Comparison of Stimulated Cycles with Low Dose r-FSH versus Hormone Replacement Cycles for Endometrial Preparation Prior to Frozen-Thawed Embryo Transfer in Young Women with Polycystic Ovarian Syndrome: A Single-Center Retrospective Cohort Study from China

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Objective: The principal purpose of this study was to compare reproductive outcomes for stimulated cycles (STC) and hormone replacement cycles (HRC) for endometrial preparation before frozen-thawed embryo transfer (FET) in young women with polycystic ovary syndrome (PCOS).

Methods: We conducted a retrospective study of 1434 FET cycles from January, 2017 to March, 2020 in our reproductive center, in which stimulated and hormone replacement cycles were used for endometrial preparation. Pregnancy outcomes of couples undergoing routine STC-FET or HRC-FET were analyzed by propensity score matching (PSM) and multivariable logistic regression analyses.

Results: Data on 1234 HRC protocols (86% of the total) and 200 STC protocols (14%) were collected. After PSM, 199 patients were included in both groups, respectively. There was no significant difference in positive pregnancy rate (52.7% vs 54.8%, p=0.763), clinical pregnancy rate (51.8% vs 52.8%, p=0.841), live birth rate (45.2% vs 43.7%, p=0.762), pregnancy loss rate (9.7% vs 16.2%, p=0.164) and ectopic pregnancy rate (1.5% vs 0.5%, p=0.615) between STC and HRC protocols. Subsequent multivariate logistic regression analysis also yielded similar results.

Conclusion: STC for endometrial preparation had similar pregnancy outcomes compared with HRC protocols. Evidence is available which shows that for young women with PCOS in preparation for FET, HRC could be a reasonable choice for patients who are unwilling to accept injections. However, STC may reduce unnecessary anxiety and operational costs and offer more flexibility for patients. Eventually, we must embrace the concepts of individualization, securitization, and optimization in the clinic.

Keywords: polycystic ovarian syndrome, stimulated cycle, hormone replacement cycle, frozen-thawed embryo transfer, endometrial preparation, propensity score matching

Introduction

The first successful frozen-thawed embryo transfer (FET) was reported in 1983 and the first live birth in 1984. Since then, elective embryo transfer and “freeze-all” strategy with segmentation of in vitro fertilization (IVF) and intracytoplasmic
sperm injection (ICSI) treatment, aiming to cryopreserve all good quality embryos produced in a fresh cycle and to transfer these embryos in subsequent endometrial prepared cycles, has been widely used in assisted reproductive technology (ART) in recent years. FET can profoundly mitigate the risk of ovarian hyperstimulation syndrome (OHSS) and its use has now been extended to include those cycles of pre-implantation genetic diagnosis/screening, late-follicular progesterone elevation and embryo-endometrial asynchrony. Compared with fresh embryo transfer, FET increase maternal safety, improve pregnancy rates, decrease ectopic pregnancy rates. In addition, FET, avoiding the negative impact of controlled ovarian stimulation (COS) on endometrial receptivity, can provide a more physiologic uterine environment for embryo implantation with a fresh start and regrowth under alternative less intensive endometrial preparation regimens.

PCOS is a heterogeneous endocrine disorder affecting reproductive aged women, with an estimated prevalence of between 8% and 13%. Patients with PCOS usually had menstrual dysfunction, infertility, hirsutism, acne, obesity, and metabolic syndrome. IVF had become an important therapeutic technique for infertility of PCOS. As known, PCOS patients refer to high responder group, hence, elective freeze-all strategy is recommended worldwide to prevent OHSS, and to alleviate the harmful effects of supra-physiologic steroid hormones on the endometrium before embryo implantation.

Chen et al reported that FET increases live birth rates (LBRs) in their RCT of women with PCOS.

Essentially, the outcomes for the FET could be affected by female age, embryo quality, endometrium and embryo synchronization, as well as endometrial receptivity, etc. If it is assumed that those factors do not differ between protocols, that endometrial preparing cycles is critical for it is assumed that those factors do not differ between cycles of pre-implantation genetic diagnosis/screening, late-follicular progesterone elevation and embryo-endometrial asynchrony. Compared with fresh embryo transfer, FET increase maternal safety, improve pregnancy rates, decrease ectopic pregnancy rates. In addition, FET, avoiding the negative impact of controlled ovarian stimulation (COS) on endometrial receptivity, can provide a more physiologic uterine environment for embryo implantation with a fresh start and regrowth under alternative less intensive endometrial preparation regimens.

