To the editor:

False-positive Epo test concerns unfounded

The brief report by Beullens et al is misleading regarding the urine test that the World Anti-Doping Agency (WADA) uses to detect recombinant human erythropoietin (rhEpo). The WADA-recommended test is based on immunoelectrophoresis and double blotting (IEF/DB), and was developed by Lasne and de Ceaurriz in 2000.

Our WADA-accredited laboratory has performed the IEF/DB test for rhEpo on more than 6800 urine samples, including more than 2600 doping control samples from athletes. Of the latter, we have reported 9 positive cases for rhEpo: 3 of these have publicly confessed to using rhEpo, 3 have accepted penalties, the physician of a seventh has been indicted for distribution of rhEpo, and 2 maintained their innocence but lost on appeal.

We take issue with Beullens’s use of the term “false positive” because, as the authors emphasize, the compound they are discussing is not rhEpo. If the compound detected can be identified as not rhEpo, then it cannot cause a false positive. This term sensationalizes an otherwise interesting case report that could in due course contribute to the body of science.

The criteria used by the WADA laboratories are well known and readily available. Beullens et al do not state the criteria they used to make the “false-positive” claim. Using the WADA criteria, the “false-positive” electropherogram is clearly negative. Moreover, they do not include a negative or a positive urine quality-control sample. Forensic testing results are normally accompanied by a comprehensive documentation package that supports the conclusion. A report such as this that raises a profound issue (false accusations against an athlete) at a minimum requires far more documentation.

Another important but unexplained issue is the nature of the compound that appears to migrate in the same general region as rhEpo and is characterized by bands. The pH range of the ampholytes is needed in order to fully interpret the data. In the left panel of their Figure 1A, the epoetin-β lane shows 3 faint bands and possibly a very weak fourth band. The bands in the darbepoetin lane are overly dense. Knowing the pH range of the ampholytes might explain why the darbepoetin region seems closer to the rhHuEPO region than we customarily observe (compare with Figure 1 here).

The right panel of their Figure 1A shows an apparent protein with bands, but it does not look like a typical rhEpo positive (Figure 1 here). Further, 1 and maybe 2 of the bands migrate more basically than the most basic epoetin-β band. Under the WADA rules, the identification criteria are not met.

Finally, the athlete has a puzzling renal disease characterized by a concentrating defect and an excessive number of casts that apparently does not interfere with his athletic prowess. He should have a full nephrology evaluation.

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References


To the editor:

No doubt about the validity of the urine test for detection of recombinant human erythropoietin

Beullens et al report the “false-positive detection of recombinant human erythropoietin in urine following strenuous physical exercise.” This report, based on observations conducted on urine from 1 single subject, relies in fact on serious errors of interpretation of poor-quality images. A first sample, collected just after exercise and analyzed by double-blotting following double-electric focusing of the retentate from ultrafiltrated urine, gives rise to a banding pattern interpreted as unrelated to Epo based on the argument that this pattern is missing in a second sample collected 1 hour later. A simple routine assay (using other antibodies than the AE7A5 used for immunoblot) of the Epo level in these 2 ultrafiltrated samples before IEF would have probably shown that a high concentration of this hormone was present in the first one but not in the second one. It is quite surprising that this basic control
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