


## Hidden in Plain Sight: Autosomal Dominant Tubulointerstitial Kidney Disease



The banner features a dark blue background with a grid pattern. On the left, it includes the Critical Path Institute logo and text: 'RARE AND ORPHAN DISEASE PROGRAMS' and 'CRITICAL PATH INSTITUTE'. Next to it is the 'RDCA-DAP' logo with the tagline 'Rare Disease Cures Accelerator Data and Analytics Platform'. The central text reads 'WEBINAR SERIES' in large white letters, followed by the title 'Hidden in Plain Sight: Autosomal Dominant Tubulointerstitial Kidney Disease' in yellow and white. A portrait of Dr. Anthony Bleyer is shown on the right. Below the portrait, his name and title are listed: 'Anthony Bleyer, MD, MS' and 'Professor, Nephrology, Wake Forest University School of Medicine'. A yellow box in the top right corner contains the date and time: 'Thurs | June 20 12 PM ET' and a 'REGISTER NOW' button.

C-Path's Rare and Orphan Disease Programs is excited to have Dr. [Anthony Bleyer](#) present it's June webinar, **“Hidden in Plain Sight: Autosomal Dominant Tubulointerstitial Kidney Disease (ADTKD).”** Dr. Bleyer is a Professor of Internal Medicine/Nephrology at Wake Forest University School of Medicine, with adjunct appointments at the Broad Institute of Harvard Medical School and Massachusetts Institute of Technology, and the Institute for Inherited Metabolic Disorders at Charles University, Czech Republic.

Dr. Bleyer led the team that identified mutations in the *UMOD* gene as a cause of ADTKD and has been leading the Wake Forest Rare Inherited Kidney Disease team for the last 25 years. His presentation will cover the genetics of ADTKD, its biology and mechanisms, a description of the Wake Forest registry, and unmet drug development needs.

Join us for this valuable opportunity to gain insights from an expert in the field.

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