Objective
To evaluate whether oral contraceptive pill (OCP) pretreatments in gonadotropin-releasing hormone (GnRH) antagonist ovarian stimulation protocols take positive effects on in vitro fertilization (IVF) outcomes in respect to retrieved oocyte number, oocyte maturation rate, fertilization rate, good quality embryo rate, cycle cancellation rate, pregnancy rate and clinical abortion rate.

Methods
A total of 194 cycles using GnRH antagonist protocol was performed at infertility clinic of our institute from September 1st, 2009 to February 28th, 2010. The medical records of GnRH antagonist protocols for IVF with or without OCP pretreatment in our IVF unit were retrospectively analyzed. We compared the IVF outcomes between OCP pretreated (n=41) and no pretreatment group (n=153).

Results
In cycles with OCP pretreated group, the total used dosage of gonadotropin (3019.38±1379.00 IU) were higher than that of no pretreatment group (2551.52 ± 1157.05 IU, P = 0.054). The duration of ovarian stimulation in OCP pretreated group (11.5 ± 2.0) was significantly longer than that of control group (9.5 ± 1.9, P = 0.000). The number of gained total embryo (2.8±0.9 vs. 2.5±1.0, P = 0.055) and fertilization rate (77.2% vs. 65.5%, P = 0.017) were significantly higher in OCP pretreated group. There is no significant difference in pregnancy rate between two groups (39.4% vs. 30.0%, P = 0.304).

Conclusion
OCP pretreatment before GnRH antagonist protocol for IVF appears not to have reliable benefit in terms of IVF outcomes. Well-controlled and large-scaled studies are needed.

Keywords: Gonadotropin-releasing hormone antagonist; Oral contraceptive pill pretreatment; Ovarian stimulation; In vitro fertilization

Gonadotropin-releasing hormone (GnRH) antagonists have been widely used after its first introduction in assisted reproductive technologies to prevent a premature luteinization [1]. GnRH antagonist protocols are preferred for poor responders because of shorter duration and use of lower amount of gonadotropins for ovarian stimulation as compared with traditional GnRH long agonist protocols [2]. However, it induces insufficient synchronization of follicular development and lack of flexibility in the starting day of ovarian stimulation, which is less likely in GnRH agonist long protocols [3]. For getting over these limitations, several pretreatments have been applied [4-7]. Among them, oral contraceptive

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Pills (OCP) pretreatment has been reported to induce higher numbers of oocytes retrieved compared to no pretreatment group in GnRH antagonist cycles [8]. However, OCP pretreated GnRH antagonist cycles presented longer duration and needed more larger amount of gonadotropin for stimulation than no pretreatment cycles [8, 9].

In this study, we aimed to evaluate the effect of OCP pretreatment in in vitro fertilization (IVF) cycles using GnRH antagonists by comparison between OCP-pretreated and non-treated groups.

**Materials and Methods**

A total of 421 cycles of GnRH antagonist protocol were performed at infertility clinic of our institute from September 1st, 2009 to February 28th, 2010. After exclusion of cycles with male factor infertility, of having uterine anomalies, and cycles for pre-implantation diagnosis, 194 GnRH antagonist cycles were included in this study. Among them, 41 cycles used OCP pretreatment in previous menstrual cycle before starting GnRH antagonist protocol for IVF (OCP pretreated group). In contrary, 153 cycles used no medicine before GnRH antagonist protocol for IVF (no pretreatment group).

We reviewed the medical records of each group and compared basal characteristics of the cycles and IVF outcomes between two groups.

