Cycle scheduling for in vitro fertilization with oral contraceptive pills versus oral estradiol valerate: a randomized, controlled trial

Erik E Hauzman\textsuperscript{1}, Azucena Zapata\textsuperscript{1}, Alfonso Bermejo\textsuperscript{1}, Carlos Iglesias\textsuperscript{1}, Antonio Pellicer\textsuperscript{2,3} and Juan A Garcia-Velasco\textsuperscript{1,4*}

Abstract

\textbf{Background:} Both oral contraceptive pills (OCPs) and estradiol (E\textsubscript{2}) valerate have been used to schedule gonadotropin-releasing hormone (GnRH) antagonist in vitro fertilization (IVF) cycles and, consequently, laboratory activities. However, there are no studies comparing treatment outcomes directly between these two pretreatment methods. This randomized controlled trial was aimed at finding differences in ongoing pregnancy rates between GnRH antagonist IVF cycles scheduled with OCPs or E\textsubscript{2} valerate.

\textbf{Methods:} Between January and May 2012, one hundred consecutive patients (nonobese, regularly cycling women 18–38 years with normal day 3 hormone levels and <3 previous IVF/ICSI attempts) undergoing IVF with the GnRH antagonist protocol were randomized to either the OCP or E\textsubscript{2} pretreatment arms, with no restrictions such as blocking or stratification. Authors involved in data collection and analysis were blinded to group assignment. Fifty patients received OCP (30 $\mu$g ethinylinE\textsubscript{2}/150 $\mu$g levonorgestrel) for 12–16 days from day 1 or 2, and stimulation was started 5 days after stopping OCP. Similarly, 50 patients received 4 mg/day oral E\textsubscript{2} valerate from day 20 for 5–12 days, until the day before starting stimulation.

\textbf{Results:} Pretreatment with OCP (mean±SD, 14.5±1.7 days) was significantly longer than with E\textsubscript{2} (7.8±1.9 days). Stimulation and embryological characteristics were similar. Ongoing pregnancy rates (46.0% vs. 44.0%; risk difference, \textminus2.0\% [95\% CI \textminus21.2\% to 17.3\%]), as well as implantation (43.5\% vs. 47.4\%), clinical pregnancy (50.0\% vs. 48.0\%), clinical miscarriage (7.1\% vs. 7.7\%), and live birth (42.0\% vs. 40.0\%) rates were comparable between groups.

\textbf{Conclusions:} This is the first study to directly compare these two methods of cycle scheduling in GnRH antagonist cycles. Our results fail to show statistically significant differences in ongoing pregnancy rates between pretreatment with OCP and E\textsubscript{2} for IVF with the GnRH antagonist protocol. Although the study is limited by its sample size, our results may contribute to a future meta-analysis. An interesting future direction would be to extend our study to women with decreased ovarian reserve, as these are the patients in whom an increase in oocyte yield—due to the hypothetical beneficial effect of steroid pretreatment on follicular synchronization—could more easily be demonstrated.

\textbf{Trial registration:} ClinicalTrials.gov NCT01501448.

\textbf{Keywords:} IVF, GnRH antagonist, Cycle scheduling, Oral contraceptives, Estrogen pretreatment

\* Correspondence: juan.garcia.velasco@ivi.es
\textsuperscript{1}IVI Madrid, Avda del Talgo, 68-70, 28023 Madrid, Spain
\textsuperscript{2}Rey Juan Carlos University, Avda del Talgo 68-70, 28023 Madrid, Spain

Full list of author information is available at the end of the article

© 2013 Hauzman et al; licensee BioMed Central Ltd. This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/2.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.
Background
In the last decade, gonadotropin-releasing hormone (GnRH) antagonist protocols have become increasingly popular for controlled ovarian hyperstimulation (COH). Their use offers several advantages over the long agonist protocol [1], including shorter overall treatment duration, absence of perimenopausal symptoms caused by pituitary desensitization, no risk of inadvertent administration at the beginning of pregnancy or of ovarian cyst formation, and a lower consumption of gonadotropins. According to the conclusions of a recent meta-analysis [2], similar live birth rates can be achieved with the use of GnRH antagonist protocols, while significantly reducing the risk of ovarian hyperstimulation syndrome, compared to the long GnRH agonist protocol.