Essentially, the outcomes for the FET could be affected by female age, embryo quality, endometrium and embryo synchronization, as well as endometrial receptivity, etc. If it is assumed that those factors do not differ between protocols, that endometrial preparing cycles is critical for frozen-thawed embryo transfer.

There are different ways for endometrial preparation, ranging from natural cycle (NC-FET) to stimulated cycle (STC-FET), or hormone replacement cycle (HRC-FET). However, elucidating which is the best option remains to be determined. The NC cycle is suitable for patients with regular menstrual periods. The endometrium is better developed and breakthrough bleeding is less likely when the NC regimen is used. However, it is reported that premature ovulation and follicular dysplasia lead to the cancellation of cycle, especially in women with PCOS. In addition, in the light of menstrual dysfunction, the natural cycle used in the preparation of endometrium is not applicable. The mild ovarian stimulation induces follicular development by generating endogenous hormones. That process of follicular development and ovulation is important to function of the corpus luteum. What calls for special attention is that the initial dosing of gonadotropins (Gn) should be low for preventing the risk of OHSS. Women also should be monitored closely. The most commonly used FET protocol for women with PCOS is the HRC. This cycle is easy to plan, thus improving patient convenience. The main reasons for canceling cycles in HRC group were related to an inadequate endometrial response.

Currently, there are few data comparing stimulated cycles with hormone replacement cycles for FET, especially in PCOS patients. A recent meta-analysis indicated that, compared with the hormone replacement cycles (HRC), the letrozole stimulation cycle may have a lower miscarriage rate (MR). No significant difference had been found between the mild ovarian stimulation (OS) cycle and AC protocols in live birth rate (LBR), ongoing pregnancy rate (OPR), clinical pregnancy rate (CPR) and embryo implantation rate (IR), some researches were opposed to this meta in artificial and stimulated cycle for FET in PCOS. The main objective of this study was to compare reproductive outcomes for stimulated and hormone replacement endometrial preparation protocols in frozen embryo transfer (FET) cycles of PCOS. To minimize potential biases, we applied the PSM method to implement post-hoc randomization.

Materials and Methods
Study Design and Participants
We performed a retrospective cohort study of 1434 FET cycles of PCOS from January 2017 to March 2020 in the fertility unit at a University Hospital. Patients in this present study had previously undergone treatment by IVF or intracytoplasmic sperm injection (ICSI) cycles. The study was approved by the Reproductive Ethics Committees of the Affiliated Hospital of Shandong University of TCM (ref approval no. SDTCM20201215). All participants provided written informed consent. Eligible patients included women with PCOS aged between 21 and 35 years, diagnosed by Rotterdam criteria: oligo-or anovulation, clinical or biochemical evidence of hyperandrogenism, and polycystic ovarian morphology on ultrasonography (defined as an ovary that either contains ≥12 antral follicles or that has a volume >10 cm³), with at least one
embryo vitrified mainly at day 3, and for whom it was the first FET performed. The exclusion criteria were: (i) Body Mass Index (BMI) ≥30Kg/m² at the time of embryo vitrification; (ii) Endometriosis; (iii) Preimplantation genetic diagnosis/screening cycle; (iv) History of recurrent pregnancy loss or recurrent implantation failure; (v) Uterine pathology; (vi) Cycles cancelled due to failure of embryo thawing and survival.

Controlled Ovarian Stimulation Protocol
All participants had undergone the IVF/ICSI treatment as clinically indicated. Furthermore, a flexible GnRH antagonist (GnRH-ant) (Cetrorelix; Merck Serono, Darmstadt, Germany) and long GnRH-a (Triptorelin, Decapeptyl, Ipsen, France) protocols were employed with 150–225 IU/day of recombinant FSH (Gonal-F, Merck-Serono, Lyon, France). Additionally, the doses of gonadotropin were determined based on the characteristics of individual patients. Thereafter, oocyte retrieval was conducted under ultrasound transvaginal guidance, 34–36 hours after triggering with 0.1mg GnRH-a or recombinant hCG (Ovitrelle®, 250μg, Merck), after which conventional IVF/ICSI were performed as previously described.22 The IVF/ICSI procedure had either been followed by a fresh embryo transfer and preservation of the redundant good embryos by vitrification or by a freeze-all strategy on clinical indication. Regular monitoring during controlled ovarian hyperstimulation (COS) treatment includes vaginal ultrasound (to assess endometrial thickness and follicle development) and blood hormone assays (including estradiol, progesterone and LH plasma levels).

