Severe Oligospermia Treatment with Testicular Sperm Using “ICSI”

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ABSTRACT

Introduction: Assisted reproductive technology has been developed significantly throughout the past few years, particularly diagnosing and treating male infertility. Many studies have been performed showing that Intracytoplasmic Sperm Injection (ICSI) is a successful method to attain clinical pregnancy and live birth through impaired spermatozoa characteristics or low sperm count, such as severe oligospermia. Severe oligospermia indicates low sperm count, which in some cases leads to azoospermia. Severe oligospermia can be caused by several factors such as genetics or medication. In search of efficient treatment for couples with Severe oligospermia, numerous retrospective and prospective researches have reported high pregnancy and live birth rates through testicular sperm for men with severe oligospermia and cryptozoospermia with or without high sperm

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DNA damage. The research showed that the use of testicular sperm in combination with ICSI yielded a high pregnancy rate and live births over another source of sperm, such as ejaculated sperms. Moreover, the use of ICSI in severe oligospermia has shown successful fertilization and pregnancy.

**Methods:** In search for effective treatment for couples with severe male factor, a number of small retrospective and prospective studies have reported high pregnancy and live birth rates using testicular sperm for men with necrozoospermia, cryptozoospermia and oligozoospermia with or without elevated sperm DNA damage. Although the data suggest that there may be some benefit in performing testicular sperm retrieval (TSR)-ICSI in select groups of non-azoospermic infertile men, there are potential risks involved with TSR. Clinicians should balance these risks prior to the recommendation of TSR-ICSI on the result of a semen analysis or sperm DNA test alone. Careful evaluation and management of male factor infertility is important. The use of TSR-ICSI in the absence of specific sperm DNA defects is still experimental.

**Discussion:** In 1992 and subsequently, several reports indicated that ICSI was a successful technique to achieve clinical pregnancy and live birth using spermatozoa with severely impaired characteristics. The initial optimism over the ability of ICSI to overcome significant sperm abnormalities was later tempered by the findings of more recent publications suggesting that some sperm deficits may not be as effectively treated with ICSI.

**Conclusion:** Severe oligospermia indicates low sperm count, which can lead to male infertility; severe oligospermia which can be overcome through ICSI. Genetic factors like microdeletions of the Y chromosome (Yq) can cause severe oligospermia or chemotherapy molecules, affecting the sperm count directly.

**Keywords:** Severe oligospermia; Medication and cancer treatment influence on oligozoospermia; ICSI for men with sever oligozoospermia; testicular sperm, sperm retrieval; ICSI; male infertility; sperm DNA.

**ABBREVIATIONS**

- GnRH: Gonadotropin-releasing hormone
- ICSI: Intracytoplasmic Sperm Injection
- GnRH: Gonadotropin-releasing hormone
- LH: Luteinizing hormone
- FSH: Follicle-stimulating hormone
- Yq: Y chromosome
- AZF: Azoospermia factor
- CFTR: Cystic fibrosis transmembrane conductance regulator
- OAT: Oligo-astheni-teratozoospermia
- ART: Assisted reproductive technology
- DFI: DNA fragmentation index
- SDF: Sperm DNA fragmentation

**1. INTRODUCTION**

Male infertility factors have been recognised in around 40 %–50 % of infertile couples; more than 90% of cases in male infertility are due to low sperm counts, poor sperm quality or both. Other factors involve anatomical disorders, hormonal imbalances and genetic defects [1]. Moreover, male factors can include ejaculatory problems such as retrograde ejaculation, premature ejaculation or anejaculation, defects in sperm count, mobility, or abnormalities [1]. Some of the most common factors that correlate to irregularities in sperm are listed in (Table 1).

Male infertility has geographical variation in the prevalence of male infertility. High fertility levels can be found in France (59 %), 26 % –32 % in the UK and Kashmir Valley in India, and about 36 % in South Africa, Indonesia and Finland. Moreover, it has been shown that mean sperm concentration has regional variation in men from different regions of the USA and France [2].

<table>
<thead>
<tr>
<th>Sperm Abnormalities</th>
<th>Descriptions</th>
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<tbody>
<tr>
<td>Aspermia</td>
<td>Male with no semen</td>
</tr>
<tr>
<td>Azoospermia</td>
<td>Male with no sperms in the semen</td>
</tr>
<tr>
<td>Oligozoospermia</td>
<td>Low sperm concentration</td>
</tr>
<tr>
<td>Oligospermia</td>
<td>Low sperm count</td>
</tr>
<tr>
<td>Teratozoospermia</td>
<td>Irregular sperm shape (96% of sperms)</td>
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There are different ways of male infertility treatment, depending on the male infertility factors, for instance, improving the quality of sperm, surgery to repair a varicocele or intracytoplasmic sperm injection (ICSI) procedures. ICSI requires injecting one sperm directly into an oocyte prior to transferring to the uterus. ICSI has the advantage of using few viable sperm to fertilise an egg and can bypass natural barriers that prevent fertilisation [1].