A potential disadvantage of GnRH antagonist protocols is that, classically, stimulation is started on day 2 or 3 of the menstrual cycle, which makes it difficult to schedule stimulation and laboratory activities. Cycle scheduling can be used to avoid weekend retrievals and to equally distribute the workload throughout the week, thereby avoiding excessive opening of incubator doors and the associated negative impact on embryo development, as well as reducing the amount of unplanned work, which can result in loss of concentration and reduced efficiency of the laboratory staff [3-5].

Both oral contraceptive pills (OCPs) and synthetic progestogens have been used for many years to schedule cycles [6-9]. More recently, the use of natural estrogens has also been advocated [10]. Estrogens primarily inhibit FSH secretion, whereas progestogens are mainly involved in the control of LH secretion.

Some smaller-scale studies on the hypothetical negative impact of OCPs on IVF cycles yielded controversial results in terms of ongoing pregnancy rates [11,12], while a recent meta-analysis, which summarized the results of six randomized controlled trials, found that ongoing pregnancy rates in normal responders were affected by OCP pretreatment [13]. However, this meta-analysis was confounded by the use of different OCPs for a varying number of days and with a varying pill-free interval. As far as the underlying mechanisms are concerned, it has been postulated that the gestagen component of OCPs could exert a negative impact on endometrial receptivity in the subsequent cycle. Alternatively, low LH concentrations after OCP pretreatment could impair oocyte competence or endometrial receptivity when ovarian stimulation is performed with recombinant FSH without LH [14]. However, in a recent randomized, controlled trial performed by our group that used one type of OCP for a relatively narrow range of days followed by a fixed pill-free period, we did not find any significant differences in ongoing pregnancy rates or live birth rates compared to the long agonist protocol [15].

The use of oral estrogens started in the midluteal phase of the cycle preceding ovarian stimulation was proposed recently for cycle scheduling purposes, based on the inhibitory effect of estradiol (E2) on follicle growth through its negative feedback on the increase in FSH during the luteal–follicular transition [16], which stops as soon as E2 is discontinued. Pretreatment with estrogens offers the advantage of a shorter duration of pretreatment than OCPs. Previous studies showed that estrogen pretreatment does not affect cycle outcome as compared to no pretreatment [17-19] or to the long GnRH agonist protocol [20], while it does promote follicular growth coordination during ovarian stimulation [21]. However, none of the studies compared treatment outcomes directly with those obtained after scheduling with OCPs. The aim of our study was to see whether these two methods of cycle scheduling are equally effective in terms of ongoing pregnancy rate and the secondary outcome measures detailed below.

Methods
Subjects
Between January and May 2012, we enrolled 118 women who were undergoing an IVF–ICSI cycle at IVI Madrid (Madrid, Spain). The inclusion criteria were age 18–38 years, regular normo-ovulatory menstrual cycles (26–35 days), body mass index (BMI) < 30 kg/m², normal cycle day-3 basal serum hormone levels (FSH < 10 IU/l and E2 < 60 pg/ml), and <3 previous IVF/ICSI attempts. Exclusion criteria were previous ovarian surgery, low ovarian response (cancellation of the cycle due to poor follicular development after at least 7 days of gonadotropin stimulation, or <5 oocytes retrieved) in a previous IVF/ICSI cycle, and polycystic ovarian syndrome according to the Rotterdam criteria [22]. A total of 100 women were included in the study, with only one cycle per patient (Figure 1). This project was approved by our institutional review board, and all patients provided written informed consent. The study was registered with ClinicalTrials.gov (identifier: NCT01501448).

