Evaluation of selected semen parameters and biomarkers of male infertility – preliminary study [version 1; peer review: 1 approved, 2 approved with reservations]

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Abstract

Background: Because the etiopathogenesis of male infertility is multifactorial our study was designed to clarify the relationship between standard semen parameters, testicular volume, levels of reproductive hormones and the fragmentation of sperm nuclear DNA (SDF).

Methods: Patients (n = 130) were clustered as subjects: 1) with an abnormal volume (ultrasoundography) of at least one testis (<12 mL) or with a normal volume of testes and 2) with abnormal levels of at least one of the reproductive hormones (FSH, LH, PRL, TSH, total T – electrochemiluminescence method) or with normal hormonal profiles and 3) with high level of SDF (>30%), moderate (>15–30%) or low (≤15%) (sperm chromatin dispersion test).

Results: In subjects with a decreased testicular volume and in subjects with abnormal levels of reproductive hormones, decreased basic semen parameters were found. Participants with abnormal testicular volume had a higher percentage of SDF and a higher level of FSH (Mann–Whitney U test). In turn, men with a high level of SDF had lower testicular volume and conventional sperm parameters than
men with a low level of SDF (Kruskal–Wallis test).

**Conclusions:** We showed that spermatogenesis disorders coexisted with decreased testicular volume and increased FSH levels. The disorders of spermatogenesis were manifested by reduced basic sperm characteristics and a high level of sperm nuclear DNA damage.

**Keywords**
male infertility, semen characteristics, testicular volume, reproductive hormones, sperm nuclear DNA integrity
Introduction
The etiopathogenesis of male infertility is a multifactorial medical problem and is correlated with many congenital and acquired defects of the urogenital tract, cancers, urogenital infections, heat stress in the scrotum, hormonal disorders, genetic abnormalities and immunological factors. It is estimated that approximately 30–50% of male infertility cases are recognized as idiopathic, very often associated with low-quality of spermatozoa.1–4 On the other hand, unexplained infertility (couples where male patients have normal basic semen parameters and female patients have normal ovulation and fallopian tube potency) is diagnosed in 15–30% of cases.1–4 Therefore, the comprehensive evaluation of male fertility status should be developed using scrotal ultrasonography (USG) and assessment of the key reproductive hormone as well as advanced seminological tests.3,5–12

Available data has suggested that it is possible that infertile men could have normal standard semen characteristics.1,13,14 Therefore, it is important to look beyond conventional semen analysis. Many authors report that among the advanced sperm tests, the assays that verify sperm nuclear DNA fragmentation (SDF) are the most clinically useful. Furthermore, evaluation of the percentage of SDF could significantly help in determining the most beneficial treatment algorithm for couples trying to have offspring.5,7,15–20 An SDF ≤15% is considered a normal value (low level of nuclear DNA damage) and correlates with high male fertility potential. In these cases, the chance of becoming pregnant naturally or by intrauterine insemination (IUI) is high. In turn, SDF >15–30% (moderate level of DNA damage) can be associated with a reduced chance of becoming pregnant through natural conception and IUI or even in vitro fertilization (IVF) treatment. This range of SDF values and history of previous unsuccessful attempts to achieve pregnancy might indicate the need to introduce intracytoplasmic sperm injection (ICSI). Finally, a high level of SDF (>30%) is strongly associated with a significantly increased risk of reproductive failure, including ICSI treatment. It should be highlighted that even if pregnancy due to assisted reproductive technology (ART) is achievable, the percentage of sperm cells with a fragmented genome >30%, especially >40%, may significantly increase the risk of pregnancy loss.4,16,19,21–30 These three ranges of sperm DNA damage (≤15%, >15–30% and >30) were primarily recommended for interpretation of the SCSA test results.16,22,25,26,29 However similar ranges also have been successfully adapted to sperm chromatin dispersion test (SCD).19,23,31 Hence, an in-depth assessment of male fertility status, including testicular ultrasound, the levels of reproductive hormones and basic and advanced semen analysis, is clinically justified. Therefore, our study was designed to (1) determine the relationship between testicular volume, levels of reproductive hormones (follicle-stimulating hormone – FSH, luteinizing hormone – LH, prolactin – PRL, total testosterone – total T, thyroid-stimulating hormone – TSH), standard semen analysis and sperm genomic integrity and 2) compare standard semen parameters and investigated biomarkers of male infertility between groups of participants with low, moderate and high levels of nuclear DNA fragmentation.

Methods
Ethical considerations
In accordance with the Declaration of Helsinki, all participants in the study indicated their written conscious and voluntary consent to participate in the scientific project. The study was conducted according to the guidelines of the Declaration of Helsinki and approved by the Ethics Committee of Pomeranian Medical University, Szczecin, Poland (KB-0012/21/18, date of approval: 5 February 2018).

Study population
The study population consisted of 130 male infertile participants (median age: 33.00 years; range: 23–51 years) who were treated in 2018–2021 in the Individual Specialist Medical Practice (Szczecin, Poland) and The Fertility Partnership Vitrolive in Szczecin (Poland) – Gynaecology and Fertility Clinic and who gave their consent to participate in the study. All patients were partners of women (n = 130; median age: 30.00 years; range: 22–46 years) who did not become pregnant during one year (median: 2 years; range: 1.00–14.00 years) of regular intercourse without contraception (Figure 1). All initially qualified participants during a medical interview reported to the Laboratory of Andrology in the Department of Histology and Developmental Biology (Pomeranian Medical University, Szczecin, Poland) for seminological analysis. Based on the performed basic semen analysis, men with azoospermia and cryptozoospermia were excluded from the study group.

The infertile status of subjects was verified based on an in-depth medical interview conducted by a specialist in urology (M. K.). The interview included information about factors that may affect fertility potential (genital injuries, cryptorchidism, varicocele, urogenital infections, chronic diseases, pharmacotherapy, use of anabolic steroids, operations and treatments, exposure to harmful factors, lifestyle, stimulants and others) (the interview form can be found as Extended data.11) Moreover, the physical medical examination included body assessment (body and hair proportions), and palpation (penis, gonads, epididymides, seminal cords, inguinal canal, prostate and mammary glands) was carried out.
Conventional semen analysis

Standard semen analysis was carried out according to World Health Organization (WHO) recommendations.32 Semen samples were collected in a sterile urine container by masturbation after a two- to seven-day sexual abstinence. After complete liquefaction of semen (at 37 °C), standard semen analysis was carried out at room temperature (22°C). The macroscopic evaluation of the semen included color, viscosity, volume and pH. In turn, the microscopic assessment (light/phase-contrast microscope DM500, Lecia, Heerbrugg, Switzerland) included the initial verification of the samples (presence of mucus bands, erythrocytes, epithelial cells, spermine crystals, residual bodies, aggregation and agglutination of sperm) as well as the assessment of the sperm concentration and the total sperm count, motility (progressive and nonprogressive motility), morphology, vitality and the concentration of inflammatory cells. Sperm concentration (analyzed in an improved Neubauer hemocytometer – Heinz Hernez Medizinalbedarf GmbH, Hamburg, Germany, ref no 1080339), sperm motility and vitality (eosin-positive cells and hypoosmotic-reactive cells: HOS test-positive cells) were assessed under a light/phase-contrast microscope using a 40xC2 objective. To evaluate sperm cell morphology (including the teratozoospermia index reflecting multiple morphological defects per spermatozoon – TZI), native sperm smears were fixed and stained according to the Papanicolaou method (Aqua-Med, Lodz, Poland) and were analyzed under a bright light microscope using a 100xC2 objective oil immersion lens. The concentration of leukocytes (peroxidase-positive cells) was calculated using the Endtz test (LeucoScreen kit, FertiPro N.V., Beernem, Belgium) and assessed in an improved Neubauer hemocytometer.

Sperm Chromatin Dispersion (SCD) test

To verify sperm nuclear DNA fragmentation, a commercial chromatin dispersion test – a Halosperm G2® kit (Halotech DNA, Madrid, Spain) – was applied. The procedure was performed strictly according to the manufacturer’s guidelines and was described in detail in our previous publications.33-36 To calculate the percentage of sperm cells with fragmented DNA, a minimum of 300 sperm cells per sample was counted under the 100x objective of a bright light microscope. According to the manufacturer’s guidelines, the following evaluation criteria were used: (1) sperm cells without nuclear DNA fragmentation (spermatozoa with a large halo – equal to or higher than the diameter of the core of spermatozoa and spermatozoa with a medium-sized halo – >1/3 of the diameter of the core of spermatozoa) and (2) sperm cells with nuclear DNA fragmentation (spermatozoa with a small halo – ≤1/3 of the diameter of the core of spermatozoa and spermatozoa without a halo but with a strongly stained core or without a halo and degraded chromatin – sperm cells showing no halo and simultaneously presenting an irregularly or weakly stained core) (Figure 2). The results of the SCD test (SDF) are presented as the sum of spermatozoa with nuclear DNA fragmentation divided by the total number of assessed sperm cells and multiplied by 100%.

Hormone profile of infertile subjects

To assess the panel of basic hormones influencing male fertility, potential vein blood was collected from participants in the morning (7.30–09.00), and the following hormones presented in Table 1 were measured. The hormone levels were determined by the electrochemiluminescence method (ECLIA) using the Cobas e801 analytical unit (Roche Diagnostics GmbH, Mannheim, Germany). The ECLIA method is based on the binding of biotinylated monoclonal-specific antibodies directed against the measured hormones and specific monoclonal antibodies labeled with a complex containing ruthenium metal (sandwich complex). In the next step, streptavidin-bound microparticles were used to bind to biotinylated antibodies directed against the measured hormones. Unbound substances were removed. The bound
microparticles were magnetically trapped on the electrode surface. The electrode voltage induced chemiluminescence emission, which was measured by a photomultiplier. The result was determined based on a two-step calibration. All hormonal analyses were assessed strictly in accordance with the manufacturer's instructions.