In OCP pretreated group, daily OCP (Myvlar®, Bayer Schering Pharma AG, Berlin, Germany) was applied from 3rd day of previous menstrual cycle. About 3-4 days after OCP discontinuation, the ovarian stimulation was done with gonadotropin as extensively described. Briefly, the patients underwent pituitary downregulation with daily GnRH antagonist (Orgalutran®, Schering-Plough, Whitehouse Station, NJ, USA or Cetrotide®, Merck-Serono, Geneva, Switzerland) from mid or late follicular period of this cycle applied when dominant follicle reached to 12 or 13 mm. When two or more follicles reached 18 mm in diameter 5,000-10,000 IU of hCG (Ovidrel®, Merck Serono, Bari, Italy) was administered. Trans-vaginal ultrasound guided oocyte pick-up (OPU) was performed 34-36 hours later and then, maturity and quality of retrieved oocytes was evaluated.

According to the quality and number of sperm, conventional insemination or microinjection (intracytoplasmic sperm injection, ICSI) was carried out 4-6 hours later. After 3-5 days of in vitro culture, less than 4 selected embryos were transferred to uterus. For the purpose of luteal support, daily intramuscular injection of progesterone (Progesterone®, Watson Pharmaceuticals Inc., Morristown, NJ, USA) was started at the time of OPU day up to first inspection of pregnancy. In case of successful pregnancy, luteal support was continued to 7th to 8th gestational weeks.

The IVF outcomes such as retrieved oocyte number, oocyte maturation rate, fertilization rate, good quality embryo rate, cycle cancellation rate, pregnancy rate and clinical abortion rate were compared between OCP pretreated and no pretreatment group. Statistical analysis was performed using SPSS ver. 16.0 (SPSS Inc., Chicago, IL, USA). Each variable was presented as mean ± standard deviation. Student’s t-test and Chi-square test were used wherever appropriate. P-value of < 0.05 was considered statistically significant.

**Results**

The mean age and body mass index in OCP pretreated and non-pretreated groups were similar (35.8 ± 3.1 vs. 36.8 ± 3.5 years and 21.1 ± 2.4 kg/m² vs. 20.5 ± 1.9 kg/m²). The basal follicle stimulating hormone (FSH) level (9.89 ± 4.0 IU/mL vs. 10.5 ± 2.1 IU/mL) was also similar between two groups. The primary infertility rate tended to be higher in OCP pretreated group (65.9% vs. 55.6%) but not statistically significant. The mean duration of infertility was longer in control group (52.4 ± 40.2 months vs. 49.7 ± 33.3 months) but not statistically significant (Table 1).

Mean amount of gonadotropins for controlled ovarian stimulation in OCP pretreated group was higher than that of control group. The mean duration of ovarian stimulation in OCP pretreated group was significantly longer than that of no pretreatment group. The mean number of retrieved oocytes was similar between two groups. The good quality embryo rate and oocyte maturation rate tended to be higher in OCP pretreated group (65.9% vs. 55.6%) but not statistically significant. The number of gained total embryo was higher in OCP pretreated group with borderline significance (P = 0.055). Fertilization rate was also higher in OCP pretreated group. The implantation and pregnancy rate were similar between two groups. The clinical abortion rate was also showed no significant difference between two groups. The cycle cancellation rate tended to higher in OCP pretreated group than OCP non-treated group but not statistically different.

**Discussion**

The retrieval of good quality oocyte is very important factor to
achieve pregnancy in infertile women, especially older women who are barely gettable many oocytes with one cycle of controlled ovarian hyperstimulation (COH). To gain good quality embryo, growth of finely matured oocyte is firstly needed. To get more matured oocytes, the synchronized growing of follicles is one of important factor.

During COH, most of the early antral follicles are required to grow coordinately in response to exogenous gonadotropins thus accomplishing simultaneous functional and morphological maturation. Marked discrepancies of follicular size at the end of COH may be counterproductive since they imply that a substantial fraction of FSH-sensitive follicles fail to undergo satisfactory maturation. This phenomenon potentially reduces the number of viable oocytes and embryos and the probability of conception. Selection of good embryos for transfer depends on embryo cohort size: implications for the ‘mild ovarian stimulation’ debate [10]. The number of embryos available for transfer predicts successful pregnancy outcome in women over 39 years with normal ovarian hormonal reserve testing [11]. During the last year, we took notice of unusually low maturation rate of oocytes, especially in GnRH antagonist cycles. Primarily, we thought it derived from rapidly rising age of patients which reached 36 years old averagely. And then we assumed that the less competent oocyte is caused by asynchronous follicular development and a limited number of dominant follicles due to ovarian stimulation without pituitary suppression in GnRH antagonist protocols.

