Creatine clearance in anorexia nervosa

We read with interest the recent article by Boag et al1 concerning abnormally low creatinine clearance in untreated patients with anorexia nervosa. From their data the authors assumed that these patients had a reversible impairment of renal function. We, however, offer an alternative explanation for the low creatinine clearance.

In normal subjects creatinine clearance calculated from the serum and 24 hour urinary creatinine measured by the Jaffe reaction closely approximates the glomerular filtration rate. This close approximation, however, is due to two compensating errors: firstly, in normal subjects 2-3 mmol (0.23-0.34 g), representing 20-30% of the urinary creatinine, is from tubular secretion rather than glomerular filtration; and secondly, 15-35 μmol/l (0.17-0.40 mg/100 ml), representing 15-30% of the serum creatinine as measured by conventional Jaffe methods, is from non-creatinine chromogens.2 Less than 5% of urinary creatinine measured by the Jaffe reaction is due to these non-creatinine chromogens. The two errors cancel each other on calculation of creatinine clearance, as one is in the numerator and the other is in the denominator of the clearance equation.

In various pathological conditions the relative magnitude of these errors no longer quantitatively compensate each other, and creatinine clearance becomes a poor predictor of glomerular filtration rate. In severe renal failure creatinine clearance gives a significant overestimate of the glomerular filtration rate. Although tubular secretion still accounts for 20-30% of urinary creatinine, serum non-creatinine chromogens still account for 15-35 μmol/l (0.17-0.40 mg/100 ml) of the serum creatinine (Jaffe reaction) and now represent <5% of the total serum creatinine.3 In patients with low muscle mass who have low serum and 24 hour urinary creatinine creatinine clearance gives a significant underestimate of the glomerular filtration rate. In these patients the Jaffe reaction still measures urinary creatinine relatively accurately, but now the non-creatinine chromogens in the serum may represent 50% or more of the serum creatinine (measured by Jaffe reaction). A particularly dramatic case, which we encountered several years ago, was a woman weighing 45 kg with congenital anterior horn cell disease and severe wasting of the muscles. Her 24 hour urinary creatinine was only 1.2 mmol/l (136 mg) and her serum creatinine was 35 μmol/l (0.040 mg/100 ml) by a Jaffe method, yielding a creatinine clearance of 25 ml/min. Her serum urea concentration was normal at 3-9 mmol/l (23 mg/100 ml). Her creatinine clearance was artefactually low because non-creatinine chromogens accounted for >60% of her serum creatinine measured with the Jaffe reaction, and her renal function was normal.

Among the principal non-creatinine chromogens that react to a Jaffe method in serum are various ketoacids.2 In patients with diabetic ketoacidosis and a normal fasting subject raised acetoacetate concentrations cause spurious increases of serum creatinine measured by the Jaffe method and thus spuriously low creatinine clearances. With prolonged starvation as seen with anorexia nervosa relatively severe ketosis would be expected.4 Consequently, the serum Jaffe creatinine would be artefactually high. In the work reported by Boag et al sera for creatinine determinations were collected at 8.00 am. How long the patients had been fed a high energy diet before the "before treatment" creatinine clearances were determined is not clear. Even if the patients had been fed for several days, however, with their severely depleted muscle mass and lack of gluconeogenic substrates they might have become reasonably ketotic with fasting lasting as little as one night. If ketosis did occur serum creatinine measured with the Jaffe reaction would have been artefactually high, and this would contribute to the artefactual lowering of the creatinine clearance. Consequently, without more reliable measurements of glomerular filtration rates, such as inulin clearance, in at least some patients, or documentation of the absence of serum ketones, low creatinine clearance measured by Jaffe methods in patients with untreated anorexia nervosa should not be assumed to indicate renal dysfunction.

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References

Drs Boag and Dandona reply as follows:

We thank Drs Eckfeldt and Freier for pointing out the limitations of indices of renal function based on creatinine clearance, especially in cases in which the turnover of creatinine is likely to fall and ketogenesis is likely to produce chromogenic reactions that mimic those of creatinine.

