Detection of Sperm Deformity by Kruger Strict Criteria using Pre-staining Slides Before and After Post Coital Test

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Abstract
Objective: There are many methods to examine the sperm morphology. One of the most important methods with high accuracy is Kruger strict criteria. The objective of this study is to: 1-simplified the method and lowering the time consumption for examination. 2-find out any changes may occurs on sperm morphology through post coital test.

Design: Prospective study
Settings: IVF Institute, Al-Nahrain University, Baghdad-Iraq.

Materials and Methods: Seventy four unselected couples were involved at the present work who attend the Infertility Clinic of the IVF Institute. Semen fluid analysis(SFA) was performed for all men as recommended by WHO(1999). According to the results of SFA, the samples were divided into normal and abnormal semen. Azoospermic, severe oligospermic, severe asthenospermic samples were excluded from this investigation. Special pre-stained slides(Testsimplet® slides, Warymart-Germany) were used for examination of sperm deformity. One drop of liquefied semen was mounted on warmed pre-stained slides and covered by standard cover slip. The slides were left for 5 minutes at room temperature and examined as described by Kruger strict criteria. When the results of men samples showed normal sperm morphology, Post coital test were done for their spouses with normal cervical mucus score. The results of sperm morphology obtained from PCT were recorded by using the Kruger criteria too.

Results: There was 28 out of 74 semen samples with normal sperm morphology. Fifty two samples revealed abnormal sperm morphology when examined by Kruger criteria and 36 samples were abnormal by using staining method alone. The percentage of morphologically normal sperm using Kruger criteria pre-coital test was 14.8% and following PCT the percentage was 13.4%. The statistical analysis showed a significant (P<0.05) differences between them.

Conclusion: It is concluded from the present study that some results of normal sperm morphology were changed in the cervix to abnormal sperm form by using Kruger criteria, even the cervical mucus showed normal score. This may interpret some unexplained infertility cases. At the same time, the study noticed that the examination by the stained slides and using Kruger criteria was so simple, easy to perform and the time required to prepare the samples is short to start reading the results.

Key words: Kruger strict criteria, testsimplet slides, male infertility, post -coital test

Introduction
Semen fluid analysis including sperm morphology remains the main pillar for male infertility work up. However, different methodologies for sperm morphology assessment have remained controversial because of the lack of universally acceptable method,(Said,2005). Meanwhile, there are many diagnostic laboratory techniques to evaluate semen quality, and utilize to interpretation infertility in both male and female, however explanation of infertility still obscure in about10-15 % of cases,(Harrison et al., 1997). Therefore, during the past decades, several methods of assessing sperm function have been introduced including the assessment of hyperactivated motility pattern of spermatozoa, the sperm zona pellucida (ZP) binding test, evaluation of the acrosome
status, zona free hamster egg penetration test and post coital test, and monitoring of reactive oxygen species generation (Liu and Baker, 1992). In vitro fertilization has provided a unique opportunity to determine how predictive these tests are of the fertilizing capacity of spermatozoa (Franken et al., 1990).

In 1986, Kruger and co-workers found that in a group of men with sperm densities greater than 20 million per milliliter and motility greater than 30% IVF rates were 37% for those with less than 14% normal sperm by strict criteria morphology (SCM) and 91% for those with greater than 14% normal sperm. Furthermore, Kruger noted that men with less than 4% normal SCM had an IVF rate of 7.6%, whereas those with 4% to 14% normal forms had a 63.9% fertilization rate (Kruger et al., 1986). Subsequent studies have confirmed the correlation between SCM and IVF rate (Ombelet et al., 1995). Thus, one of the most important methods to examine the sperm morphology with high accuracy is Kruger strict criteria. Therefore, the objective of this study is 1- simplified the method and lowering the time consumption for examination. 2- find out any changes may occurs on sperm morphology through post coital test.

