



Relationship between morphological grading of day five and day six embryos and clinical pregnancy rate

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Abstract

Objective (s): assessment of embryo grades on day five/six? And the potential of slow developing embryos

Design: prospective analysis of single embryo transfer (SET).

Setting: private IVF center *Giza, Egypt*.

Subject: women who had fresh (SET) on day five/six. From January 2016 to December 2017. Age 20 – 35 years old were included. Embryo scoring was done using Gardner grading system, besides, 5 grades were added. Only non-compacting embryos were excluded.

Intervention(s): chromosomal analysis of each grade may add more prediction.

Result(s): Analysis of cycles (n=326) revealed positive clinical pregnancy (42.3%). Analysis of difference resulted single embryo with expansion or hatching (EH) stages or 1, 5, or 6 were significantly associated with higher clinical pregnancy rates.

Conclusion: EH and TE are significantly correlated with clinical pregnancy, however, no specific grade should be excluded except for arrested embryos, as no golden grade. Prediction of pregnancy might be calculated according to our findings.

Keywords: embryo grading, pregnancy outcome, embryo quality, clinical pregnancy

1. Introduction

Embryo quality is the only parameter independently associated with ongoing pregnancy and mainly examined by microscope at certain times using morphological scoring systems, and it is still the best strategy for the selection of embryos [1-2]. Assessment of embryos using morphological and developmental criteria is the conventional method and inefficient indicator for pregnancy potential [3]. Placido and his coworkers postulated other factors to predict outcome as zygote quality, and in-vitro development because of embryo morphology alone isn't a predictive tool for IVF outcome [4]. On the other hand, both cytogenetic and molecular analysis could be performed, and the result may be used to score embryos. Since, chromosomal investigation of the embryo could provide embryologists with an additional marker for embryo quality which is partially independent of embryo morphology or using both morphological and chromosomal [5-7].

Relationship between embryo morphology and pregnancy occurrence after IVF was assured, from 1993 a study of morphological assessment of embryos was directly correlated with implantation rate [8]. Morphological assessment leads to reduce the potential of multiple pregnancies through selecting fewer embryos at embryo at ET [9].

Top-quality embryos could be chosen by measurement of the cleavage rate and degree of fragmentation [10]. Early cleavage has shown to correlate with embryo quality, with the like hood to reach to blastocyst stage [11].

For day five/six embryos, we have two choices to transfer embryos, firstly; active, which means morphological and chromosomal assessment secondly; indirect by culturing the

embryos as long as possible, in which unsuitable embryos will be arrested [5].

Many stages and grades of embryos were identified, dealing with this enormous numbers of forms make it difficult if not treated. There are many systems identified to make scoring easier. Embryo grading systems are useful in the prediction of embryo implantation [12].

Thus, the main aim of this study is to investigate the relationship between individual morphology parameters and clinical pregnancy outcome. In addition to adopt logistic regression model for prediction of the probability of which embryo could be transferred based on blastocyst morphology scoring.

2. Patients and Methods

2.1 Study Design

A prospective analysis, observational study of SET was conducted in a private IVF center *Giza, Egypt* during January 2016 to December 2017. Inclusion criteria were women who had fresh SET on day five/six, Age 20 – 35 years old. While Cases of well-known poor endometrial receptivity, uterine anomalies, and who had systemic conditions that may predispose to miscarriage were excluded. Regarding data documentation all demographic and initial characteristics of women included, type of the protocol used for ovarian stimulation, number of oocytes retrieved, number of embryos transferred, and their grading were documented daily during work. Women included were contacted directly in the center to record results.