The choice of embryos for vitrification was expected to focus on the inclusion of no less than six blastomeres with ≤20% fragmentation. Embryos that presented a fragmentation rate between 20% and 50% were vitrified only when they had reached the 8-cell stage on Day 3. The applied vitrification procedure has been described in detail before.22

Endometrial Preparation Protocols
Women with PCOS were instructed to wait for spontaneous menses or prescribed with progestin to induce menses before endometrial preparation.23 The two endometrial preparation protocols used before the FET were the following:

Hormone Replacement Cycles
In hormone replacement cycles, 4 mg of oral estradiol valerate was administered starting on the second or third day of the menstrual cycle and continuing for five days. This was followed by 6 mg of oral estradiol for 6–8 days. When the endometrial thickness reached 7 mm and the serum progesterone level was below 1.5 ng/mL, we added vaginal supplementation with progesterone 90 mg daily (8% Crinone, Merck-Serono, Switzerland) prior to FET. The embryo was transferred according to its development stage at the time of freezing. The supplementation continued until a pregnancy test was performed. In case of a positive test, the patients were instructed to continue treatment until the 10th gestational week.24

Stimulated Cycles
In stimulated cycles, patients received a daily subcutaneous injection of Gonafen (Merck Serono SA Aubonne Branch) (37.5–75 IU) from day 5 of the cycle onwards. The dose was adjusted according to the BMI, the ovarian reserve and any previous ovarian response to stimulation. A subcutaneous injection of hCG (5000 IU) or recombinant hCG (250 μg) was administered to induce oocyte ovulation, when the ovulation criteria were met (one dominant follicle ≥16 mm and peak plasma estradiol level ≥200 pg/mL). These patients had no intercourse on ovulation day. The adequacy of the luteal phase was evaluated by measuring blood progesterone levels three days after ovulation had been triggered. If the progesterone level 3 days after ovulation triggering exceeded 3 ng/mL, FET was implemented (depending on the embryo’s development stage at the time of freezing). STC protocols for endometrial preparation were not supplemented with progesterone.23

Study Endpoints and Definitions
Positive pregnancy was defined as a serum β-hCG level greater than 10 IU/L in the 14 days after cleavage embryo transfer.26,27 The patient underwent ultrasonographic monitoring to determine the number of gestational sacs and fetal viability at the 6th-7th week of gestation, ie, clinical pregnancy, if the β-hCG assay yielded a positive result. Pregnancy loss was defined as clinically recognized spontaneous loss of pregnancy before the completion of twenty gestational weeks. Ectopic pregnancy, defined as a pregnancy in which implantation takes place outside the uterine cavity, diagnosed by ultrasound, surgical visualization or histopathology. Live birth, defined as the birth
of at least one child with breath and heartbeat, irrespective of the duration of gestation. A birth weight of 3500 g or more can be used if gestational age is unknown. Furthermore, this pathological state in which the death of a fetus prior to the complete expulsion from its mother after 20 completed weeks of gestational age was diagnosed as stillbirth. As opposed to live birth, the fetus does not breathe or show any other evidence of life.

Statistical Analysis
All data are evaluated using version 26.0 of SPSS program (SPSS Inc., Chicago, USA). Shapiro–Wilk test was used to assess data normality. Quantitative variables are expressed as means ± standard deviations (SD) or median (range) and were analyzed using Student’s t-test (normal distribution), Independent-Samples Mann–Whitney U-Test (when normal distribution was not obeyed). Qualitative variables are expressed as frequencies and percentages and were analyzed using the χ²-test. P < 0.05 was considered statistically significant for the two groups of data tested. Furthermore, a propensity score matching (PSM) model was established to balance differences in baseline characteristics between the two groups.20 The propensity scores were calculated using binary logistic regression analyses based on the following patients’ characteristics: female age, infertility duration, body mass index (BMI), infertility type (primary or secondary), AMH, protocol of COS (Long GnRH-a protocol or GnRH-ant protocol), initial treatment (IVF or ICSI), Gn usage time, Gn dosage, oocytes retrieved, total number of embryos, good quality embryos, transferred embryos.28 Patients undergoing STC were matched with the HRC group using the nearest-neighbor random matching algorithm in a ratio of 1:1. In our study, we also used binary logistic regression analysis to assess the association between endometrial preparation protocols and pregnancy outcomes. We calculated crude odds ratios (OR) and adjusted OR with 95% confidence interval (CI).