Materials and Methods
A total number of seventy four unselected male patients who attended the infertility clinic of the Institute of Embryo Researches and Infertility Treatment - University of Al-Nahrain, Baghdad-Iraq, were included in this study during the period from November 2005 to October 2006. The mean age of these patients was 32 years and the majority of them (82.4%) had history of primary infertility. The duration of their infertility was ranging between 1.5-18 years with a mean of 4.6 years.

The standard semen analysis was performed according to the WHO, (1999) guidelines. Semen was collected in a clean, sterile, dry, plastic and warm disposable wide mouth dish, labeled with patient’s name and the time of collection (Silverberg and Turner, 2001). Semen sample was then immediately transferred to the laboratory and placed in an incubator at 37°C, waiting for liquefaction.

One drop (10µl) of liquefied thoroughly mixed semen was, mounted between warmed slides and covered with a standard cover slip. At least two hundred spermatozoa were counted and percent of normal morphology was reported (Acosta et al., 1986).

\[ \text{Morphologically Normal Sperm%} = \frac{\text{Number of normal sperms}}{\text{Total number of sperms}} \times 100 \]

1- Modified Kruger Strict Criteria (MKSC):
In this study it has been used especial pre-stained slides for examination of sperm morphology. The stain contains two dyes, 2.1µg/cm² crystal violet acetate and 1.0µg/cm² new methylene blue. One drop (10µl) of semen was mounted between warmed slides covered with standard cover slip and left for 5 minutes at room temperature then examined under light microscope using oil-emersion power 1000X and graduated optic. Ten fields or 100 sperms were examined by measuring the length and width of the head axis for each sperm in the slide by the graduated optic and labeled in a table. The normal head axis length is 5-6 µm and normal head axis width 2.5-3.5 µm as reported by Kruger, et al. (Kruger et al., 1987).
2. Post Coital Test (PCT):-

After evaluation of 74 unselected male patients who attended the infertility clinic in IVF Institute by routine seminal fluid analysis with examination of sperm morphology by MKSC; 20 couples out of the number of men had normal semen parameters including normal sperm morphology by MKSC were done PCT. The female partners of those patients were examined physically and by vaginal ultrasound. All of them had normal reproductive hormonal profile and tubal potency. The PCT was done according to WHO recommendations (WHO, 1999).

Post Coital Sperm Morphological Examination By Using MKSC:-

A drop of 10µl of collected cervical mucus (which was placed in a micro tube in an incubator) was placed on the especial pre-stained slides (mentioned previously), covered with standard cover slip and left for 5 minutes at room temperature then examined under light microscope using oil-emersion power 1000X and graduated optic. 100 sperms were examined by measuring the length and width of the head axis for each sperm in the slide. (Kruger et al., 1987).

Statistical analysis

The data was shown as mean ± SEM. The results were statistically analyzed using the Student's t-test and Chi-square depending on the nature of the data. When P value reaches the 0.05, the result was considered significant (Sorlie, 1995).

Results

There was a significant (P<0.05) different between all the semen criteria that examined of fertile and infertile men. The sperm concentration and the percentage of active sperm motility in fertile men were significantly (P<0.001) higher than that of infertile men. There was 28 out of 74 semen samples with normal sperm morphology. Fifty two samples revealed abnormal sperm morphology when examined by Kruger criteria and 36 samples were abnormal by using staining method alone. Using Kruger strict criteria, the percentage of MNS in fertile men was 15.5% ±0.14 and in the infertile men was 10.3% ± 0.25. A significant (P<0.05) different was found in the percentage of MNS between fertile and Infertile semen samples (Table-1) by using High power field method or Kruger criteria (Figure-1).

There was a significant (P<0.001) increase in sperm agglutination(%) between fertile and infertile semen. The number of round cells(cell/HPF) of infertile semen was significantly (P<0.001) higher than that of fertile semen (8.94 vs. 3.0, respectively) as shown in table-1.

In table-2, all the spouses (n=50) were shown normal cervical score (11.9 ± 0.176), normal endometrial thickness (8.76 ± 0.269) and follicle size of preovulatory period (19.53± 0.393).