2.2 Ovarian Stimulation

Hormonal profile including estradiol (E2), luteinizing

hormone (LH), prolactin (PRL), and follicular stimulating hormone (FSH) were monitored. Doses of initial gonadotropin used for ovarian stimulation is 75-300 IU/ml adjusted according to patient's age, Body Mass Index (BMI), previous response to stimulation and FSH concentrations on day 2 or 3 of menstruation. Patients underwent ovarian stimulation by rFSH (Fostimon; Serono Laboratories Inc, Norwell, Maine, USA) with gonadotrophin-releasing hormone (GnRH) agonist pituitary down-regulation by menotrophen (Merional, IBSA International) or (GnRH) antagonist by ectromelia acetate (Cetrotide, EMD Serono, USA), and trigger human chorionic gonadotrophin (HCG) (Choriomon, IBSA International). It was given when two or more follicles reach a diameter of ≥ 18 mm. Egg retrieval was done by transvaginal ultrasound 36 h after HCG administration. After ET patients were asked to administer 80 mg/day of progesterone (Prontogest, Marcyrl, Egypt) till clinical pregnancy.

2.3 Egg Retrieval and Semen Preparation

Oocytes were retrieved from the follicular fluid and poured in a Falcon dish (Falcon, USA) with 3 ml of HEPES-MOP'S buffer medium (Multipurpose Handling Medium, MHM® with Gentamicin, Irvin Scientific). Once all oocytes were retrieved, they were washed in 3 ml of same buffer. Semen was prepared discontinuous density gradients (Isolate, Irvin Scientific), and simple washing by (Sperm wash, Irvin Scientific).

2.4 ICSI and Embryo Culture

One hour after egg retrieval, oocytes were denuded by two alternative steps; Biochemical in which buffered Hyaluronidase Solution 80 IU (90101, Irvin Scientific) in 100 μ l for 30-45 seconds. Secondly; mechanical in which flexible pipettes used for denuding by gently multiple movements with appropriate diameter 170 μ l (Cook® Flexipet® Pipette, Cook® medical). Upon completion, oocytes were graded for maturity then placed into the incubator (37°C) for 1–2 h until ICSI. 38-39h post hCG ICSI was started, the first polar body either at the 6 o'clock or 12 o'clock position. A single spermatozoon was aspirated into the cytoplasm and then back into the needle, the spermatozoon and aspirated cytoplasm were expelled to the oocyte. The oocytes were incubated in single step culture media (Global® TOTAL, life global group) at 37°C in 6.9% CO₂, 5.0% O₂, 98% humidity, PH 7.24 to 7.28 in (Forma water jacket incubator, Thermo fisher scientific, USA).

2.5 Fertilization and Embryo Assessment

Oocytes were evaluated for pronucleus PN 16–18 hours after injection. Fertilization was considered normal when only two PN were seen, abnormal were excluded. Embryo assessment on day two post-insemination, the evaluation considered the number of blastomeres, degree of fragmentation, multi-nucleation, and symmetry of blastomeres. On day five, grading was conducted according to Gardner grading system/ Gardner – Schoolcraft grading system [13]. Fourteen days later, biochemical test for β -HCG level was measured. Seven weeks later female was asked for ultrasound to record clinical pregnancy.

2.6 Statistical Analysis

These results were statistically analyzed using Statistical

Package for Social Science (SPSS®) software (Statistical Package for Social Science, version 23, Illinois, USA). results were expressed as means \pm SD. Analysis of difference using Mann-Whitney's U-Test and Chi-Squared Test. Differences were statistically significant when $p < 0.05$.

3. Results

Age range 20 – 35 years with Mean \pm SD (28.57 \pm 4.08). The major of included indications was male factor (44.8%), while the minor indication was gender selection. Table (1) shows the characteristics of inclusion.