Results
Demographic and ART Characteristics of Patients
One thousand four hundred and thirty-four cycles undergoing IVF or ICSI who had been performed the first freeze-thaw embryo transfer were studied. In detail, A total of 200 (14%) patients received STC, and 1234 (86%) underwent HRC before FET. Simultaneously, the 199 cycles were matched after PSM. Patient characteristics before and after PSM for STC and HRC groups are presented in Table 1. There were significantly different between two groups in initial treatment, good quality embryos, transferred embryos of D3 before PSM. Furthermore, no significant differences were found regarding patient characteristics between two groups after PSM.

Pregnancy Outcome Measures
Pregnancy outcomes reflected by matched FET method are shown in Table 2. However, no statistical significance was detected between STC and HRC groups in terms of positive pregnancy rate (PPR), clinical pregnancy rate (CPR), live birth rate (LBR), pregnancy loss rate (PLR) and ectopic pregnancy rate (EPR). (all P > 0.05)

A binary logistic regression model was also used to assess the association between endometrial preparation protocols and pregnancy outcomes while adjusting for potential confounders (Table 3). In the crude and adjusted models, the STC group was comparable to the HRC group in terms of PPR, CPR, and LBR.

Discussion
To our best knowledge, few studies have evaluated the different ways in which endometrium is prepared in young women with PCOS. In comparison to previous research, our practice can provide evidence-based guidance to select suitable endometrium preparation protocols for FET based on post-hoc randomization and large sample. In the current retrospective cohort analysis, we compared two different endometrial preparation protocols for FET with STC and HRC. Our findings showed that there was no statistical significance in the pregnancy outcomes between two groups.

PCOS resulted in infertility could have been attributed to anovulation as well as endometrial dysfunction which affect endometrial receptivity.16 In particular, hyperandrogenism and high level of LH during the follicular phase may decrease the rate of conception, the latter may lead to poor oocyte quality and embryo quality. Tomas et al noticed that the hormone replacement therapy (HRT) population has a higher pregnancy loss risk, which could be correlated with a higher prevalence of PCOS.29 Few studies have compared OS with HRT of PCOS patients in the reproductive outcomes. Most literature focuses on live birth rates and clinical pregnancy rates. In accordance with our outcomes, some literatures had the similar conclusion in HRT versus OS.17,30 In Yu et al retrospective study, the
two protocols resulted in LPR (30.0% vs 31.7%), CPR (41.0% vs 41.6%), OPR (36.6% vs 34.7%), which were not statistically different. In addition, there is a relatively high cycle cancellation rate in stimulated cycle. A systematic review and meta-analysis in 2016 similarly found STC and HRC endometrial preparation protocols are equally effective, despite the low quality of evidence, for women with PCOS. A systematic review and meta-

