PROJECT REVIEW

“Analysis of 19-Norsteroids, Testosterone and Precursors Metabolites in Human Urine”

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Detecting the use of androgenic anabolic steroids, potentially endogenous in humans, and which are prohibited substances in sport doping control programmes, still represents a major challenge to the analysts. These steroids include testosterone, its precursors androstenedione and dehydroepiandrosterone, and the 19-norsteroids equivalents, some of which are commercially available for oral self-administration.

This project is aimed at applying the IRMS (isotope ratio mass spectrometry) to the detection of the “natural” testosterone and 19-nortestosterone anabolic agents. Complement of the existing GC/MS methods, the GCIC/IRMS permits the differentiation of the exogenous or endogenous origin of urinary androgens metabolites, by measuring the isotopic content of their carbon atoms. That novel application of an otherwise well-known technique, now requires its validation in different experienced laboratories, the determination of international reference ranges of the metabolites of endogenous origin and the documentation of the changes commonly observed following the administration of these steroids. Ultimately, one of its most direct outcomes will be to verify that although different methodological approaches and different equipment are used, the same determination is made from the analysis of common specimens.

Testosterone and 19-nortestosterone (nandrolone) are potent androgenic anabolic steroids of known abuse in sport. Anabolic steroids are banned in Olympic sports for more than 20 years and since 1986, the highest number of positive cases reported are due to these two steroids. The administration of testosterone and its precursors, androstenedione and DHEA has been described to significantly alter the parameters of the urinary androgens steroid profile measured by GC/MS. The administration of testosterone is first detected in human urine by the GC/MS measurement of a testosterone/epitestosterone (TIE) value higher than 6, which is caused by the relative increase of excreted testosterone glucuronide (Donike, 1983)). The oral intake of androstenedione and DHEA was shown to transiently increase the excreted T/E value in females and in some male volunteers, from whom T/E values slightly higher than one were measured (Uralets (1999); Lévesque (2000); Bowers (1999); Garle (1998)). Other alterations of the urinary steroid profile, such as abnormally high concentration of androsterone and etiocholanolone and the presence of the characteristic hydroxylated metabolites glucuro- and sulfoconjugated, 6c~ androstenedione, 6P-epiandrosterone, had permit to report positive findings (Lévesque, 1999).

Disruption of the normal urinary profiles of androgens metabolites can be
demonstrated by comparison to the described population reference ranges and to
the individual’s norm (Donike (1993); Ayotte (1997) and reference cited therein).
That requires the investigation of the athlete’s previous or subsequent tests results
in order to exclude the few individuals who naturally produce urine samples in
which elevated $TIE$ values are systematically measured. Although successfully
applied in many cases, this method is time-consuming and complex. Considering
only the T/E values above 6 also leads to false negative results since it is known
that the limit will not be exceeded following the administration of testosterone and
precursors, when the basal values are lower than one, which is a characteristic but
not exclusively, of the Asian population (Shackleton, (1997)). The level of
androgens in female samples is generally very low and the uncertainty of the
measurements may represent a problem to which must be add reports of alteration
of the normal values attributed to other sources than the administration of
androgens.

The administration of 19-nortestosterone and of its precursors, 19-
norandrostenedione and 19- norandrostenediol, which are available for oral self-
administration, results in the excretion of 19- norandrosterone and 19-
noretiocholanolone, mostly found in the glucuroconjugated form. The period, during
which the metabolites can be detected, is drastically reduced when the oral
preparations are taken (Engel (1958); Masse (1985); Ayotte (1996); Schanzer
(1996); Kintz (1999)). In the last years, many positive cases were reported with
low levels of the urinary metabolites. Extremely low levels of 19-norandrosterone
can be endogenously excreted in human urine, and that has prompted the IOC to
safely recommend a threshold in males and females. However, as it is the case with
the androgens, natural factors are systematically invoked to challenge the positive
test results.
Analysis of Testosterone and Precursors Metabolites in Human Urine by GC/C/IRMS: a comparative inter-laboratory study.

Results and Conclusions

The three laboratories participating to the project have years of experience in testing urinary steroids by GC/MS and by GC/C/IRMS. For the latter, they have developed and validated techniques which make use of different instruments, different sample preparation and analytical methods. The internal urinary reference steroids also differ. Nothing has been changed to the laboratory validated protocols. However, having observed in the early phases of the project that one laboratory had values differing significantly from the two others when authentic standards were analysed, the project was halted until the necessary verifications and adjustments were made.

The results indicate that similar conclusions are reached by the three laboratories when sharing urine samples. It further confirms the need to consider and compare the difference of delta values between the intact and altered urinary metabolites for each sample and not the absolute individual values which were found to vary. This does not limit in any way the applicability of the technique since already only the difference of $^{13}$C/$^{12}$C values (delta values per mil) is significant in individual samples, allowing for individual variations that could be due to external environmental factors.

As an example, absolute mean $^{13}$C/$^{12}$C values of alcanes of know and certified isotopic content and authentic standards of different steroids were found to vary in the three laboratories by less than 0.53 and up to 1.6 0/00 respectively. When urine samples were shared, we observed that the absolute delta values of some steroids could vary up to 2.3 0/00. However, when the differences of the values between the metabolites and the reference steroids were compared, before and after the administration of the testosterone precursor, coefficients of variation of less than 21% were obtained for the main urinary metabolites. In all three laboratories, significant alterations of the $^{13}$C/$^{12}$C values of androsterone and etiocholanolone (after the hydrolysis of the glucuronide) were recorded in samples collected following the administration of testosterone, androstenedione and DHEA. The values measured in reference steroids e.g., pregnandiol, pregnantriol and cholesterol remained stable, as expected.