Raman spectrum: A potential biomarker for embryo assessment during in vitro fertilization

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Abstract. The aim of the study was to investigate whether Raman spectrum is consistent with the morphological scoring of the embryo of day 3 during in vitro fertilization (IVF). The spent culture media of embryo of day 3 from 10 patients were collected and analyzed. The samples were analyzed using Raman spectroscopy and graded according to the standard embryo scoring system simultaneously. Data showed that the Raman spectra obtained from the droplet of media were useful, as they can act as the characteristic signature for protein and amino acids. The Raman biospectroscopy-based metabonomics profiling of spent media was consistent with the result of conventional morphological evaluation. In conclusion, this technology offers great potential for the development of tools allowing rapid non-invasive assessment of the quality of embryo of day 3 during IVF.

Introduction

Infertility is a serious public health issue, estimated to affect 15% of the reproductive aged population. It has gained much attention worldwide. In vitro fertilization (IVF) has appeared as a powerful solution in the aspect of infertility treatment (1). The process of IVF involves monitoring and stimulating a woman's ovulatory process and further removing an ovum or ova (egg or eggs) from the woman's ovaries for IVF with sperm in a laboratory. The fertilized egg (zygote) is cultured for 2 to 6 days in a growth medium and then implanted in the woman's uterus with intention of establishing a successful pregnancy (2). Accordingly, morphological assessment was the primary method used in the fertility clinics to select \textit{in vitro} generated embryo(s) for the transfer into the uterus (3). However, it was widely known that the correlation between embryo morphology and implantation potential is also elusive (4). Indeed, embryo morphological grading methods mainly depends on the observer experience and the assessment criteria is rather subjective. As a milestone technique in reproductive medicine, the first time-lapse microscopy system for IVF was approved in 2009 for clinical use. Embryo morphokinetic scoring system has been shown to further improve the rates of pregnancy (5). The images were compiled using specialized software to create a time-lapse sequence of embryo development. Consequently, the study negated the need for the embryologist to remove the embryos from the incubator for morphological assessment. Additionally, widely using this technique also addresses some of the encountered obstacles presently due to its high cost. Therefore, in order to increase the rates of implantation, the improved assessment methodology was required. Accordingly, many different approaches have been suggested to reflect the embryo's function or physiological state. Several non-invasive methods assessing embryo viability and the potentiality of development have been established, such as proteomics (6), measurement of respiration rate (7), soluble human leukocyte antigen-G (8), pyruvate uptake (9) and glucose uptake (10). However, all of these methods have not been widely used in clinic because they were either expensive and required dedicated equipment, technical staff or do not provide results quickly enough to be used within the time frame of clinical IVF.

However, the Raman spectroscopy technique, used to observe vibration of bonds, rotational, and other low-frequency modes in a system, has been commonly used in chemistry to provide a fingerprint by which molecules can be identified (11) and serves as an are of interest. Indeed, over the past few years, Raman spectroscopy has become a powerful diagnostic tool in the life sciences (12). Currently, in the field of assisted reproduction technology, there are few studies using the Raman technique to investigate the correlation between the metabonomics profile and the embryo quality as well as the developmental potential and the outcome of pregnancy. Seli \textit{et al} reported that the non-invasive metabonomics profiling of embryo culture media using Raman and near infrared (NIR) spectroscopy were correlated with the pregnancy outcome in women undergoing IVF (13). They also carried out two studies demonstrating that metabonomics models developed using NIR or Raman may predict embryo viability (14,15). The same group conducted an analogous
drops of sequential culture media. Fertilized oocytes were individually cultured for 3 days in 25 µl pre-equilibrated medium (SAGE In-Vitro Fertilization, Inc., Trumbull, CT, USA) supplemented with 10% serum substitute supplement (Irvine Scientific, Santa Ana, CA, USA) to create the culture conditions. After 3 days of culture, the cumulus complexes were isolated, the majority of the cumulus cells were stripped mechanically and the oocytes were placed into individual 25 µl droplets of media (Quinn’s advantage cleavage medium). Patients who did not undergo embryo transfer were selected to our study. Oocyte retrieval was performed 36 h after the human chorionic gonadotropin injection. The patients’ treatment was performed according to the long gonadotropin-releasing hormone (GnRH) agonist protocol or a flexible GnRH antagonist protocol. All the patients enrolled in our study were treated at the Department of Reproductive Medicine of Nantong Maternity and Child Health Hospital. Each couple whose embryos were enrolled into the study signed a written informed consent. Ethical approval for the protocol was received from the Medical Ethics Committee of Nantong Maternity and Child Health Hospital, in Nantong, China. Informed consent was obtained from patients and ethics approval was granted by working in clean air hoods.

