Clinical significance of the low normal sperm morphology value as proposed in the fifth edition of the WHO Laboratory Manual for the Examination and Processing of Human Semen

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Abstract

The very low cut-off value for sperm morphology of 4% morphologically normal spermatozoa, as proposed in the new edition of the World Health Organization (WHO) manual on semen analysis, is in agreement with recently published values and reflects the trend of a decline in reported mean values for normal sperm morphology. The reduced value for morphologically normal spermatozoa over the years may be due to several factors. The first is the introduction of strict criteria for the evaluation of sperm morphology. Other reasons may include the introduction of additional criteria for sperm morphology abnormalities and the suggested decrease in semen parameters because of increasing negative environmental influences. Although on its own the newly proposed very low normal value may not provide the strong predictive value for a males’ fertility potential, as originally reported for sperm morphology evaluated according to strict criteria, a good predictive value can still be obtained if the holistic, strict approach for sperm morphology evaluation is followed together with additional sperm morphology parameters now available, because certain morphology patterns and sperm abnormalities are now known to be of strong prognostic value. In addition, better international standardization of the technical methodology, consensus on the interpretation of sperm morphology evaluation criteria and standardized international external quality control (EQC) schemes, are of utmost importance to maintain the good predictive value of sperm morphology.

Keywords: male fertility potential, normal cut-off values, semen analysis, sperm morphology evaluation, strict criteria, WHO manual

1 Introduction

Reaching a professional consensus always involves a difficult balance of divergent views, and no consensus has been more difficult to reach than that regarding the evaluation and reporting of sperm morphology. Thus, there are sharply differing views on how to assess sperm morphology. This difficulty is reflected in the introduction (Chapter 1) of the 5th edition of the World Health Organization (WHO) Laboratory Manual for the Examination and Processing of Human Semen [1], which says, on the topic of assessment of sperm morphology, ‘Some laboratories assess only normal forms, while others consider the type, location and extent of abnormality to be more important. Whether these or differential or semi-quantitative assessments increase the value of the semen analysis remains contentious. Evidence to support the relationship between the percentage of ‘normal’ forms, (as defined by strict categorization or computer-aided assessment of morphology) with fertilization rates in vivo, justifies the approach of trying to determine a morphologically distinct sub-population of spermatozoa within semen.’

The introduction goes on to note, ‘In the 4th edition, an obvious omission, but inevitable given the lack of data, was the absence of a reference value for the percentage of
morphologically normal forms; this has been rectified in this manual with the inclusion of data on the percentage of normal forms in a fertile population.’ For this purpose, the lower fifth percentile was used, resulting in a very low cut-off value of 4% morphologically normal spermatozoa. At first impression, this lower reference limit of the percentage of normal spermatozoa in fertile men is so low that it is unlikely that a lower population (from infertile men) could be distinguished from it.

The aim of this article is, therefore, to discuss the usefulness of this very low reference value as well as the biological meaning of such tight categorization of a small minority of spermatozoa, and available tools and data to improve the diagnostic value of the suggested cut-off value.

2 History of sperm morphology evaluation

The interest in sperm morphology as a tool in the evaluation of a man’s fertility potential started in the early 1900s, when it became accepted that normal and pathological sperm forms could appear simultaneously in a semen sample [2]. Although the importance of normal sperm morphology has been stressed since that time—for example, with statements by Cary in 1916 [3], Williams and Savage in 1925 [4] and Moench and Holt in 1931 [5] that sperm morphology was the single most important source of information on the reproductive potential of spermatozoa—the manner in which the normality or abnormality of the spermatozoa should be evaluated has been controversial and remains a field of continuous debate.

The history of sperm morphology evaluation consists of two approaches or phases, which are also reflected in the evolution of sperm morphology evaluation criteria in the consecutive WHO manuals, from the first in 1980 to the soon-to-be-published fifth edition [1, 6–9]. The two approaches are the early, or liberal, approach [10] and the strict (Tygerberg) approach, as introduced by Menkveld in 1987 [11] and described in detail by Menkveld et al. in 1990 [12].

2.1 Liberal approach

Human males exhibit extreme heterogeneity, or polymorphism, of sperm morphology among [13] and even within specific males [14]. This varied morphology contrasts with that of most domestic animals, which generally reveals a very homogeneous sperm population in individual species. This homogeneity for animal sperm morphology made it possible to use the appearance of the modal spermatozoon form, as seen in proven fertile animals, to describe the morphologically normal spermatozoon form for that specific species. Owing to the heterogeneity of human sperm morphology, however, it was not possible to apply this concept successfully in human males. In the typical approach, investigators would describe different obvious abnormalities and use these abnormalities as criteria to evaluate sperm morphology. All spermatozoa not considered abnormal, were regarded as normal, and morphologically normal spermatozoa were, therefore, identified by default. These so-called (remaining) normal spermatozoa as well as specific spermatozoal abnormalities were often depicted by schematic and inaccurate drawings [15]. The liberal approach is illustrated by the words of MacLeod and Gold [16], who stated, ‘we are not prepared at this time to classify any but the most distorted forms as truly abnormal.’ Owing to the fact that on the basis of this approach no specific criteria were put forward for a morphologically normal spermatozoon, poor correlations could theoretically be expected with normal sperm functions and fertilization rates, as reported by Page and Holding [17].