Randomization procedure
A study nurse randomized and assigned patients at the time of cycle scheduling to either the OCP or E2 pretreatment arms in 1:1 ratio, based on a computer-generated random number list, with no restrictions such as blocking or stratification. The sequence was concealed in opaque, consecutively numbered envelopes until an intervention was assigned. Patients meeting all inclusion criteria were recruited consecutively. Authors involved in data collection and data analysis were blinded to group assignment.

Protocols
Patients randomized to the OCP group started with the pill (30 μg of ethinyl E2 plus 150 μg of levonorgestrel [Microgynon; Bayer Schering Pharma, Berlin, Germany])
(n = 50) on day 1 or 2 of menses of the cycle prior to the scheduled IVF/ICSI procedure, and they took it for 12–16 days. As suggested by Cédrin-Durnerin et al. [23], ovarian stimulation was initiated 5 days after discontinuation of the pill, regardless of the specific day of the onset of menses, with a daily dose of recombinant FSH (Puregon; MSD, Oss, The Netherlands, or Gonal F; Merck Serono, Madrid, Spain) ranging from 200 to 225 IU on the first 5 days. From day 6 of stimulation, gonadotropin doses were adjusted according to serum E2 levels and ovarian response, which was assessed by vaginal ultrasound, every 2 days.

In patients allocated to the E2 arm of the study (n = 50), pretreatment with E2 valerate (Meriestra; Novartis Farmacéutica, Barcelona, Spain) was started on day 20 of the cycle preceding the IVF/ICSI cycle at a daily dose of 4 mg (2 mg twice a day) orally for 5–12 days, until the day before the initiation of ovarian stimulation, independent of the specific day of onset of menstruation, as it has been shown that the number of days by which E2 pretreatment is extended beyond the menses does not have a significant effect on cycle outcome [24].

The number of pretreatment days in both groups was chosen with the aim of starting stimulation between Wednesday and Sunday and thus equally distributing oocyte pickup among working days and minimizing weekend retrievals, as coordinated by a designated person at the centre.

In both groups, the GnRH antagonist ganirelix (Orgalutran; MSD) was introduced at a daily dose of 0.25 mg when the leading follicle reached 13 mm in mean diameter. Ovarian triggering was performed with 250 μg of recombinant human chorionic gonadotropin (hCG) (Ovitrelle; Merck Serono), which was administered as soon as two leading follicles reached ≥17 mm mean diameter. Ovum pickup was performed 36 hours later. IVF or ICSI was used to fertilize oocytes, according to individual requirements. A maximum of two embryos were transferred on day 3, under ultrasound guidance. Luteal support was started with 200 mg of micronized vaginal progesterone (Progeffik 200; Laboratorios Alcalá Farma, Alcalá de Henares, Spain) every 12 hours, beginning the night after ovum pickup. Serum βhCG was evaluated 12 days after embryo transfer; a result of > 10 IU/l was
considered positive. Clinical pregnancy was confirmed by transvaginal ultrasound 2 weeks later if βhCG was positive.

Clinical pregnancy was defined as an intrauterine sac with heartbeat 4 weeks after embryo transfer, whereas ongoing pregnancies were characterized by the presence of a developing embryo at >12 weeks of gestation.

**Sample size estimate**

The primary outcome of the study was ongoing pregnancy rate. Secondary outcome variables were implantation rate, clinical pregnancy rate, miscarriage rate, and live birth rate.

Sample size calculations were based on the results of our recently published previous study [15], in which ongoing pregnancy rates in GnRH antagonist cycles pretreated with OCP (47.8%) were compared with the GnRH agonist protocol. Based on a two-sided significance level of 0.05 and a power of 80%, at least 1,555 cycles in each group would have been necessary to detect a 5% difference in ongoing pregnancy rates. However, reaching a sample size large enough for an adequately powered investigation is not feasible for a single-centre study. Therefore, we arbitrarily chose the number of patients to be recruited in this study to provide clinically useful data that could be incorporated into a future meta-analysis.

