

Standards in semen examination: publishing reproducible and reliable data based on high-quality methodology

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ABSTRACT: Biomedical science is rapidly developing in terms of more transparency, openness and reproducibility of scientific publications. This is even more important for all studies that are based on results from basic semen examination. Recently two concordant documents have been published: the 6th edition of the *WHO Laboratory Manual for the Examination and Processing of Human Semen*, and the International Standard ISO 23162:2021. With these tools, we propose that authors should be instructed to follow these laboratory methods in order to publish studies in peer-reviewed journals, preferable by using a checklist as suggested in an Appendix to this article.

Key words: reproducibility / basic semen examination / standardized laboratory procedures / andrology / reproductive medicine / laboratory training / quality control / patient security / science development / journal requirements

Appeal to the scientific society involved in Andrology and Reproductive Medicine

As scientists are aware, there has been much discussion about the transparency, openness and reproducibility of science. This is not a new issue. Ten years ago, Begley and Ellis proposed a series of recommendations to improve the reliability of studies in preclinical cancer research (Begley and Ellis, 2012) that helped to initiate a series of developments to address and improve reproducibility. These have included more detailed reporting and transparency of methods such as the STAR Methods for *Cell Press* journals <https://www.cell.com/star-authors-guide>. Concomitant with these developments, national programmes, such as The MDAR (Materials Design Analysis Reporting) Framework, for transparent reporting in the life sciences have been launched (Macleod et al., 2021) and specific consortia have been developed to repeat key published experiments, e.g. Reproducibility Project: Cancer Biology (RP: CB) (<https://elifesciences.org/collections/9b1e83d1/reproducibility-project-cancer-biology>) (Rodgers and Collings, 2021). Furthermore, there are significant resources available such as EQUATOR guidelines (<https://www.equator-network.org/>). The clear direction of travel is to improve standards and have transparent reporting of research (Amara, 2022). There are challenges, however. For example, in the RP:CB project, insufficient information was available to repeat selected experiments published in high impact journals. Furthermore, in the experiments that could be repeated (50/193), fewer than half yielded similar results. As such, the final report of the RP: CB consortia suggested that 'it is hard to assess whether reported findings are credible' (Errington et al., 2021).

In our own discipline of Andrology and Reproductive Medicine, there is a plethora of evidence to show that using non-standardized methods produces unreliable data including, for example, for human sperm concentration and sperm motility assessments. This has created significant problems for the field, making it difficult to determine the scientific accuracy of many studies and ultimately establish their real

clinical and public health impact. A recent example of this is the study of Campbell et al. where they updated the World Health Organization (WHO) semen analysis distribution values (Campbell et al., 2021). The authors reported several challenges in obtaining key information of the quality of the semen examination methods used across the studies being considered for inclusion. Standardization of semen examination has been a long-standing issue that the profession has collectively failed to address, despite the availability of proven accurate methods and robust training approaches (Björndahl et al., 2002, 2016; Barratt et al., 2011; Carrell and De Jonge, 2016; Cairo Consensus Workshop, 2020). Too many studies depending on semen analysis derived data continue to demonstrate poor robustness and rigour in semen analysis methodology (Serrano et al., 2014). When methods with a high degree of uncertainty are used, differences between normal and pathological conditions are likely to be impossible to discover since each observation, burdened by large variability due to measurement uncertainty, will have a more-or-less random result. This will cause considerable overlap in results from the different populations, making them practically inseparable.

The question for all of us working in Andrology, including Editors of journals publishing research in this field, is: What can be done to improve the situation? We believe there is currently a window of opportunity for action. The recent publication of ISO Standard 23162 for the basic examination of human semen (International Organization for Standardization, 2021) finally means that the field has *de facto* reference methods. These methods form the basis of those recommended in the new 6th edition of the WHO andrology laboratory manual (World Health Organization, 2021), which contains simple to follow and proven high-quality methods for semen examination. We propose that authors should be instructed to follow these laboratory methods in order to publish studies in peer-reviewed journals. To facilitate this, we present in the Appendix an author checklist, modified from Björndahl et al. (2016), which authors can complete and submit with their manuscript, making it simple for the journals, reviewers and readers alike to assess the quality of the semen assessment methods used,

and hence of the data being reported. We suggest that any deviation from the checklist, for purposes of testing a new reagent, different method or procedure for improving on the performance of a current recommendation, should be specified and measured against those in the checklist. If authors did not follow these methods, a separate section of the Materials and Methods should specify what differed and why, and how the variations might have impacted the accuracy of results. In other disciplines, checklists have assisted with improving the reporting of results (Nature, 2018; NPQIP Collaborative Group, 2019). This approach is consistent with the TOP Guidelines (Transparency and Openness Promotion; Centre for Open Science, <https://www.cos.io/initiatives/top-guidelines>; Nosek et al., 2015).

This is an important initiative. We suggest it be implemented by all journals in our discipline to help improve the transparency, openness and reproducibility of science.

