Abstract. Background/Aim: Studies on the impact of intrauterine human Chorionic Gonadotropin (hCG) administration in order to improve the In Vitro Fertilization (IVF) outcome have yielded conflicting results. The aim of the present systematic review and meta-analysis is to investigate whether timing of intrauterine hCG administration prior to embryo transfer affects its efficiency. Materials and Methods: A systematic search of the literature on Pubmed/Medline, Embase and Cochrane databases was performed. Only Randomized Control Trials were included in this meta-analysis. Results: Live birth rates were not improved following hCG administration (RR=1.13, 95%CI=0.88-1.46, p=0.34) in the pooled results. Combined live birth and ongoing pregnancy rates were borderline statistically significant following hCG administration (RR=1.27, 95%CI=1.00-1.62, p=0.05). Following subgroup analysis regarding live birth and ongoing pregnancy rates, only the 5-12 minutes prior to the embryo transfer group reported a statistically significant improvement. Conclusion: Intrauterine infusion of hCG within an IVF-Intracytoplasmic Sperm Injection (ICSI) cycle improves outcome only when administered 5-12 min prior to embryo transfer.

The success rate of in vitro fertilization (IVF) has been reported to be at 29.1% on the first cycle when considered irrespectively of age and infertility etiology (1). Despite its 40 years of practice, numerous conundrums still remain to be delineated in an effort to identify a foolproof approach towards achieving a clinical pregnancy. Implantation of an embryo into the endometrial cavity depicts the product of a constructive, complex, molecular dialog between the embryo and the endometrium (2). Thus, a potential implantation failure may originate from the equal contribution of embryonic, endometrial and maternal parameters (3). Clinicians are often called to identify which parameter is the culprit of a failed implantation. In practice, it may often be unrealistic to reveal the true origin of the failure, which would enable the design of a future treatment accordingly. The process of implantation is still considered to be a “black box” in the IVF treatment, while years of efforts have focused on how to understand it and subsequently enhance implantation rates. The key-molecule that intervenes and principally controls the implantation procedure, in line with the subsequent pregnancy that it entails, is the human chorionic gonadotropin (hCG) (4).

HCG is an heterodimeric hormone, secreted by the blastocyst prior to implantation (5, 6). Its initial role appears to be the stimulation of progesterone production by the corpus luteum (7, 8). Thus, hCG’s role shifts towards the maintenance of the upcoming pregnancy, principally through endorsing uterine angiogenesis, assuring maternal tolerance of the semi-allograft embryo, assisting uterine enlargement, which goes hand in hand with fetal development (5).
In light of the fact that the main driver of research in the assisted reproduction field is to improve pregnancy rates, scientists have tackled that issue aiming to address questions such as: “How can we improve embryo culture conditions?”, or “How can we achieve optimal receptivity of the endometrium?” It was not long before a hypothesis was formed questioning whether hCG administration via intrauterine infusion prior to the embryo transfer (ET) procedure may offer a noteworthy solution to implantation failure, resulting in superior pregnancy outcomes (9).

The novel approach of intrauterine hCG infusion was first brought up by Mansour and colleagues. The study concluded that 500 IU of hCG in 1 mL of culture media mirrors the ideal concentration to be injected into the uterine cavity, about 7 minutes prior to ET, promising enhanced pregnancy rates (10). Hitherto, several prospective studies have advocated the benefit of hCG administration as a valid option for improvement of pregnancy rates during IVF/ICSI cycles (6, 11-13). In contrast, other studies have reported no positive effect of this approach with respect to pregnancy rates, and particularly for cases involving blastocyst ET (14, 15).