Ultrasonography of the scrotum

To verify testicular volume, USG of the scrotum (ultrasound system Z-5 with a 75L38EA linear head; frequency range of 5–10 MHz, Mindray, Shenzhen, China) was performed by a senior urologist. The measurements were calculated using the following formula: length × width × height × 0.71. Furthermore, they were expressed in mL. According to the most commonly accepted criterion in clinical practice, the hypotrophic gonad was considered when the volume of the testis was less than 12 mL. Additionally, the homogeneity and echogenicity of the gonadal parenchyma as well as the presence of possible focal lesions and microcalcifications were assessed.

Statistical analysis

Because the Shapiro–Wilk test showed that the data were not normally distributed, a nonparametric Mann–Whitney U test and Kruskal–Wallis test were applied to compare quantitative variables between two or more studied groups, respectively. Quantitative variables are expressed as the median with the range and mean ± standard deviation (SD). Additionally, to verify the relationships between study parameters, the Spearman’s rank (r_s) correlation coefficient was calculated (Figure 1). To interpret the strength of dependence between parameters, the following levels of correlation

Table 1. Reproductive hormones evaluated in study population.

| Hormone | Short description of biological function | Reference value |
|---------|------------------------------------------|-----------------|
| FSH     | FSH levels are considered a marker of impaired spermatogenesis | 1.5–12.4 μIU/mL |
| LH      | LH levels are helpful for differentiating primary (testicular) hypogonadism from secondary (nontesticular) hypogonadism | 1.7–8.6 μIU/mL |
| total T | Total T levels influence spermatogenesis through Sertoli cells | 8.0–41.7 nmol/L |
| PRL     | PRL levels may result in decreased libido and erectile function | 4.04–15.02 ng/mL |
| TSH     | Both hyperthyroidism and hypothyroidism can negatively affect spermatogenesis | 0.27–4.2 μIU/mL |

FSH – follicle-stimulating hormone, LH – luteinizing hormone, PRL – prolactin, TSH – thyroid-stimulating hormone, total T – total testosterone.

Figure 2. Visualization of sperm chromatin dispersion test (SCD). Micrographs obtained by light microscopy, 100 ×. Scale bar = 5 μm. Raw micrographs were edited in Corel Photo-Paint 2019 (Corel Corporation, Ottawa, Canada, RRID:SCR_014235). Editing included only: cropping (to center the presented sperm cells), rotation, brightening, contrast enhancing and enlargement.
were presumed: <0.2 – lack of linear dependence (regardless of the p value), 0.2–0.4 – weak dependence, >0.4–0.7 – moderate dependence, >0.7–0.9 – strong dependence and >0.9 very strong dependence. For all statistical analyses, a p value < 0.05 was considered significant. Data analysis was performed using Statistica version 13.3 (StatSoft, Krakow, Poland, RRID:SCR_014213) and MedCalc version 18.2.1 (MedCalc Software, Ostend, Belgium, RRID:SCR_015044). Open source statistical software which can be used in the study – GNU PSPP.

Results
Seminological characteristics of study population
Of 130 obtained semen samples, 26 were classified as normozoospermia (total sperm count ≥39 × 10⁶ cells, sperm progressive motility ≥32%, normal sperm morphology ≥4%), 36 as teratozoospermia (abnormal sperm morphology), 36 as oligoasthenoteratozoospermia (simultaneously abnormal total sperm count, progressive motility and morphology), 19 as oligoteratozoospermia (simultaneously abnormal total sperm count and morphology), nine as asthenoteratozoospermia (simultaneously abnormal progressive sperm motility and morphology), three as oligozoospermia (abnormal sperm total count) and one as asthenozoospermia (abnormal sperm progressive motility). Moreover, 51 men had abnormal levels of at least one of the assessed reproductive hormones (FSH, LH, PRL, total T, TSH), and 37 men had an abnormal volume of at least one testis (<12 mL). The descriptive statistics of the investigated parameters are provided in Table 2.

Comparison of study parameters between infertile subjects with an abnormal volume of at least one testis (<12 mL) and subjects with a normal volume of testes (each ≥12 mL)
Compared groups did not differ in age. The subjects from the group with abnormal testicular volume (n = 37) had significantly higher levels of FSH than the reference group (n = 91) (median: 8.05 mIU/mL vs. 5.29 mIU/mL), whereas the levels of other study hormones (LH, PRL, total T, TSH) did not differ significantly (Table 3). On the other hand, in case of seminological parameters, patients with decreased testicular volume had a significantly reduced sperm concentration (medians: 6.25 × 10⁶ cells/mL vs. 19.00 × 10⁶ cells/mL), total sperm count (medians: 15.35 × 10⁶ cells/mL vs. 29.48 × 10⁶ cells/mL), total sperm motility (%) (medians: 48.79 ± 22.32 vs. 58.67 ± 22.25), and sperm nonprogressive motility (%) (medians: 5.56 ± 4.40 vs. 5.92 ± 4.40).

Table 2. Descriptive statistics of study parameters in the infertile group of men (n=130).

| Study parameters                  | median (range) | mean ± SD |
|-----------------------------------|----------------|-----------|
| Age (y)                           | 33.00 (23.00–51.00) | 33.77 ± 5.28 |
| LTV (mL)                          | n = 128 15.00 (2.40–25.00) | 14.99 ± 4.81 |
| RTV (mL)                          | n = 128 15.00 (4.30–25.00) | 15.50 ± 4.95 |
| FSH (mIU/mL)                      | n = 128 6.33 (1.17–21.35) | 6.84 ± 3.90 |
| LH (mIU/mL)                       | 5.51 (1.21–17.10) | 5.66 ± 2.63 |
| total T (nmol/L)                  | 16.60 (3.37–39.48) | 17.53 ± 6.22 |
| PRL (ng/mL)                       | 13.22 (5.55–46.56) | 14.25 ± 5.90 |
| TSH (μU/mL)                       | n = 129 1.78 (0.63–14.40) | 2.17 ± 1.50 |
| Semen volume (mL)                 | 3.00 (0.50–8.50) | 3.34 ± 1.63 |
| Sperm concentration (×10⁶/mL)     | 13.82 (0.40–251.00) | 21.60 ± 28.54 |
| Total number of spermatozoa (×10⁶) | 42.70 (0.84–426.70) | 67.76 ± 79.57 |
| Morphologically normal spermatozoa (%) | 1.00 (0.00–11.00) | 1.80 ± 2.49 |
| TZI                               | 1.75 (1.34–2.50) | 1.80 ± 0.29 |
| Sperm progressive motility (%)    | 44.00 (0.00–87.00) | 43.22 ± 22.25 |
| Sperm nonprogressive motility (%) | 5.00 (0.00–29.00) | 5.56 ± 4.40 |
| Total sperm motility (%)          | 50.00 (1.00–98.00) | 48.79 ± 22.32 |
| Eosin-negative spermatozoa – live cells (%) | 76.50 (30.00–96.00) | 74.33 ± 13.29 |
| HOS test-positive spermatozoa – live cells (%) | n = 107 78.00 (26.00–92.00) | 75.60 ± 10.51 |
| Peroxidase-positive cells (%)     | 0.25 (0.00–10.25) | 0.66 ± 1.32 |
| SDF (%)                           | 20.00 (3.00–58.00) | 22.06 ± 12.04 |

FSH – follicle-stimulating hormone, HOS test – hypo-osmotic swelling test, LH – luteinizing hormone, LTV – left testis volume, SDF – sperm DNA fragmentation, PRL – prolactin, RTV – right testis volume, TSH – thyroid-stimulating hormone, total T – total testosterone, TZI – teratozoospermia index, n – number of subjects, SD – standard deviation.
Table 3. Descriptive statistics and comparison of age, testicular volume and levels of reproductive hormones between infertile subjects with an abnormal volume of at least one testis (<12 mL) and subjects with a normal volume of testes (each ≥12 mL).

| Study parameters | Subjects with abnormal volume of at least one testis (<12 mL) n = 37 | Subjects with normal volume of testes (each ≥12 mL) n = 91 | p     |
|------------------|-------------------------------------------------|-------------------------------------------------|-------|
| Age (y)          | 33.00 (23.00–49.00) 33.34 ± 4.89               | 33.00 (22.00–48.00) 33.60 ± 5.06               | NS    |
| LTV (mL)         | 10.00 (2.40–18.00) 9.81 ± 3.27                  | 16.00 (12.00–25.00) 17.10 ± 3.58               | <0.000000 |
| RTV (mL)         | 10.00 (4.30–24.00) 10.82 ± 4.10                 | 16.70 (9.00–25.00) 17.40 ± 3.90                | <0.000000 |
| FSH (mIU/mL)     | 8.05 (2.10–21.35) 8.66 ± 4.23                   | n = 90 5.29 (1.17–17.00) 6.05 ± 3.30           | 0.002677 |
| LH (mIU/mL)      | 5.86 (2.30–17.10) 6.37 ± 3.32                   | 5.49 (1.21–11.20) 5.41 ± 2.22                  | NS    |
| total T (nmol/L) | 16.12 (4.16–27.07) 4.76 ± 1.62                  | 17.75 (3.37–39.48) 5.18 ± 1.87                 | NS    |
| PRL (ng/mL)      | 13.12 (6.80–25.52) 13.84 ± 5.28                 | 13.20 (5.55–46.56) 14.39 ± 6.18                | NS    |
| TSH (μU/mL)      | 1.61 (0.63–14.40) 2.13 ± 2.21                   | n = 90 2.01 (0.71–6.70) 2.19 ± 1.09            | NS    |

FSH = follicle-stimulating hormone, LH = luteinizing hormone, LTV = left testis volume, PRL = prolactin, RTV = right testis volume, TSH = thyroid-stimulating hormone, total T = total testosterone. n = number of subjects; NS = no statistical significance; SD = standard deviation. Statistical significance in the Mann-Whitney U test was reached when p < 0.05.

vs. 6.01.2 × 10^6 cells), sperm morphology (medians: 0.00% vs. 1.00%), progressive motility (medians: 37.00% vs. 51.00%), total motility (medians: 43.50% vs. 57.00%) and vitality – eosin-negative sperm cells (medians: 73.00% vs. 79.00%) and hypoosmotic (HOS) test-positive sperm cells (medians: 73.50% vs. 78.00%). It should be highlighted that in the group of men with abnormal testicular volume, a significantly higher percentage of SDF was found (medians: 27.00% vs. 17.00%) (Table 4).