Asynchronous multi-follicular growth during COH may be a direct consequence of size heterogeneities of early antral follicles during controlled ovarian hyperstimulation.

Table 1. Comparison of baseline characteristics and ART outcomes

<table>
<thead>
<tr>
<th></th>
<th>OCP pretreated group (n = 41)</th>
<th>No pretreatment group (n = 153)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age of female (yr)</td>
<td>35.8 ± 3.1</td>
<td>36.8 ± 3.5</td>
<td>NS</td>
</tr>
<tr>
<td>Age of husband (yr)</td>
<td>36.9 ± 4.0</td>
<td>38.3 ± 4.0</td>
<td>0.039</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>21.1 ± 2.4</td>
<td>20.5 ± 1.9</td>
<td>NS</td>
</tr>
<tr>
<td>Basal serum FSH (mIU/mL)</td>
<td>9.9 ± 4.0</td>
<td>10.5 ± 2.1</td>
<td>NS</td>
</tr>
<tr>
<td>Primary infertility (%)</td>
<td>65.9</td>
<td>55.6</td>
<td>NS</td>
</tr>
<tr>
<td>Duration of infertility (mon.)</td>
<td>49.7 ± 33.3</td>
<td>52.4 ± 40.2</td>
<td>NS</td>
</tr>
<tr>
<td>Dosage of gonadotropin (IU)</td>
<td>3019.4 ± 1379.0</td>
<td>2551.5 ± 1157.1</td>
<td>0.054</td>
</tr>
<tr>
<td>Duration of COH (day)</td>
<td>11.5 ± 2.1</td>
<td>9.5 ± 1.9</td>
<td>0.000</td>
</tr>
<tr>
<td>E₂ on hCG day (pg/mL)</td>
<td>1170.8 ± 1267.3</td>
<td>1086.0 ± 877.5</td>
<td>NS</td>
</tr>
<tr>
<td>Em thickness on hCG day (mm)</td>
<td>10.4 ± 2.6</td>
<td>10.0 ± 2.4</td>
<td>NS</td>
</tr>
<tr>
<td>Number of retrieved oocyte</td>
<td>7.2 ± 5.1</td>
<td>7.0 ± 5.2</td>
<td>NS</td>
</tr>
<tr>
<td>Number of matured oocyte</td>
<td>5.6 ± 4.3</td>
<td>5.1 ± 4.1</td>
<td>NS</td>
</tr>
<tr>
<td>Oocyte maturation rate (%)</td>
<td>81.1</td>
<td>75.1</td>
<td>NS</td>
</tr>
<tr>
<td>Number of total gained embryo</td>
<td>2.8 ± 0.9</td>
<td>2.5 ± 1.0</td>
<td>0.055</td>
</tr>
<tr>
<td>Good quality embryo rate (%)</td>
<td>26.8</td>
<td>19.1</td>
<td>NS</td>
</tr>
<tr>
<td>Number of transferred embryo</td>
<td>2.6 ± 0.9</td>
<td>2.3 ± 1.0</td>
<td>NS</td>
</tr>
<tr>
<td>Implantation rate (%)</td>
<td>10.6</td>
<td>10.5</td>
<td>NS</td>
</tr>
<tr>
<td>Fertilization rate (%)</td>
<td>77.2 ± 0.2</td>
<td>65.5 ± 0.3</td>
<td>0.017</td>
</tr>
<tr>
<td>ICSI rate (%)</td>
<td>75.8</td>
<td>83.1</td>
<td>NS</td>
</tr>
<tr>
<td>Cycle cancellation rate (%)</td>
<td>19.5</td>
<td>14.5</td>
<td>0.468</td>
</tr>
<tr>
<td>Pregnancy rate/embryo transfer (%)</td>
<td>39.4</td>
<td>30.0</td>
<td>0.304</td>
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<tr>
<td>Clinical abortion rate (%)</td>
<td>41.7</td>
<td>42.1</td>
<td>1.000</td>
</tr>
</tbody>
</table>

