Screening for Chronic Kidney Disease: Clinical Benefit and Burden

LAURENCE E. CARROLL, M.D., F.A.S.N.
Hypertension and Kidney Specialists

BACKGROUND
I have previously written in this Journal about the role of the estimated Glomerular Filtration Rate (eGFR) calculated from the Modification of Diet in Renal Disease (MDRD) equation, and the associated classification of Chronic Kidney Disease (CKD) into Stages 1-4. That article was presented in response to the 2002 Kidney Disease Outcomes Quality Initiative (KDOQI), a project of the National Kidney Foundation and the broad-based "kidney community," which defined those Stages.

In January 2005, Lancaster General Hospital added a creatinine-based calculated eGFR value to its blood chemistry panels (BMPs & CMPs). Since then, Pennsylvania and other states have mandated that all clinical laboratories must provide eGFRs, with the result that a national initiative has begun for a "standardized creatinine," to try to provide eGFR results that have greater accuracy and reproducibility.

BENEFIT
All this activity has raised awareness of CKD and has helped to improve care by allowing earlier referral and the development of new CKD Clinics. Aggressive management of anemia, hyperparathyroidism, and Vitamin D deficiency has become more common, and use of arteriovenous fistulas (AVF) has increased. AVFs are the optimal vascular access for hemodialysis, and their use now exceeds the Healthy People 2010 goal of 50%. (Our local use is >70% - the highest in the Regional Kidney Network.) We're hoping these achievements will pay dividends in future years by lowering mortality rates, especially in Cardiovascular Disease (CVD) for which CKD is now accepted to be an independent risk factor.

The controversy over Stage 3 CKD starts with its definition: an eGFR of 30-60 cc/min, with no requirement for an abnormal imaging study (usually Kidney Ultrasound) or urinary abnormality (albuminuria/proteinuria, hematuria, or both). Critics point out that age related decreases in GFR and kidney function have been known for over 50 years, and that previous population studies have suggested a 10% decrease in GFR every 10 years after the third decade of life. This means that by eGFR alone, 38% of individuals over 70 years will be classified as having CKD.

Since only 24% of Stage 3 CKD patients have albuminuria/proteinuria, 76% of these patients are in this category only because of an eGFR of 30-60 cc/min – a finding that may be more the result of aging than of an actual "disease."

THERAPEUTIC FOCUS
The absence of albuminuria/proteinuria is an important consideration, because it appears to be a critical marker for the risk of progressive CKD in most diabetics and other patients. A recent Chinese study showed that when Stage 1-2 CKD (GFR >60 cc/min) was associated with albuminuria the risk of developing progressive CKD was increased; without albuminuria, the risk was not increased even in Stage 3 CKD. Previous studies in diabetics (the RENAAL trial - Reduction ofEndpoints in Non-Insulin Dependent Diabetes Mellitus with Angiotensin II Antagonist Losartan) have shown that proteinuria, especially above 1 gram urinary protein...
per gram of urinary creatinine, indicates a high risk for progressive CKD. Reduction of proteinuria in diabetic kidney disease has been shown to be beneficial. In the non-diabetic population, the use of angiotensin converting enzyme (ACE) inhibitors, angiotensin receptor blockers (ARBs), and aldosterone blockers (alone or in combination), is a matter of ongoing studies.

**RECOMMENDATIONS**

The protocol that accompanies this article, “Primary Care in Early Chronic Kidney Disease,” is based on the Kidney Early Evaluation Program of the National Kidney Foundation’s (KEEP-NKF) (Table 1); the 2008 Guidelines for CKD of the Royal College of Physicians; and some of my personal opinions. It designates specific populations to screen, and the survey tools to use: blood pressure (BP), serum creatinine, eGFR, urinalysis, and urine albumin/creatinine ratio (ACR). An appendix will be made available at the LGH website for normal values and further information.

Although some have proposed that only stage 4 CKD patients should be referred to a nephrologist to prepare them for dialysis or transplant, such a plan could delay therapy for potentially reversible glomerulonephritis or vasculitis. The protocol therefore focuses on two factors that differentiate patients who need to be referred for specialty care from those who only need to be followed in the primary care office. First, the protocol suggests that all patients with “glomerular” hematuria (dysmorphic RBCs in the urine and RBC casts), should be referred for evaluation, at which time kidney biopsy should be considered. Those with significant proteinuria should be seen with particular urgency.

Second, referral to nephrology is not necessary in older patients with Stage 3 CKD if they have no structural abnormality on imaging studies, have a normal urinalysis, a normal albumin/creatinine ratio (ACR) or normal microalbuminuria (abnormal >30 mcg/mg creatinine). They can be followed with measurements of blood pressure and serum creatinine at 3–6 month intervals along with the other primary care goals stated.

**CONCLUSIONS**

Nephrology needs to partner with primary care providers to deal with the burden of CKD in our society. Using the tools available such as this protocol, we should be able to provide appropriate and effective care to enhance outcomes. These measures will slow progression to the more advanced Stages of CKD, where costly and somewhat limited renal replacement therapies become the only alternative.
REFERENCES


Neither Dr. Carroll nor any member of his immediate family have any relevant relationships to disclose with any corporate organizations associated with the manufacture, license, sale, distribution or promotion of a drug or device.

Laurence E. Carroll, M.D., F.A.S.N.
Hypertension Kidney Specialists
2112 Harrisburg Pike - Suite 2112
Lancaster, PA 17604
717-544-3232
lecjtc@comcast.net