Comparison of study parameters between infertile subjects with abnormal levels of at least one of the assessed hormones and subjects with normal hormonal profiles

There were no significant differences in age between the compared groups, whereas higher levels of FSH, LH, PRL and TSH were noted in infertile men with abnormal hormonal profiles (n = 51) than in infertile men with normal hormonal profiles (n = 79). Unexpectedly, the study groups did not differ in the level of total T (Table 5). Regarding the semen parameters, infertile men with hormonal disorders had significantly lower total sperm count (medians: 30.25 × 10^6 cells vs. 54.00 × 10^6 cells), sperm morphology (medians: 0.00% vs. 1.00%), progressive motility (medians: 35.00% vs. 50.00%) and total motility (medians: 41.00% vs. 57.00%). Furthermore, the percentage of SDF was increased in the group with hormonal abnormalities, but the difference was not statistically significant (medians: 22.00% vs. 18.00%). Additionally, the compared groups did not differ in testicular volume (Table 6).

Comparison of study parameters between infertile subjects with SDF >30%, >15–30% and ≤15%

Based on the publications of other authors,16,19,22,23,25,26,29,31 the study group was divided into three subgroups: 1) with a high level of sperm nuclear DNA damage (SDF >30%, low fertility potential), 2) with a moderate level of sperm nuclear DNA damage (SDF >15–30%, moderate fertility potential) and 3) with a low level of sperm nuclear DNA damage (SDF ≤15%, high fertility potential) (Tables 7, 8).

Statistical analysis revealed some significant differences between men with SDF >30% (n = 28) and men with SDF ≤15% (n = 43). The first group had significantly lower left testis volume (medians: 13.00 mL vs. 16.00 mL) and right testis volume (medians: 12.00 mL vs. 16.00 mL), sperm concentration (medians: 5.65 × 10^6 cells/mL vs. 21.75 × 10^6 cells/mL), total sperm count (medians: 20.02 × 10^6 cells vs. 70.76 × 10^6 cells), sperm morphology (medians: 0.00% vs. 3.00%), progressive motility (medians: 26.00% vs. 63.00%), total motility (medians: 33.00% vs. 68.00%) and vitality – eosin-negative sperm cells (medians: 67.00% vs. 86.00%) and HOS test-positive sperm cells (medians: 67.00% vs. 83.00%) (Tables 7, 8).

In addition, in contrast to men with SDF ≤15%, infertile men with SDF >15–30% (n = 59) had a significantly lower total sperm count (medians: 41.25 × 10^6 cells vs. 70.76 × 10^6 cells), sperm morphology (medians: 0.00% vs. 3.00%), progressive motility (medians: 40.00% vs. 63.00%), total motility (medians: 48.00% vs. 68.00%) and sperm vitality – eosin-negative sperm cells (medians: 74.00% vs. 86.00%) and HOS test-positive sperm cells (medians: 72.00% vs. 83.00%) (Table 7).
Table 4. Descriptive statistics and comparison of semen parameters between infertile subjects with an abnormal volume of at least one testis (<12 mL) and subjects with a normal volume of testes (each ≥12 mL).

| Study parameters                      | Subjects with abnormal volume of at least one testis (<12 mL) n = 37 | Subjects with normal volume of testes (each ≥12 mL) n = 91 | p        |
|---------------------------------------|---------------------------------------------------------------|------------------------------------------------------|----------|
| median (range) mean ± SD              |                                                               |                                                      |          |
| Semen volume (mL)                     | 3.00 (0.75–6.50) 3.04 ± 1.25                                   | 3.00 (0.50–8.50) 3.48 ± 1.76                        | NS       |
| Sperm concentration (×10⁶/mL)         | 6.25 (0.40–57.25) 11.26 ± 12.63                                | 19.00 (0.45–251.00) 26.05 ± 32.17                    | 0.000111 |
| Total number of spermatozoa (×10⁶)    | 15.35 (0.88–169.00) 33.28 ± 39.25                              | 60.12 (0.84–426.70) 82.64 ± 87.72                    | 0.000046 |
| Morphologically normal spermatozoa (%)| 0.00 (0.00–10.00) 1.05 ± 1.95                                  | 1.00 (0.00–11.00) 2.14 ± 2.63                        | 0.028614 |
| TZI                                   | 1.71 (1.36–2.50) 1.79 ± 0.29                                   | 1.75 (1.34–2.48) 1.81 ± 0.28                         | NS       |
| Sperm progressive motility (%)        | 37.00 (0.00–87.00) 37.36 ± 20.47                               | 51.00 (2.00–86.00) 45.98 ± 22.49                      | 0.032661 |
| Sperm nonprogressive motility (%)     | 5.00 (0.00–18.00) 5.50 ± 3.98                                  | 5.00 (0.00–29.00) 5.63 ± 4.59                        | NS       |
| Total sperm motility (%)              | 43.50 (1.00–90.00) 42.86 ± 20.70                               | 57.00 (4.00–88.00) 51.62 ± 22.42                      | 0.026757 |
| Eosin-negative spermatozoa – live cells (%) | 73.00 (41.00–95.00) 70.07 ± 13.08                             | 79.00 (30.00–96.00) 76.16 ± 13.10                    | 0.003811 |
| HOS test-positive spermatozoa – live cells (%) | n = 26 73.50 (54.00–88.00) 72.38 ± 9.60 | n = 80 78.00 (26.00–92.00) 76.73 ± 10.66 | NS      |
| Peroxidase-positive cells (%)         | 0.25 (0.00–10.25) 0.65 ± 1.66                                  | 0.25 (0.00–9.50) 0.66 ± 1.17                         | NS       |
| SDF (%)                               | 27.00 (3.00–58.00) 29.00 ± 14.30                               | 17.00 (4.00–46.00) 19.16 ± 9.75                      | 0.000127 |

HOS test – hypo-osmotic swelling test, SDF – sperm DNA fragmentation, TZI – teratozoospermia index. n – number of subjects, NS – no statistical significance, SD – standard deviation. Statistical significance in the Mann–Whitney U test was reached when p < 0.05.

Table 5. Descriptive statistics and comparison of age, testicular volume and levels of reproductive hormones between infertile subjects with abnormal hormone profiles (abnormal levels of at least one evaluated hormone) and subjects with normal hormone profiles.

| Study parameters                      | Subjects with abnormal hormone profile n = 51 | Subjects with normal hormone profile n = 79 | p        |
|---------------------------------------|------------------------------------------------|------------------------------------------------|----------|
| median (range) mean ± SD              |                                                               |                                                      |          |
| Age (y)                               | 33.00 (25.00–47.00) 33.43 ± 4.45                        | 33.00 (22.00–49.00) 33.48 ± 4.44                    | NS       |
| LTV (mL) n = 49                       | 14.00 (4.30–24.00) 13.98 ± 5.02                        | 15.60 (4.00–25.00) 15.62 ± 4.59                     | NS       |
| RTV (mL) n = 50                       | 14.00 (4.30–24.00) 14.61 ± 4.98                        | 15.00 (4.60–25.00) 15.60 ± 4.78                     | NS       |
| FSH (mIU/mL) n = 78                   | 7.70 (1.17–21.35) 7.90 ± 4.23                          | 7.80 (1.80–24.17) 7.80 ± 4.33                       | 0.11483 |
| LH (mIU/mL) n = 50                    | 6.82 (1.45–17.10) 6.94 ± 3.07                          | 4.49 (1.21–8.55) 4.83 ± 1.91                        | 0.00024  |
| total T (nmol/L)                      | 15.60 (3.37–28.77) 16.08 ± 6.37                        | 17.95 (8.32–39.48) 5.31 ± 1.74                      | NS       |
| PRL (ng/mL) n = 80                    | 19.10 (6.42–46.56) 18.85 ± 6.35                        | 11.79 (5.55–17.90) 11.29 ± 3.00                      | < 0.00000|
| TSH (μU/mL) n = 50                    | 16.25 (0.63–14.40) 2.49 ± 1.98                         | 1.65 (0.70–6.70) 1.97 ± 1.05                        | 0.028703 |

FSH – follicle-stimulating hormone, LH – luteinizing hormone, LTV – left testis volume, PRL – prolactin, RTV – right testis volume, TSH – thyroid-stimulating hormone, total T – total testosterone. n – number of subjects, NS – no statistical significance, SD – standard deviation. Statistical significance in the Mann–Whitney U test was reached when p < 0.05.
Table 6. Descriptive statistics and comparison of semen parameters between infertile subjects with abnormal hormone profiles (abnormal levels of at least one evaluated hormone) and subjects with normal hormone profiles.