Values are presented as mean ± standard deviation.
ART, assisted reproductive technology; OCP, oral contraceptive pill; NS, not statistically significant; BMI, body mass index; FSH, follicle stimulating hormone; COH, controlled ovarian hyperstimulation; E₂, estradiol; EM, endometrial thickness; hCG, human chorionic gonadotropin; ICSI, intracytoplasmic sperm injection.

*Statistically significant (P-value < 0.05).
ing the early follicular phase [12]. Luteal estradiol administration strengthens the relationship between day 3 FSH and inhibin B levels and ovarian follicular status [13]. Some follicles are able to respond to lower FSH levels than others by their intrinsic sensitivity to FSH, and, start their development during the late luteal phase [14]. Since larger follicles are more FSH responsive than are smaller follicles, exogenous gonadotrophin administration is likely to intensify further size discrepancies of growing follicles during COH [15]. Follicular development begins during the luteal phase of the human menstrual cycle. Hence, COH protocols such as mid-luteal long protocol, suppression of luteal FSH secretion could prevent untimely and uncoordinated development of FSH-sensitive follicles during the luteal-follicular transition and faster follicular growth synchronization during COH can be obtained [16]. However, this luteal suppression of FSH cannot be achieved in GnRH antagonist COH protocols. Therefore, marked follicular size discrepancies would be occurred in GnRH antagonist COH cycles. In previous study, OCP pretreatment in GnRH antagonist cycles in low responders resulted in improving ovarian response by intrinsic gonadotropins before COH [17]. Based on that result, we tried to apply OCP pretreatment in GnRH antagonist IVF cycles bearing in mind their insufficient action of follicular synchronization.

In our study, the baseline characteristics of IVF cycles and outcomes between both groups were comparable regardless of age. In OCP pretreated group, as shown in Fig. 1, it presented improvement of fertilization rate and gained more number of fertilized embryos than that of OCP non-pretreated group even if longer duration and larger used dose of gonadotropin for ovarian stimulation. According to previous meta-analysis regarding OCP pretreatment in GnRH antagonist cycles [18], OCP pretreatment was associated with an increased gonadotropin consumption and increased duration of stimulation without improvement of ongoing pregnancy rate. There were many other studies which concerned to OCP pretreatment and IVF outcomes in GnRH antagonist cycles. Among them, Kolibianakis et al. [19], reported that OCP pretreated GnRH antagonist COH cycles have no significant benefit in ongoing pregnancy rates and moreover results in a significantly higher early pregnancy loss of compared to non-OCP cycles. In another systemic review and meta-analysis analyzed by Griesinger et al. [20], OCP pretreatment in GnRH antagonist for COH have no significant benefit in increasing ongoing pregnancy rates. Those results were in accordance with our current data. A recent study focused on compromised group like as low responders [21]. The study showed higher number of retrieved and matured oocytes, and fertilized oocytes in OCP pretreatment group in low responders which was defined as elevated basal FSH level (>8.5 mIU/mL), and/or antral follicle count <5. In the present study, the number of gained embryo and oocyte fertilization rate were higher in cycles of OCP pretreatment.

Old age and high basal FSH level are predictors of poor responder in controlled ovarian stimulation for IVF. However, these two parameters do not always show consistent result, and moreover basal FSH level is variable by each cycle. The other study focused on women's age which divided into young (less than 36 years old) and old group (over 36 years old); they reported that OCP pretreatment in GnRH antagonist cycles just results in increased use of gonadotropin and increased time of stimulation without any benefit, regardless of age [22].