Supplementary data

Supplementary data are available at *Human Reproduction* online.

Data availability

No new data were generated or included in the manuscript.

Authors' roles

L.B., C.B. and D.M. outlined the first manuscript and contacted all other authors for comments. L.B., C.B. and D.M. summarized all suggestions and finalized the manuscript that all authors have received and confirmed their participation in.

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Conflict of interest

This is an Opinion article based on the necessity for improvement of standards in the field of Andrology and Reproductive Medicine, based on the two non-profit publications, by the World Health Organization and the International Organization for Standardization. No conflicts of interest are declared. CB, as an employee of the University of Dundee, serves on the Scientific Advisory board of ExSeed Health (from October 2021, financial compensation to the University of Dundee) and is a scientific consultant for Exscientia (from September 2021, financial compensation to the University of Dundee). CB has previously received a fee from Cooper Surgical for lectures on scientific research methods outside the submitted work (2020) and Ferring for a lecture on male reproductive health (2021). CB is Editor for *Reproductive Bio Medicine Online*. DL, as an employee Weill Cornell Medicine, declares: American Board of Bioanalysis (Secretary-Treasurer:

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Appendix

Semen analysis methodology checklist for authors, reviewers and editors (modified from Björndahl et al., 2016).

An accessible version of the checklist is available as a [supplementary file](#). This checklist is based on the ISO Standard on basic semen examination and the current WHO recommendations ("WHO6") [1,2], and on general scientific standards. Full compliance requires that all boxes are ticked.

A deviation from this checklist does not necessarily mean that the study cannot be published, but all deviations must be declared in the Materials and Methods section of the manuscript, including their impact on accuracy and measurement uncertainty of the data, in order to allow the reader to evaluate the quality of the analyses performed. For studies not reporting all characteristics of a basic semen examination, the checklist includes the option 'Not applicable to the study'.

Investigations that would be subject to the requirements in this checklist can roughly be classified as clinical (evaluating patient treatment, diagnostic classification or predictive powers of certain assessments), experimental (e.g. exposure of spermatozoa to different compounds or *in vitro* treatments [3,4]) or epidemiological (evaluating variations in semen characteristics or effects of exposure populations to certain compounds or other circumstances).

Any scientific rationale for not complying with the guidelines, which is not included in the Materials and Methods section of the manuscript, must be substantiated to the Editor and Reviewers.

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I. Patients

☐ Not applicable to the study

- ☐ I.1 The studied population (e.g. patients or volunteers) has been declared in the manuscript, together with the recruitment method and inclusion and exclusion criteria. In a study concerning couples being investigated for infertility, the following is specified in the manuscript: fertility status of female partner; and primary, secondary or other level of investigation of the man.
- ☐ I.2 If used in the manuscript, the term 'male factor' is completely defined.

- ☐ I.3 Reference limits provided in WHO5 or 5th percentile of distribution of semen examination results in WHO6 have not been used to characterize subjects as infertile, subfertile or fully fertile.

2. General aspects

☐ Not applicable to the study

- ☐ 2.1 Patients were instructed to maintain 2–7 days of sexual abstinence before collecting an ejaculate for investigation.
- ☐ 2.2 Patients were informed about the importance of reporting any missed early ejaculate fractions, and their responses were noted on the laboratory record.
- ☐ 2.3 For specimens not collected at the laboratory, patients were instructed to avoid cooling (under 20 °C) or heating (above 37 °C) of the semen specimen during transport to the laboratory.
- ☐ 2.4 In the laboratory, specimens were kept at 37 °C before initiation of and during the analysis in case of sperm motility assessment.
- ☐ 2.5 For specimens collected adjacent to the laboratory, analysis was initiated after completion of liquefaction and within 30 min after ejaculation. If some of the specimens were collected at the laboratory and others collected at home the influence on the data is declared and discussed in the manuscript.
- ☐ 2.6 Liquefaction was first checked within 30 min after ejaculation.
- ☐ 2.7 Volume was determined either by weighing or using a wide-bore volumetric pipette.
- ☐ 2.8 Viscosity was measured using either a wide-bore pipette or a glass rod.
- ☐ 2.9 All staff members who performed the analyses have been trained in basic semen analysis (ESHRE Basic Semen Examination Course—or equivalent—with further in-house training to establish competency), and participate regularly in internal quality control.
- ☐ 2.10 When more than one method is recommended for a particular characteristic (e.g. to measure volume), only one was used in the study. For a multicentre study, all laboratories used the same method or variable methods are declared in the manuscript.