| Study parameters | Subjects with abnormal hormone profile n = 51 | Subjects with normal hormone profile n = 79 | p       |
|------------------|-----------------------------------------------|---------------------------------------------|---------|
|                  | median (range) mean ± SD                      |                                             |         |
| Semen volume (mL)| 3.00 (0.75–7.90) 3.07 ± 1.49                  | 3.50 (0.50–8.50) 3.52 ± 1.70                | NS      |
| Sperm concentration (× 10⁹/mL) | 12.87 (0.40–118.50) 17.42 ± 20.70 | 15.62 (0.45–251.00) 24.31 ± 32.46 | NS      |
| Total number of spermatozoa (× 10⁶) | 30.25 (0.88–412.77) 52.53 ± 69.77 | 54.00 (0.84–426.70) 77.60 ± 84.27 | 0.018847 |
| Morphologically normal spermatozoa (%) | 0.00 (0.00–8.00) 1.17 ± 1.95 | 1.00 (0.00–11.00) 2.21 ± 2.72 | 0.013886 |
| TZI               | 1.75 (1.35–2.50) 1.82 ± 0.32                  | 1.75 (1.34–2.45) 1.79 ± 0.27                | NS      |
| Sperm progressive motility (%) | 35.00 (0.00–86.00) 37.19 ± 21.44 | 50.00 (1.00–87.00) 47.11 ± 22.03 | 0.011304 |
| Sperm nonprogressive motility (%) | 5.00 (0.00–18.00) 4.88 ± 3.19 | 5.00 (0.00–29.00) 6.01 ± 4.99 | NS      |
| Total sperm motility (%) | 41.00 (1.00–88.00) 42.07 ± 21.56 | 57.00 (2.00–90.00) 53.12 ± 21.84 | 0.005652 |
| Eosin-negative spermatozoa – live cells (%) | 74.00 (44.00–96.00) 73.47 ± 11.54 | 79.00 (30.00–95.00) 74.88 ± 14.35 | NS      |
| HOS test-positive spermatozoa – live cells (%) | n = 41 76.00 (54.00–91.00) 75.12 ± 8.66 | n = 66 78.50 (26.00–92.00) 75.90 ± 11.56 | NS      |
| Peroxidase-positive cells (%) | 0.25 (0.00–10.25) 0.94 ± 2.00 | 0.25 (0.00–2.15) 0.48 ± 0.51 | NS      |
| SDF (%)           | 22.00 (7.00–58.00) 23.54 ± 12.05              | 18.00 (3.00–54.00) 21.10 ± 12.02            | NS      |

HOS test – hypo-osmotic swelling test, SDF – sperm DNA fragmentation, TZI – teratozoospermia index. n – number of subjects, NS – no statistical significance, SD – standard deviation. Statistical significance in the Mann–Whitney U test was reached when p < 0.05.

Table 7. Descriptive statistics and comparison of age, testicular volume and levels of reproductive hormones between infertile subjects with SDF >30%, >15–30% and ≤15%.

| Study parameters | Subjects with SDF >30% (1) n = 28 | Subjects with SDF >15–30% (2) n = 59 | Subjects with SDF ≤15% (3) n = 43 | p 1 vs. 2 | p 1 vs. 3 | p 2 vs. 3 |
|------------------|-----------------------------------|------------------------------------|-----------------------------------|-----------|-----------|-----------|
|                  | median (range) mean ± SD          |                                    |                                   |           |           |           |
| Age (y)          | 31.00 (27.00–51.00) 33.00 ± 5.63   | 34.00 (25.00–49.00) 34.35 ± 5.90   | 34.00 (23.00–43.00) 33.48 ± 4.04 | NS        | NS        | NS        |
| LTV (mL)         | 13.00 (4.00–23.00) 13.48 ± 4.76    | n = 57 14.00 (2.40–25.00) 14.55 ± 5.02 | 16.00 (9.00–25.00) 16.58 ± 4.15 | NS        | 0.024251  | NS        |
| RTV (mL)         | n = 27 12.00 (8.00–23.00) 13.78 ± 4.86 | n = 58 15.00 (4.30–25.00) 15.38 ± 5.12 | 16.00 (8.00–24.00) 16.73 ± 4.52 | NS        | 0.032806  | NS        |
| FSH (mIU/mL)     | 6.32 (1.80–14.50) 6.60 ± 3.55      | 6.70 (1.93–21.35) 7.51 ± 4.49      | n = 42 5.44 (1.17–13.40) 6.06 ± 3.06 | NS        | NS        | NS        |
| LH (mIU/mL)      | 5.07 (2.10–9.30) 5.45 ± 2.47       | 5.77 (1.45–17.10) 5.93 ± 3.09      | 5.20 (1.21–11.20) 5.43 ± 2.01    | NS        | NS        | NS        |
| total T (nmol/L) | 18.16 (4.16–28.08) 5.10 ± 1.51     | 16.19 (3.37–37.20) 17.05 ± 6.76    | 16.12 (8.66–39.48) 18.09 ± 6.10  | NS        | NS        | NS        |
| PRL (ng/mL)      | 11.89 (6.42–25.52) 13.89 ± 5.48    | 14.16 (6.42–28.30) 14.68 ± 4.95    | 12.14 (5.55–46.56) 13.91 ± 7.30  | NS        | NS        | NS        |
| TSH (μU/mL)      | 2.01 (0.63–14.40) 2.48 ± 2.52      | n = 58 1.73 (0.84–6.70) 2.07 ± 1.06 | 1.68 (0.71–5.48) 2.11 ± 1.07     | NS        | NS        | NS        |

FSH – follicle-stimulating hormone, LH – luteinizing hormone, LTV – left testis volume, PRL – prolactin, RTV – right testis volume, TSH – thyroid-stimulating hormone, total T – total testosterone. n – number of subjects, NS – no statistical significance, SD – standard deviation. Statistical significance in the Kruskal–Wallis test was reached when p < 0.05.
Table 8. Descriptive statistics and comparison of semen parameters between infertile subjects with SDF >30%, >15–30% and ≤15%.

| Study parameters                                      | Subjects with SDF >30 (1) | Subjects with SDF >15–30% (2) | Subjects with SDF ≤15% (3) | p 1 vs. 2 | p 1 vs. 3 | p 2 vs. 3 |
|-------------------------------------------------------|---------------------------|--------------------------------|-----------------------------|-----------|-----------|-----------|
|                                                        | median (range) mean ± SD  |                                 |                              |           |           |           |
| Semen volume (mL)                                     | 3.50 (1.50–8.50) 3.66 ± 1.78 | 3.00 (0.50–8.00) 3.06 ± 1.47 | 3.00 (1.50–8.00) 3.52 ± 1.72 | NS        | NS        | NS        |
| Sperm concentration (× 10^6/mL)                       | 5.65 (0.40–60.25) 10.90 ± 13.50 | 13.04 (0.80–104.50) 20.40 ± 21.46 | 21.75 (2.50–251.00) 30.22 ± 39.93 | NS        | 0.000175  | NS        |
| Total number of spermatozoa (×10^6)                   | 20.02 (0.88–221.00) 42.42 ± 59.41 | 41.25 (0.84–365.75) 63.75 ± 78.07 | 70.76 (8.10–426.70) 89.77 ± 88.40 | NS        | 0.000841  | 0.025102  |
| Morphologically normal spermatozoa (%)                | 0.00 (0.00–3.00) 0.46 ± 0.83 | 0.00 (0.00–8.00) 1.20 ± 1.74 | 3.00 (0.00–11.00) 3.51 ± 3.11 | NS        | 0.000011  | 0.000497  |
| TZI                                                   | 1.79 (1.36–2.48) 1.82 ± 0.26 | 1.69 (1.35–2.50) 1.82 ± 0.33 | 1.75 (1.34–2.30) 1.77 ± 0.24 | NS        | NS        | NS        |
| Sperm progressive motility (%)                        | 26.00 (6.00–70.00) 28.82 ± 16.00 | 40.00 (0.00–70.00) 38.54 ± 19.72 | 63.00 (15.00–87.00) 59.02 ± 19.86 | NS        | <0.00001  | 0.000016  |
| Sperm nonprogressive motility (%)                     | 4.50 (0.00–10.00) 4.39 ± 2.97 | 6.00 (0.00–29.00) 6.74 ± 5.50 | 4.00 (0.00–13.00) 4.72 ± 2.89 | NS        | NS        | NS        |
| Total sperm motility (%)                              | 31.00 (6.00–71.00) 33.21 ± 17.18 | 48.00 (1.00–77.00) 45.28 ± 20.01 | 68.00 (15.00–90.00) 63.74 ± 19.47 | NS        | <0.00001  | 0.000103  |
| Eosin-negative spermatozoa – live cells (%)           | 67.00 (30.00–86.00) 62.71 ± 15.82 | 74.00 (40.00–86.00) 73.13 ± 9.49 | 86.00 (47.00–96.00) 83.53 ± 8.87 | NS        | <0.00001  | <0.000001 |
| HOS test-positive spermatozoa – live cells (%)        | n = 17 67.00 (26.00–84.00) 65.82 ± 13.67 | n = 49 72.00 (53.00–85.00) 72.97 ± 7.44 | n = 41 83.00 (67.00–92.00) 82.80 ± 7.01 | NS        | <0.00001  | 0.000001  |
| Peroxidase-positive cells (%)                         | 0.25 (0.00–0.25) 0.81 ± 1.90 | 0.25 (0.00–9.50) 0.76 ± 1.41 | 0.25 (0.00–2.00) 0.41 ± 0.51 | NS        | NS        | NS        |

