HARMONIZATION OF ANALYSIS AND REPORTING OF 19-NORSTEROIDS RELATED TO NANDROLINE

1.0 Introduction

This document has been established to harmonize the analysis and reporting of 19-norsteroids related to nandrolone as Adverse Analytical Findings by Laboratories.

The detection of the use of nandrolone and other 19-norsteroids is based primarily upon the identification of the main urinary metabolite, 19-norandrosterone (19-NA) in an amount greater than 2 ng/mL. More than one metabolite (e.g., 19-noretiocholanolone (19-NE)) of administered norsteroids may be detected and reported but the identification and quantification of the 19-NA metabolite only (derived from hydrolysis with β-glucuronidase) is sufficient to report an Adverse Analytical Finding.

2.0 Analysis

2.1 Identification and Quantification

In addition to meeting the identification criteria described in the IDCR Technical Document, the Laboratory shall demonstrate that the concentration of 19-NA is above the Decision Limit (DL) as set out in the DL Technical Document.

The quantification method used to calculate the concentration of 19-NA shall include or have the following characteristics:

- A deuterated internal standard (d4-19-NA-glucuronide is the preferred internal standard since it corrects for both the hydrolysis and other analytical steps);
- A calibration curve at an appropriate range bracketing the estimated concentration of the analyte;
- The use of appropriate quality control samples. For example, a negative control (without the presence of 19-NA or at a concentration ≤ 1ng/mL of 19-NA) and a positive control in the range of 3 to 5 ng/mL of 19-NA may be used. Alternatively, a freeze-dried urine reference material with approximately 2 ng/mL of 19-NA may be used (e.g., NMI Reference number MX002);
- The combined standard measurement uncertainty established at the threshold by the Laboratory shall be less than the $u_{c_{\text{Max}}}$ specified in the DL Technical Document.

2.2 Additional mandatory tests

The Laboratory shall also perform methods to test for pregnancy (e.g. hCG) and detection of tetrahydronorethisterone in urine Samples from female Athletes that have 19-NA concentrations greater than the threshold.
In extremely rare circumstances, 19-NE, and to a lesser extent 19-NA, can be formed by 19-demethylation of abundant endogenous steroids. These “unstable” Samples show the presence of low and comparable levels of 19-NA and 19-NE where the ratio of 19-NA/19-NE is less than the ratio of A/E (androsterone/etiocholanolone).

When concentrations less than 10 ng/mL of 19-NA are measured and the Sample shows the above feature of instability, the Laboratory shall perform a stability test. The stability test, which incorporates deuterated androsterone and etiocholanolone, has been described by Grosse et al [1].

- If the stability test is positive (i.e. Sample is “unstable”), the 19-NA result shall not be reported.
- If the stability test is negative, the Sample shall have a GC/C/IRMS analysis performed on the 19-NA [2]. The criterion for an unstable urine by GC/C/IRMS shall be a difference of less than 3.0‰\(^1\) between the measured δ values (\(\Delta\delta = \delta_{\text{Androsterone}} - \delta_{19-\text{NA}}\)) of 19-NA and androsterone. The GC/C/IRMS analysis should include the confirmation of the 19-NA peak identity (for example by GC/MS analysis).
- If the stability test is negative and the GC/C/IRMS test shows that the 19-NA is endogenous, the 19-NA result shall not be reported.
- If the stability test is negative and GC/C/IRMS test shows that the 19-NA is exogenous, the result shall be reported as an Adverse Analytical Finding.

### 3.0 Reporting

The Laboratory shall report the finding of 19-NA in a urine Sample from a male or a female Athlete at a concentration above the DL after adjustment of the threshold for specific gravity (see section 3.1 and 3.2 below) and meeting the requirements of IDCR TD as defined below:

A. Samples from male or non-pregnant female Athletes for which 19-NA concentrations are determined to be greater than 25 ng/mL shall be reported as an Adverse Analytical Finding simply as “>25 ng/mL” without the need for reporting concentration.

B. Samples from male or non-pregnant female Athletes for which 19-NA concentrations are determined to be between 10 ng/mL and 25 ng/mL shall be reported as an Adverse Analytical Finding as the mean concentration from triplicate determinations to not more than two significant figures\(^2\) as well as the measurement uncertainty in compliance

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1 The difference between δ values need not be corrected for any derivatizing agent since both androsterone and norandrosterone would have the same contribution from added carbon atoms.

2 A result between 2 and 10 would be reported, for example, as "5.1 ng/mL". A result between 10 and 25 would be reported as "17 ng/mL".
with the DL Technical Document. A comment or opinion shall be added according to 3.3 below as applicable.

