ERYTHROPOIETIN (Epo) may have effects on exercise capacity and physiological regulation beyond a simple increase in red cell mass and the associated improvement in oxygen transport (4). In the context of a larger study on this topic, Lundby and colleagues (11) also asked questions about the reliability of urine testing for recombinant human Epo (rHuEpo). They studied eight healthy male subjects during a 4-wk “loading” and 2-wk “boosting” phase of Epo use followed by a 2-wk maintenance phase. In the parent study they showed that the effects of Epo on exercise performance were confined to its impact on red cell mass and not to other physiological effects of the hormone. These results were consistent with ideas about the relationship between maximal oxygen uptake and red cell mass or total body hemoglobin that emerged in the 1950s. The findings are timely and have implications for public policy relating to the control of doping practices. In this short report a number of challenges related to urine testing for Epo are highlighted.

Testing for recombinant Epo in urine may seem practical at first sight but appears to be a very difficult task. The amount of endogenous Epo in urine is extremely low (5). The physiological background for testing Epo in urine is complex and the handling of Epo by the renal tubules is poorly understood (16). Furthermore, exercise-induced renal ischemia and the accompanying postexercise proteinuria may affect the clearance of this 32- to 39-kDa protein and the quality of the urine matrix.

The Epo test that has been adopted in World Anti-Doping Agency (WADA)-accredited laboratories is based on isoelectric focusing (2, 10). Since the introduction of the test in 2000, Epo-abusing athletes have altered their dosing schemes, proving the initial efficiency of the test. By injecting microdoses of rHuEpo, the window of detection can be reduced to as little as 12–18 h postinjection (8).

Several problems (e.g., the lengthy sample preparation, the low sample load capacity, difficulties with interlaboratory standardization, non-specific binding of the secondary antibody to urinary proteins, sensitivity issues) were already identified in a WADA-commissioned report in 2003 (14). The test requires a 700- to 1,000-fold concentration of the specimen before analysis can be carried out, and the concentrated urine forms a pellet that is difficult to solubilize.

Despite this enormous concentration factor, up to 20% of the investigated samples do not show detectable Epo (14). The non-specific interaction of the used monoclonal antibody with human, bacterial, and yeast proteins is worrisome (4, 6). Epo test results are clearly not always interpreted identically (4). The use of the software processing has been criticized (6).

The American WADA-accredited laboratory has performed the direct Epo test on more than 2,600 samples, only nine of them were found to be positive (3). The low numbers of athletes caught by the test are somewhat contradictory to the overall increase of mean hematocrit values since rHuEpo became available (12). Additionally, in some high profile legal cases in the United States, athletes who were clearly doping with a variety of compounds including Epo “passed” hundreds of individual drug tests.

Along these lines, Lundby et al. (11) convincingly demonstrated that the performance of the urinary Epo test is somewhat disappointing. Although the judgment process of “real” doping cases differs from the one applied in the present study, the high number of false-negative results imply a risk that athletes doping with Epo will avoid detection and damage the fundamental goal of fair competition. The earlier reported flaws of the test help to understand the relatively low efficiency of the direct Epo test and the current results emphasize the need for improving Epo testing.

The detection window for Epo shows an interindividual variation because the actual positivity criteria take into consideration the endogenous Epo production rate, which varies enormously between individuals. In subjects with a naturally elevated or stimulated Epo production rate (altitude training, hypoxic tent, etc.), there is a reduced detection window. The positivity criteria used by anti-doping laboratories are strict and (in the case of the Epo analog darbepoietin) could be adapted by only taking into account the position and the specific distribution of the bands in the most acidic area of the gel and no longer the intensity of the bands. This would rule out the differences in interpretation of the test due to different endogenous Epo levels in individual athletes (8). While the performance of the existing test can likely be improved by paying more attention to the preanalytical care after prelevation of urine specimens (4), there also is concern about protease-treated urine specimens that could mask Epo abuse (9).

On the other hand, blood-based indirect Epo tests have a better physiological basis and offer the advantage that other kinds of blood doping can also be detected (1, 13, 15). This approach focuses on consistent tracking and establishes upper limits of normal permitted for competition. The so-called “passport” approach will be further facilitated by the increased availability of well-validated mobile hematological analysis equipment. Longitudinal monitoring of blood profiles and comparing an athlete’s individual hematological values against his or her own historical baseline rather than a population-derived threshold further enhances the potency of indirect testing (1). In the Union Cycliste International (UCI), the governing body of cycling, anti-doping program “100% Against Doping,” an individual hematological profile is created. However, indirect Epo/blood doping detection methods require blood sampling, which is a practical disadvantage. Additionally, various dietary or saline infusion strategies for the purposes of short-term hemodilution are likely being practiced to circumvent the “passport” approach prior to important compe-
tions (7). Does this mean that a combination of regular tracking tests with random elements along with a marker of whole body fluid status is needed?

In summary, the data provided by Lundby et al. demonstrate that an improvement in the current Epo test is necessary or that a different strategy to detect Epo use and blood doping should be considered. Blood-based indirect Epo tests offer an interesting alternative. The rapidly changing blood doping landscape will definitely encourage the use of indirect Epo/blood doping testing with a much broader application.

REFERENCES