Table 1: initial characteristics

Age (years)	
Range	20 – 35
Mean \pm SD	28.57 \pm 4.08
Duration of Infertility (years)	
Range	2 – 21
Median (IQR)	7.95 \pm 3.86
Indication for ICSI	
Male Factor	146 (44.8%)
Tubal Factor	30 (9.3%)
Endometriosis	31 (9.6%)
Anovulation	26 (8.4%)
Unexplained Infertility	41 (12.6%)
Combined Factors	36 (11.1%)
Gender Selection	14 (4.2%)

SD standard deviation –IQR interquartile range - IVF in vitro fertilization - ICSI intracytoplasmic sperm injection Data presented as range, mean \pm SD; or number (percentage) Cycles stimulated using GnRH α Protocol represented (87%), and GnRHant Protocol were (12.8%). Wide range of number of retrieved oocytes (1-40) median 10 (6-14) as shown in table (2).

Table 2: ICSI Cycle Characteristic

Protocol of COH	
GnRH α Protocol	284 (87.2%)
GnRHant Protocol	42 (12.8%)
Source of Sperm Harvesting	
Fresh Ejaculate	262 (80.4%)
Frozen Ejaculate	15 (4.5%)
Fresh TESE	6 (2%)
Frozen TESE	30 (9.2%)
Fresh FNA	9 (2.7%)
Frozen FNA	4 (1.3%)
No. of Retrieved Oocytes	
Range	1 – 49
Median (IQR)	10 (6 – 14)

ICSI intracytoplasmic sperm injection – COH controlled ovarian hyperstimulation
 GnRH α gonadotropin releasing hormone agonist
 GnRHant gonadotropin releasing hormone antagonist
 TESE testicular sperm extraction – FNA fine needle aspiration
 Data presented as number (percentage); or range, median (IQR)
 Analysis of SET (n=326) revealed positive clinical pregnancy 138 (42.3%). Analysis of difference using Mann-Whitney's U-Test and Chi-Squared Test resulted that single embryo with expansion or hatching EH stages or 1, 5, or 6 were significantly associated with higher clinical pregnancy

rates. On the contrary, transfer of a single embryo with stages of morula (20%), compacting cells (30.9%), and cavitating morula (25%). There was no significant association between clinical pregnancy outcome and inner cell mass ICM (A) grade (63.6%) P=0.372 (NS), (B) grade (40%) P=0.134 (NS).

Otherwise, trophoectoderm TE grade (A) showed significantly positive clinical pregnancy (67.7%) P=0.027 (S). Table (3) shows the difference between women who had positive clinical pregnancy outcome and those who did not regarding EH Stage.

Table 3: Difference between Women who had Positive Clinical Pregnancy Outcome and Those who did Not among Women who had Single Embryo Transfer regarding EH Stage, ICM and TE Grades

Single Embryo Transfer (n=326)	Positive Clinical Pregnancy (n=138)	Negative Clinical Pregnancy (n=188)	RR (95% CI)	P
EH Stage				
1 (n=28)	18 (64.3%)	10 (35.7%)	1.6 (1.17 to 2.17)	0.014 (S)
2 (n=28)	16 (57.1%)	12 (42.9%)	1.4 (0.99 to 1.98)	0.097 (NS)
3 (n=16)	10 (62.5%)	6 (37.5%)	1.51 (1.01 to 2.26)	0.094 (NS)
4 (n=20)	6 (30%)	14 (70%)	0.7 (0.35 to 1.38)	0.249 (NS)
5 (n=40)	28 (70%)	12 (30%)	1.82 (1.42 to 2.34)	<0.001 (HS)
6 (n=4)	4 (100%)	0 (0%)	2.4 (2.11 to 2.73)	0.019 (S)
Contracted (n=6)	4 (66.7%)	2 (33.3%)	1.59 (0.89 to 2.84)	0.223 (NS)
Cavitating Morula (n=32)	8 (25%)	24 (75%)	0.57 (0.31 to 1.04)	0.037 (S)
Abnormal Cavity (n=2)	2 (100%)	0 (0%)	2.38 (2.1 to 2.71)	0.098 (NS)
Morula (n=40)	8 (20%)	32 (80%)	0.44 (0.23 to 0.83)	0.002 (S)
Compacted cells (n=110)	34 (30.9%)	76 (69.1%)	0.64 (0.47 to 0.88)	0.003 (S)
ICM Grade				
A (n=66)	42 (63.6%)	24 (36.4%)	1.27 (0.7 to 2.31)	0.372 (NS)
B (n=10)	4 (40%)	6 (60%)	0.62 (0.28 to 1.35)	0.134 (NS)
C (n=2)	2 (100%)	0 (0%)	1.65 (1.38 to 1.98)	0.257 (NS)
TE Grade				
A (n=62)	42 (67.7%)	20 (32.3%)	1.81 (0.94 to 3.48)	0.027 (S)
B (n=6)	2 (33.3%)	4 (66.7%)	0.52 (0.17 to 1.64)	0.139 (NS)
C (n=10)	4 (40%)	6 (60%)	0.62 (0.28 to 1.35)	0.134 (NS)