In summary, OCP pretreatment for women in GnRH antagonist protocol is a valid option in terms of better fertilization rate and higher number of fertilized embryos. The oocyte maturation rate appears to improve by OCP pretreatment. But, it also has a weak point in respect to longer stimulation duration and increased gonadotropin consumption. The OCP pretreated cycle had no ad-
vantages in respect to pregnancy rate, cycle cancellation rate, and clinical abortion rates.

In conclusion, OCP pretreatment before GnRH antagonist protocol for IVF seems to have no reliable benefit in respect to final IVF outcomes. Our present study have a some different features from other previous studies because we focused on new aspects of the effects, such as maturation and fertilization rate in OCP pretreatment in GnRH antagonist protocols although it was retrospective small. Well controlled, large scaled studies are needed to support ineffectiveness of OCP pretreatment before starting GnRH antagonist ovarian stimulation protocol for IVF.

References

18. Bodri D, Sunkara SK, Coomarasamy A. Gonadotropin-releasing hormone agonists versus antagonists for controlled ovarian hyperstimulation in oocyte donors: a systematic review and
생식샘자극호르몬 분비호르몬 길항제를 이용한 난소자극 투여법에서의 경구복합피임제의 전처치: 비교 연구

판동대학교 의과대학 제일병원 산부인과
추연실, 한혜라, 양승헌, 성나영, 차선화, 김혜옥, 박찬우, 송인옥, 궁미경, 양광문

목적
생식샘자극호르몬 분비호르몬 길항제를 이용한 난소자극 투여법에서 경구복합피임제의 전처치가 체외수정시술의 결과에 미치는 효과에 대해 평가해 보고자 하였다.

연구방법
2009년 9월 1일에서 2010년 2월 28일까지 본원 불임센터에서 체외수정시술을 위해 총 194회의 생식샘자극호르몬 분비호르몬 길항제 요법을 시행한 환자 중 경구복합피임제를 전처치한 경우와 그렇지 않은 경우의 체외수정 및 배아이식술의 진료 기록을 후향적으로 분석하였다.

결과
경구복합피임제 전처치군의 경우 사용된 생식샘자극호르몬의 사용량이 전처치하지 않은 군보다 많았다(3019.38±1379.90 vs. 2551.52 ± 1157.05 IU, P=0.054). 난소 자극 시작부터 난소 채취를 위한 마지막 생식샘자극호르몬의 투여일까지 기간 또한 경구복합피임제 전처치 군(11.5 ± 2.0)에서 전처치하지 않은 군(9.5 ± 1.9)에 비해 통계학적으로 의미 있게 길었다(P=0.000). 배아 총 획득 갯수(2.8 ± 0.9 vs. 2.5 ± 1.0, P=0.055) 및 난자수정률(77.2% vs. 65.5%, P=0.017) 또한 경구복합피임제 전처치 군에서 난자 수정률에 비해 통계학적으로 의미 있게 높았다. 한편, 임신율에 있어서는 두 그룹 간에 통계학적으로 유의한 차이가 없었다(39.4% vs. 30.0%; P=0.0304).

결론
체외수정시술을 위한 생식샘자극호르몬 분비호르몬 길항제 사용 시 경구복합피임제를 전처치 하였을 때, 그렇지 않은 군에 비해 최종적인 체외수정시술의 결과는 총에서 현저한 차이는 없는 것으로 보인다. 하지만 좀 더 명확한 결론을 도출하기 위해서는 더 많은 연구군을 포함한 전량적인 연구가 필요할 것으로 생각된다.

중심단어: 생식샘자극호르몬 분비호르몬 길항제, 경구복합피임제 전처치, 난소자극투여법, 체외수정시술