conventional IVF cycles, and those authors believed that the source of embryos transferred did not have a significant impact on clinical pregnancies and live birth rates[4]. Capablo et al. reported the analysis of eight zygotes scored as 0PNs in ICSI cycles by PGT-A and found that all were diploid bi-parental[15]. Destouni et al. also reported genome-wide haplotyping embryos developing from 0PN zygotes increased transferrable embryos in PGT-M[16]. These results suggest that 0PN-embryos should not be eliminated from further studies since they are probably misclassified zygotes due to the prokaryotes form, or they may disappear too early or too late and miss the prokaryotic observation point [3, 4, 17].

In conventional IVF and ICSI, a fertilization check is usually performed 16–20 hours after insemination on day 1 after oocyte pick-up (OPU) [1, 3, 4]. Therefore, in Guangzhou, China, insemination is often performed at about 1:00–3:00 p.m. on OPU day and the prokaryotic observation is conducted at 7:00–8:00 a.m. on the next morning. Thus, L-R-ICSI is carried out after discovery of non-fertilization, usually at 8:30 – 9:00 a.m., so the time of reinsemination in L-R-ICSI is 5–6 hours earlier than normal insemination. The observation point is delayed to more than 22 hours in L-R-ICSI, making it difficult to observe the prokaryotes at regular fertilization check (about 2–3 o’clock midnight) during work time, since going to work 5–6 hours earlier than normal work is unrealistic. Therefore, we speculate that the origin of the "0PN" zygote is primarily due to the disappearance of the prokaryot in L-R-ICSI, and the criteria for the available embryos of conventional IVF and ICSI that must be derived from 2PN may need to be changed. Based on this deduction, we required patients’ signed informed consent for use of 0PN embryo and used it as backup, which not only increased the embryo reserve for patients, but also brought substantial benefits to the patients. The birth of five healthy babies confirmed the feasibility of this decision. Chian et al. (2003) also reported successful healthy live births from 0PN embryos after L-R-ICSI following failed fertilization. Their results combined with our cases presented in this report suggest that "0PN"-derived available embryos can produce healthy babies in L-R-ICSI cycles.

Another highlight of our work was the transfer of embryos during thawing cycles. The pregnancy rates after L-R-ICSI are lower than normal ICSI due to either poor quality of embryos or asynchrony between endometrial secretory pattern and embryo development. Aging of oocytes in in vitro culture and the long interval between oocytes retrieval and L-R-ICSI may result in poor quality of embryos [18, 19]. Also, an asynchrony between endometrial secretory pattern and embryo development may influence implantation and pregnancy. Though there had been previous reports of live births of fetuses in fresh embryo transfer cycles in L-R-ICSI [1, 20], prior to 2013, we had not yet had a successful pregnancy with 10 fresh transplants in spite of the embryo appearing to meet the requirements of good embryos, either D3 or D4 after egg retrieval. So, we considered that the dyssynchrony of the developmental stage of the embryo and endometrial receptivity may be the main cause. All available embryos were frozen during the fresh cycle and thawed and transplanted 2–3 months later. Since then, eight clinical pregnancies and live births have been derived through FET in our center between 2015 and 2017. Results of the present case
study emphasize the importance of synchronization of embryonic development and endometrium for embryo transfer and that the FET cycle is the preferred strategy in L-R-ICSI.

In this case report, we describe four successful full-term pregnancies and five births after the transfer of L-R-ICSI embryos derived from "0PN". This result indicates that we should give attention to "0PN" embryos in L-R-ICSI and consider that they are probably misclassified and zygotes are discarded due to the absence of pronuclei. We suggest that the conventional method of discarding embryos because zygotes originally lacked pronuclei at the fertilization check should be reconsidered in L-R-ICSI. Transferring "0PN-derived" embryos/blastocysts in the FET cycles with subsequent expected pregnancies can benefit families where the number of transferrable embryos is very limited, and can be performed as the last ray of hope in case of failed fertilization. Therefore, it is reasonable to suggest that "0PN" embryos more likely to originate from 2PN in L-R-ICSI. Certainly, confirming this viewpoint requires more cases. In addition, it’s worth considering that our successful patients were only 26–32 years old. Whether 0PN embryos from older women have developmental competency after L-R-ICSI remains unknown.

It is worth emphasizing, there are multiple publications on apronuclear and unipronuclear embryos in which better technologies have been utilized to assess these embryos such as time lapse system (non-invasive) and chromosomal/ploidy analysis (invasive) [21–22]. In this instance, incubation in a time lapse incubator may have provided additional important information regarding the true origin of these abnormal zygotes, but that is currently missing from this study because of having no time lapse incubator. We checked the PN morphology more than 22 hours post-insemination and classified the "0PN" zygotes at this time. it is most likely that syngamy and PN fading already occurred and the transferred embryos could be regular 2PN. Our center does not have the qualification of PGT, so we cannot do biopsy to analyse chromosomal/ploidy. The PN status of these '0PN' embryos were may wrongly classified due to late fertilization check in L-R-ICSI zygotes. It is for this reason, we appeal that "0PN" embryos can be reserved for those patients who do not have 2PN embryos in L-R-ICSI.