2.2 Strict (Tygerberg) criteria

The strict approach for sperm morphology evaluation was conceptualized in the late 1970s and early 1980s, as described by Menkveld in 1987 [11] and Menkveld et al. in 1990 [12]. In contrast to the liberal approach, the definition for a morphologically normal spermatozoon was based on biological evidence obtained from spermatozoa found at the level of the internal cervical os. Spermatozoa found in such samples present with a mostly homogeneous appearing sperm population. Small biological variations in the head morphology can be observed in such samples, but these variations were considered normal. One of the most important aspects of the strict evaluation criteria is that the range allowed for these minute normal biological variations has to be kept as small as possible. For this reason, the so-called ‘borderline’ or ‘slightly abnormal’ head forms are regarded as abnormal.

By strict evaluation, the description of a morphologically normal spermatozoon is furthermore supported by the morphological appearance of spermatozoa found tightly bound to the human zona pellucida [18, 19].

2.3 Evolution of WHO criteria for sperm morphology evaluation

There was no clear textual description for morphologically normal spermatozoa in the 1st WHO manual published in 1980 [6], except in some legends for the color plates in which it is mentioned that ‘the germinal cell classifications are based on those proposed by MacLeod,’ thus following the liberal approach. Although slightly improved, the 2nd (1987) WHO manual [7] followed the same liberal approach [13] and provided only a little additional information on sperm morphology evaluation criteria.

In the 3rd (1992) WHO manual [8], the scheme for
the morphological classification of human spermatozoa was described carefully. The manual stated, ‘strict criteria should be applied when assessing the morphological normality of the spermatozoon.’ The strict evaluation definition for a morphologically normal spermatozoon is then given, as well as the statement that all ‘borderline’ forms should be considered abnormal. The 4th (1999) edition of the WHO manual [9] completely adopted the strict (Tygerberg) approach.

2.4 Evolution of the WHO cut-off point for normal morphology

Through the progression of the first four WHO manuals, some minor changes occurred in the cut-off values with respect to most semen variables, such as concentration, motility and viability. However, for normal sperm morphology, the values were lowered dramatically from the first edition of the manual published in 1980 to the fourth edition in 1999, and now especially in the fifth edition (Table 1). The first edition [6] included a table with different sperm morphology variables, such as normal, large oval heads and tapering heads, along with mean values and ranges. These values were based on the results of 602 ejaculates from 73 men who were fathers. The mean normal morphology was 80.5%, with a range of 48.0%–98.0%. In a footnote to the table, it is stated that, ‘Analysis of ejaculate (multiple samples) from at least 50–70 men of recently proven fertility, needs to be carried out at each centre to establish the normal values for a given population, in terms of sperm counts, germinal cell types and their percentages.’ Readers were referred to publications by MacLeod [20] and Eliasson [21] for more information. Eliasson [21] considered the morphology cut-off value to be ≥60% morphologically normal spermatozoa.

In the second edition of the WHO manual [7], the normal sperm morphology value was lowered to 50% morphologically normal forms. In the third edition [8], the value for normal sperm morphology was changed to ≥30% morphologically normal spermatozoa. A note added to this value stated that, ‘Although no clinical studies have been completed, experience in a number of centres suggest that the percentage of normal forms should be adjusted downwards when more strict criteria are applied. An empirical reference value is suggested to be 30% or more with normal forms.’ In the fourth edition, issued in 1999 [9], no specific value was indicated for normal sperm morphology, but a note was added stating that ‘Multicentre population-based studies utilising the methods of morphology assessment in this manual are now in progress. Data from assisted reproductive technology programmes suggest that, as sperm morphology falls below 15% normal forms using the methods and definitions described in this manual, the fertilisation rate in vitro decreases.’

In the fifth edition of the WHO manual [1], a new approach has been adopted with regard to the establishment of so-called ‘normal,’ or reference, values. This new direction is long overdue: many authors have been advocating the change for many years. One of the first groups to advocate these changes was that of the Andrology Unit at Tygerberg Hospital, in the publications by Van Zyl et al. [22, 23], Menkveld and Kruger [24–26], and Menkveld [27].

