



Genomics Education Programme

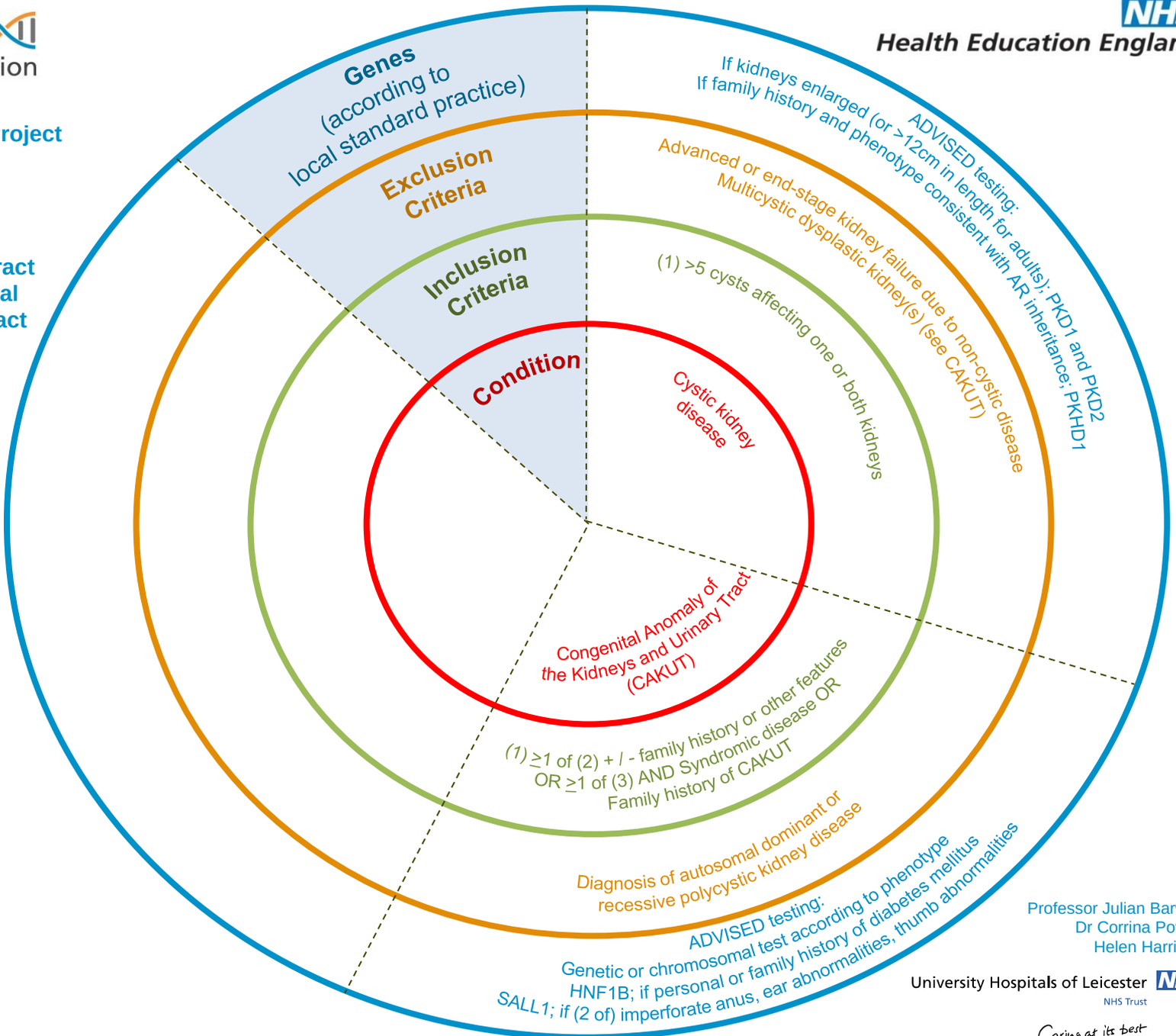
100,000 Genomes Project Eligibility wheels A tool for clinicians

Rare Diseases: Renal and urinary tract disorders - Structural renal and urinary tract disease

(1) Recruitment should NOT include unaffected relatives (normal screening) unless part of a TRIO/DUO. Multiple affected relatives are preferable to singleton cases.

- (2)
- Renal hypodysplasia or agenesis
 - Congenital multicystic kidney
 - Congenital, persistent, severe hydroneurter
 - Congenital, persistent, severe congenital hydronephrosis
 - Bladder exstrophy
 - Posterior urethral valves

- (3)
- Duplex kidney
 - Vesicoureteric reflux
 - Vesicoureteric or ureteropelvic junction obstruction



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Caring at its best