C. **Samples** from male or non-pregnant female *Athletes* not using norethisterone for which 19-NA concentrations are determined to be between the DL and 10 ng/mL and for which the features of an “unstable” urine are NOT met shall be reported as an *Adverse Analytical Finding* as the mean concentration from triplicate determinations to not more than two significant figures as well as the measurement uncertainty in compliance with the DL Technical Document. A comment or opinion shall be added according to 3.3 and 3.4 below as applicable.

D. **Samples** from male or non-pregnant female *Athletes* not using norethisterone for which 19-NA concentrations are determined to be between the DL and 10 ng/mL and for which the feature of an “unstable” urine is met and the stability test is positive shall be reported as Negative. If the stability test is negative, then the *Sample* shall be submitted to GC/C/IRMS analysis and reported according to section 2.2 above.

E. **Samples** from non-pregnant female *Athletes* using norethisterone shall be reported as *Adverse Analytical Finding* but adding a comment or opinion about the use of norethisterone as described below under section 3.3.3. The presence of tetrahydronorethisterone should be confirmed.

F. **Samples** from pregnant female *Athletes* for which 19-NA concentrations are determined to be greater than the DL shall be reported as Negative unless IRMS testing is consistent with exogenous administration. The Laboratory shall include in the report a comment indicating the test results that support the opinion of the pregnancy as described below under section 3.3.

### 3.1 Adjusted Threshold

Only in the case of urine *Samples* measured with a specific gravity above 1.020 (in the Laboratory), an adjustment to the threshold shall be made to take into account the specific gravity (SG) of the *Sample* (since higher 19-NA concentrations have been associated with higher specific gravities (e.g. [3])), using the following formula:

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\text{Threshold}_{\text{adjusted}} = \left[ \frac{(\text{Specific gravity of the } \text{Sample} - 1)}{(1.020 - 1)} \right] \times 2\text{ng/mL}
\]

### 3.2 Decision limit for 19-NA

The DL for 19-NA is established in the DL Technical Document. In cases where the SG is greater than 1.020, the guard band represented by the difference between the value of the DL and the value of the threshold, as specified for 19-NA in the DL Technical Document, shall be added to the adjusted threshold to determine the DL
for an individual 19-NA test result\(^3\). The DL (after adjustment of the threshold for SG) shall be included on the Laboratory test report. The DL shall be used to determine whether an Adverse Analytical Finding is reported for the Sample.

The result from a Sample shall be reported as an Adverse Analytical Finding if the measured mean concentration is greater than the DL (adjusted for SG, if necessary) unless a Sample meets one of the conditions discussed in sections 3.3 and 3.4 below. If the Sample does meet one of the conditions discussed in 3.3 and 3.4 below, then it shall be reported as an Atypical Finding and both the Testing Authority and WADA shall be notified of the results as a comment in the test report.

### 3.3 Female Athlete’s Samples

When 19-NA exceeds the DL in the urine Sample of a female Athlete, the Laboratory shall include in the test report the results of tests to determine whether the 19-NA is due to pregnancy or to the intake of a medication containing norethisterone.

3.3.1 If hCG is present at a concentration less than 1000 IU/L, a comment or opinion shall be added to the test report indicating that the test was performed and that the 19-NA result was not consistent with pregnancy. An example would be:

"The Sample was analyzed for hCG and the concentration was less than 1000 mIU/mL; indicating that the 19-NA finding is not the result of pregnancy."

3.3.2 If another compound, such as pregnanediol, is used to detect pregnancy, a comment or opinion shall be added to the test report indicating that the test was performed and that the result was not consistent with pregnancy.

3.3.3 If tetrahydronorethisterone (metabolite of norethisterone contraceptive) is not detected in the urine Sample, a comment or opinion shall be added to the test report indicating that the test was performed and that norethisterone was not present. An example would be:

"The Sample was found not to contain tetrahydronorethisterone by GC/MS analysis; indicating that the finding is not the result of an administration of a norethisterone contraceptive."

### 3.4 19-demethylation

In the rare event that a Sample is found to meet the common features of an “unstable” urine and therefore tested as in Section 2.2 above, the Laboratory shall include the results of the stability test(s) in the test report for the 19-NA Adverse Analytical Finding. Therefore, a comment shall be on the test report indicating that the stability test was performed and that the Sample is stable. In addition, the results of the GC/C/IRMS analysis, including the δ values for 19-NA and androsterone and the Δδ value shall be included in the test report.

\(^3\) Thus the Decision Limit for the 19-NA test result for a Sample with a specific gravity of 1.020 or less is 2.5 ng/mL. When the specific gravity of a Sample is 1.030, for example, the adjusted threshold is 3.0 ng/mL and DL shall be 3.5 ng/mL (according to TD2010DL).
4.0 References