The percentage of morphologically normal sperm of infertile semen using Kruger criteria pre-coital test was 14.8% and following PCT the percentage was 13.4%. The statistical analysis showed a significant (P<0.05) differences between them (Figure-2).
Table-1: Certain microscopic examination of the semen of fertile and infertile patients using two methods to measure the percentage of morphologically normal sperm

<table>
<thead>
<tr>
<th>Sperm parameters</th>
<th>Fertile patients N=7*</th>
<th>Infertile patients N=67</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sperm concentration (millionx10⁶)</td>
<td>49±2.14</td>
<td>17.57±0.53</td>
</tr>
<tr>
<td>Active sperm motility(%)</td>
<td>55.7±1.1</td>
<td>33.12±1.86</td>
</tr>
<tr>
<td>morphologically normal sperm By /HPF(%)</td>
<td>56.78±1.63</td>
<td>29.37±1.02</td>
</tr>
<tr>
<td>morphologically normal sperm By Kruger strict criteria(%)</td>
<td>15.5±0.14</td>
<td>10.3±0.25</td>
</tr>
<tr>
<td>Sperm agglutination(%)</td>
<td>6.6±2.3</td>
<td>23.3±0.8</td>
</tr>
<tr>
<td>Round cells/HPF</td>
<td>3.0±.21</td>
<td>8.94±0.7</td>
</tr>
</tbody>
</table>

Values are mean± SEM
*P<0.05: significantly different from other group

Table-2: Some female reproductive system characters of pre-ovulatory period by using ultra-sound vaginal examination

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Mean</th>
<th>SEM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cervical score</td>
<td>11.90</td>
<td>0.176</td>
</tr>
<tr>
<td>Endometrial thickness</td>
<td>8.76</td>
<td>0.269</td>
</tr>
<tr>
<td>Follicle size</td>
<td>19.53</td>
<td>0.393</td>
</tr>
</tbody>
</table>

Values are mean ± SEM
No. females=50
Figure-1: The percentage of morphologically normal sperms of fertile (control, n = 7) and infertile (treated, n=67) men.

Figure-2: Morphologically normal sperm in infertile patients pre-post coital test using Kruger strict criteria. +P<0.05: Significantly different from post-coital test.
Discussion

In this study there was a highly significant correlation (P<0.01) between pre-and-PCT sperm morphology which was examined by MKSC using the special pre-stained slides mentioned previously. This data suggest that more sperm with abnormal morphology were present after migration through cervical mucus than before. Under normal condition the cervical mucus act as the first selective fluid encountered by sperm after entering the female genital tract. It selects sperm according to their kinetic efficiency and morphology (Al-Dujaily et al.,2006 ; Ole et al.,2003). According to this hypothesis, the percentage of live, progressively motile sperm and morphologically normal sperm must be higher after migration through cervical mucus than before (Ole et al.,2003). The explanation for this controversial may be the presence of abnormal cervical mucus (providing that partner’s semen were normal) either due to the presence of congenital cervical anomalies (e.g lack of endo-cervical glands, absence of hormonal receptors), silent pelvic inflammatory disease, low pH, destruction of the endocervical mucosa by previous operations, presence of anti-sperm antibodies, (Kremer and Jager, 1988 and 1992) and the treatment with anti-estrogen (e.g. Clomphen citrate ).

Although, the semen characters and cervical mucus score in some couples had shown normal values, there was an increase of abnormal sperm morphology following PCT. This finding indicates that these couples classified as unexplained infertility cases, whereas our result proved that the cervix may altered the sperm morphology due to changes in cervix secretion or the sperm may had negative response to cervical secretion affecting the permeability of cell membrane causing abnormality of sperm shapes. This in turn will decrease the possibility of sperm-oocyte interaction or sperm fusion process (Rodrigues-Martined et al., 2001; Aziz et al., 1996) , there will be no chance for fertilization or pregnancy.

References