HOS test – hypooptic swelling test, SDF – sperm DNA fragmentation, TZI – teratozoospermia index. n – number of subjects, NS – no statistical significance, SD – standard deviation. Statistical significance in the Kruskal–Wallis test was reached when p < 0.05.
Table 9. Spearman’s rank correlation coefficient ($r_s$) between age, testicular volume and levels of reproductive hormones.

| Study parameters | LTV (mL) | RTV (mL) | FSH (mIU/mL) | LH (mIU/mL) | total T (nmol/L) | PRL (ng/mL) | TSH (μU/mL) | SDF (%) |
|------------------|----------|----------|---------------|-------------|-----------------|-------------|-------------|---------|
| Age (y)          | n = 128  | rs = -0.0320 NS | rs = -0.0767 NS | rs = -0.1203 NS | rs = 0.0657 NS | rs = -0.0599 NS | rs = -0.0783 NS |         |
|                  |          |          |               |             |                 |             |             |         |
| LTV (mL)         | - n = 127 | rs = **0.8259** p < 0.000001 | - n = 127 | rs = **-0.2491** p = 0.004745 | n = 128 | rs = **0.1529** NS | n = 128 | rs = **0.1225** NS |         |
|                  |          |          |               |             |                 |             |             |         |
| RTV (mL)         | n = 127  | rs = **0.2491** p = 0.006525 | - n = 127 | rs = **-0.2402** p = 0.006525 | n = 128 | rs = **0.0524** NS | n = 128 | rs = **0.0720** NS |         |
|                  |          |          |               |             |                 |             |             |         |
| FSH (mIU/mL)     | n = 127  | rs = **0.2491** p = 0.006525 | - n = 127 | rs = **-0.2402** p = 0.006525 | - n = 127 | rs = **0.5018** p < 0.000001 | n = 129 | rs = **0.0524** NS |         |
|                  |          |          |               |             |                 |             |             |         |
| LH (mIU/mL)      | n = 128  | rs = **-0.1529** NS | n = 128 | rs = **-0.1596** NS | rs = **0.5018** p < 0.000001 | rs = **-0.0568** NS | rs = **0.1673** NS | rs = **0.0533** NS |         |
|                  |          |          |               |             |                 |             |             |         |
| total T (nmol/L) | n = 128  | rs = **0.1225** NS | n = 128 | rs = **0.1418** NS | n = 128 | rs = **0.0524** NS | rs = **-0.0703** NS | rs = **0.0102** NS |         |
|                  |          |          |               |             |                 |             |             |         |
| PRL (ng/mL)      | n = 128  | rs = **-0.0195** NS | n = 128 | rs = **-0.0759** NS | rs = **0.1673** NS | rs = **-0.0703** NS | rs = **-0.0892** NS | rs = **0.0802** NS |         |
|                  |          |          |               |             |                 |             |             |         |
| TSH (μU/mL)      | n = 127  | rs = **0.0832** NS | n = 127 | rs = **0.0564** NS | n = 128 | rs = **0.0533** NS | n = 129 | rs = **-0.0892** NS | rs = **0.0321** NS |         |

FSH – follicle-stimulating hormone, LH – luteinizing hormone, PRL – prolactin, RTV – right testis volume, TSH – thyroid-stimulating hormone, total T – total testosterone. n – number of subjects, NS – no statistical significance, SD – standard deviation. Statistical significance was reached when $p < 0.05$. 

Table 9. Spearman’s rank correlation coefficient ($r_s$) between age, testicular volume and levels of reproductive hormones.
| Study parameters | LTV (mL) | RTV (mL) | FSH (mIU/mL) | LH (mIU/mL) | total T (nmol/L) | PRL (ng/mL) | TSH (μU/mL) | SDF (%) |
|------------------|---------|---------|-------------|-------------|-----------------|-------------|-------------|---------|
| **Semen volume (mL)** | n = 128 | n = 128 | n = 129 | n = 129 | n = 129 | n = 129 | n = 129 | n = 129 |
| r_s = 0.0376 | r_s = -0.0524 | r_s = -0.1285 | r_s = -0.0994 | r_s = -0.0010 | r_s = -0.1236 | r_s = -0.0940 | r_s = -0.0010 | r_s = -0.0265 |
| NS | NS | NS | NS | NS | NS | NS | NS | NS |
| **Sperm concentration (× 10^9/mL)** | n = 128 | n = 128 | n = 129 | n = 129 | n = 129 | n = 129 | n = 129 | n = 129 |
| r_s = 0.4345 | r_s = 0.4019 | r_s = -0.2205 | r_s = -0.0564 | r_s = -0.0760 | r_s = -0.3461 | r_s = -0.3343 | r_s = -0.000055 | r_s = p < 0.00001 |
| p = 0.000001 | p = 0.000603 | p = 0.011703 | p = 0.035873 | p = 0.000005 | p = 0.000101 | p = 0.00001 | p = 0.00000055 | p = 0.0000001 |
| **Total number of spermatozoa (× 10^6)** | n = 128 | n = 128 | n = 129 | n = 129 | n = 129 | n = 129 | n = 129 | n = 129 |
| r_s = 0.4191 | r_s = 0.3452 | r_s = -0.2350 | r_s = -0.1067 | r_s = -0.1011 | r_s = -0.3461 | r_s = -0.3343 | r_s = -0.000055 | r_s = p < 0.00001 |
| p = 0.000001 | p = 0.000603 | p = 0.007108 | p = 0.000005 | p = 0.000101 | p = 0.000101 | p = 0.00000055 | p = 0.0000001 | p = 0.0000001 |
| **Morphologically normal spermatozoa (%)** | n = 128 | n = 128 | n = 129 | n = 129 | n = 129 | n = 129 | n = 129 | n = 129 |
| r_s = 0.1634 | r_s = 0.1579 | r_s = -0.0681 | r_s = -0.0269 | r_s = -0.1368 | r_s = -0.0102 | r_s = -0.0553 | r_s = -0.0470 | r_s = -0.1966 |
| p = 0.000001 | p = 0.000003 | p = 0.035873 | p = 0.000005 | p = 0.000101 | p = 0.000001 | p = 0.000001 | p = 0.000001 | p = 0.000001 |
| **TZI** | n = 128 | n = 128 | n = 129 | n = 129 | n = 129 | n = 129 | n = 129 | n = 129 |
| r_s = 0.0062 | r_s = 0.0549 | r_s = 0.0235 | r_s = -0.0940 | r_s = -0.1121 | r_s = 0.0066 | r_s = -0.1199 | r_s = -0.5476 | r_s = p < 0.000001 |
| NS | NS | NS | NS | NS | NS | NS | NS | NS |
| **Sperm progressive motility (%)** | n = 128 | n = 128 | n = 129 | n = 129 | n = 129 | n = 129 | n = 129 | n = 129 |
| r_s = 0.2048 | r_s = 0.1313 | r_s = -0.0940 | r_s = -0.0095 | r_s = -0.1121 | r_s = -0.1199 | r_s = -0.5476 | r_s = p < 0.000001 | r_s = p < 0.000001 |
| p = 0.020393 | p = 0.000603 | p = 0.035873 | p = 0.000005 | p = 0.000101 | p = 0.000101 | p = 0.00000055 | p = 0.0000001 | p = 0.0000001 |
| **Sperm nonprogressive motility (%)** | n = 128 | n = 128 | n = 129 | n = 129 | n = 129 | n = 129 | n = 129 | n = 129 |
| r_s = 0.0562 | r_s = -0.0665 | r_s = 0.0059 | r_s = -0.0287 | r_s = 0.0462 | r_s = 0.1629 | r_s = -0.0553 | r_s = 0.0235 | r_s = -0.0102 |
| NS | NS | NS | NS | NS | NS | NS | NS | NS |
| **Total sperm motility (%)** | n = 128 | n = 128 | n = 129 | n = 129 | n = 129 | n = 129 | n = 129 | n = 129 |
| r_s = 0.2115 | r_s = 0.1236 | r_s = -0.1126 | r_s = -0.0096 | r_s = -0.1008 | r_s = -0.1180 | r_s = -0.5374 | r_s = p < 0.000001 | r_s = p < 0.000001 |
| p = 0.016524 | p = 0.016524 | p = 0.016524 | p = 0.016524 | p = 0.016524 | p = 0.016524 | p = 0.016524 | p = 0.016524 | p = 0.016524 |
| **Eosin-negative spermatozoa – live cells (%)** | n = 128 | n = 128 | n = 129 | n = 129 | n = 129 | n = 129 | n = 129 | n = 129 |
| r_s = 0.1956 | r_s = 0.0945 | r_s = -0.0470 | r_s = 0.1652 | r_s = -0.0577 | r_s = -0.0247 | r_s = 0.0102 | r_s = 0.0102 | r_s = 0.0102 |
| p = 0.026909 | p = 0.026909 | p = 0.026909 | p = 0.026909 | p = 0.026909 | p = 0.026909 | p = 0.026909 | p = 0.026909 | p = 0.026909 |
| **HOS test-positive spermatozoa – live cells (%)** | n = 105 | n = 105 | n = 106 | n = 107 | n = 107 | n = 107 | n = 107 | n = 107 |
| r_s = 0.1904 | r_s = 0.0934 | r_s = -0.0359 | r_s = -0.0147 | r_s = -0.0147 | r_s = -0.0147 | r_s = -0.0147 | r_s = -0.0147 | r_s = -0.0147 |
| NS | NS | NS | NS | NS | NS | NS | NS | NS |
| **Peroxidase-positive cells (%)** | n = 128 | n = 128 | n = 129 | n = 129 | n = 129 | n = 129 | n = 129 | n = 129 |
| r_s = 0.1248 | r_s = 0.1629 | r_s = 0.0387 | r_s = 0.1138 | r_s = -0.0247 | r_s = 0.1441 | r_s = -0.0553 | r_s = -0.0553 | r_s = -0.0553 |
| p = 0.026503 | p = 0.026503 | p = 0.026503 | p = 0.026503 | p = 0.026503 | p = 0.026503 | p = 0.026503 | p = 0.026503 | p = 0.026503 |

FSH – follicle-stimulating hormone, HOS test – hypo-osmotic swelling test, LH – luteinizing hormone, LTV – left testis volume, SDF – sperm DNA fragmentation, PRL – prolactin, RTV – right testis volume, TSH – thyroid-stimulating hormone, total T – total testosterone, TZI – teratozoospermia index, n – number of subjects, NS – no statistical significance, SD – standard deviation. Statistical significance was reached when p < 0.05.
On the other hand, we did not observe any significant differences between men with SDF >30% and men with SDF >15–30% in any study parameters. Additionally, in the case of age, hormone levels (FSH, LH, PRL, total T, TSH), TZI index, sperm nonprogressive motility and concentration of peroxidase-positive cells in semen, no significant differences between the compared three groups were recorded (Tables 7, 8).