### Table 1 Patient Characteristics for HRC and STC Groups

| Variables                              | Before Propensity Score Matching |          | P-value |          |          |          |          |
|----------------------------------------|----------------------------------|----------|---------|----------|---------|---------|---------|
|                                        | STC (N=200)                      | HRC (N=1234) |        |          | STC (N=199) | HRC (N=199) |        |
| Female age (years)                     | 29.7±2.9                         | 29.6±3.1 | 0.725"  |          | 29.7±2.9 | 29.9±3.1 | 0.560"  |
| Infertility duration (years)           | 3 (1.11)                         | 3 (1.13) | 0.800"  |          | 3 (1.11) | 3 (1.13) | 0.397"  |
| BMI (kg/m²)                            | 23.7±3.9                         | 23.7±3.9 | 0.786"  |          | 23.8±3.9 | 23.8±3.5 | 0.975"  |
| Infertility type (n, %)                |                                  |          |         |          |         |         |         |
| Primary infertility                    |                                  |          |         |          |         |         |         |
| Secondary infertility                  |                                  |          |         |          |         |         |         |
| AMH (ng/mL)                            | 6.3±2.0                          | 6.5±2.1 | 0.319"  |          | 6.3±2.0 | 6.4±2.0 | 0.837"  |
| Protocol of COS (n, %)                 |                                  |          |         |          |         |         |         |
| Long GnRH-a protocol                  | 119/200 (59.5%)                  | 712/1234 (57.7%) | 0.632"  |          | 119/199 (59.8%) | 80/199 (40.2%) | 0.760"  |
| GnRH-ant protocol                     | 81/200 (40.5%)                   | 522/1234 (42.3%) |          |          | 80/199 (42.7%) | 83/199 (41.7%) |
| Initial treatment (n, %)               |                                  |          |         |          |         |         |         |
| IVF                                    | 115/200 (57.5%)                  | 427/1234 (34.6%) | <0.001" |          | 114/199 (57.4%) | 85/199 (42.7%) | 0.227"  |
| ICSI                                   | 85/200 (42.5%)                   | 807/1234 (65.4%) |          |          | 85/199 (42.7%) | 97/199 (48.7%) |
| Gn usage time (days)                   | 11.2±1.7                         | 11.0±1.6 | 0.102"  |          | 11.2±1.7 | 11.1±1.6 | 0.319"  |
| Gn dosage (IU)                         | 2326.6±547.6                     | 2274.0±582.5 | 0.232"  |          | 2330.0±546.9 | 2239.7±558.2 | 0.104"  |
| Oocytes retrieved (n)                  | 17 (2, 60)                       | 18 (1, 64) | 0.384"  |          | 17 (2, 60) | 18 (5, 57) | 0.897"  |
| Total number of embryos (n)            | 6 (1, 16)                        | 6 (1, 14) | 0.190"  |          | 6 (1, 16) | 6 (1, 14) | 0.629"  |
| Good quality embryos (n)              | 2 (0, 15)                        | 1 (0, 16) | 0.010"  |          | 2 (0, 15) | 1 (0, 12) | 0.310"  |
| Transferred embryos of D3 (n)          | 2±0.3                            | 1.9±0.3 | 0.002"  |          | 2±0.3 | 2.0±0.2 | 0.101"  |

Notes: Data are presented as mean ± SD, median (min, max) or n (%). "t-test for equality of means. t In independent-Sample Mann–Whitney U-test. χ²-test.

Abbreviations: STC, stimulated cycle; HRC, hormone replacement cycle; BMI, body mass index; IVF, in vitro fertilization; ICSI, intracytoplasmic sperm injection; AMH, anti-Müllerian hormone; COS, controlled ovarian stimulation; Gn, gonadotropin.

### Table 2 Pregnancy Outcomes for HRC and STC Groups

| Outcomes                             | Before Propensity Score Matching |          | P-value |          |          |          |          |
|--------------------------------------|----------------------------------|----------|---------|----------|---------|---------|---------|
|                                      | STC (N=200)                      | HRC (N=1234) |        |          | STC (N=199) | HRC (N=199) |        |
| Positive pregnancy rate, PPR (n, %)  | 106/200 (53.0%)                  | 630/1234 (51.1%) | 0.609  |          | 105/199 (52.7%) | 109/199 (54.8%) | 0.763  |
| Clinical pregnancy rate, CPR (n, %)  | 104/200 (52.0%)                  | 619/1234 (50.2%) | 0.630  |          | 103/199 (51.8%) | 105/199 (52.8%) | 0.841  |
| Live birth rate, LBR (n, %)           | 91/200 (45.5%)                   | 505/1234 (40.9%) | 0.223  |          | 90/199 (45.2%) | 87/199 (43.7%) | 0.762  |
| Pregnancy loss rate, PLR (n, %)       | 10/104 (9.6%)                    | 97/619 (15.7%) | 0.108  |          | 10/103 (9.7%) | 17/105 (16.2%) | 0.164  |
| Ectopic pregnancy rate, EPR (n, %)    | 3/200 (1.5%)                     | 17/1234 (1.4%) | 1.000  |          | 3/199 (1.5%) | 1/199 (0.5%) | 0.615  |

Notes: χ²-test.

Abbreviations: STC, stimulated cycle; HRC, hormone replacement cycle.
Table 3 Relationship Between Endometrial Preparation and Pregnancy Outcomes in Different Models

| Pregnancy Outcomes | Endometrial Preparation | Crude Model a | Adjusted Model b |
|---------------------|-------------------------|---------------|-----------------|
|                     | OR (95% CI)             | P value       | OR (95% CI)     | P value         |
| Positive pregnancy  |                        |               |                 |                 |
| HRC                 | Reference               | 1.08 (0.80 to 1.46) | 0.609          | Reference       | 1.05 (0.77 to 1.44) | 0.767 |
| STC                 | Reference               | 1.08 (0.80 to 0.45) | 0.630          | Reference       | 1.05 (0.76 to 1.43) | 0.190 |
| Clinical pregnancy  |                        |               |                 |                 |
| HRC                 | Reference               | 0.83 (0.61 to 1.12) | 0.223          | Reference       | 0.83 (0.61 to 1.14) | 0.246 |
| STC                 | Reference               |                 |                 |                 |
| Live birth          |                        |               |                 |                 |
| HRC                 | Reference               |                 |                 |                 |
| STC                 | Reference               |                 |                 |                 |