Further to the current discrepancies from reported results on efficiency, several studies have conducted intrauterine hCG administration using different protocols of infusion. These differences refer to the optimal concentration, and time points of administration, along with the employment/ or not of sonography. The scenario of conflicting studies concerning the benefits and drawbacks of novel approaches in IVF practice is well argued and documented in literature. Three meta-analyses have been conducted on the effect of intrauterine administration of hCG on the day of ET with conflicting results (16-18). The hitherto heterogeneous cohort of results, constitutes a dilemma for clinicians, and reflects the unmet need for a universal standardized protocol regarding the application of this innovative approach. It is this need, that renders this new meta-analysis timely and essential towards reaching a consensus regarding the true place of hCG administration. Other contributions have been submitted on the matter of hCG administration, with a recent contribution assessing hCG administration efficiency from the point of view of the developmental stage of the embryos transferred (16). It is noteworthy, that the current study brings a different “angle” to the investigation, enriching the point of view regarding the optimal timing of hCG administration.

This metanalysis aims to address the need for designing a specific protocol on hCG administration. Besides, the ultimate

### Table I. Search strategy employed in all databases.

|   |   |
|---|---|
| 1 | IVF |
| 2 | In vitro fertilization |
| 3 | ICSI |
| 4 | Intracytoplasmic sperm injection |
| 5 | Assisted reproduction |
| 6 | ART |
| 7 | MAR |
| 8 | #1 OR #2 OR #3 OR #4 OR #5 OR #6 OR #7 |
| 9 | Cleavage stage embryo |
| 10 | Morula |
| 11 | Blastocyst |
| 12 | Day 2 |
| 13 | Day 3 |
| 14 | Day 4 |
| 15 | Day 5 |
| 16 | Day 6 |
| 17 | #9 OR #10 OR #11 OR #12 OR #13 OR #14 OR #15 OR #16 |
| 18 | Embryo transfer |
| 19 | #17 AND #18 |
| 20 | Human chorionic gonadotropin |
| 21 | hCG |
| 22 | #20 OR #21 |
| 23 | Intrauterine |
| 24 | Infusion |
| 25 | Administration |
| 26 | #24 OR #25 |
| 27 | #23 AND #26 |
| 28 | #22 AND #27 |
| 29 | #8 AND #19 AND #28 |

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Table II. Characteristics for studies included in this meta-analysis.

| Study                        | Number of participants | Intervention group Volume of Intrauterine infusion of culture media with hCG | Control group | Dosage (IU) | Time point Prior to ET |
|------------------------------|------------------------|--------------------------------------------------------------------------------|---------------|-------------|------------------------|
| Aaleyasin et al., 2015       | 483                    | 50 µl                                                                           | Culture mediaa | 500         | 5-7 min                |
| Cambiaghil et al., 2013      | 44                     | Volume not specified                                                            | No placebo    | 500         | 6 h                    |
| Eskandar et al., 2016        | 240                    | Volume not specified                                                            | No placebo    | 500         | 10 min                 |
| Firouzabadi et al., 2016     | 159                    | 0.04 µl                                                                         | No placebo    | 500 or 1000 | 7 min                  |
| Hafezi et al., 2018          | 180                    | 40 µl                                                                           | Culture media & no placebo | 500       | 7-10 min               |
| Hong et al., 2014            | 300                    | 20 µl                                                                           | Culture mediaa | 500         | <3 min                 |
| Hosseini et al., 2016        | 100                    | 40 µl                                                                           | No placebo    | 500         | 7 min                  |
| Huang et al., 2016           | 161                    | 1 ml                                                                            | Saline & no placebo | 1000   | 3 days                 |
| Kokkali et al., 2014b        | 194                    | Volume not specified                                                            | Culture mediaa | 500         | 4 min                  |
| Leao et al., 2013            | 36                     | Volume not specified                                                            | No placebo    | 500         | 6 h                    |
| Mansour et al., 2011         | 212                    | 40 µl                                                                           | Culture mediaa | 500         | 7 min                  |
| Mostajeran et al., 2017      | 100                    | 5 ml                                                                            | No placebo    | 700         | 5-10 min               |
| Navali et al., 2014          | 138                    | 0.5 ml                                                                          | Saline        | 500         | 3 days                 |
| Santibañez et al., 2014      | 210                    | 20 µl                                                                           | Culture mediaa | 500         | 4 min                  |
| Singh et al., 2014           | 216                    | 40 µl                                                                           | Culture mediaa | 500         | 5 min                  |
| Wirleitner et al., 2015c      | 1186                   | 40 µl                                                                           | Culture mediaa | 500         | 3 min or 2 days        |
| Zarei et al., 2014           | 182                    | Volume Not specified                                                           | Unidentified volume of saline | 6500b   | 12 min                 |

