Arguments for an age-adapted definition of chronic kidney disease

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Defining a disease or a pathological status is not easy. For many pauci-symptomatic diseases, biological results are of outstanding importance. Most clinicians are scarcely aware how much laboratories have to do to establish such reference or “normal” values. Laboratories have to follow different complex procedures. In the same vein, laboratory specialists well know that every biological result given to the clinicians is vitiated by an inevitable error, known as the measurement uncertainty. On the other hand, clinicians too often consider laboratory results as Gospel words carved in stone. The current definition of chronic kidney disease (CKD) is no exception. Indeed, Kidney Disease: Improving Global Outcomes (KDIGO) guidelines are recommending that all subjects with estimated glomerular filtration rate eGFR below 60 mL/min/1.73m² should be considered as a CKD patient. The drawbacks arising from the use of fixed thresholds must be considered. An estimation of 59 mL/min/1.73m² will be too frequently considered as CKD, whereas a not-so-different value of 61 mL/min/1.73m² will not. Only the uncertainty measurement surrounding the analytical determination of creatinine could be enough to argue that these two values are actually not different (because sex and gender, the other variables in eGFR equations, are quite solid variables, most of the time). To help in the interpretation of an eGFR result, it is important to underline that the KDIGO also recommend that a low eGFR value must be confirmed at three months. This recommendation is important, and often neglected in epidemiological studies, to exclude patients with acute kidney injury, but repeating the measurement also helps to better diagnose these patients with an eGFR at the limit of the threshold.

Beyond the inherent uncertainty of any biological measurement, one must bear in mind that, regarding a CKD diagnosis, an estimation, eGFR, with an equation is considered. And this estimation can be of low precision. An eGFR equation is considered as accurate when the relative difference with measured GFR is within 30%. This means that, for a threshold at 60 mL/min/1.73m², a value of 42 or 78 mL/min/1.73m² will be too frequently considered as CKD, whereas a not-so-different value of 61 mL/min/1.73m² will not. Only the uncertainty measurement surrounding the analytical determination of creatinine could be enough to argue that these two values are actually not different (because sex and gender, the other variables in eGFR equations, are quite solid variables, most of the time). To help in the interpretation of an eGFR result, it is important to underline that the KDIGO also recommend that a low eGFR value must be confirmed at three months. This recommendation is important, and often neglected in epidemiological studies, to exclude patients with acute kidney injury, but repeating the measurement also helps to better diagnose these patients with an eGFR at the limit of the threshold.

The choice of a fixed threshold of 60 mL/min/1.73m² has the advantage of simplicity. That is undeniable. However, beyond the limitations inherent in any threshold, the choice of a fixed threshold for CKD definition has other specific limitations that might be strongly challenged. The first is the very well-known normal physiological decline in GFR with aging. Several studies have shown that after 40 years, GFR declines with aging. This means that a large proportion (up to 25%) of healthy subjects over 75 years of age could be diagnosed as having CKD. Importantly, this decline in GFR has been illustrated with measured GFR and with estimating GFR (using different equations) in the world’s population. Percentiles of eGFR or measured GFR do thus exist and are available for clinical use7,8. One main argument of the KDIGO for keeping the fixed GFR threshold at 60 mL/min/1.73m² is an epidemiological one. Many large-scale studies from the Chronic Kidney Disease epidemiology (CKD-EPI) consortium have claimed that mortality was significantly higher in general or high-risk populations when eGFR was below 60 mL/min/1.73m², whatever the age considered9. However, all these epidemiological studies shared the same limitation, as they have a control group with an eGFR around 90 mL/min/1.73m². An analysis of the same database according to age categories showed that a different group control should be considered according to age. Indeed, the “normal” GFR range, corresponding to the GFR associated with the lowest mortality rate, is observed in higher GFR ranges in young people than in old populations. Calculation of risk with such a methodology confirm that a higher mortality risk is observed when eGFR is below 45 mL/min/1.73m² in subjects older than 65 years; below 60 mL/min/1.73m² in subjects between 40 and 65 years, and below 75 mL/min/1.73m² in subjects younger than 40 years3. Several authors claim for an age-adapted CKD definition3,10. Using GFR percentiles is the best scientific solution for such an age-adapted definition. Using percentiles is less impacted by the measurement uncertainty. Percentiles are also very useful for early detection of a “brake” in the slope of GFR of a given patient (even if the GFR value is in the normal range). Percentiles could potentially help in the detection of hyperfiltration. It is important to note that such percentiles are useful in the early detection of CKD in young subjects, as the threshold at 60 mL/min/1.73m² seems totally unsuitable and much too low in this specific population3,11. At the population level, the use of such percentiles will profoundly impact the epidemiology of CKD, with a large decrease of CKD prevalence (up to 30 to 50%) in elderly populations, but an increase of CKD prevalence in young populations. However, the impact of this increase prevalence in young populations will have a limited impact on the global crude CKD prevalence because, hopefully, the prevalence of CKD in young people is low12,13.

In this perspectives article, we focused on GFR and its interpretation. We must keep in mind that there is another key parameter, namely the measurement of urine albumin-to-creatinine ratio (ACR), for the diagnosis of ACR. Interpretation of ACR has its own caveats, but it remains very important to measure ACR in patients at risk. As
an example, ACR is still too frequently forgotten in diabetic patients, whereas the presence of abnormal albuminuria level is enough for a CKD diagnosis, whatever the GFR level.

Interpretation of GFR percentiles is thus easy and can help the nephrologists in many situations\(^{11}\). The results of percentiles must ideally be interpreted with an ACR result to establish the renal risk and propose a dedicated therapy\(^{11}\).

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