Genetics in chronic kidney disease: conclusions from a Kidney Disease: Improving Global Outcomes (KDIGO) Controversies Conference

KDIGO Conference Participants

Numerous genes for monogenic kidney diseases with classical patterns of inheritance, as well as genes for complex kidney diseases that manifest in combination with environmental factors, have been discovered. Genetic findings are increasingly used to inform clinical management of nephropathies, and have led to improved diagnostics, disease surveillance, choice of therapy, and family counseling. All of these steps rely on accurate interpretation of genetic data, which can be outpaced by current rates of data collection. In March of 2021, Kidney Diseases: Improving Global Outcomes (KDIGO) held a Controversies Conference on “Genetics in Chronic Kidney Disease (CKD)” to review the current state of understanding of monogenic and complex (polygenic) kidney diseases, processes for applying genetic findings in clinical medicine, and use of genomics for defining and stratifying CKD. Given the important contribution of genetic variants to CKD, practitioners with CKD patients are advised to “think genetic,” which specifically involves obtaining a family history, collecting detailed information on age of CKD onset, performing clinical examination for extrarenal symptoms, and considering genetic testing. To improve the use of genetics in nephrology, meeting participants advised developing an advanced training or subspecialty track for nephrologists, crafting guidelines for testing and treatment, and educating patients, students, and practitioners. Key areas of future research, including clinical interpretation of genome variation, electronic phenotyping, global representation, kidney-specific molecular data, polygenic scores, translational epidemiology, and open data resources, were also identified.


KEYWORDS: genetic kidney disease; genetic testing; genome-wide association studies; monogenic; polygenic; single-nucleotide polymorphism

Chronic kidney disease (CKD) affects approximately 10% of the global adult population. Multiple genetic and environmental risk factors contribute to kidney diseases, making identification of the underlying pathophysiologic mechanisms difficult. However, the advent of high-throughput genotyping and massively parallel sequencing, combined with the availability of large datasets of genomic and health information, has led to rapid advances in our understanding of the genetic basis of kidney function and disease.

To date, more than 600 genes have been implicated in monogenic kidney diseases, and known single-gene disorders account for up to 50% of nondiabetic CKD in pediatric cohorts, and 30% in adult cohorts. In addition, genetic variation plays an important role for kidney function in the normal range, and common genetic variants account for approximately 20% of the estimated genetic heritability of estimated glomerular filtration rate (eGFR). Common genetic variants also have been shown to contribute to disorders, such as IgA nephropathy, membranous nephropathy, and nephrotic syndrome. Hence, the pathogenesis model for many kidney diseases has expanded to include multiple genetic and environmental factors that together contribute to the pathology, commonly referred to as “complex disease.”

Genetic findings increasingly are used to inform clinical management of many nephropathies, enabling more precise diagnostics, targeted disease surveillance, and better-informed choices for therapy and family counseling. Clinical management relies on accurate interpretation of genomic data, a labor-intensive process that can be outpaced by the speed of discovery. To realize the promises of genomic medicine for kidney disease, many technical, logistical, ethical, and scientific questions must be addressed. In March of 2021, Kidney Diseases: Improving Global Outcomes (KDIGO) held a Controversies Conference on the topic of “Genetics in CKD” to review the current state of understanding of monogenic and complex kidney diseases, processes for applying genetic findings in clinical medicine, and use of genomics for defining and stratifying CKD. Participants identified areas of consensus, gaps in knowledge, and priorities for research (Table 1). The conference agenda, discussion questions, and plenary session presentations are available on the KDIGO website: https://kdigo.org/conferences/genetics-in-ckd/.

Definitions and epidemiology of genetic kidney diseases

The familial aggregation and substantial heritability of CKD are well described across the world. Recent large-scale analyses of electronic medical records estimated observational

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The KDIGO Conference Participants are listed in the Appendix.
Table 1 | Summary points from the Genetics in CKD Controversies Conference

Consensus

- Monogenic and complex kidney diseases exist on a continuum, but dichotomous categories are useful for practical distinction.
- There is no upper age-limit for monogenic CKD.
- Actionable genes in kidney diseases refers to genes in which the identification of pathogenic variants can lead to specific clinical actions for treatment or prevention, following recommendations based on evidence.
- There is a need for development of a reference kidney disease gene list and standardization of gene/variant reporting for kidney diseases.
- A larger workforce with expertise in kidney genetics, genomics, and computational research is needed.
- Education of the workforce is necessary for successful implementation of genetic testing in clinical nephrology.
- New studies are needed that involve divergent populations worldwide to ensure equitable and generalizable implementation of genetic testing, obtain evidence of causality, establish global prevalence, and facilitate variant discovery.
- Interdisciplinary expert boards (including nephrologists, clinical geneticists, molecular biologists, genetic counselors) should be assembled to discuss potential genetic diagnostic findings and counsel primary and secondary care centers.
- Genomics should be integrated into clinical trials on kidney diseases.
- Estimates of the prevalence of monogenic CKD are important, but they are currently imprecise due to selection bias.