*aVolume of the culture media infused for the control group was equal to the volume infused for the intervention group. bBoth published and unpublished data are included following communication. cWirleitner study included two arms. In the first arm the administration was performed 3 minutes prior to ET, whereas in the other arm hCG administration was performed 2 days prior to ET. dThe dosage corresponds to recombinant hCG.*

goal of a metanalysis is to conclude on an optimal practice and assist clinicians in decision making regarding the potential inclusion of a novel approach as a Standard Operation Procedure (SOP). The purpose of this study is to produce a systematic review and a meta-analysis that delineates and clarifies -for the first time- which is the optimal timing of hCG intrauterine infusion in order to adequately cement its application in clinical practice and efficiently assist the clinicians towards deciding on optimal practice.

**Materials and Methods**

**Search strategy.** A systematic search of the literature was performed in Pubmed/Medline, Embase and Cochrane Central databases on July 2018 (Table I). The keywords employed and combined for the search strategy were: “In Vitro Fertilization”, “IVF”, “Assisted Reproduction”, “Assisted Reproduction Techniques”, “Medical Assisted Reproduction”, “Intracytoplasmic Sperm Injection”, “ICSII”, “human chorionic gonadotropin”, “hCG”, “Intrauterine Infusion”, “Randomized control trial”, “prospective study”. The original search yielded 1054 studies from the three databases. Following the removal of duplicate studies (n=184), all records were screened and full-text was sought and obtained for relevant articles. Relevant articles (n=47), were identified following title and abstract screening, employing the flow chart of Preferred Reporting Items for Systematic Reviews and Meta-analysis (PRISMA) as presented in Figure 1. A total of 17 randomized controlled trials (RCTs) (6, 10-15, 19-28) were employed in the present meta-analysis. Screening and selection of literature was performed independently by three authors. Citation mining was performed where the reference lists of all included articles and relevant reviews and metanalyses were reviewed to identify other articles of relevance. The search was limited to full-length manuscripts published in English in peer-reviewed journals up to July 2018. No protocol was submitted to the Prospero International Prospective Register of Systematic Reviews, providing details on conducting this study.

**Excluded studies.** The first arm of the study up to citation (10) was excluded as the low dosage levels were considered ineffective by the authors. The studies by Ye et al., 2015, Osman et al., 2016, Craciunas et al., 2016 (17, 18, 29) were not included in the quantitative synthesis as they were meta-analyses. The study by Janati et al., 2014 (30) was excluded as it was identical to (14), with the latter including more outcomes. The update of the study by Craciunas et al., 2016 was published in October 2018 (16), following the completion of the literature search performed for this metaanalysis which was concluded in July 2018.

**Study selection.** Only RCTs were selected for the present study. The population included women undergoing IVF, while the intervention was defined as the administration of hCG prior to the ET procedure. The primary outcome measure was the live birth rate (LB). The secondary outcome measures were: i) biochemical pregnancy (BP), ii) clinical pregnancy (CP), iii) live birth plus ongoing pregnancy (LB+OP), and iv) miscarriage rate. Ongoing pregnancy was coupled to live birth in a single measure, as performed in other meta-analyses (31, 32). It has been voiced that when performing RCTs on interventions regarding fertility/infertility, ongoing pregnancy is considered to be the desired outcome (33). Nonetheless, this point of view raises considerable controversy (34).
Bias assessment. Assessment of bias was performed on the selected studies regarding: i) selection bias (randomization), ii) allocation concealment, iii) selective reporting, iv) blinding of patients and personnel, v) blinding of outcome assessment, vi) incomplete outcome data and vii) other possible sources of bias. The bias assessment was performed independently by three authors.