Materials and methods

Patients and ethics approval. All the patients enrolled in our study were treated at the Department of Reproductive Medicine of Nantong Maternity and Child Health Hospital, in Nantong, China. Ethical approval for the protocol was received from the Medical Ethics Committee of Nantong Maternity and Child Health Hospital. Each couple whose embryos were enrolled into the study signed a written informed consent.

Patient treatment. Ten patients who underwent ovarian stimulation using either a long gonadotropin-releasing hormone (GnRH) agonist protocol or a flexible GnRH antagonist protocol were selected to our study. Oocyte retrieval was performed 36 h after the human chorionic gonadotropin injection by transvaginal ultrasound-guided aspiration. The oocyte cumulus complexes were isolated, the majority of the cumulus cells were stripped mechanically and the oocytes were placed into individual 25 µl droplets of media (Quinn’s advantage cleavage medium). Patients who did not undergo embryo transfer were selected to our study. Oocyte retrieval was performed 36 h after the human chorionic gonadotropin injection. The patients’ treatment was performed according to the long gonadotropin-releasing hormone (GnRH) agonist protocol or a flexible GnRH antagonist protocol. All the patients enrolled in our study were treated at the Department of Reproductive Medicine of Nantong Maternity and Child Health Hospital. Each couple whose embryos were enrolled into the study signed a written informed consent. Ethical approval for the protocol was received from the Medical Ethics Committee of Nantong Maternity and Child Health Hospital, in Nantong, China. Informed consent was obtained from patients and ethics approval was granted by working in clean air hoods.

Materials and methods

Culture medium specimens. The embryo selection for implantation was based on embryo morphology at day 3. Following embryo transfer, the spent culture media were collected individually and stored at -80°C. The equilibrated culture media without embryos was also collected and used for normalization. During each experimental step, laboratory personnel wore gloves and coat, and the physical isolation was guaranteed by working in clean air hoods.

Raman spectroscopy. The spent culture media were thawed at room temperature (25°C) for 30 min before analysis. The mineral oil on the surface of the spent media was removed by capillary siphon until there was no visual stratification. After which the media were vortexed for 10 sec and centrifuged for 10 min at 11,000 x g. Next, 20 µl of embryo culture medium, taken from the cryovials by pipette gun, were dropped into a round glass with a thin layer of gold film on the surface, and the glass was placed onto a slide. Ten points were selected and Raman spectroscopy was measured for each sample. Raman analysis was conducted using a Raman spectrometer (BWTEK; B&W Tek, Inc. Newark, DE, USA). The spectra were recorded from 600 to 2,000 cm⁻¹. The signal acquisition time was 60 sec. Raman spectroscopy was collected at 10-15 points/sample. Laser Raman spectroscopy of the samples from 10 patients was conducted under the parameters above. The original spectra were preprocessed automatically by Origin 6.0 software.

Table I. Embryo assessment criteria.

| Classification | Grade | Cell no. | Fragmentation (%) | Symmetry | Multi-nucleation | Vacuoles | Zona pellucida |
|----------------|-------|----------|-------------------|----------|-----------------|----------|--------------|
| High quality   | A     | 4(d2) → 7-8(d3) | <10 | Even | No | No | Normal |
| 3-4            | B     | 4(d2) → 7-8(d3) | 11-25 | Even | No | No | Normal |
|                |       | 4(d2) → ≥9(d3) | 11-25 | Even | No | No | Normal |
| Poor quality   | C     | 2,4,6(d2) → >7(d3) | 26-35 | Uneven | No | Few | Abnormal |
| 4-5            |       | 6(d2) → >8(d3) | <35 | Uneven | No | Few | Abnormal |
|                |       | 2 or 4(d2) → 6(d3) | <35 | Uneven | No | Few | Abnormal |
|                |       | 3(d2) → >6(d3) | <35 | Uneven | No | Few | Abnormal |