The proposed value in the fifth edition is very low, 4% morphologically normal spermatozoa, based on the fifth centile of combined data from several reports in recent publications, in which semen analyses were performed according to known and standardized methodologies [28].

| Table 1. Cut-off values for semen variables as published in consecutive WHO manuals [6–9] and as proposed in the fifth World Health Organization (WHO) manual [1]. |
|-------------------------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Semen variable                | 1980            | 1997            | 1999            | 2010            |                  |
| Volume (mL)                   | –               | ≥2.0            | ≥2.0            | ≥2.0            | 1.5             |
| Concentration (10⁶ mL⁻¹)      | 20–200          | ≥20             | ≥20             | ≥20             | 15              |
| Total sperm number (10⁹/ejaculate) | –             | ≥40             | ≥40             | ≥40             | 39              |
| Motility (%) motile          | ≥60             | ≥50 (a + b)     | ≥50 (a + b)     | ≥50 (a + b)     | 40 (a + b + c)  |
| Forward progression (for 1980 only) | ≥2⁴             | ≥25 (a)         | ≥25 (a)         | ≥25 (a)         | 32 (a + b)      |
| Morphology (% normal)        | 80.5⁴           | ≥50             | ≥30³            | (14)³           | 4               |
| Viability/vitality (% live)  | –               | ≥50             | ≥75             | ≥75             | 58              |
| White blood cells (10⁶ mL⁻¹)  | <4.7            | <1.0            | <1.0            | <1.0            | <1.0            |

¹Lower reference limit. Obtained from the lower fifth centile value. ²Grade a = rapid progressive motility (> 25 µm s⁻¹); Grade b = slow/ sluggish progressive motility (5–25 µm s⁻¹); Grade c = no-progressive motility; Grade d = immotility; Normal = ≥50% motility (grades a + b) or ≥25% progressive motility (grade a) within 60 min of ejaculation. ³Forward progression on a scale of 0–3, in which 0 = no forward progression. ⁴Mean of a fertile population (range = 48%–98%); modified Bryan Leishman staining. ⁵Arbitrary value. ⁶No actual value given. Multicentre studies in progress refer to >14% for in vitro fertilization (IVF).
3 Comparison of the proposed WHO value with recently published normal sperm morphology values

As early as 1975 and 1976, Van Zyl et al. [22, 23] published data proposing much lower normal values on the basis of interval values above which spontaneous pregnancies showed no statistically significant increases with an increase in a particular semen variable value (Table 2). Below these normal cut-off values, two additional cut-off points could be established, one for a grey zone, in which occasional spontaneous pregnancies could still be expected, and another below which a spontaneous pregnancy could only very occasionally be expected [29]. For normal sperm morphology, the normal cut-off value was 20% morphologically normal spermatozoa, compared with the 60% morphologically normal spermatozoa value proposed by MacLeod and Gold [16] and Eliasson [30]. The cut-off value for sperm morphology for the grey zone, as proposed by Van Zyl et al. [29], was 10% morphologically normal spermatozoa and 3% was the low cut-off value, below which spontaneous pregnancies were seldom seen [29, 31]. These three categories are in agreement with the differentiation of semen analysis results into prognostic classes as published by Eliasson [30], although he used the terms normal, doubtful, pathological and severe pathological.

Similarly, in 1986 and 1988, Kruger et al. [32, 33] published data suggesting that three zones could be distinguished for sperm morphology with regard to the prediction of a males’ fertility potential for in vitro fertilization (IVF) on the basis of percentage of the total number of oocytes fertilized per morphology interval group. The three cut-off values for these zones were ≥15% morphologically normal spermatozoa for a normal prognosis group, 5%–14% for the good prognosis or G-pattern group, and ≤4% for the poor prognosis or P-pattern group.

More recently, a number of papers have been published comparing sperm morphology cut-off values to in vivo pregnancy results in infertile populations or so-called fertile and subfertile populations (Table 2). Eggert-Kruse et al. [34] published data from a prospective study including 89 randomly selected couples presenting for infertility investigations, for which the female partners were apparently normal. A spontaneous pregnancy rate of 20.2% (18/89) was observed after 1 year without treatment. An analysis of sperm morphology, as evaluated by

| Author                  | Population | Cut-off value | Fertile | Subfertile | Infertile | TZI | AI |
|-------------------------|------------|---------------|---------|------------|-----------|-----|----|
| Van Zyl et al. [23, 29, 31] | In vivo pregnancies | 20 | 10 | 3 |
| Kruger et al. [32, 33] | IVF fertilization rates | Initial published intervals | $\geq 15$ | 14–4 | $\leq 3$ |
|                         |            | Intervals as used in practice | $\geq 15$ | 14–5 | $\leq 4$ |
| Eggert-Kruse et al. [34] | In vivo pregnancies | 14 | 7 | 4 |
| Omelet et al. [35] | Tenth percentile | Fertile population | 10 |
|                          | ROC curve analysis | Fertile vs. subfertile | 5 |
| Zinaman et al. [36] | Healthy couples | 8 |
| Günalp et al. [38] | Fertile vs. subfertile | 12 | 5\(^2\) |
| Menkveld et al. [37] | ROC curve analysis | Fertile vs. subfertile | 4 | 1.64 | 8 |
|                          | Adjusted | Fertile vs. subfertile | 3 | 2.09 | 3 |
|                          | Tenth (upper) percentile | Fertile population | 2 | 1.82 | 5 |
| Guzick et al. [39] | Fertile vs. subfertile | $> 12$ | 9–12 | $< 9$ |
| Haugen et al. [40] | Tenth percentile | Fertile population | 4 |
|                          | Fifth percentile | 3 | 1.72 |
| Fifth WHO manual [1] | Fifth percentile | Recent fathers | 4 |