Ongoing controversies

Definitions/terminology

- Two-part names (clinical condition PLUS gene name) are preferred for more-precise disease terminology.
- The term CKD of unknown etiology is not clear and is in need of consensus.
- There is no clear consensus on which VUS are to be reported in the framework of diagnostic testing.

Processes for improving data capture and analysis

- Improve phenotyping, including methods for electronic phenotyping.
- Improve the quality of genomic studies, including analytical and computational methods.
- Improve data access while protecting the privacy of research participants.
- Create processes for transferring genetic information obtained through clinical testing to research.
- Study health–economic impacts of genetic testing in nephrology.
- Establish a process for periodic reanalysis of unsolved cases with kidney disease.
- Implement high-throughput techniques for in silico and in vitro variant characterization.
- Identify and characterize rare variants, structural variants, and functional variants using functional genomic, epigenetic, and other multi-omic approaches.
- Employ new approaches to identify more homogeneous CKD phenotypes and subclassifications for genetic studies, such as using nontraditional omics biomarkers, electronic health record data, imaging, or machine learning.
- Assemble larger cohorts with genetically defined kidney disease for both research and clinical trials; collaborate internationally if possible.
- Reduce measurement errors in eGFR and misclassification in the resulting CKD definition; for example, reassess coefficients based on race, sex, and chronological age in eGFR equations.
- Conduct large-scale genetic studies on specific kidney sub-phenotypes, such as CKD progression, acute kidney injury, cause-specific disease severity, and manifestations.
- Integrate genetic studies with biomarker and multi-omic profiling to leverage findings and increase power for both variant and pathway identification.
- Generate comprehensive molecular maps of kidney tissue/cells as well as in vitro and animal models to enable mechanistic studies of genes identified in GWAS of kidney traits.
- Encourage broad data sharing (FAIR principles; findable, accessible, interoperable, reusable) and transparent protocols for data generation, quality control, and analyses.
- Use federated networks to standardize key data elements across platforms and countries.
- Use portals (cloud-based) to “safely” share individual data and allow for democratization and a broader scale of integrative in silico analyses.
- Extend discovery analyses to nonadditive genetic models (e.g., recessive) and include nonautosomal regions (e.g., chromosome X, mitochondrial).
- Improve imputation reference panels.
- Apply and develop approaches specific to admixed populations.
- Conduct Mendelian randomization analysis to elucidate causal mechanisms.

Priorities for Implementation

- Increase genetic and genomic resources in underrepresented populations with kidney disease.
- Investigate the use of polygenic scores in clinical settings.
- Develop guidelines for nephrologist to establish core competencies in genetics, develop evaluations to test them, and identify the educational gaps of general nephrologists (some need to be country-specific).
- Develop and test the impact of dissemination tools to spread the basic knowledge required for all nephrologists.
- Measure the quality of existing or to-be-established genetic subspecialty training for nephrologists as well as training in nephrology for genetic counselors and molecular geneticists (variant interpretation side).
- Develop guidelines for the referral of nephrology patients to genetic counseling/genetic testing/reproductive counseling.
- Analyze the impact of genetic testing on clinical outcomes of nephrology patients.
- Analyze the cost-effectiveness and longitudinal clinical utility of genetic testing.
- Analyze the impact of centers of expertise on quality of care and patient outcomes.

CKD, chronic kidney disease; eGFR, estimated glomerular filtration rate; GWAS, genome-wide association studies; VUS, variants of uncertain significance.
Table 2 | Characteristics of monogenic versus complex genetic diseases

<table>
<thead>
<tr>
<th>Allele/variant frequency</th>
<th>Monogenic (Mendelian)</th>
<th>Polygenic (complex)</th>
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<tr>
<td>Effect size of major gene driving disease</td>
<td>Rare</td>
<td>Can be common</td>
</tr>
<tr>
<td>Penetrance</td>
<td>High</td>
<td>Low</td>
</tr>
<tr>
<td>Role of environment</td>
<td>Limited</td>
<td>Strong</td>
</tr>
<tr>
<td>Inheritance model</td>
<td>Mendelian</td>
<td>None apparent</td>
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heritability of CKD to be in the range of 25%–44%, with higher estimates for patients of African ancestry.12 These estimates are generally consistent with traditional family-based heritability studies of CKD and glomerular filtration rate.27–29 The relatively high heritability of CKD is likely attributable to both monogenic causes as well as complex or polygenic factors.