Spearman’s rank correlations between the study parameters

Correlations between SDF, male age, basic semen parameters, testicular volume and hormone levels

Analysis of the Spearman’s rank correlation coefficient showed a linear relationship between SDF and selected parameters. SDF was negatively correlated with sperm concentration ($r_s = -0.3461$; weak dependence), total sperm count ($r_s = -0.3343$; weak dependence), sperm morphology ($r_s = -0.4482$; moderate dependence), progressive motility ($r_s = -0.5476$; moderate dependence), total motility ($r_s = -0.5374$; moderate dependence) and vitality – eosin-negative sperm cells ($r_s = -0.6389$; moderate dependence) and HOS test-positive sperm cells ($r_s = -0.5811$; moderate dependence). In turn, there were no significant correlations between SDF and age, ejaculate volume, nonprogressive sperm motility or peroxidase-positive cell concentration. Moreover, a negative correlation between SDF and the volume of the left testis was found ($r_s = -0.2055$; weak dependence), whereas there were no significant correlations between SDF and the volume of the right testis or study hormone levels (Tables 9, 10).

Correlations between testicular volume, hormone levels and basic semen parameters

In the examined group, the left and right testis volumes were negatively correlated with the level of FSH ($r_s = -0.2491$ and $r_s = -0.2402$, respectively; weak dependences) but was not correlated with other hormones (LH, PRL, total T, TSH). Moreover, the volumes of the left and right testes were positively correlated with sperm concentration ($r_s = 0.4345$ and $r_s = 0.4019$, respectively; moderate dependences) and total sperm number ($r_s = 0.4191$ and $r_s = 0.3455$, respectively; moderate and weak dependences). Additionally, positive correlations between left testis volume and sperm progressive motility ($r_s = 0.2048$) as well as total motility ($r_s = 0.2115$; weak dependence) were found. Furthermore, the LH level was negatively correlated with sperm concentration ($r_s = -0.2205$; weak dependence) and total sperm count ($r_s = -0.2350$; weak dependence), but there were no other significant correlations between the levels of assessed hormones and conventional semen parameters (Tables 9, 10).

The raw data can be found as Underlying data.109,110

Discussion

Reduced basic semen parameters can result from impaired spermatogenesis

Based on the obtained data, it can be suggested that the failure to become a biological father could be due to disorders of spermatogenesis manifested by reduced standard seminological parameters. It is worth noting that in our study, the median morphologically normal sperm was only 1%, and as many as 100 out of 130 infertile men had teratozoospermia (isolated or coexisting with other semen disorders). Additionally, studies conducted by other authors confirm the relationship between standard sperm parameters and male fertility.38–40 Slama et al.40 proved a significantly shorter time to pregnancy (TTP) in women whose partners had a higher percentage of sperm with normal morphology. Moreover, it was found that the percentage of morphologically normal sperm was decreased in men from couples with a history of recurrent miscarriage.31–44 Additionally, morphologically normal sperm cells play an important role not only in the case of natural conception but also in medically assisted conception (IUI, fertilization in vitro),45 and it has been shown that sperm morphology may also influence embryo development.43 On the other hand, reproductive success might be achieved even when morphologically normal sperm cells are not observed in the semen. Shabtaie et al.44 emphasize that in the case of only abnormal sperm morphology (assuming no female infertility factor), first-line therapy should not assist ART without undertaking a sufficiently long attempt at natural conception. Therefore, opinions about the predictive value of sperm morphological assessment for both natural conception and medically assisted conception are controversial.45–49

Additionally, in our study, 46 cases of asthenozoospermia (isolated or coexisting) and 58 cases of oligozoospermia (isolated or coexisting) were observed. Many authors confirm that progressive motility is one of the most important parameters influencing reproductive success both in terms of natural conception and medically assisted conception.38,40,50,51 Furthermore, Lotti et al.52 revealed a negative correlation between sperm vitality and TTP. Also, analyzing a large group of infertile men and men from the control group (candidates to be sperm donors), Li et al.53 showed that in the first group, there were significantly more men with azoospermia, asthenozoospermia and oligoasthenozoospermia, whereas surprisingly isolated oligozoospermia were detected more often in men from the control group. Some authors even suggest greater clinical implications of the total sperm count in relation to the sperm concentration.
However, it should be emphasized that low standard seminological parameters are not always synonymous with infertility status. Not all authors recognize the arbitrary division of men into fertile and infertile groups based only on the basic semen characteristics according to the WHO. Therefore, except for azoospermia, necrozoospermia, and globo- and macrozoospermia, it is difficult to determine male fertility potential considering only standard seminological parameters. This thesis is also confirmed by our previous studies, in which 19 cases of reduced basic semen parameters were found in a group of men with proven fertility (n = 64). Therefore, in this research, the standard seminological assessment was not only one criterion for qualifying a man as infertile. The patient's infertility was established by an interview indicating unsuccessful attempts for offspring during one year of regular intercourse without the use of contraception.

Relationships between testicular volume, hormone levels, basic semen parameters and sperm genome integrity

The results of our research suggest the coexistence of spermatogenesis disorders with a reduced testicle volume and a higher FSH level. It is known that the process of spermatogenesis, reflecting testicular function, depends on a hypothalamic–pituitary–gonadal axis function, in which gonadotropins LH and FSH play a key role in maintaining testosterone biosynthesis and the function of the seminiferous epithelium, respectively. In addition, it has been proven that the function of the male gonad is influenced by thyroid hormones and prolactin. Importantly, a significant decrease in testicular volume can be associated with both reduced hormonal activity (lower levels of androgens) and reproductive activity manifested by seminiferous tubule atrophy. Therefore, our study included an evaluation of not only standard seminological parameters but also testicular volume and reproductive hormone levels (FSH, LH, PRL, total T, TSH). It should be noted that the selection of the assessed hormones was based on data from the literature.

Unfortunately, to date, there have been no strict guidelines regarding the hormonal test profile that should be determined in the routine diagnosis of male infertility. On the other hand, the European Academy of Andrology (EAA) in guidelines from 2018 postulates evaluation of total T, FSH and LH in every case of an infertile man with oligoasthenoteratozoospermia (OAT). These recommendations are in line with the guidelines of the European Society Urology (EAU) from 2021. However, it is believed that the remaining hormonal tests should be performed based on an individual assessment of the patient. The levels of commonly recognized markers of spermatogenesis and Sertoli cell function (FSH, inhibin B) have been most frequently studied in the available literature. In addition, the authors of the study also paid attention to the analysis of the levels of SHBG, prolactin, estradiol, TSH, cortisol, growth hormone (GH) and insulin-like growth factor 1 (IGF-1).

Based on testicular volume measurement in our study, two groups of infertile participants were distinguished: men with a volume of at least one testis below the norm (<12 mL) and men with a normal volume of both testes. Our results showed that infertile men with a reduced volume of at least one testis had a significantly higher FSH level and a lower sperm count, sperm morphology, motility and vitality. Moreover, it should be especially highlighted that we found a significantly increased fragmentation of sperm nuclear DNA in the first group. These results were confirmed by correlation analysis. The testicular volume was negatively correlated with the level of FSH and the SDF value and positively correlated with sperm morphology, motility and vitality. Moreover, it should be especially highlighted that we found a significantly increased fragmentation of sperm nuclear DNA in the first group. These results were confirmed by correlation analysis. The testicular volume was negatively correlated with the level of FSH and the SDF value and positively correlated with the number and motility of sperm. Surprisingly, we did not find an association between testicular volume and total T level.

The obtained findings were partially consistent with the data published by other authors. Numerous researchers have reported relationships between testicular volume, conventional semen parameters, gonadotropin and testosterone levels as well as the results of advanced sperm tests (chromatin status, mitochondrial potential, apoptosis). The coexistence of reduced standard semen parameters, decreased testosterone levels and testicular volume presented by Bahk et al. and Condorelli et al. suggest that the reduction of testicular volume may be associated not only with impaired spermatogenesis but also with decreased hormonal function of male gonads. Condorelli et al. recommend periodic assessment of testosterone levels for patients with hypotrophic gonads. On the other hand, the obtained results presented by other authors are not always unambiguous. For example, in contrast to our results, Condorelli et al. revealed a relationship between testicular volume and testosterone levels, but they did not find a correlation between testicular volume and gonadotropin levels.