Statistical analysis. Data extraction was performed by three authors. Risk Ratio (RR) with 95% Confidence Intervals (95%CI) was employed for the analyses of the included studies. Either the fixed-effects or the random effects model was employed for pooling the results according to heterogeneity. Heterogeneity of the exposure effect was evaluated employing the I^2 statistic. If the I^2 value was 80% or greater, the meta-analysis was not performed due to high heterogeneity. If the I^2 value was 60% or greater, indicating significant heterogeneity, the random effects model was employed. A chi-squared test for heterogeneity was also performed and the p-values were presented. Funnel plots for a possible publication bias were performed. Sensitivity analysis was performed for the studies where culture media were administered as placebo in control groups. All statistical analyses were performed using the Review Manager (RevMan) software (built 5.3).

Results

Study characteristics. The evaluated characteristics for each study included in the present meta-analysis are presented in Table II. Characteristics include: i) number of participants, ii) intervention protocol, iii) dosage, and iv) time point of administration. In Table I, the “Intervention” column refers to the volume of intrauterine infusion of culture media enriched with hCG, while the “Control” column refers to the volume of the placebo (plain culture media or saline). In cases where the study proceeded straight to ET serving as a control group, this is indicated as “no placebo”. Only one study employed recombinant hCG (28). Assessment of bias was performed and the results are presented in Figures 2 and 3. With regards to the subgroup analysis, all the studies that have described the timing of the intervention as “straight before the ET”, report performing the infusion in less than 5 minutes prior to the ET procedure. Based on the above, the possible time frames for infusion for subgroup analysis were the following: i) “less than 5 minutes prior to ET” describing the “straight before the ET” group, ii) “5 to 12 minutes prior to ET” describing the “minutes before the ET” group, where patients were asked to wait for a few minutes between the intervention and ET, iii) “hours prior to ET” and iv) “days prior to ET” describing the “hours and days prior to the ET” groups, respectively. There are no studies that report hCG administration between 12 minutes and 6 hours prior to ET.

Live birth rates. Six studies have reported results on LB rates (10, 15, 19, 22, 24, 27). Heterogeneity was statistically significant (I^2=79%, p<0.0001), thus the random effects model was employed. Pooled results failed to reveal any statistically significant difference (RR=1.11, 95%CI=0.85-1.45, p=0.34). Performing a subgroup analysis for the group of studies in which hCG administration was performed <5 minutes prior to ET no statistically significant difference was observed (RR=0.81, 95%CI=0.57-1.15, p=0.24). Live birth rate was significantly higher following hCG administration 5 to 12 minutes prior to ET (RR=1.42, CI=1.10-1.84, p=0.007). Only one study has reported on live birth rates following hCG administration days prior to ET (15) and no study has reported on live birth rates following hCG administration hours prior to ET (Figure 4).
Live birth/ongoing pregnancy rates. Six studies have reported results on LB rates (10, 15, 19, 22, 24, 27), and 3 studies have reported on ongoing pregnancy rates (12,26,28). Heterogeneity was statistically significant ($I^2=79\%$, $p<0.0001$), thus the random effects model was employed. Pooled results revealed a trend that marginally did not reach statistical significance (RR=1.27, 95% CI=1.00-1.62, $p=0.05$), favoring hCG administration. Subgroup analysis did not reveal any statistically significant difference neither in the <5 minutes prior to ET group (RR=0.81,95% CI=0.57-1.15, $p=0.24$) nor in the days prior to ET group (RR=1.52, CI=0.87-2.66, $p=0.14$). The addition of the outcome of ongoing pregnancy strengthened the beneficial effect of hCG when administered 5-12 minutes prior to hCG (RR=1.47, CI=1.19-1.83, $p=0.0004$) (Figure 5).