**Statistical analysis**

Categorical data are expressed as number and percentage, and continuous data as mean ± standard deviation (SD). For categorical variables, the χ² test was used with continuity correction, whereas continuous variables were analyzed with Student’s t-test for independent samples, after testing for normality. A P value of <0.05 was considered to be statistically significant. All statistical analyses were performed with the SPSS 13.0 package (SPSS, Inc., Chicago, IL, USA).

**Results**

As presented in Table 1, both groups of patients (n = 50 in both groups) were comparable in terms of age, BMI, and number of previous IVF attempts.

Due to the difference between the two pretreatment protocols, significantly more days of pretreatment with OCP compared to E2 (14.5 ± 1.7 vs. 7.8 ± 1.9 days, P < 0.001) were necessary before starting stimulation. All parameters of stimulation that we analyzed (duration of stimulation, total amount of FSH used, and peak E2 and progesterone levels) were comparable between the groups (Table 2).

Stimulation was cancelled in three patients receiving OCP pretreatment and two patients pretreated with E2 due to low response in all cases. Of those patients who did undergo oocyte pickup (47 and 48 patients, respectively), four and seven women did not reach the stage of embryo transfer in the OCP and E2 groups, respectively, because no embryos were available on day 3 (Figure 1).

No significant differences were observed in the number of retrieved oocytes, fertilization rate, number of top-quality embryos, or number of transferred embryos. With regard to the outcome of the cycles, implantation rates, total and clinical pregnancy rates, early clinical miscarriage rates, and ongoing pregnancy rates and live birth rates were all comparable between the two groups (there were two late miscarriages in both groups) (Table 3). All analyses were based on the intention-to-treat principle. However, even an “as treated” analysis of the primary outcome measure, based on 23/43 vs. 22/41 ongoing pregnancies, would give a difference that remains far from statistical significance (P = 0.84).

The proportion of oocyte retrievals performed on weekend days was similar between the groups: 8.5% (4/47) with OCP and 10.4% (5/48) with E2 pretreatment (P = 0.97) (Figure 2). Both frequencies were significantly lower than 28.6%, which would be expected to occur by pure chance (i.e., on 2 out of 7 days) (P = 0.03 for OCP and P = 0.04 for E2).

**Discussion**

Our results failed to show statistically significant differences for any measures of IVF treatment outcome between pretreatment with OCP and E2 for COH with the GnRH antagonist protocol. To the best of our knowledge, this is the first study to directly compare these two methods of cycle scheduling in GnRH antagonist cycles.

**Table 2 Stimulation cycle parameters**

| Characteristics | OCP (n = 50) | E2 (n = 50) | P value |
|-----------------|-------------|-------------|---------|
| Pretreatment days | 14.5 ± 1.7 | 7.8 ± 1.9 | <0.001 |
| Stimulation days | 10.0 ± 1.7 | 10.6 ± 1.5 | 0.09 |
| Total FSH dose (IU) | 1,627 ± 565 | 1,692 ± 488 | 0.54 |
| Peak E2 (pg/ml) | 1,527 ± 729 | 1,596 ± 774 | 0.84 |
| Peak P (ng/ml) | 0.6 ± 0.3 | 0.5 ± 0.2 | 0.52 |
| Cancellation rate | 6% (3/50) | 4% (2/50) | 0.99 |
| Retrieved oocytes | 9.6 ± 4.9 | 10.2 ± 6.0 | 0.61 |
| Fertilization rate (%) | 64.0 ± 19.4 | 61.3 ± 20.8 | 0.52 |
| Top-quality embryos | 3.0 ± 2.2 | 3.0 ± 2.4 | 0.93 |
| Transferred embryos | 1.6 ± 0.4 | 1.4 ± 0.6 | 0.07 |