As mentioned above, we discovered that a group of men with at least one testis volume <12 mL had significantly reduced integrity of the sperm genome. The data could suggest that spermatogenesis disorders coexist with decreased testicular volume and are manifested not only by reduced conventional sperm parameters but also by molecular disorders of sperm chromatin. The relationship between testicular volume and sperm DNA strand brakes was also confirmed by our other findings. The participants with a high level of SDF (>30%) had significantly smaller testes than men with a low level of SDF (≤15%). Moreover, we noted a negative correlation between testicular volume and sperm chromatin fragmentation. Similar results were obtained by other authors who observed a negative correlation between the fragmentation of nuclear sperm DNA (verification using the TUNEL method), its denaturation (verification using acridine orange), sperm chromatin density (verified using propidium iodide) and the volume of testes.
In the next step of our research, we compared two groups of subjects: men with abnormal levels of at least one of the assessed hormones and men with normal hormonal profiles. The obtained findings provided nonobvious data. We noted that in the first group, sperm count, morphology and motility were reduced, but testicular volume did not differ significantly between the two groups. Moreover, the LH level was negatively correlated with the total sperm count. Additionally, other authors have confirmed the statistical relationship between the level of selected hormones and standard seminological parameters.65,79,80,81,82,83,84 Wei et al.80 showed that in patients with OAT, total T was positively correlated with sperm morphology, whereas PRL was correlated with sperm concentration and motility. Moreover, Lu et al.79 and Uhler et al.83 revealed a negative correlation between FSH level and semen volume, sperm concentration, morphology and motility as well as between LH level and sperm concentration in infertile men or healthy volunteers.

It should be pointed out that we did not find significant differences in the percentage of sperm cells with fragmented DNA between participants with abnormal levels of at least evaluated hormones and those with normal hormonal profiles. This comparative analysis was consistent with Spearman’s rank correlation test, which did not show a significant correlation between the SDF value and the level of the assessed hormones. However, other authors’ data indicated statistical relationships between sperm chromatin quality and the hormonal profile.79,81,82 The coexistence of sperm DNA fragmentation with abnormally high or low levels of gonadotropins was shown in research published by Wdowiak et al.82 These results were partially consistent with the studies of Lu et al.83 and Smit et al.81 who showed a negative correlation between sperm DNA fragmentation and elevated levels of FSH and LH. In turn, the association between sperm nuclear DNA damage and testosterone level is not always unequivocal. Some researchers Wdowiak et al.82 have found a negative correlation between these parameters, whereas others did not confirm these findings.81,83

The open question is why there was no statistically significant difference in our research in the percentage of SDF between the groups of men differing in the level of at least one of the assessed hormones. There is no doubt that the obtained results could have been influenced by the limited number of infertile men (n = 130) enrolled in our study and the hormonal heterogeneity of the group of men with abnormal levels of at least one of the verified hormones. Disturbances in the level of hormones can be both a factor influencing infertility and a consequence of such a state. In other words, an abnormal hormonal profile can be responsible for reduced semen quality or may be only a secondary effect of pathological processes in testes. In addition, it should be emphasized that there are many potential factors (e.g., obesity, occupational exposure, comorbidities, age, pharmacotherapy, stress) that may affect the interrelationship between spermatogenesis and hormone levels.84–90

**Sperm genome integrity is a key point for male fertility**

It can be assumed that in our investigated group of infertile patients, one of the major factors that limited the ability of male gametes to fertilize was probably an increased level of sperm nuclear DNA fragmentation. It was found that the group of men with SDF >30% had a significantly reduced sperm count, morphology, motility and vitality in comparison to infertile men with a normal SDF rate of ≤15%. Similarly, Erenpreiss et al.77 showed that if males had diagnosed astheno- and teratozoospermia, the odds ratio (OR) for >20% DFI or for >30% DFI was 1.9–4.0-fold higher or 2.8–6.2-fold higher, respectively, than in subjects with normal sperm motility and morphology. Additionally, Vinnakota et al.29 observed a decrease in sperm motility in participants with SDF >30%. Moreover, we showed that the level of SDF was negatively correlated (Spearman’s rank correlation test) with sperm count, morphology, motility and vitality, and our findings have been confirmed by the research of other authors.30,83,92,93 However, some researchers have not always found a correlation between SDF and basic sperm parameters.94–96

Importantly, it should be highlighted that in our study, the median SDF was 20%. In fact, according to the manufacturer of the Halosperm G2® kit, these results are in the normal range (SDF below 30%). It seems that the threshold of 30% SDF is too high (risk of a false-negative result). In our previous publications, we demonstrated that the median SDF in the group of men with confirmed fertility and/or with high reproductive potential (healthy volunteers with normozoospermia) ranged from 12% to 14%.33–36 In addition, these studies also showed a significantly satisfactory predictive value of the sperm chromatin dispersion (SCD) test to discriminate males with normal reproductive potential from those with reduced reproductive potential (based on receiver operating characteristic [ROC] analysis). The cut-off value was 18% and 20% SDF.34–36 Moreover, we obtained a threshold of 18% SDF to distinguish infertile men from fertile men (unpublished data).

These observations were in agreement with other authors who also clearly showed that the level of sperm nuclear DNA fragmentation was correlated with male infertility and that the acceptable threshold for sperm genome fragmentation was not below 30% but rather below 20%.34,36,45,46,103,104,105,106,107 For example, Bungum et al.37 showed that in the case of subjects with sperm nuclear DNA fragmentation in the range of 0–20%, the chance of spontaneous pregnancy is constant, but an increase in sperm DNA fragmentation >20% is associated with a reduced ability to achieve pregnancy. Moreover, Majzoub et al.108 estimated that the mean value of SDF for fertile subjects was 15.68%, whereas in the infertile group,
it was 27.60%. In turn, comparing the groups of fertile and infertile men, Wiweko and Utami found not only significant differences in the SDF value between the study groups (medians: 19.90% vs. 29.95%, respectively) but also reported that SDF at the cutoff point of 26.1% had a higher diagnostic value. Similar results were presented by Javed et al. (the SDF at the cutoff point was 24.47%). Moreover, Evenson emphasized that the percentage of sperm with damaged chromatin >15–25 could increase the risk of male infertility and that >20–35% spermatozoa with damaged DNA could significantly reduce the chances of becoming pregnant using in vitro fertilization. Therefore, based on own research and analysis of the results of other researchers, Evenson concluded that when the percentage of sperm with abnormal chromatin status was >20, male fertility was decreased, and in vitro fertilization as first-line therapy should be considered. These conclusions were also confirmed by Giwercman et al., who performed a comparison of ORs for the occurrence of infertility depending on the percentage of DFI. The authors showed that in the group of men with DFI 10–20%, the risk of reproductive failure was higher (OR = 2.5) than that in men with DFI <10%. In addition, Giwercman et al. observed a significant increase in the risk of infertility (OR = 8.4) in men with DFI> 20% compared to men with DFI <10%. Finally, in two most recent publications both Esteves et al. and Agarwal et al. reported that cut-off point of 20% sperm cells with fragmented DNA (verified both by SCSA, TUNEL and SCD assay) is the best criterion to discriminate fertile men from infertile.

Additionally, the influence of DFI on the fertilization process has been confirmed. Simon et al. revealed a higher risk (OR = 9.5) of a low percentage of fertilized oocytes (<40% fertilized oocytes) when men had DFI >40% compared to men with DFI ≤40%. Therefore, we can assume that sperm chromatin abnormalities may be accompanied by lowered standard sperm parameters synergistically affecting male fertility.

Conclusions

Our comprehensive assessment of male infertility factors allowed us to conclude that in the study clinical cases, spermatogenesis disorders coexisted with decreased testicular volume and increased FSH levels. Moreover, they were manifested not only by reduced basic sperm characteristics but also, very importantly, by a high level of sperm nuclear DNA damage, which has great clinical utility both in terms of natural conception and in terms of ART (Figure 1). Furthermore, our current and previous findings suggest that the cut-off value of 30% SDF given by manufacturer of the Halosperm G2® kit seems too high and should be revised/downgraded to 20%, for better prognosis of male fertility. What’s more, clarification of the relationship between standard semen parameters, testicular volume, levels of reproductive hormones, SDF and clinical features might help to develop new personalized strategies for therapeutic interventions. In the case of infertile men, a complete andrological examination including in-depth medical interview, physical examination, standard semen analysis, scrotal ultrasound, assessment of reproductive hormones and integrity of sperm genome is justified. This medical approach is necessary not only due for verification of the causes of infertility but also due to the need to detect serious health disorders that may be life-threatening. For example, it has been proven that infertile men have an increased risk of testicular cancer, which determines the recommendation of periodic ultrasound examinations of the scrotum and gonadal self-examination. Therefore, the introduction of a complex diagnosis of male infertility factors is justified and needed.

Limitations of the study

Some limitations of our study should be addressed. One of the most important factors influencing our results is the limited number of participants. In total, 130 men from couples with confirmed infertility were included to this project. It is known that the most reliable data are obtained from well-designed studies on large cohorts of patients. Due to the limited number of participants in our research, the presented results should be approached critically, and it should be borne in mind that studies conducted on a larger group could provide different results and conclusions. Moreover, the number of compared men in particular groups was not equal, which may affect the obtained statistical differences between groups. In the assessed hormonal profile, we did not include the determinations of some markers which could also be important for assessing the status of male fertility (i.a. inhibin B, SHBG, GH, estradiol, cortisol). Finally, sperm chromatin dispersion (SCD) test was performed to assess SDF. This test is a standardized diagnostic method but often not considered the gold standard for sperm DNA assessment because it does not directly evaluate breaks of DNA.