EH expansion or hatching – ICM inner cell mass – TE trophoectoderm

Data presented as number (percentage)

RR risk ratio and its 95% confidence interval

1 Analysis of Difference using Mann-Whitney’s U-Test

2 Analysis of Difference using Chi-Squared Test

S significant – HS highly significant – NS non-significant

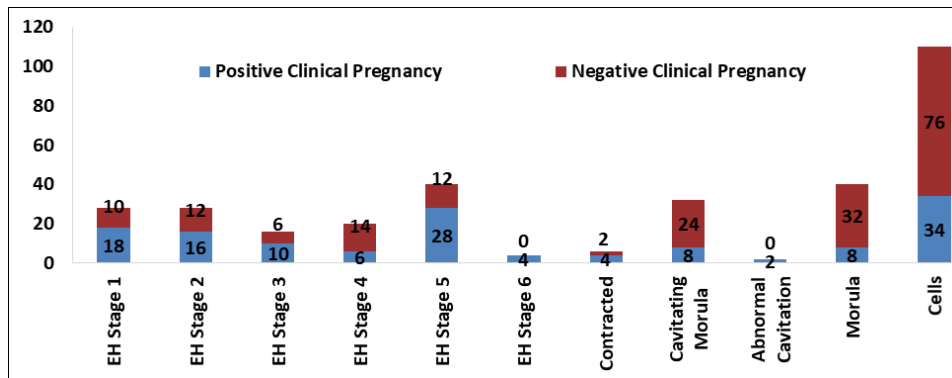


Fig 1: Bar-Chart showing Difference between Women who had Positive Clinical Pregnancy Outcome and Those who did not

4. Discussion

On day five, blastocyst has three independent factors which may affect outcome. EH, TE, and ICM were independently measured if any influences implantation. Assessment of shape and size of embryo may provide more insights to predict outcome. A lot of scoring methods have been developed. Gardner’s system is widely using in IVF laboratories. This system, which has been shown to provide higher implantation rates and better selection than other systems [14].

4.1 Blastocyst Stage

Previous results of day five embryos recorded high

implantation rate of expanded and Hatched blastocysts they achieved (64%) and (63%) respectively [15]. In 2013, a retrospective analysis on 1860 blastocysts, EH 5 showed the highest pregnancy rate among other grades (41.8%) [16]. But it was thawed embryos which may explain low pregnancy rate, however, the highest between all other EH grades. Other study recommended that the most potential parameter in blastocyst is EH stage, and EH 5 of SET showed (50%) clinical pregnancy [17].

On other hand, in 2012 a frozen embryo transfer FET research revealed lowest pregnancy of EH 5, then 6 among included grades [18]. And one more recent showed poor outcome of EH 5 also, it achieved (33.3%) compared to EH

4 (57.8%) [19]. In this study, it's noted that only 3 embryos were included in EH 5, moreover, probability of damage after thawing because of grading was done before vitrification.

Early blastocyst is the smallest EH between all blastocysts. It was more likely to achieve low pregnancy compared to EH 5. It was the lowest pregnancy [20], and [17]. Other studies reported higher results [15]. EH2 achieved higher in euploid embryos [19].