Clinical pregnancy rates. Seventeen studies (6, 10-15, 19-28) have reported results on clinical pregnancy rates. Heterogeneity was statistically significant ($I^2=65\%$, $p<0.0001$), thus the random effects model was employed. Pooled results revealed a statistically significant difference (RR=1.27, 95% CI=1.11-1.44, $p=0.0005$), favoring hCG administration. Subgroup analysis revealed a statistically significant difference only in the 5-12 minutes prior to ET group (RR=1.33, CI=1.12-1.57, $p=0.001$). Statistically significant differences were not reported on neither of the other three groups, namely i) the <5 minutes prior to ET group (RR=1.11, 95% CI=0.92-1.35, $p=0.28$), ii) the hours prior to ET group (RR=1.12, 95% CI=0.76-1.65, $p=0.56$) and iii) the days prior to ET group (RR=1.52, CI=0.86-2.69, $p=0.15$). It should be mentioned that heterogeneity in the days prior to ET group was very high, rendering the results of this particular subgroup lacking robustness ($I^2=83\%$) (Figure 6).

Biochemical pregnancy rates. Eleven studies (6, 10, 14, 15, 19, 20, 22-24, 26, 28) have reported results on biochemical pregnancy rates. Heterogeneity was statistically significant ($I^2=63\%$, $p<0.002$), thus the random effects model was employed. Pooled results revealed statistically a significant difference (RR=1.20, 95% CI=1.07-1.36, $p=0.003$), favoring hCG administration. Subgroup analysis revealed a statistically significant difference with respect to the 5-12 minutes prior to ET group (RR=1.21, CI=1.06-1.38, $p=0.006$). Statistically significant differences were not revealed neither in the <5 minutes prior to ET group (RR=1.16, 95% CI=0.92-1.47, $p=0.22$), nor in the days prior to ET group (RR=1.34, 95% CI=0.70-2.56, $p=0.15$). It should be mentioned that heterogeneity in the days prior to ET group was significantly high, resulting in obscure results ($I^2=85\%$). Only one study reported results in the hours prior to ET group thus analysis could not be performed (Figure 7).

Miscarriage rates. Ten studies (10, 12, 14, 15, 19, 22, 24, 26-28) have reported results on miscarriage rates. Heterogeneity was statistically insignificant ($I^2=0\%$, $p=0.82$) thus the fixed effects model was employed. Pooled results did not reveal a statistically significant difference (RR=1.10, 95% CI=0.83-1.45, $p=0.5$). Subgroup analysis also did not reveal any statistically significant difference neither in the 5-
12 minutes prior to ET group (RR=1.09, CI=0.80-1.48, p=0.59) nor in the days prior to ET group (RR=0.90, CI=0.45-1.80, p=0.77). The <5 minutes prior to ET group also did not present with a statistically significant difference (RR=1.20, 95%CI=0.78-1.83, p=0.41), even though the heterogeneity was significantly high (I²=86%) (Figure 8).

Sensitivity analysis. In the sensitivity analysis, only studies including culture media as placebo were employed (11, 12, 15, 19, 22, 27). Regarding the live birth rate outcome measure, excluding studies not employing culture media as placebo did not affect our results in a statistically significant fashion. Nonetheless, heterogeneity remained statistically very high (RR=1.22, 95%CI=0.94-1.60, p=0.18, five studies, I²=72%, n=2116). Subgroup analysis revealed a statistically significant beneficial effect of hCG when administered 5-12 minutes prior to ET (RR=1.64, 95%CI=1.31-2.05, p<0.0001, three studies, I²=0%, n=819). It should be emphasized that all three studies are reporting on live birth rates. No statistically significant difference was observed in the days prior to ET group (RR=1.05, 95%CI=0.79-1.38, two studies, I²=0%, n=293). Regarding the miscarriage rates outcome measure, the sensitivity analysis was performed in the same manner and did not affect our results in a statistically significant fashion (RR=0.88, CI=0.62-1.24, p=0.60, five studies, I²=0%, n=2116).