This report will be discussing and evaluating the application of ICSI, in treating men with significant oligospermia.

2. SEVERE OLIGOSPERMIA

Oligozoospermia is a disorder where sperm concentration is below the lower reference limit of 15 million sperm/ml of ejaculate. Oligozoospermia can be further classified as mild (between 10 and 15 million sperm/ml), moderate oligozoospermia (between 5 and 10 million sperm/ml), and severe oligozoospermia (less than 5 million sperm/ml) [3]. Oligozoospermia diagnosing can be challenging, as several conditions are recognised to result in oligozoospermia. Therefore, a formal and precise diagnosis can be relevant in giving patients a reason for their sterility and decreasing confusion, fear, and stigma [3].

Disturbance of the regulated hormonal axis can induce the increase to several of the endocrine-based aetiologist of oligozoospermia. Usually, the hypothalamus produces and secretes gonadotropin-releasing hormone (GnRH) into the hypothalamic-hypophyseal portal circulation; GnRH influences the anterior pituitary to induce the gonadotroph cells to produce and release luteinizing hormone (LH) and follicle-stimulating hormone (FSH) into the systemic circulation. LH excites the Leydig cells of the testes to deliver testosterone, while FSH stimulates the Sertoli cells to induce spermatogenesis (Fig. 1). Generally, any destruction of gonadotropin secretion will result in a downstream outcome of disrupting intratesticular testosterone biosynthesis and spermatogenesis [3].

3. GENETICS FACTORS AND OLIGOZOOSPERMIA

Males with infertility have higher chromosomal irregularities than fertile men and usually do not present any phenotypic features. In the oligozoospermic, men of autosomal translocations have a rate of inversions predominate (3%) over sex chromosome anomalies (1.6%). Fertility may be expected, but interference with spermatocyte chromosomal pairing results in azoospermia or, more frequently, oligozoospermia [4].

Fig. 1. The hypothalamic-pituitary-testicular axis (Reynolds-Wright and Anderson, 2019)
Yq microdeletions are the most well-known genetic condition of spermatogenic failure. However, Microdeletions of the Y chromosome (Yq) include the azoospermia factor (AZF) regions with several copies of spermatogenic genes. The high oligozoospermic rate of approximately 4% of men with sperm densities of 0.1–5 million/ml. The AZFc deletion accounts for about 60% of Y microdeletions, and of these men, one-third have severe oligozoospermia. Moreover, the high homology within palindromic sequences induces intrachromosomal recombination and deletions of 0.8–7.7 megabase segments that arise in embryonic life [4].

AZF is divided into regions a, b, and c in the Y chromosome. Complete AZF deletions rates in the general population are rare (1 in 4,000); however, they happen in 10% of patients with idiopathic nonobstructive azoospermia and 5% of men with severe oligozoospermia (mainly with a male who has <2 million spermatozoa per ml). Another factor of AZF mutation is the carriage of NAHR substrates inside the AZFc region, which can give rise to the development of various partial AZFc deletions. The most clinically relevant NAHR deletion is the gr/gr deletion for its frequency. NAHR deleted half of the AZFc region gene content, which resulted in a substantially increased risk of a male having oligozoospermia [5].

Moreover, constitutional chromosome irregularities are also a common cause of male infertility, detected in up to 20% of infertile men with oligozoospermia. The aberrations involve numerical defects, like XYY karyotype in Klinefelter syndrome, its alternatives and structural rearrangements, Robertsonian translocations, balanced reciprocal translocations, and inversions [4]. The Men with 47, XYY syndrome can also be a factor of oligozoospermia; it has been recognised that germ cells with an extra Y chromosome from a male with the 47 XYY karyotypes have odd meiotic pairing, indicating disturbed meiosis, oligozoospermia and infertility. Furthermore, structural chromosomal irregularities are found in patients with azoospermia and oligozoospermia, as autosomal defects are more common in oligozoospermia. (SCAs) include deletions, duplications, translocations, and inversions [4].

Cystic fibrosis transmembrane conductance regulator (CFTR) gene mutations can induce oligozoospermia, associated with inherited infertility. Mutations in the (CFTR) gene frequently lead to male infertility. Most frequently, CFTR mutations point to a congenital bilateral loss of the vas deferens and severe oligozoospermia. Mutations in the CFTR gene is associated with obstruction of the male genital tract rather than primary testicular failure.