Conclusions

The PN status of these '0PN' embryos were may wrongly classified due to late fertilization check in L-R-ICSI zygotes, since the observation point is delayed to more than 22 hours in L-R-ICSI. Embryos from "0PNs" such as 2PN can develop into healthy babies in L-R-ICSI. This finding indicates that "0PN"-derived embryos may be most likely 2PN embryo in L-R-ICSI cycles and the conventional discard of embryos in c-IVF and ICSI cycles that originated from 0PN zygotes at fertilization check-up may no longer apply to L-R-ICSI. We suggest that embryos from "0PNs" may be used in those infertile couples who lack 2PN embryos, instead of discarding them.

Declarations

Ethics approval and consent to participate
This study was approved by the Ethical Committee of Guangdong Provincial People's Hospital, Guangdong Academy of Medical Sciences. All methods were performed in accordance with the relevant guidelines and regulations and the approved protocol was followed throughout the study period. For each couples entering the treatment cycle, we already signed an informed consent form to state the success rate and possible complications of IVF/ICSI, and informed them that their data might be used for scientific research but their personal information is completely confidential.

Consent for publication

We all authors' agreement to submission of the manuscript.

Availability of data and material

All data and material of this study is true and reliable.

Competing interests

All authors have no conflicts of interest to report.

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Authors' contributions

CL and DH conceived and designed the study. JC and YL carried out female patients' ovulation plan, counselled the patients and followed pregnancy outcomes. DH and CZ performed ICSI and embryo freezing procedures. CL and DH performed thawing and embryo transfer procedures. JL and YZ performed semen processing and embryo/blastocysts scoring. DL performed male physical examinations and recording. CL and JC performed data and statistical analysis. CL drafted the original version of the article. CL, JC, DL and YL revised the article. All co-authors read and approved the final version of the article.

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**Tables**

| Characteristic                   | Case1  | Case2  | Case3  | Case4  |
|----------------------------------|--------|--------|--------|--------|
| Age (years)                      | 30     | 32     | 28     | 26     |
| Duration of infertility (years)  | 7      | 6      | 6      | 3.4    |
| Type of infertility              | Primary| Secondary| Primary| Primary|
| Cause of infertility             | Pelvic inflammatory disease | Ovulation factor | Pelvic inflammatory disease and tubal factor | Polycystic ovary syndrome (PCOS) |
| Protocol                         | long-acting GnRH agonist | long-acting GnRH agonist | long-acting GnRH agonist | GnRH Antagonist |
| Number of oocytes                | 14     | 10     | 14     | 12     |
| Oocytes of L-R-ICSI              | 6      | 7      | 10     | 12     |
| Number of frozen D3 Embryos      | 5(0PN) | 1(2pn) | 2(0PN) | 2(2pn) |
| Number of frozen blastocysts     | 0      | 2(0PN,D5) | 2(0PN,D6) | 2(0PN,D5) |

GnRH gonadotropin-releasing hormone
Table 2
Data of FET cycles

| Characteristic                        | Case1 | Case2 | Case3 | Case4 |
|---------------------------------------|-------|-------|-------|-------|
| Protocol for endometrial preparation  | HRT   | NC    | NC    | HRT   |
| Endometrial pattern                   | 8     | 8.5   | 12    | 7.6   |
| Endometrial thickness (mm)            | A     | A-B   | A-B   | A-B   |
| Number of embryo transferred          | 2     | 2     | 2     | 2     |
| Day of embryo transferred             | 3     | 5     | 3     | 5     |
| Grade of embryo or blastocyst         | 8.21 /13.21 | 4AC/4CB | 7.21 /7.21 | 4BC/4CB |
| Sac                                   | 2     | 1     | 1     | 2     |
| fetal heartbeat                       | 1     | 1     | 1     | 2     |
| Weeks of delivery                     | 41 + 3 | 39 + 6 | 40 + 2 | 35 + 6 |
| Type of delivery                      | Cesarean section | Cesarean section | uneventful delivery | Cesarean section |
| Number of infant                      | 1     | 1     | 1     | 2     |
| Gender                                | female | female | Male | male | female |
| Weight(kg)                            | 3.75  | 3.45  | 3.77  | 2.95  | 2.30  |
| Length(cm)                            | 50    | 50    | 52    | 48    | 46    |

**FET** frozen–warmed embryo transfer;

**HRT** hormone replacement treatment

**NC** natural cycle