Data availability

Underlying data

Zenodo: Evaluation of selected semen parameters and biomarkers of male infertility – preliminary study. https://doi.org/10.5281/zenodo.6536196.

This project contains the following underlying data:

- Kups et al. for database.xlsx (raw data)
Zenodo: Evaluation of selected semen parameters and biomarkers of male infertility – preliminary study. [10]

This project contains the following underlying data:

- Raw microphotographs

Extended data

Zenodo: Evaluation of selected semen parameters and biomarkers of male infertility – preliminary study. [11]

This project contains the following extended data:

- Urological and andrological medical interview Michal Kups.pdf (Patient card used during the medical interview)

Data are available under the terms of the Creative Commons Attribution 4.0 International license (CC-BY 4.0).

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111. Kups M, Gill K, Rosiak-Gill A, et al.: Evaluation of selected semen parameters and biomarkers of male infertility - preliminary study. [Data set]. Zenodo. 2022. Publisher Full Text
Open Peer Review

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Version 1

Reviewer Report 20 February 2024

https://doi.org/10.5256/f1000research.133505.r238146

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Eva Tvrdá
Slovak University of Agriculture, Nitra, Slovakia

This is a nicely written and appropriately designed study evaluating and interconnecting standard semen parameters in infertile men. While I appreciate papers that turn "back to the basics" and re-evaluate/re-verify the importance of traditional parameters of sperm quality, I am missing the originality of the study. The authors should place more emphasis on the originality and scientific rigour of their experiments as well as provide a solid justification as to why such types of papers on generally well-accepted knowledge are necessary.

Finally, the title of the article hints evaluation of biomarkers of male infertility. If so, what are the biomarkers and why should these be more of a center focus for the andrologists?

Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Yes
**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Sperm quality, oxidative stress, antioxidants, molecular andrology.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

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**Reviewer Report 20 February 2024**

https://doi.org/10.5256/f1000research.133505.r238141

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Nafisa Balasinor

ICMR-National Institute for Research in Reproductive and Child Health, Parel, India

In the present manuscript the authors have studied the association between standard semen parameters, testicular volume, reproductive hormones profile and the sperm nuclear DNA fragmentation.

Overall the study is well done and has clinical utility. However, there are few points which needs to be addressed:

1. How was the sample size of 130 calculated?
2. Was infertility due to known female factor ruled out?
3. The authors should mention how they categorized abnormal hormonal levels.
4. Table 3: volume in the right testis of normal volume group ranged between 9 to 25ml. However, testicular volume of > or equal to 12 was taken as normal testicular volume. Please check the data.
5. Table 5: Men with abnormal hormonal profile group had increase in all hormones except Testosterone. However, the values do not indicate so. Hence need to check if correct statistical tool was used.
6. The authors need to check testosterone levels in this infertile group as compared to fertile group or normal range.
7. SDF in men with normal semen and sperm parameters should be analyzed. If SDF is found in this group too than the importance to including SDF in male infertility work out will be evident.
8. It would be interesting to see the sperm parameters in individuals in which all 3 parameters, namely, Volume, hormone profile and SDF.

Is the work clearly and accurately presented and does it cite the current literature?

Yes

Is the study design appropriate and is the work technically sound?

Partly

Are sufficient details of methods and analysis provided to allow replication by others?
Partly

**If applicable, is the statistical analysis and its interpretation appropriate?**
I cannot comment. A qualified statistician is required.

**Are all the source data underlying the results available to ensure full reproducibility?**
Yes

**Are the conclusions drawn adequately supported by the results?**
Partly

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Reproductive endocrinology, male fertility and epigenetics

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

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Abdul S Ansari  
Department of Zoology, University of Rajasthan, Jaipur, Rajasthan, India

In the present manuscript the author(s) evaluated etiopathogenesis of male infertility by investigation of the relationship between standard semen parameters, testicular volume, levels of reproductive hormones and the fragmentation of sperm nuclear DNA (SDF). A total of 130 subjects, divided into three groups, with at least one abnormal testis volume, minimum low level among any reproductive hormones and with SDF. Results obtained from the study revealed that the subjects with decreased testicular volume and abnormal levels of hormones were observed with decreased basic semen parameters. Abnormal testicular volume also had, respectively, higher percentage of SDF and enhanced levels of FSH. Subjects with high SDF level had low testicular volume and seminal parameters. Authors concluded that spermatogenesis disorders coexisted with decreased testicular volume and increased FSH levels which manifested by reduced seminal characteristics and high sperm nuclear damage. However, the research article contains several lacunae in terms of writing the manuscript, language, presentation of data, etc. Moreover, the title of the article does not match the contents of the article. It correlated several parameters.

**Specific comments:**

**Abstract:**
1. Page 1: Methods: The participants of the study were not patients, instead they are subjects.
Modify the sentence as “Participant subjects (n=130) were divided into”.

**Introduction:**
1. Page 3, Introduction: Write full form of SCSA test followed by its abbreviation SCSA in parentheses.
2. Page 3, Methods (Study population): The study population, i.e., 130 male infertile subjects is very small. If the number would have been more, it is easy to draw a conclusion.
3. Page 3, Methods (Study population): Replace “All patients…” with “All subjects…”.
4. Page 3, Methods (Study population): How many subjects with azoospermia and cryptozoospermia were excluded? Mention their number.
5. Page 3, Methods (Study population): Replace the phrase “seminal cords” with “spermatic cords”.
6. Page 4, Methods (Conventional semen analysis): Room temperature is not -20 °C. Correct the sentence.
7. Page 4, Methods (Conventional semen analysis): Add “liquefaction time” of semen as a macroscopic semen analysis parameter.
8. Page 4, Methods (Conventional semen analysis): The terminology total sperm count has changed by WHO as “sperm density” (million/ejaculate). Replace the same with new terminology.
9. Page 4, Methods (Conventional semen analysis): The sperm vitality or viability is carried out the mixing of a well mixed semen sample with eosin-nigrosin stain and and spreaded on a slide and dried. The unstained spermatozoa are regarded as alive/vital, while pink coloured sperms are considered dead. The HOS test is done with the mixing of a semen sample with a hypoosmotic solution and incubated. After incubation spermatozoa with coiled tails are counted, and calculated in percent (%), termed as HOS positive sperms. How vitality indicating both eosin-positive cells and hypoosmotic-reactive cells (HOS test-positive cells can be assessed?
10. Page 4, Methods (Sperm Chromatin Dispersion (SCD) test): The chromatin dispersion test carried out in the study with Halosperm G2 kit is sort of In vitro Nuclear Chromatin Decondensation (NCD) Test of WHO routinely carried out for infertility diagnosis for fertilizing ability. In the test Sperm heads decondensed which results into swelling of heads, appears as, halos. Whereas, Non-fertilizable sperm heads remain condensed due to defective nuclear chromatin. Decondensed sperms heads are counted versus condensed sperm heads and represented as percent. I am surprised that why so expensive SCD test was carried out when other several simple tests are available. A detailed comment is required from author(s) on this point.
11. Page 4, Methods (Hormone profile of infertile subjects): Add Inter-assay and Intra-assay coefficient of variation of each hormone assayed.

**Results:**
1. Page 6, Seminological Characteristics of study population: Replace the word “Seminological” with “Seminal”.
2. Page 6, Comparison ... (each $\geq$ 12 mL): Write P (significant) value of FSH.

**Discussion:**
1. Page 14, Para 1, Line 5: This is not a thesis, instead a study or investigation. Change the phrase “This thesis” with an appropriate phrase.
2. Page 14, Para 3, Line 2: Complete the word “norm” as “normal”.
3. The write-up of discussion is too lengthy, reduced the same to half length.

4. Conclusions:
5. A brief conclusion of the study carried out similar with Abstract is required.

6. **Data availability:**

7. Replace the word “project” with “research article”.

8. **References:**

9. Reference Nos. 27, 51: Italicize the words “in Vitro”.

10. Reference 59: The title of the reference in upper (capital) case. Change the same as per Instruction to Authors.

11. References 109 to 111 are related to the present research article, thus should be deleted from the list of References.

12. **Tables:**

13. Table 1: This table contains all assayed hormones in the study and well described in the text, therefore, should be deleted.

14. Tables 2, 4, 6, 8, 10: In all these tables data on Sperm nonprogressive motility (%) have been depicted. When data on Sperm progressive motility (%) are presented, there is no use of depicting this. The rest of Sperm progressive motility (%) will obviously related to the Sperm nonprogressive motility (%).

15. The present study contains a total 10 tables. Presentation of data in the present research article merely only in tables is not so impressive and very hard to understand by readers. Prepare histograms or bar diagrams for depiction of data.

16. Table 10: The data of this table should be depicted with Scatter diagrams.

**References**

1. Kups M, Gill K, Rosiak-Gill A, Harasny P, et al.: Evaluation of selected semen parameters and biomarkers of male infertility – preliminary study. *F1000Research*. 2022; 11. [Publisher Full Text]

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**Is the work clearly and accurately presented and does it cite the current literature?**

Yes

**Is the study design appropriate and is the work technically sound?**

Yes

**Are sufficient details of methods and analysis provided to allow replication by others?**

Yes

**If applicable, is the statistical analysis and its interpretation appropriate?**

I cannot comment. A qualified statistician is required.

**Are all the source data underlying the results available to ensure full reproducibility?**

Yes

**Are the conclusions drawn adequately supported by the results?**

Partly

**Competing Interests:** No competing interests were disclosed.
Reviewer Expertise: Reproductive physiology, Male contraception, infertility diagnosis, cancer diagnosis, endocrinology

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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