Discussion

In the last four decades, respectful efforts focus on improving IVF success rates via launching novel trends (35). These range...
from techniques employed to support the IVF laboratory performance, namely metabolomics for optimal embryo assessment and selection (36), to approaches regarding protocols in clinical practice. Emphasizing on improving implantation rates, the scientific community has brought to light a wide variety of innovative options in order to enhance the endometrial receptivity and contribute towards a successful implantation procedure. Studies have reported that: i) endometrial injury or scratching (37), ii) the employment of stem cells (38, 39), iii) intrauterine platelet-rich plasma (PRP) infusion (40, 41), and iv) intrauterine hCG infusion prior to ET (17) could result in improving endometrial receptivity and implantation rates. The aforementioned options are highly promising, though still may be considered as ground-breaking and controversial, meriting further RCTs to cement their true role in optimal practice. Various perils could be associated with the employment of highly promising novel approaches. Hence, even if studies support the safety and effectiveness of clinical application, still the leap to adoption in clinical practice should be preceded by exhaustive trials. An example of this is the widely and routinely employed technique of Intracytoplasmic Sperm Injection (ICSI) that is still believed, by a part of the scientific community, to be the cause of epigenetic alterations (42). Research and development should involve studies on animal models, followed by in vitro tests on embryos that are donated for research, advancing to RCTs (43).

The scientific community has a responsibility towards accepting novel trends that leave conflicting impressions and are not part of established guidelines in clinical practice. The ideal scenario would be for the highly-esteemed Societies of Reproductive Medicine, to evaluate alternative trends and report on specific guidelines, regarding the clinical application of IVF. However, the accurate establishment of guidelines, accompanied by universally standardized
protocols, is anticipated to require a considerable time. The final step prior to the admission of novel approaches in the clinical routine setting should be the completion of meta-

analyses with favorable and cohesive results.

The process of implantation still remains as a conundrum in the physiology of reproduction. In the context of assisted reproduction, intrauterine infusion of hCG emerges as one of the promising trends in promoting successful implantation fueling this metaanalysis. Different mechanisms have been described throughout the literature, regarding the underlying basis of hCG’s fundamental function to adjust the process of the embryo’s implantation. In 1998, results on delineating unknown mechanisms revealed the paracrine function of hCG influencing various growth factors and cytokines of paramount importance for the implantation window, such as the intrauterine insulin-like growth factor binding protein 1 (IGFBP-1), the macrophage colony-stimulating factor (M-MSCF), the leukemia inhibitory factor (LIF), the vascular endothelial growth factor (VEGF) and the matrix metalloproteinase 9 (MMP-9) (44). Other studies have indicated that hCG could stimulate the production of the transforming growth factor beta (TGF-β) or the receptor of interleukin-1. The growth factors and the cytokines secreted as a response to hCG are required in order to constitute the endometrium receptive towards the embryo (45). Last but not least, hCG has been documented to enable attraction of immune cells, namely neutrophils, monocytes and lymphocytes in vitro (4, 6, 46).

The primary aim of this systematic review and meta-

analysis study is to highlight the contradictions on the matter of hCG intrauterine infusion, and concur on its employment and effectiveness in clinical IVF routine. Our primary results indicate that hCG administration provides beneficial results regarding live birth rates only when administered 5-12 minutes prior to ET. As a secondary outcome, our results indicate that biochemical and clinical pregnancy rates, along with ongoing pregnancy and live birth rates, are also improved following hCG administration 5-12 minutes prior to ET. With respect to intrauterine hCG infusion and the

Figure 6. Risk ratio and Forest plot regarding clinical pregnancy rates.
possibility of associating it with miscarriage rates, no statistically significant difference was identified neither in the pooled results nor in any subgroup, extrapolating safety of the practice in that perspective. HCG administration performed during any other time frame did not convey any statistically significant improvement on live birth rates. When combining live birth rates with ongoing pregnancy rates no statistically significant difference was observed neither when hCG administration was performed in less than 5 minutes, or in the very opposite, days prior to ET. Biochemical and clinical pregnancy rates were not improved at any other point except during the 5-12-minute time frame prior to ET. With regards to limitations of our analysis, heterogeneity was very high regarding the clinical pregnancy rates in the 2-3 days prior to ET group, thus rendering the results of this subgroup less robust. Nonetheless, no statistically significant effect was observed in this subgroup regarding the ongoing pregnancy/live birth rates.