Continuous data are expressed as mean ± SD.
Cédrin-Durnerin et al. showed a significantly higher consumption of gonadotropins in OCP-pretreated cycles as compared to either pretreatment with E₂ or no pretreatment, with no significant differences in treatment outcomes [23]. However, their study was mainly aimed at examining changes in the hormonal environment and antral follicles during the washout period after discontinuation of different methods of steroid pretreatment, and it included an even lower number of patients per treatment arm. Their data showed that steroid pretreatments differently affect the hormonal environment before the start of stimulation. According to their findings, it took 5 days after stopping OCP for FSH and LH to return to baseline levels from a strong suppression, suggesting this was an optimal washout period in cycles pretreated with OCP. In contrast, E₂ pretreatment did not reduce FSH levels significantly but rather avoided the increase of FSH during the luteal-follicular transition, and the rapid FSH rebound after stopping natural estrogen intake argues for a short, 1–2 day washout interval. Indeed, we administered E₂ until the day before the initiation of ovarian stimulation in order to optimize the synchronization of endogenous and exogenous FSH stimuli.

In a more recent study [19], the same group of researchers hypothesized that a 1-day washout period was too short to allow for complete recovery of baseline FSH levels, and that this was responsible for the increased gonadotropin consumption as compared to cycles without any pretreatment. However, no clinical studies have been performed with COH started 2 days after stopping E₂ administration.

Although the length of steroid exposure varied considerably both within each pretreatment group and between the two groups, previous studies have shown that the number of pretreatment days – at least within the range of days used in our study – has no impact on COH outcomes in cycles pretreated with either E₂ [24] or OCPs [25]. There is, however, considerable difference between OCPs in terms of their suppressive effect, depending on their progestin component [26]. A recent, small-scale, retrospective cohort study, performed in egg donors, suggested that OCPs containing androgenic (estrane- and gonane-derived) progestins lead to a more profound suppression of follicular development than those with anti-androgenic activity, resulting in lower oocyte yields [27]. The authors hypothesized that OCPs including an androgenic progestin would inhibit gonadotropin support for the growing follicle but maintain androgen exposure, which would lead to initial androgen-driven follicle growth but, in the end, to atresia of growing follicles because of

| Parameters                              | OCP (n = 50) | E₂ (n = 50) | Risk difference (%) | P value |
|-----------------------------------------|--------------|-------------|---------------------|---------|
| Implantation ratea                      | 43.5% (30/69) | 47.4% (27/57) | 3.9 (–13.4 to 21.1) | 0.79    |
| Pregnancy rate per cycle                | 56.0% (28/50) | 52.0% (26/50) | –4.0 (–23.1 to 15.4) | 0.84    |
| Clinical pregnancy rate per cycle       | 50.0% (25/50) | 48.0% (24/50) | –2.0 (–21.3 to 17.4) | 0.99    |
| Early clinical miscarriageb rate per pregnancy | 7.1% (2/28)   | 7.7% (2/26)   | 0.6 (–16.4 to 18.3)  | 0.66    |
| Ongoing pregnancy rate per cycle        | 46.0% (23/50) | 44.0% (22/50) | –2.0 (–21.2 to 17.3) | 0.99    |
| Live birth rate per cycle               | 42.0% (21/50) | 40.0% (20/50) | –2.0 (–21.0 to 17.1) | 0.99    |

*Implantation rate is the number of intrauterine sacs with heartbeat / number of transferred embryos.

bClinical pregnancies with miscarriage at <12 weeks of gestation.

CI confidence interval.
Moreover, extending E2 pretreatment beyond the menses cycle scheduling by reducing weekend pickups [18,24].

ment on follicular synchronization to the hypothetical beneficial effect of steroid pretreatment in women with decreased ovarian reserve, as these are results to a future meta-analysis on the subject. An interesting approach to other stimulation protocols.