It's evident from current analysis and [15, 17, 21-22] that EH grade correlated with pregnancy rate. EH 5 is the best for transfer, EH 6 and 1 have high clinical pregnancy. On contrary, some papers postulated that EH grade didn't affect pregnancy [18, 19, 20]. In our findings, women who received SET of EH 5 and 6 were the highest but EH 6 was only 6 embryos.

TE morphology may be the most important parameter when selecting a single blastocyst for transfer, it is the strongest morphological predictor for outcome A grade was the highest then B and C [18, 15]. In contrary [19-21] didn't find relation. TE morphology may affect the rates of ongoing pregnancy. In our findings, TE with grade A was significantly the highest clinical pregnancy. This may be due to high numbers of cells and compaction of grade A which help in implantation.

For ICM, The higher ongoing pregnancy embryos those had A grade compared to C [18, 15]. But it wasn't significantly linked with pregnancy [15]. Among poor quality or average quality embryos, ICM morphology priority should be chosen with better grade because it is a better predictor of pregnancy outcomes than other blastocyst components [19]. The relation between ICM and predicting pregnancy has been linked [17, 21] also. On the other hand, studies found that no significant association between ICM grade and clinical pregnancy outcome [15, 18, 20]. In current study, ICM wasn't important parameter when selecting SET. It's found that A grade associated (not significantly) with clinical pregnancy.

4.2 Morula and Cavitating Morula (CM)

It is generally accepted that morula on day 4 is normal according to the healthy developing embryo [23]. Therefore, when single elective morula was transferred on day 4, it was an alternative to single elective day 5 blastocyst [24]. Low pregnancy (15.8% for morula, and 21.1% for CM) was recorded on day 5 [23]. Cavitating morula, morula and unexpanded early blastocyst in one group achieved 30% implantation rate [25].

According to our results, transfer of a single embryo Gardner not included stages; cavitating morula, morula, or compacting was significantly associated with lower clinical pregnancy rates. Clinical pregnancy of morula was only (20%).

In a simplified grading system morula and CM reported implantation, but live birth rate was higher in cavitating embryos [26]. But included cavitating and morula groups were limited as mentioned, morula or cavitating weren't significantly different from they poor blastocyst also. Similar result was reached, about 36% of morula implanted successfully, but higher miscarriage rate occurred [15]. No pregnancies were observed following transfers of 9 embryos non-cavitating on day five [27]. But only 3 transfers were considered. Clinical pregnancy of cavitating morula was slightly higher than morula of included embryos in the previous study. In comparison of transferring of fresh

embryos and culturing for one more day, higher blastulation rate of CM was recorded than morula, however, low pregnancy achieved after freezing/thawing [23].

4.3 Compacting Cells

Grading in many studies added compacting cells to morula or compacting morula [25-27] but it was separated here to assess their potential. Slow developing embryos might be by the action of chromosomal abnormality, it was stated that auto-correction of late developed embryos was done, the progression of normal cells exceeded from (12.5%) on day six to (47.8%) on day twelve [28].

In our study, 110 single embryos were transferred, 30% clinical pregnancy was achieved, and there were transferred early day five. Their potential was low, however, they should be considered after patients counseling about low outcome instead of excluding if they were the only available embryos. Non-compact; cell embryos on day 5 were excluded from current study. It was considered as arrested embryos [25].

5. Conclusion

No a specific grade should be excluded except for arrest embryos, as no golden grade, however, EH and TE were the most blastocyst variables may influence outcome. Otherwise, ICM grade was not significant. Relationship between morphological grading of day five and/or day six embryos and clinical pregnancy rate might be calculated according to previous finding. Chromosomal study of each different grade might add more prediction.

Conflicts of Interest: Authors declare that no conflict of interest regarding the publication of this article.

6. References

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