None of our patients had ketonuria when we began to measure their plasma creatinine concentration and urinary creatinine excretion.1 Whether this was the result of the initial feeding and "stabilisation" for 24 to 72 hours, which may suppress lipolysis and ketogenesis, or whether such patients have switched off lipolysis and ketogenesis through some other metabolic mechanism is not clear at present. In this context it is worth noting that patients with anorexia nervosa do not have raised free fatty acid concentrations.3 This was also true of a series of patients with anorexia nervosa whom we investigated, their free fatty acid concentrations (565 μmol/l) were comparable with those in controls (654 μmol/l).4 Insulin receptor population is known to be increased in anorexia nervosa,5 possibly, enhanced sensitivity to insulin may somehow "protect" such patients from lipolysis and ketogenesis. The concentrations of acetoacetate required to cause spurious increases of plasma creatinine concentrations are far higher than those expected in patients with anorexia nervosa.

Finally, we now have data to show that glomerular filtration rate as measured by excretion of edetic acid labelled with 54chromium is also greatly diminished in...
these patients and that it increases with weight gain.

In conclusion, Eckfeldt and Freier have raised an interesting theoretical point regarding the low creatinine clearance in patients with anorexia nervosa. Our patients were, however, not ketotic and therefore Eckfeldt's and Freier's objection is not directly relevant. Furthermore, we are all too aware of the fallacies of basing indices of renal function on creatinine clearance, and have taken care to show that the true glomerular filtration rate based on clearance of edetic acid labelled with $^{51}$chromium is also diminished in these patients.

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References


Electron Microscopy in Human Medicine.  

This is the penultimate book in this massive 12 volume series (for some curious reason the last two volumes are to be called 11 (a) and 11 (b) rather than 11 and 12), the format being the same as the earlier volumes. Professor Hashimoto is the editor of this volume and there are 11 other contributors, mainly from Japan and the United States. The material of the book consists of well illustrated ultrastructural descriptions of skin pathology under the headings of neoplastic conditions; bullous dermatoses; hereditary ichthyoses; hair abnormalities; and viral infections, together with a chapter on the freeze-fracture technique as applied to the skin. No doubt, any institution that has already collected the previous volumes will want to add this one to the set and it will be a valuable reference work for pathologists interested in dermatology. What is missing, however, is a clear indication as to the conditions for which electron microscopy is likely to be diagnostically useful and which ultrastructural features are simply of academic interest.

JULIE CROW

Laboratory Manual of Histochemistry.  

This is traditional histological technique in modern guise. Loose leaved, spiral bound, the manual is set in typescript of differing styles and pitch that change haphazardly within the same section or the same page. The illustrations are restricted to line diagrams and a few photographs that lack lustre. Some may feel the title to be pretentious; only one tenth of this book is concerned with enzyme histochemistry, and immunohistochemistry is briefly described in 23 pages. The rest of this book is a manual on tissue preparation, section cutting, and dye staining. Lillie and Puchter influenced the work of the author and their papers dominate the bibliography.

Overall, this manual is comprehensive and seems to be useful and reliable, but its success on this side of the Atlantic will depend on its ability to rise above the lecture note presentation and suspect standing in laboratory use.

RAB DRURY

Arthritis and Allied Conditions.  

This new edition to one of the best and most well known major rheumatology textbooks is published six years after its predecessor. Its length, over 1700 pages, permits a comprehensive review of virtually all rheumatological conditions. There are 104 chapters written by over 100 contributors who comprise the best of North American rheumatologists. Unfortunately, the absence of any British, or indeed European contributors, gives a rather too severe American flavour. By any standards, however, this is an excellent synopsis of rheumatic diseases.

The book is well indexed and contains key references within the text. Its outline is similar to the previous edition. It comprises an introductory section on the epidemiology and differential diagnosis of arthritis; a relatively detailed account of some of the scientific pillars of current rheumatological thought; a brief resume on antirheumatic drugs; and then sections on the major rheumatic diseases. The book concentrates on clinical aspects rather than giving detailed accounts of laboratory techniques. For the practising pathologist it will provide an ever ready source of clinical information to place in the context of pathological findings.

It is a book to dip into rather than read exhaustively. I found it simple to review rapidly diverse subjects; for example, can