Abbreviations: IVF, in vitro fertilization; ROC, receiver operating characteristic; TZI, teratozoospermia index; AI, acrosome index. \(^2\)See text for more detailed descriptions of populations investigated. \(^3\)Based on the positive and negative predictive values (see Günalp et al. [38]).
strict criteria, indicated three morphology cut-off points for the occurrence of a pregnancy, namely, ≥ 4% with a pregnancy rate of 21.5% (17/79), ≥ 7% with a pregnancy rate of 27.4% (17/62) and ≥ 14% with a pregnancy rate of 34.3% (12/35).

Ombelet et al. [35] compared the semen parameters of 144 males from couples, in whom the wives had achieved a recent pregnancy within 12 months of unprotected intercourse. The subfertile group consisted of 143 consecutive men attending the infertility clinic during the same period, who did not achieve a pregnancy. Using the lower tenth percentile of the fertile population, the cut-off value for sperm morphology was 10% morphologically normal spermatozoa; using the receiver operating characteristic (ROC) curve analysis method, the cut-off value to distinguish between the fertile and subfertile group was 5% morphologically normal spermatozoa.

Zinaman et al. [36] also conducted a prospective study with 210 healthy couples. The authors followed each couple for 12 menstrual cycles, during which the couple tried to conceive, and performed multiple semen analyses in each man over the period of exposure with known abstinence intervals. The couples were instructed to have intercourse as frequently as possible on and around the day of predicted ovulation. Semen analyses results were divided into pregnant and non-pregnant groups. The mean (SD) percentage of morphologically normal spermatozoa for the pregnant group was 6.2 (3.7)% and that for the non-pregnant group was 4.1 (3.5)%. A decline in pregnancy rate was suggested below 8% morphologically normal spermatozoa.

Like Ombelet et al. [35], Menkveld et al. [37] compared the semen parameters of a fertile and subfertile population. The fertile population comprised a group of 107 healthy males without any history of fertility problems, the partners of whom had a spontaneous pregnancy within 1 year of unprotected intercourse and were pregnant at the time the husband was included into the study. A total of 103 males of couples attending the infertility clinic were included in the subfertile group. Using the ROC curve analysis, the cut-off point values for sperm morphology, teratozoospermia index (TZI) and the acrosome index (AI) were 4% morphologically normal spermatozoa, 1.64 and 8% normal acrosomes, respectively. Taking into consideration that the male contributes to the infertility problem in about 50% of cases, these values were recalculated in terms of false positive and false negative predictive values. The recalculations resulted in cut-off point values of ≤ 3% morphologically normal spermatozoa, a TZI value of ≥ 2.09 and an AI value of ≤ 3% normal acrosomes. Taking the lower tenth percentile values of the fertile population, the cut-off points were 2% morphologically normal spermatozoa, 5% for the AI and a TZI value of 1.82.

Gunalp et al. [38] also published ROC curve analysis data of a prospective study that included 61 fertile males whose wives were at a gestational age of 8–23 weeks and 62 males attending the infertility clinic, resulting in a cut-off value of 12% morphologically normal spermatozoa. After an adjustment for the assumption of an infertility prevalence of 15% in the population, the cut-off point was calculated at 5% morphologically normal spermatozoa.

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4 Possible reasons for the decline in reference values

The reason for the drastic decrease in normal sperm morphology cut-off values over the years, as illustrated in Figure 1, are mainly three-fold, namely, (i) the implementation of strict evaluation criteria with the unfortunate result that sperm morphology evaluation became over-critical with regard to normality, (ii) the fact that over the years more criteria for sperm morphological abnormalities were identified and introduced into the evaluation system, and (iii) a true decline because of negative environmental factors.

4.1 Introduction of strict criteria

The statement that, with strict evaluation, borderline spermatozoa should be regarded as abnormal, which is against the principle for the liberal approach that borderline spermatozoa should be regarded as normal, automatically placed a large number of spermatozoa, which would have been regarded as normal by the liberal approach, into the abnormal category when using the strict criteria approach,
leading to an overall lower percentage of morphologically normal spermatozoa. This has been illustrated in several articles [37, 41].