HCG is widely considered as the pregnancy hormone. The mRNA of hCG is firstly produced at the cleavage developmental stage of the embryo, and the mature mRNA at the blastocyst stage (47, 48). The cross-talk between the embryo and the endometrium may require the constant presence of hCG along with the respective effect it exerts, as it is proposed by a recent review (49). Therefore, it may be a safe hypothesis that the presence of hCG 5-12 minutes prior to the ET procedure may provide a beneficial effect on implantation potential contributing towards a successful pregnancy sustenance that results in a live birth.

The time frame 5-12 minutes prior to the ET appears to be optimal. Nonetheless, with regards to limitations, no study has reported hCG administration in the time frame defined as 13 minutes to <6 hours prior to ET, or in the time frame defined as ≥6 hours to <2 days. With regards to hCG administration at 6 hours and onwards, our results indicate that it may be ineffective. This may be attributed to the fact

| Study or Subgroup | Experimental | Control | Risk Ratio | Risk Ratio |
|-------------------|--------------|---------|------------|------------|
|                  | Events | Total | Events | Total | Weight | M-H, Random, 95% CI | M-H, Random, 95% CI |
| 1.4.1 4.5 minutes prior to ET | Kokkali et al., 2014 | 68 | 97 | 56 | 97 | 10.9% | 1.21 [0.68, 1.50] |
|                  | Saridakezi et al., 2014 | 53 | 101 | 39 | 109 | 8.3% | 1.47 [1.07, 2.00] |
|                  | Wirtzinger et al., 2015 | 264 | 510 | 261 | 494 | 13.6% | 0.97 [0.66, 1.09] |
|                  | Subtotal (95% CI) | 708 | 700 | 32.7% | 1.16 [0.92, 1.47] |
| Total events | 36 | 382 | 36 | 380 | 24.7% |
| Heterogeneity: Tau² = 0.03; CH² = 7.93, df = 2 (P = 0.22); P = 75% Test for overall effect: Z = 1.24 (P = 0.22) |

| 1.4.2 5-12 minutes prior to ET | Alesyusin et al., 2015 | 131 | 240 | 87 | 243 | 11.2% | 1.52 [1.24, 1.87] |
|                  | Firouzbabadi et al., 2016 | 35 | 108 | 18 | 53 | 5.3% | 0.97 [0.61, 1.54] |
|                  | Hafezi et al., 2018 | 21 | 60 | 46 | 120 | 6.2% | 0.91 [0.60, 1.38] |
|                  | Hosseini et al., 2016 | 14 | 50 | 10 | 50 | 2.8% | 1.40 [0.68, 2.85] |
|                  | Mansour et al., 2011 | 84 | 107 | 65 | 105 | 11.9% | 1.27 [1.06, 1.53] |
|                  | Zarei et al., 2014 | 31 | 54 | 22 | 96 | 5.4% | 1.64 [1.04, 2.61] |
|                  | Subtotal (95% CI) | 647 | 669 | 42.8% | 1.29 [1.09, 1.53] |
| Total events | 316 | 248 | 248 |
| Heterogeneity: Tau² = 0.01; CH² = 7.14, df = 5 (P = 0.17); P = 35% Test for overall effect: Z = 2.97 (P = 0.003) |

| 1.4.3 Hours prior to ET | Cambiagli et al., 2013 | 19 | 22 | 18 | 22 | 9.7% | 1.06 [0.82, 1.37] |
|                  | Subtotal (95% CI) | 22 | 22 | 9.7% | 1.08 [0.82, 1.37] |
| Total events | 19 | 18 | 18 |
| Heterogeneity: Not applicable Test for overall effect: Z = 0.41 (P = 0.68) |