Notes: *No adjustments for other covariates. *Adjusted for female age, infertility duration, infertility type (primary, secondary), BMI (<24 kg/m², ≥24 kg/m²), AMH, protocol of COS (Long GnRH-a protocol, GnRH-ant protocol), number of oocytes retrieved, total number of embryos, good quality embryos, number of transferred embryos, and initial treatment (IVF, ICSI).

Abbreviations: BMI, body mass index; AMH, anti-Müllerian hormone; OR, odds ratio; CI, confidence interval; STC, stimulated cycle; HRC, hormone replacement cycle.

analysis including pooled results of only two studies of PCOS patients found letrozole produces similar CPR, LBR, and birth defect rates as NC and HRC; there were similar CPR and lower LBR in letrozole-stimulated cycle compared to HMG stimulation. Added to Chen et al conclusions, a recent meta-analysis comparing OS using letrozole or HMG with HRT for FET in patients with PCOS found no difference between mild OS cycles and HRT groups for OPR and embryo implantation rate (IR); the letrozole-stimulated cycle may lower the miscarriage rate more than the HRT cycle. Multiple retrospective cohort studies pointed out a different view, letrozole-stimulated cycle had significantly higher LBR and lower PLR compared with HRT after adjusting for possible confounding factors. Therefore, whether letrozole has an advantage in preparing FET requires more high-quality researches for confirmation. In contrast with our findings, in a recent historical cohort analysis on women with PCOS, OS protocol achieves a better pregnancy outcome than the HRT protocol. In detail, that LBR with HRT requires more transvaginal ultrasound examinations, urine LH measurement, precise hCG injection timing, and subsequent FET in order to prevent missing the implantation.
window. As a result, HRC might be a viable option for young PCOS patients undergoing IVF-ET and being unable to take injections. STC may reduce unnecessary anxiety and operational costs, and offer more flexibility for patients. Eventually, we must embrace the concepts of individualization, securitization, and optimization in the clinic.

Although this study has the advantage of using PSM to balance the variables that potentially affect the outcomes. Nevertheless, there are some drawbacks to this study. Every retrospective nature may not allow completely introduced selection or information bias. First of all, we cannot investigate other confounding factors, including exercise, nutritional supplements and diet, which may add information bias. Second, since this is not an RCT, according to professional experience, patients are assigned to multiple groups, which can add selection bias.

**Conclusion**

In conclusion, STC for endometrial preparation had similar PPR, CPR, LBR, PLR, EPR compared with HRC by excluding heterogeneous factors after PSM. For young women with PCOS who were undergoing in-vitro fertilization-embryo transfer (IVF-ET), HRC could be a reasonable choice for patients who are unwilling to accept injections. Additionally, STC may reduce unnecessary anxiety and operational costs, and offer more flexibility for patients and IVF centres. We should follow the principles of individualization, securitization and optimization. To validate the obtained results, broader analyses, as well as an economic assessment of the costs involved, are required.

**Data Sharing Statement**

The data generated and analyzed from the current study will be availed by the corresponding author upon request.

**Ethics Approval and Consent to Participate**

This study was conducted in accordance with the Declaration of Helsinki. The study was approved by the Reproductive Ethics Committees of the Affiliated Hospital of Shandong University of TCM (ref approval no. SDTCM20201215). All participants provided written informed consent.

**Consent for Publication**

Written informed consent for publication was obtained from all participants.

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**Author Contributions**

Zhen-Gao Sun and Jing-Yan Song conceived and designed the study. All authors made a significant contribution to the work reported, whether that is in the conception, study design, execution, acquisition of data, analysis and interpretation, or in all these areas; took part in drafting, revising or critically reviewing the article; gave final approval of the version to be published; have agreed on the journal to which the article has been submitted; and agree to be accountable for all aspects of the work.

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**Disclosure**

None of the authors have a conflict of interest to declare with regard to this study.

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