Furthermore, the term ‘strict’ introduced a trend of over-criticality in the application of the definition of normality. An analysis of our own laboratory data indicated that the decline, as illustrated in Figure 1, was partly because of a real decline in normal sperm morphology and also because of stricter application of the criteria for normality of spermatozoa [42]. On the basis of the re-evaluation of 80 randomly selected semen smears from 1988 in 1995, the mean (SD) percentage of the 80 smears as evaluated in 1988 was 12.4 (7.8)% morphologically normal spermatozoa. The mean (SD) of the same 80 smears evaluated in 1995 was 9.6 (5.5)% morphologically normal spermatozoa, a decrease of 2.8 percentage points because of stricter application of the criteria over the period of 8 years. The overall mean percentage of morphologically normal spermatozoa in 1988 was also 12.4 (7.6)% (n = 219) and decreased to a mean of 7.8 (5.1)% (n = 227) in 1995, showing overall decrease of 4.6 percentage points from 1988 to 1995. Of the overall decrease of 4.6 percentage points from 1988 to 1995, at least 2.8 percentage points could be attributed to a stricter approach in the evaluation of sperm morphology, and it can be postulated that the remaining 1.8 percentage points may be because of negative environmental factors.

This tendency toward a stricter approach over time was also illustrated in a publication by Horte et al. [43], in which a series of sperm morphology smears from 1980 to 1994 were re-evaluated. The results showed that the observed decline in the percentage of morphologically normal spermatozoa observed over the 15-year period was mainly because of a stricter evaluation approach over the years. On the basis of the predictive values from a multiple regression analysis to calculate the end points of regression lines, the proportion of spermatozoa with normal morphology declined from 42.8% to 36.7% over the 15-year period. The re-evaluation of the sampled smears only showed a minimal change, from 13.7% in 1980 to 13.3% in 1994. The effect of the application of stricter criteria, as suggested by the consecutive WHO manuals, can be very clearly seen in the substantially lower means obtained with the re-evaluation of the sampled morphology smears.

4.2 Introduction of additional criteria for sperm morphology evaluation

The original description of sperm abnormalities, as described, for example, by MacLeod and Gold [16], and Freund [44], concerned head abnormalities only. In 1971, Eliasson [30] introduced the concept that for a spermatozoon to be classified as normal, the whole sperm must be normal, including abnormalities of the neck/midpiece and tail, as well as the presence of excess cytoplasmic residue material. This automatically decreased the possible number of normal spermatozoa in a sample. As more sperm functions in the fertilization pathway [45] became known and these sperm functions could be associated with certain sperm morphology structures, the probability of sperm morphology abnormalities increased for a specific semen sample with the corresponding decrease in the mean percentage of morphologically normal spermatozoa. One of the more recently identified sperm morphology parameters based on sperm functionality includes that of the sperm acrosome. Acrosome size, such as too big or too small, and acrosome staining abnormalities are now important criteria for sperm morphology evaluation that had not been recognized until recently [46]. Thus, as more sperm functional abnormalities are recognized, a corresponding decrease in the percentage of morphologically normal spermatozoa will likely be observed.

4.3 Negative environmental factors

Carlson et al. [47] suggested a declining trend in semen quality worldwide. One of the main reasons for the suggested decline was the increasing effect of endocrine disruptors and other environmental polluters,
such as insecticides and pesticides [48]. The entire issue of declining sperm parameters, including sperm morphology, has gained importance because of the recognition of several other trends that reflect a decline not only in semen production, but also in overall reproductive health, including the increase of testicular cancers and higher trends of cryptorchidism, together with a more general increase in testicular dysfunction [49], which also has a negative effect on normal sperm morphology. Another contributing factor may be a wider adoption of a more promiscuous lifestyle in some communities, leading to an increase in sexually transmitted diseases and male accessory gland infections (MAGI). The higher prevalence of MAGI, as identified by the presence of leukocytospermia, can also have a negative effect on spermatogenesis by reducing the percentage of morphologically normal spermatozoa and increasing the percentage of elongated spermatozoa [50].

5 Additional tools to improve/refine sperm morphology evaluation

By itself, the very low normal sperm morphology cut-off value proposed in the fifth edition of the WHO manual and also in the recently published articles by Menkveld et al. [37] and Haugen et al. [40] may be of limited prognostic value. However, over the years, the evaluation of sperm morphology has been refined and relies on more than just the percentage of morphologically normal spermatozoa. Additional sperm morphology parameters have been developed to improve the diagnostic abilities of sperm morphology parameters in the prediction of males’ fertility potential. In males presenting with normal spermatozoa below the proposed WHO cut-off value of ≤ 4%, two groups, or prognostic categories, can be identified from the morphology patterns and morphological abnormalities present.