| 1.4.4 Days prior to ET | Navali et al., 2016 | 42 | 71 | 21 | 67 | 6.3% | 1.69 [1.26, 2.23] |
|                  | Wirtzinger et al., 2015 | 42 | 69 | 45 | 63 | 8.5% | 0.86 [0.62, 1.02] |
|                  | Subtotal (95% CI) | 168 | 269 | 14.8% | 1.34 [0.79, 2.26] |
| Total events | 84 | 66 | 66 |
| Heterogeneity: Tau² = 0.19; CH² = 8.62, df = 1 (P = 0.01); P = 85% Test for overall effect: Z = 0.88 (P = 0.36) |
| Total (95% CI) | 1537 | 1551 | 100.0% | 1.22 [1.07, 1.39] |
| Total events | 801 | 688 | 688 |
| Heterogeneity: Tau² = 0.03; CH² = 30.8, df = 11 (P = 0.001); P = 64% Test for overall effect: Z = 2.89 (P = 0.004) Test for subgroup differences: CH² = 1.84, df = 3 (P = 0.61), P = 0% |

Figure 7. Risk ratio and forest plot regarding biochemical pregnancy rates.
that the half-life of exogenous hCG is 5-6 hours during the rapid phase (50). Further to that, the dosage of 500 IU that was administered in this study may not be efficient to enhance clinical pregnancy and live-birth rates. On the other hand, trying to understand why hCG administration in less than 5 minutes prior to ET appears to be ineffective, one may hypothesize that the volume of fluid introduced in the uterine cavity so close to the ET procedure may contribute to increased volume following ET, and subsequent embryo “floating”. Interestingly, embryo “floating” has been associated with a lower implantation potential (51). Moreover, the procedure of introducing a catheter prior to ET, similarly to the practice of mock ETs, may harbor perils depending on standards of practice. Previous studies have shown that low pregnancy rates may be associated with traumatic catheter introduction into the uterine cavity, as evident through direct visualization of endometrial lesions assessed immediately following catheter introduction (52). Thus, the possibility of causing minor trauma so close prior to the actual ET procedure may be involved in the lower rates corresponding to the hCG administration in less than 5 minutes prior to ET. Of course, such a hypothesis would stand independently as a confounder regarding any of the time frames. Nonetheless, no study has been conducted to assess the response of the endometrium with regards to the pathways activated during certain timeframes following minor injury.

The present meta-analysis may assist the clinicians in decision-making regarding the optimal time of hCG administration prior to ET. A seemingly simple time shift of hCG administration in the patients’ schedule with respect to the ET procedure, could potentially exert a considerable
effect on the clinical outcome and effectiveness of the IVF. Therefore, it is of pivotal importance to design and implement a specific ET SOP in cases where hCG infusion is included in the protocol. This may be of particular importance, especially in a highly demanding IVF laboratory setting. In addition to that, the results of the present meta-analysis could be considered in line with the recent meta-analysis of Craciunas and colleagues (16). It may be possible that the optimal practice includes hCG administration at minimum dosage of 500 IU, 5 to 12 minutes prior to the transfer of cleavage stage embryos. The current study highlights a significant improvement on ongoing pregnancy and live birth rates following intrauterine administration of hCG 5-12 minutes prior to ET. Further to that, the data sourced herein are reassuring with respect to miscarriage rates not being affected by this intervention.

Conflicts of Interest

The Authors declare that no conflicts of interest exist in regards to this study.

Authors’ Contributions

MS contributed to conceiving and designing the review, drafting and editing of the manuscript and assessment of bias; KS contributed to conceiving and designing the review; EM in assessment of bias, data extraction and statistical analysis; PT in study selection, assessment of bias and data extraction; PG in study selection and drafting; AP in study selection and literature search; SG in statistical analysis and drafting of the manuscript; PC in literature search and drafting; GA in data extraction and editing of the manuscript; KP in study selection and editing of the manuscript; MK in reviewing and editing the manuscript.

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