3. Sperm concentration assessment

☐ Not applicable to the study

- ☐ 3.1 Semen aliquot to be diluted for sperm concentration assessment was taken with a positive displacement pipette (i.e. a 'PCR pipette') using a recommended diluent (state which diluent: _____).
- ☐ 3.2 Only standard dilutions were used (1:50, 1:20 or 1:10, i.e. 1 + 49, 1 + 19 or 1 + 9).
- ☐ 3.3 Sperm concentration was assessed using haemocytometers with improved Neubauer ruling.
- ☐ 3.4 Haemocytometers were allowed to rest for 10–15 min in a humid chamber to allow sedimentation of the suspended spermatozoa onto the counting grid before counting.
- ☐ 3.5 Sperm counting was done using phase contrast microscope optics (200–400×).
- ☐ 3.6 Comparisons were made between duplicate counts, and counts were re-done when the difference exceeded the acceptance limits.
- ☐ 3.7 Typically at least 200 spermatozoa were counted in each of the duplicate assessments.

4. Sperm motility assessment

☐ Not applicable to the study

- ☐ 4.1 Motility assessments were performed at $37^{\circ}\text{C} \pm 0.5^{\circ}\text{C}$.
- ☐ 4.2 Motility assessments were initiated within 30–60 min after sample collection.
- ☐ 4.3 Motility assessments were performed using phase contrast microscope optics (200–400 $\times$).
- ☐ 4.4 Sperm motility was classified using a four-category scheme: rapid progressive, slow progressive, non-progressive, and immotile.
- ☐ 4.5 Motility assessments were done in duplicate and compared; counts were re-done on new preparations when the difference between duplicates exceeded the acceptance limits.
- ☐ 4.6 The wet preparation was made using a drop of ____ μl and a ____ $\times$ ____ mm coverslip to give a depth of ____ μm (must be at least 10 μm depth, but not too deep so as to allow spermatozoa to move freely in and out of focus; typically ca. 20 μm).
- ☐ 4.7 At least 200 spermatozoa were assessed in each duplicate motility count.
- ☐ 4.8 At least five microscope fields of view were examined in each duplicate count.

5. Sperm vitality assessment

☐ Not applicable to the study

- ☐ 5.1 A validated supravital stain, appropriate to the type of microscope optics employed, was used to assess sperm vitality.
- ☐ 5.2 At least 200 spermatozoa were evaluated.
- ☐ 5.3 Assessments were done under high magnification ($\times 1000$ –1250) using a 100 $\times$ high resolution oil immersion objective and bright field microscope optics (Köhler illumination).

6. Sperm morphology assessment

☐ Not applicable to the study

- ☐ 6.1 Tygerberg Strict Criteria were used for the evaluation of human sperm morphology. (Another classification could be used for scientific studies with specific aims if the classification is described or referenced. Depending on the aim of the study, the evaluation of particular abnormal forms might be useful.)
- ☐ 6.2 Abnormalities are recorded for the four defined regions of the spermatozoon (head, neck/midpiece, tail and cytoplasmic residue).
- ☐ 6.3 The Papanicolaou staining method adapted for the assessment of human sperm morphology was used. For specific aims, other staining methods could be used but must then be declared and explained.
- ☐ 6.4 At least 200 spermatozoa were assessed in each ejaculate.
- ☐ 6.5 Assessments were done under high magnification ($\times 1000$ –1250) using a 100 $\times$ high resolution oil immersion objective and bright field microscope optics (Köhler illumination).

7. External Quality Assessment (EQA)

- ☐ 7.1 The laboratory participated in EQA for the semen examination methods used to obtain data for the study.
- ☐ 7.2 Name of the EQA scheme: _____

8. Other findings

☐ Not applicable to the study

- ☐ 8.1 The presence of abnormal clumping (aggregates and agglutinates) was recorded.
- ☐ 8.2 Abnormal viscosity was recorded.

9. Analysing data

☐ Not applicable to the study

- ☐ 9.1 The actual duration of sexual abstinence (in 'hours' or 'days') was recorded for each specimen and included in the data reported in the manuscript.
- ☐ 9.2 As a minimum in clinical studies, semen volume, sperm concentration, total number of spermatozoa per ejaculate and abstinence time are given to reflect sperm production and output; only samples identified as having been collected completely were included in the study.
- ☐ 9.3 Confounding factors have been considered for statistical analysis: e.g. abstinence time and age, to consider secular or geographical variations in sperm concentration or sperm count.
- ☐ 9.4 If appropriate, optional biochemical markers for prostatic, seminal vesicular and epididymal secretions were analysed and reported, both as concentration and total amount.
- ☐ 9.5 Signs of active infection/inflammation were noted and considered in the analysis of data in the study (e.g. presence of non-germ line round cells, inflammatory cells, impaired sperm motility, possibly also anti-sperm antibodies or reduction of secretory contributions).

10. Data Repository

- ☐ 10.1 For the sake of transparency, all data without identification of individual patients or research participants have been saved to a trusted online repository, and there is a statement of this in the Results section of the manuscript.

Declaration by the corresponding author:

The information provided in this checklist is solemnly declared to be true.

Signature: _____ Date: _____

Name: _____

Affiliation: _____