The first group will have a poor prognosis because of mostly genetically determined sperm pattern defects, such as globozoospermia; short tail syndrome; spermatozoa with small heads, in most cases combined with very small acrosomes; cases of large-headed spermatozoa, mostly combined with the presence of large acrosomes; and the presence of large or small acrosomes alone. All of these types of abnormal spermatozoa have been associated with abnormal sperm functions and, thus, with poor in vivo and IVF rates.

The second group will include males with unspecified or non-genetically determined sperm defects or patterns caused by environmental factors, including medication.

5.1 Genetically determined sperm defects

5.1.1 Globozoospermia

One of the best-known genetically determined sperm sterilizing defects or patterns is globozoosperma. These spermatozoa are easily recognizable by their small, round-head form and the absence of acrosomes. Owing to the absence of acrosomes, the sperm head does not contain acrosin and is unable to bind and penetrate the zona pellucida. When couples are treated by intracytoplasmic sperm injection (ICSI), pregnancies can be achieved in some cases [51, 52].

5.1.2 Short tail syndrome

In some semen samples, the spermatozoa present with abnormalities of the spermatozoon tail and neck region. The most common defects are the stump or short tail syndrome, spermatozoa with dysplasia of the fibrous sheath, acaephic spermatozoa and spermatozoa with defects of the head–neck attachments [53]. In spermatozoa with neck attachment defects, the heads attach either to the tip or to the sides of the midpiece without an alignment with the sperm axis. This misalignment ranges from a complete lack of connection (difficult to observe) to a lateral positioning of the nucleus at a 90–180° angle [53]. These patients have a poor prognosis, even with ICSI, and in cases in which fertilization has resulted, the pronuclei do not fuse, and zygotes fail to cleave and, thus, degenerate [53]. This sperm defect can be because of defective sperm aster formation, syngamy and cleavage. Even when sperm are carefully selected for the absence of these abnormalities using ICSI and positive human chorionic gonadotropin (βHCG) levels are achieved, preclinical abortions are common [54].

5.1.3 Small-headed spermatozoa

Small-headed spermatozoa in men occur more frequently in couples complaining of infertility than currently diagnosed because of the unfortunate fact that this condition is not commonly recognized and reported as a severe sperm abnormality. This abnormality is stable over time. When recognized, small-headed spermatozoa are easy to observe, because most spermatozoa are small (head length < 3.5 µm and head width < 2.5 µm with Diff-Quik staining) [55]. These spermatozoa may also present with very small, abnormally formed acrosomes. Individuals who are not diagnosed with this condition will achieve poor results with IVF and even with ICSI, but by selecting the more normally appearing spermatozoa, ICSI results can be improved. Even when larger spermatozoa with small acrosomes are present, with < 30% of normal sperm head size, the prognosis for the patient with IVF is poor, because these spermatozoa have a low vitality and are not able to undergo the acrosome reaction [56].

5.1.4 Large-headed spermatozoa

These spermatozoa present with severely abnormal
megalone heads and multiple tails, and are observed in the ejaculate and testicular biopsies [57].  Low fertilization and pregnancy rates are achieved with ICSI, which may be because of the high incidence of chromosomal aberrations in the ejaculated megalo-headed spermatozoa [58].

5.2 Non-genetic sperm morphology aberrations

The second group, with unspecified or non-genetically determined sperm defects or patterns caused by environmental or other stress factors, including medication, differs from the group with genetically determined permanent sperm defects, which will not respond to treatment.  Sperm defects in the second group can return to normal or disappear when the source of the alternations or stress is removed or treated.  Males with these types of sperm abnormalities still have a good chance for normal in vivo conception or, alternatively, a good prognosis for IVF without ICSI.  To further enhance the diagnosis of men with a potentially good IVF chance, several additional morphology tools are available, such as the TZI, AI, the presence of cytoplasmic residues and sperm head measurements.

5.2.1 Megalo heads

Megalo heads can also be caused by the use of medicine, such as sulfasalazine, for the treatment of ulcerative colitis [59] and Crohn’s disease [60].  When sulfasalazine treatment is stopped, the semen parameters can return to normal values, and the megalo heads can disappear.  It is not clear whether this will result in fertilization and viable pregnancies.  However, the substitution of sulfasalazine with mesalazine has resulted in improved semen quality and the reduction of the megalo heads, subsequently resulting in viable pregnancies [61].