A major limitation of our study is its sample size. In fact, with 50 patients in each arm of the study, only a difference of >26% could have been detected with 80% power, at a 0.05 significance level. However, as mentioned earlier, our aim was to contribute our clinical results to a future meta-analysis on the subject. An interesting future direction would be to extend our study to women with decreased ovarian reserve, as these are the patients in whom an increase in oocyte yield—due to the hypothetical beneficial effect of steroid pretreatment on follicular synchronization—could more easily be demonstrated. As far as the use of OCPs in poor responders is concerned, a modification to the GnRH antagonist protocol has been proposed by Orvieto et al. [31] The so-called “ultrashort GnRH agonist/GnRH antagonist protocol” is supposed to combine the beneficial effects of OCP pretreatment with that of the gonadotropin flare induced by the GnRH agonist, which is administered at the beginning of COH. The latter could circumvent the alleged detrimental effect of the OCP pretreatment on endogenous LH levels, which in turn could impair oocyte competence or endometrial receptivity. This protocol has been used with success in poor responder patients [32]. However, no prospective studies have compared so far the efficacy of this combined approach to other stimulation protocols.

Conclusions
In conclusion, we could not demonstrate differences in clinical results by using either OCP or E2 pretreatment for COH, if differences in washout periods after stopping pretreatment are accounted for; admitting, however, that the study was underpowered for the detection of small differences. Our results support those of earlier studies demonstrating that endogenous FSH suppression before starting ovarian stimulation is an efficient way to schedule ovarian stimulation in GnRH antagonist cycles. While observing comparable clinical outcomes, the use of E2 might offer several practical advantages compared to OCP: (1) Because E2 pretreatment is started in the midluteal phase, preparation for a programmed stimulation with a GnRH antagonist protocol can be scheduled in the same cycle even if the patient expresses a desire to start treatment after the early follicular phase. (2) Pretreatment is shorter with E2 than with OCPs (5–12 vs. 12–16 days, respectively, in our practice). (3) By using E2, GnRH antagonist cycles can be started in a scheduled manner even in patients who have objections to or present contraindications for taking OCPs even for a short period. Furthermore, by avoiding OCP pretreatment, we can give them one more chance to get pregnant spontaneously in the cycle preceding IVF.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
The study was designed by AB, AP and JAGV. Data were collected by AZ and CI, analyzed and interpreted by EEH and JAGV. The manuscript was drafted by EEH and revised critically by all other authors. The study was supervised by AP and JAGV. All authors read and approved the final manuscript.

Author details
1IVI Madrid, Avda del Talgo, 68-70, 28023 Madrid, Spain. 2IVI Valencia, Plaza de la Policía Local, 3, 46015 Valencia, Spain. 3Universidad de Valencia, Avda de Blasco Ibáñez, 13, 46010 Valencia, Spain. 4Rey Juan Carlos University, Avda del Talgo 68-70, 28023 Madrid, Spain.

Received: 3 August 2013 Accepted: 24 September 2013
Published: 28 September 2013
References

1. Devroye P, Aboughar M, Garcia-Velasco J, Griesinger G, Hurnaidan P, Kolibianakis EM, Ledger W, Tomás C, Fauser BC: Improving the patient’s experience of IVF/ICSI: a proposal for an ovarian stimulation protocol with GnRH antagonist co-treatment. Hum Reprod 2009, 24:764–774.

2. Alinyay HC, Youssef MAHM, Aboughar M, Broekmans F, Steinberg M, Smit J, Abou-Serri AM: Gonadotrophin-releasing hormone antagonists for assisted reproductive technology. Cochrane Database Syst Rev 2011, 5:CD001750.

3. Mortimer D, Mortimer ST: Quality and risk management in the IVF laboratory. Cambridge: Cambridge University Press; 2005.

4. González RM, Canales E, García R, Martín C, Rollán M, Fernández M, Prados N: Recuperación real de la temperatura y porcentaje de CO2 en los incubadores de fecundación in vitro. In Proceedings of the XIXth national congress of AEFT. Madrid: AEFT, 2008.

5. Janssens R, Souffreau R, Haentjens P, Van de Velde H, Verheyen G: New approach to follicular stimulation for in vitro fertilization: programmed oocyte retrieval. Fertil Steril 1986, 46:657–662.