Embryo assessment. A standard embryo scoring system was used for the evaluation of the quality of embryo based on the rate of cleavage and morphology (17). The morphological criteria considered include: i) The number of pronuclei and polar bodies (zygotes); ii) cell number; iii) evenness of mitotic divisions; iv) presence of micronuclei; and v) amount of cellular fragmentation (cleavage embryos). The embryo grade A and B were defined as high-quality embryo and embryo grade C was considered as poor quality embryo (Table I). The same embryologist preformed all embryology and embryo scoring in this study to ensure the consistency.

Results and Discussion

Although numerous studies have verified the feasibility of metabolomics profiling of spent media to select the high quality embryo, the validation from other laboratories is needed for the consistency of results on the diversity of the
medium, Raman instrument and protocol of analysis. In the present study, we analyzed spent culture media of embryos using Raman spectroscopy, trying to evaluate the embryo, and compare with the conventional morphological methods. The IVF spent culture media were thawed at room temperature for 30 min. The mineral oil layer on the surface, which act as the protecting shell to prevent moisture volatilizing were taken away with physical siphon until there was no visual stratification. Our preliminary results showed that there is no effect on spectral signals of embryo media for mineral oil. To verify the consistency of measurement, different grade embryos from the same couple were analyzed. Figs. 1 and 2 show the typical Raman spectrograms and microphotography of embryos of day 3 in different grade by morphological scoring, respectively. The peak area of at Raman shifting 755 cm\(^{-1}\) from the high quality embryo was obviously different to the poor one (Fig. 1). Raman shifting 750 cm\(^{-1}\) was the characteristic peak for tryptophan, an essential amino acid which cannot be synthesized by the organism and must be provided by the exterior nutrition (15). In the fresh media for embryo culture, tryptophan is 0.02 mM in general. More tryptophan was consumed by the high quality embryo for their development comparing to the poor embryo. Thus, the metabolism of a healthy embryo may alter the surrounding environment differently from one that is less healthy and thus possessing less reproductive potential. It may produce a profile that may more clearly be associated with embryo viability.

Amino acids have numerous roles in early embryo development in addition to protein synthesis. Preimplantation embryos can consume and produce amino acids in a manner dependent on the stage of development that may be predictive of subsequent viability. Tryptophan has a variety of metabolic functions within the cell. It is incorporated into the polypeptide chains of enzymes and proteins, and it is a biosynthetic precursor of the cofactor NAD, the siderophore quinolobactin, and the neuron transmitters serotonin and melatonin (18).

Numerous studies have analyzed the remaining media to determine novel assessing and screening methods to select the greatest potential embryo (16,19). Different from the former studies, the spent media from conventional IVF were analyzed with Raman spectroscopy in our research. Although to the best of our knowledge, there is no research or epidemiology study indicating that intracytoplasmic sperm injection could impair the embryo development and the pregnancy outcome to this day, the different protocol of fertilization may have led to the different metabolites. Thus, only the remaining conventional IVF media were collected and analyzed in this study.

There are several studies trying to understand the correlation between Raman spectrum of spent media and the outcome of pregnancy of a woman implanted with the embryo (20). It is clear that the quality of the embryo is not an independent factor contributing to the pregnancy outcome. There are other reasons which may impair the outcome of the embryo transfer, such as the endometrium situation, psychological state, or hormone level. Therefore, we only focused on the correlation between the Raman spectrograms and morphological grading. In the present study, we compared the metabolic profile with the morphological classification. However, we did not calculate the correlation index, because of the tiny number of samples in the study. In subsequent investigations, we intend to include a larger number of samples.

In conclusion, the profile of Raman spectrum of the spent embryo culture media was presented comparing the morphological assessment in our study. Without any doubt, an appropriate evaluation method can improve the rates of embryo implantation and pregnancy. To substantiate the results in our study, additional investigations of the correlation between the Raman spectrum and rate of pregnancy should be conducted in the near future.

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