5.2.2 Elongated spermatozoa

Elongation is generally recognized as a stress-induced sperm morphology aberration, and is prevalent especially in MAGI and in the presence of a varicocele [62].  The classic tapering or narrow form has been described in detail by Eliasson [30] as being a sperm head longer than 5 µm with a width < 3 µm or a length of < 5 µm and a width of < 2 µm.  Pyriform heads are also included under elongation of spermatozoa.  Sperm elongation is accompanied by severe structural as well as DNA damage.  The increased sperm head length results from an abnormally elongated nucleus, which also presents particular membranous layers between the outer and inner leaves of the nuclear envelope.  The sperm nuclear anomalies are also associated with anomalies of the neck region, persistence of cytoplasmic residual material and an increased frequency of chromosomal aneuploidies, together with impaired chromatin compaction, possibly because of meiotic non-disjunction during spermatogenesis [63].  Low ICSI fertilization rates have also been found in men with severely elongated spermatozoa compared with other sperm morphology abnormalities [64].  However, when the MAGI is treated with long-term antibiotics or the varicocele is treated with a varicocelectomy, semen characteristics, including sperm morphology, can improve and pregnancies can result [65].  In case of varicocelectomies, in which no substantial improvement in semen quality is observed, pregnancies do still occur because sperm DNA quality improves because of the reduction of ROS production following the varicocelectomy [66].

5.2.3 Cytoplasmic residues

In addition to abnormalities of the head, neck/midpiece and tail, the fourth major sperm abnormality is the presence of cytoplasmic material at the neck/midpiece region and sometimes on the tails of spermatozoa.  The presence, size and terminology of the cytoplasmic droplets or cytoplasmic residues are controversial.  Originally, Eliasson [30], in the 1980 and 1987 WHO manuals [6, 7], and Menkveld et al. [12] indicated that a normal cytoplasmic droplet present on spermatozoa should be < 50% of a normal sperm head.  This was changed to < 30% in the 1999 WHO manual [9].  Recently, Cooper et al. [67] and Cooper [68] addressed the issue of the presence and size of cytoplasmic bodies, as well as the correct terminology to be used.  According to Cooper [68], it is clear that retention of cytoplasmic material on spermatozoa, as seen in air dried and stained semen smears, can be associated with impaired sperm function.  It is clear from this article and from our own experience [69] that no amount of cytoplasmic material should be present on a normal spermatozoon and, if observed, it should be regarded as an abnormality, regardless of the size or amount of cytoplasmic material present.  Cooper [68] also suggested that the correct term to be used for the presence of cytoplasmic material should be ‘excess cytoplasmic residues’ or cytoplasmic residues.  The presence of cytoplasmic residues is associated with sperm immaturity and especially the production of ROS, with subsequent sperm DNA damage; although IVF pregnancies may occur, decreased pregnancy rates are reported [70].  Therefore, much more attention should be given to the presence of cytoplasmic residues with the routine evaluation of sperm morphology.

5.2.4 Sperm measurements

The earlier sections showed that sperm size, and therefore sperm measurement, is a very important aspect of the sperm morphology evaluation process.  However, the measurements proposed by Eliasson [30], and adopted in the WHO manuals [6–9] and other publications [12] are in need of re-evaluation, because the range allowed,
especially the normal head length of 3.0–5.0 μm, is too wide. Our own experience indicates that the head length for normal spermatozoa may vary between 4.0 and 4.5 μm, with a mean of 4.07 ± 0.19 μm and a mean width of 2.98 ± 0.14 μm, as measured with a built-in microscope eye-piece micrometer [27]. We have shown in several publications that males presenting with large-headed spermatozoa of > 5.0 μm in length, and proportional increases in width and/or large acrosomes, as seen in Papanicolaou-stained smears, are associated with poor IVF results [32] and decreased sperm functional abilities [46].

Alternative measurements have been proposed by several authors, such as Katz et al. in 1986 [71] and Garrett and Baker in 1995 [72], who reported length/width measurements of 4.37 μm by 2.83 μm and 4.35 μm by 2.89 μm for Papanicolaou- and Shorr-stained smears of donors, respectively. The new WHO manual no longer refers to the ‘old’ sperm measurements, but states that head dimensions of 77 Papanicolaou-stained semen smears measured with a computerized system, have a median length of 4.1 μm (3.7–4.7 μm, 95% CI) and width of 2.8 μm (2.5–3.2 μm) and a length-to-width ratio of 1.5 (1.3–1.8). Unfortunately, the manual also states that ‘once a normally shaped spermatozoa is identified, an eye-piece micrometer may be useful for distinguishing between the normal and abnormal (size – own word) spermatozoa, but with this technique the form of the sperm head is much more important than its dimension unless grossly abnormal.’ However, sperm measurements are of utmost importance for the normal oval form. Sperm measurement is an area deserving much more attention in today’s sperm morphology assessment methodology, and more research is needed to establish more accurate sperm measurements.

5.2.5 Detection of leukocytes

Although not actually a sperm morphology parameter, determination of the presence of leukocytes, especially polymorphonuclear leukocytes, is an important aspect of the sperm morphology evaluation procedure [65]. Polymorphonuclear leukocytes are the main source of ROS production in semen, leading to increased sperm DNA damage. Although sometimes reported as part of the morphology evaluation procedure, much more emphasis should be placed on the presence of polymorphonuclear leukocytes in semen and the importance of antibiotic treatment for leukocytospermia and the subsequent possible improvement in pregnancy outcome [65, 70].