4. MEDICATION AND CANCER TREATMENT INFLUENCE ON OLIGOZOOSPERMIA

Medications are identified to affect the hypothalamic-pituitary-testicular axis functioning and could lead to infertility from oligozoospermia. For example, endocrine influences of long-term opioid therapy include suppression of the hypothalamic-pituitary-gonadal axis via suspension of hypothalamic pulsatile GnRH secretion, which results in hypogonadism, sexual dysfunction, and infertility [6]. Opioid high doses and long duration of opioid therapy are correlated with an increased rate of opioid-induced hypogonadism, though the impact points to be reversible following opioid discontinuation; as a result, this distribution can lead to oligozoospermia [6].

Cancer treatments effectiveness increase if a mixture of molecules is used, such as mechloethamine, vincristine, procarbazine, and prednisone) and ABVD (adriamycin, bleomycin, vinblastine, and dacarbazine) to treat cancer such as Hodgkin’s lymphoma [7]. Moreover, the medication regimen taken by children, adolescents, and young adults include alkylating agents (e.g., chlorambucil, cyclophosphamide, cisplatin, busulphan). Most chemotherapy molecules are associated with a high risk of infertility, that is, according to the degree of spermatogenesis damage ranging from oligozoospermia to non-obstructive azoospermia after recovery [7].

5. ICSI FOR MEN WITH SEVERE OLIGOZOOSPERMIA

New researches have investigated the potential of ICSI to overcome oligozoospermia; in cases of male subfertility, ICSI treatment results in high fertilization rates per oocyte compared with conventional IVF treatment. Also, it has been shown that severe patients of oligo-asthenoteratozoospermia (OAT) can now be successfully treated with ICSI [8].
Furthermore, investigate the association between prewash progressive sperm motility and pregnancy rate in severely oligoasthenozoospermic patients in (ICSI) cycle. The patients were classified into two groups depending on prewash progressive sperm motility [9]. Group one had progressive sperm motility under 10%, and group two had progressive sperm motility equal to or greater than 10%. The result shows no significant differences among the two groups regarding the total number of oocytes retrieved, number of mature oocytes, fertilization rate, or the number of transferred embryos. In group two, the clinical pregnancy rate was significantly higher where sperm motility was higher (62.5% [25/40] than in group I (32.5% [13/40]), with P = 0.014 [9].

Evaluate assisted reproductive technology (ART) in treating male infertility; patients were treated using different types of assisted reproductive techniques such as IUI, IVF or ICSI. IVF had the highest pregnancy rate among other ART treatments (46.4%) compared to ICSI and IUI, 35.5 and 16%, respectively. (Mukhtar et al., 2017) Moreover, in a patient with oligospermia, the pregnancy outcome among male partners was 47.8%, 36.4% and 20% in IVF, ICSI, and IUI, respectively [10] (Fig. 2).

Sperm parameters, such as morphology and motility impact fertilization and pregnancy rate, in addition to sperm DNA integrity, which is considered to be a major factor of the quality of the injected sperm. Male infertility with high sperm DNA fragmentation index (DFI) has result in poor ICSI outcomes with lower fertilization rate, pregnancy rate, live birth rate, and higher abortion rate [11]. HM, 2019, conducted a study to compare (ICSI) results in patients with severe oligospermia using testicular vs ejaculated spermatozoa. As testicular sperm has been shown to have considerably less DNA fragmentation than other sperm sources; as a result, the use of the testicular sperm with its higher DNA integrity can generate higher success rates in oligospermia men [11] The clinical pregnancy percentage was significantly higher in group B, where testicular spermatozoa were injected (49.50% vs 35.57%, p=0.044*). Still, there was no significant difference in the fertilization rate between group A and B (group A 67.93%, group B 68.01%, p=0.960). Moreover, high-quality embryos were found to be higher in the testicular sperm group [11].

Testicular sperm's strength relies on robust chromatin integrity since the main pathways leading to sperm DNA fragmentation are induced through sperm transport into the seminiferous tubules or epididymis transit [12]. The validity of this biological aspect relies on, firstly, chromatin compaction is continuous throughout epididymal transit. Secondly, high reactive oxygen species can be produced in the epithelial cells of epididymis under physicochemical stressors like high temperature and environmental circumstances. Finally, the DNA of mature live sperm can be cleaved by endonucleases. Consequently, sperm DNA fragmentation can occur within various pathways, revealing the high rate of SDF in live ejaculated sperm (Fig. 3) [12].

![Fig. 2. ART outcomes to treat male infertility, where IVF had the highest pregnancy rate (Mukhtar, Shaman, Mirghani and Almasalmah, 2017)](image-url)
Fig. 3. Origin of sperm and DNA fragmentation rate, as sperm DNA damage, can arise only in the testis and increase post testicular site (Muratori, 2006).