6. Wardle PG, Foster PA, Mitchell JD, McLaughlin EA, Williams JAC, Corrigan E, Ray BD, McDermott A, Hull MG: Norethisterone treatment to control timing of IVF cycle. Hum Reprod 1986, 1:1453–457.

7. Zorn JR, Boyer P, Guichard A: Never on a Sunday: programming for IVF-ET and GIFT. Lancet 1987, 1:385–386.

8. Géreli S, Remohi J, Pizzocro P, Bontero C, Balmaceda JP, Silk JS, Asch RH: Programming of ovarian stimulation with norethisterone acetate in IVF/GIFT cycles. Hum Reprod 1989, 4:746–748.

9. de Ziegler D, Jääskeläinen AS, Brioschi PA, Fanchin R, Bulletti C: Synchronisation of endogenous and exogenous FSH stimuli in controlled ovarian hyperstimulation (COH). Hum Reprod 1988, 13:561–564.

10. Rombaud L, Neay D, Norman RJ: Comparative randomized trial to assess the impact of oral contraceptive pretreatment on follicular growth and hormone profiles in GnRH antagonist-treated patients. Hum Reprod 2006, 13:235–245.

11. Kolibianakis EM, Papanikolaou EG, Camus M, Tournaye H, Van Steirteghem A, Devroye P: Effect of oral contraceptive pill pretreatment on ongoing pregnancy rates in patients stimulated with GnRH antagonists and recombinant FSH for IVF. A randomized controlled trial. Hum Reprod 2006, 21:352–357.

12. Griesinger G, Kolibianakis EM, Venetsis CA, Diedrich K, Tzatzizis B: Oral contraceptive pretreatment significantly reduces ongoing pregnancy likelihood in gonadotropin-releasing hormone antagonist cycles: an updated meta-analysis. Fertil Steril 2010, 94:2382–2394.

13. Griesinger G, Venetsis CA, Marx T, Diedrich K, Tzatzizis BC, Kolibianakis EM: Oral contraceptive pill pretreatment in ovarian stimulation with GnRH antagonists for IVF: a systematic review and meta-analysis. Fertil Steril 2008, 90:1055–1063.

14. Garcia-Velasco JA, Bermejo A, Ruiz F, Martínez Salazar J, Requena A, Pellicer A: Cycle scheduling with oral contraceptive pills in the GnRH antagonist protocol vs the long protocol: a randomized, controlled trial. Fertil Steril 2011, 96:900–903.

15. Le Nestroy E, Marroniu J, Lahlou N, Roger M, de Ziegler D, Bouchard P: Role of estradiol in the rise in follicle-stimulating hormone levels during the luteal-follicular transition. J Clin Endocrinol Metab 1993, 77:439–442.

16. Fanchin R, Salomon L, Castelo-Branco A, Olivennes F, Fauser B, Devroye P: Luteal estradiol pre-treatment coordinates follicular growth during controlled ovarian hyperstimulation with GnRH antagonists. Hum Reprod 2003, 18:2698–2703.

17. Blockeel C, Engels S, De Vos M, Haentjens P, Polyzos NP, Stoop D, Camus M, Devroye P: Oestradiol valerate pretreatment in GnRH-antagonist cycles: a randomized controlled trial. Reprod Biomed Online 2012, 24:272–280.

18. Cédrin-Durnerin I, Guivarch-Leveque A, Hugues JN: Pretreatment with estrogen does not affect IVF-ICSI cycle outcome compared with no pretreatment in GnRH antagonist protocol: a prospective randomized trial. Fertil Steril 2012, 97:1359–1364.

19. Ye H, Huang GN, Zeng PH, Pei L: IVF/ICSI outcomes between cycles with luteal estradiol (E2) pre-treatment before GnRH antagonist protocol and standard long agonist protocol: a prospective and randomized study. J Assist Reprod Genet 2009, 26:105–111.