5.3 TZI

The two most recent WHO manuals [8, 9] recommended that spermatozoa should only be classified as normal or abnormal and that a note should be made if a specific abnormality occurs with a frequency of > 20%. However, an abnormal spermatozoon may have only one specific abnormality or any combination of two to four abnormalities, as described above. The TZI was introduced to reflect the mean number of abnormalities per abnormal spermatozoon.

The usefulness of the TZI seems to be limited, and it is seldom used in the literature as a tool in decision-making with regard to possible treatment of patients because it does not indicate a specific dominant abnormality present in a specific semen sample. However, a high TZI value of ≥ 1.90 can be regarded as a poor prognosis for normal IVF, and these patients should be taken directly to ICSI [73]. Following a literature search and on the basis of an analysis of our own data, we concluded that the TZI has little predictive value for in vivo and IVF outcomes [74].

5.4 Acrosome index (AI)

The AI is an indication of the percentage of spermatozoon with morphologically normal acrosomes present in a semen sample and can be regarded as a reflection of the functional ability of the spermatozoa, because the AI is based on the morphological appearance of the acrosomes with regard to size (too small or too large), staining properties and form (oval or amorphous shaped).

Incorporation of the AI in the sperm morphology evaluation process is, therefore, an important tool in the prediction of fertilization outcome, especially in cases of severe teratozoospermia (now ≤ 4% normal forms). The AI was found to be a more sensitive predictor of expected IVF rates compared with normal sperm morphology, whereas the TZI was a less sensitive predictor compared with normal sperm morphology, on the basis of an ROC curve analysis for the prediction of a > 37% IVF rate with areas under the curve of 0.927, 0.712 and 0.653 for the AI, percentage of normal morphology and TZI values, respectively [75].

6 Future aspects of sperm morphology evaluation

The results of sperm morphology evaluations can be further improved by better international standardization for the whole sperm morphology evaluation procedure, better international training and improvement, and standardization of international external quality control (EQC) schemes.

6.1 International standardization of morphology evaluation procedures

According to the literature, although many authors mention in their articles that strict criteria/1999 WHO criteria have been used for sperm morphology evaluation in their laboratories, many may still use wet preparations, Diff-Quik-stained smears or other staining methods,
contributing to confusion in the literature and diverse results obtained for sperm morphology evaluation in different laboratories throughout the world [76]. Different staining procedures can lead to different sperm morphology evaluation results [77].

Therefore, before comparisons of the results of different laboratories can be made, large-scale standardization should be implemented in laboratories worldwide as far as preparation and staining techniques are concerned, focusing on well-prepared smears and staining according to the Papanicolaou method only.

6.2 Standardization of international external quality control schemes

The lack of standardization in the EQC schemes is illustrated in the article by Cooper et al. [78], in which the results of three EQC programs for the same semen smears were compared, namely, those of the ESHRE SIG-A, the EAA, and the UKNEQAS schemes. Assessment by the SIG-A EQC was stricter than that of the other two programs, and the results of the UKNEQAS were much more widely distributed compared with the EAA scheme results.

6.3 International cooperation in standardization

The users of different international EQC programs should start to work together by interchanging standardized QC material and then, after a trial period, come together to discuss the results and to try to standardize their interpretations for sperm morphology evaluations on the basis of strict evaluation principals.

7 Conclusions

The very low cut-off value for sperm morphology of 4% morphologically normal spermatozoa, as proposed in the new WHO manual [1], is in agreement with recently published values [37, 40] and with the trend of declining mean normal sperm morphology values reported in the literature [34–40].

On its own, the new proposed very low normal value may not provide a strong predictive value for a male’s fertility potential, as originally reported for sperm morphology evaluated according to strict criteria [79]. However, a good predictive value can still be obtained if the holistic strict approach for sperm morphology evaluation is followed and used with the additional sperm morphology parameters available in cases in which severe teratozoospermia (≤ 4% normal) is present. In these cases, it becomes of utmost importance that clinicians receive more detailed data on abnormal sperm patterns and specific abnormalities so they may use the additional information to make a final decision regarding the treatment of the patient, because certain morphology patterns and sperm abnormalities are known to be of strong prognostic value.

The normal WHO value for sperm morphology is based on Papanicolaou-stained smears. Another set of normal cut-off values needs to be established for rapid staining methods because the literature indicates that higher normal values are obtained in Diff-Quik-stained semen samples compared with the same samples stained with the Papanicolaou method [77].

A high percentage of morphologically abnormal spermatozoa can be associated with the presence of poor sperm functionality. On the other hand, morphologically normal-appearing spermatozoa should also be further investigated for normal sperm function and DNA content because morphological normality does not necessarily imply normal sperm function [80, 81].

In addition, better international standardization of the technical methodology, consensus on the interpretation of sperm morphology evaluation criteria, and standardized international EQC schemes are of the utmost importance.

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