Fig. 4. Overview of TESA procedure. A) Percutaneous Testicular Sperm Aspiration where a small biopsy is aspirated of the testis tissue using a needle attached to a syringe (A1, A2). In B, conventional Testicular Sperm Extraction (TESE) is performed to open scrotal layers down to the albuginea. In C1, Testicular biopsy is flushed into the sperm media tube, then the testicular biopsy is washed free from blood clots, and seminiferous tubules are mechanically dispersed (C2); finally, the testicular homogenates are examined under the microscope to prove the presence of spermatozoa [13]

Table 2. Fertilization and embryo development after ICSI with ejaculated and testicular spermatozoa [16-19]

<table>
<thead>
<tr>
<th>Sperm Source</th>
<th>Attempts</th>
<th>Oocytes injected</th>
<th>Normal zygotes a</th>
<th>Fertilization rate b</th>
<th>Cleaved embryos c</th>
<th>Good-morphology embryos d</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ejaculate</td>
<td>18</td>
<td>185</td>
<td>131</td>
<td>70.8% e</td>
<td>124(94.7%) e</td>
<td>59(47.6%) e</td>
</tr>
<tr>
<td>Testis</td>
<td>18</td>
<td>187</td>
<td>140</td>
<td>74.9% e</td>
<td>133(95.0%) e</td>
<td>68(51.1%) e</td>
</tr>
</tbody>
</table>
Therefore, testicular sperm collected through testicular sperm aspiration (TESA) or extraction (TESE) have the potential of using and selecting spermatozoa free of DNA damage for ICSI (Fig. 4).

Besides, oocyte fertilization by a Genomically intact testicular spermatozoon will increase the possibilities of forming a normal embryonic genome that will eventually improve the likelihood of pregnancy and live birth [14].

Furthermore, as mentioned before, differences observed in fertility outcomes in male infertility treatment are due to the sperm DNA fragmentation level between ejaculated and testicular sperm. For instance, the results of terminal deoxynucleotidyl transferase-mediated dUTP nick-end labelling (TUNEL) positive sperm with DNA fragmentation is lower in the testicular sample compared with ejaculated samples. The use of testicular sperm for ICSI was associated with a 50% pregnancy and live birth rate [15].

Evaluate the potential of ICSI using testicular sperm to treat infertility in men with high sperm DNA fragmentation (SDF) and oligozoospermia. Sperm DNA fragmentation rate was almost 5-fold less in testicular sperm than in ejaculated sperm, where the usage of testicular sperm was associated with improved ICSI results in men with oligozoospermia and high SDF [15].

Evaluate the potential of ICSI using testicular sperm to treat infertility in men with high sperm DNA fragmentation (SDF) and oligozoospermia. Sperm DNA fragmentation rate was almost 5-fold less in testicular sperm than in ejaculated sperm, where the usage of testicular sperm was associated with improved ICSI results in men with oligozoospermia and high SDF [15]. Testicular sperm retrieval produced motile sperm for injections in all cases; accordingly, the sperm DNA fragmentation index was 8.3% in testicular sperm, compared to 40.7% in ejaculated sperm. For the testicular sperm - ICSI group, the clinical pregnancy rate was 51.9% versus 40.2% in the ejaculated sperm - ICSI group, the miscarriage rate was 10.0% for testicular sperm retrieval 34.3% in ejaculated sperm [15].

In the Greco et al., 2005 study, the DNA fragmentation rate in the testicular sperm samples was 3.6%, significantly lower than the ejaculated sperm samples from the same individuals (23.6%). Moreover, the use of testicular spermatozoa for ICSI decreases the reproductive disadvantage associated with the use of ejaculated spermatozoa for ICSI in oligozoospermia patients (Table 2) [16-19].

6. CONCLUSION

Severe oligospermia indicates low sperm count, which can lead to male infertility; severe oligospermia which can be overcome through ICSI. Genetic factors like microdeletions of the Y chromosome (Yq) can cause severe oligospermia or chemotherapy molecules, affecting the sperm count directly. The research results indicate that ICSI, effectively treat infertility due to severe oligospermia, mainly if testicular sperms are used through TESA. Since testicular sperms have a robust chromatin integrity, which can save sperms from DNA fragmentation in cases of severe oligospermia.

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Objective of the Association for Scientific Research of the IRIFIV-AISRG Reproductive Medicine and Endometriosis Research Group (IRIFIV-AISRG), Research foundation in Casablanca, Maintaining consistent and reliably high success rates is a monthly challenge for in IVF labs, the IRIFIV Fertility Center in Casablanca – Morocco Department of Reproductive Medicine and Reproductive Biology and Embryology, advocacy of interdisciplinary Department of Reproductive Medicine and Reproductive Biology and Embryology study, encompassing the areas of research, collections and publishing Articles.

COMPETING INTERESTS

Authors have declared that no competing interests exist.

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