Monogenic (also termed “Mendelian”) CKD generally refers to diseases caused by rare, pathogenic variants in a single gene (Table 2); the genotype-to-phenotype relationship is strong, and environmental factors have limited influence. Oligogenic disorders are determined by rare variants in a few genes. Complex or polygenic diseases lack simple patterns of inheritance (e.g., dominant, recessive, or sex-linked) and instead are influenced by the aggregate effect of many common genetic variants in multiple genomic regions, as well as environmental factors.30 Such aggregate effects of common variants (or single-nucleotide polymorphisms [SNPs]) can be quantified by SNP-based heritability, which has been estimated for various types of kidney disorders to range from 14% for renal cancer among individuals of European ancestry to 43% for membranous nephropathy among individuals of East Asian ancestry. The proportions of variance explained by known loci of these diseases are smaller, ranging from <1% for urinary albumin-to-creatinine ratio to 32% for membranous nephropathy among individuals of East Asian ancestry (Figure 1).13,14,17,19,31,39 However, common genetic factors also may influence the age of onset, severity, rate of progression, and associated extrarenal complications of monogenic diseases, which often have variable expression.32,33 In addition to CKD attributed to specific etiologies, genetic studies also use phenotypic readouts, such as measures of kidney function or damage (e.g., eGFR, albuminuria), kidney histology classification, and molecular injury markers to define CKD (Table 3).34,35

Monogenic variants account for approximately 30%–50% of cases of CKD stages G3b–G5 in children,5–56,36,37 and 10%–30% in adults.3–30 Diagnostic yields vary between 12% and 65% among studies, with selection bias likely contributing to the variability. However, prevalence estimations for genetic diseases are likely to change over time as genetics-first approaches to diagnosis (in which sequence data are obtained first, followed by characterization of associated phenotypes) become more common.38 Many common variants associated with specific kidney function measures or complex kidney diseases have been identified through genome-wide association studies (GWAS) and exome- or genome-sequencing studies of large population samples—usually of European or East Asian ancestry (Figure 2).13,14,17,19,31,39–41 The largest number of loci—genomic regions containing associated SNPs—were discovered for the continuous kidney function measure eGFR, with studies based on data from >1 million individuals reporting more than 250 such loci.12–14,17,19,22,23,31,40,42–66

Although the distinction of monogenic versus polygenic disorders provides a useful practical framework, genetic risk variants for kidney diseases occur on a spectrum from rare variants with large effects to common variants with small effects, and many diseases do not fit neatly into either category. For example, apolipoprotein L1 (APOL1)—associated kidney risk variants are common among some populations of African ancestry and impart a relatively high risk under a recessive mode of inheritance, but these variants are not considered monogenic. The magnitude of the risk associated with APOL1 variants varies significantly for different forms of nephropathy. For example, Black South Africans with untreated human immunodeficiency virus (HIV) and 2 APOL1 risk alleles have been reported to have a more than 80-fold increased risk of developing HIV-associated nephropathy, but the magnitude of the risk conferred by the same risk alleles ranged between 1.2 and 2 for CKD or nondiabetic kidney failure (Figure 3).67–85 Similarly, the combination of 2 common variants in the HLA-DR and PLA2R1 loci imparts a high risk of the complex disease membranous nephropathy, defying the common variant/small effect paradigm.20
Table 3 | Disease definitions for genetic studies based on kidney function, kidney histology, and molecular markers: advantages and disadvantages

<table>
<thead>
<tr>
<th>Kidney function markers (e.g., eGFR, albuminuria)</th>
<th>Disadvantages</th>
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<tr>
<td>• Readily attainable and standardized information in low- and high-income settings</td>
<td>• Descriptive nature of disease categorization</td>
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<tr>
<td>• Deployed routinely in clinical care and interventional trials</td>
<td>• Agnostic to underlying kidney pathology and pathophysiology and disease heterogeneity</td>
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<tr>
<td>• Allows the identification of genetic determinants of kidney function and factors impacting the progression of kidney disease</td>
<td>• Urinary albumin excretion is underutilized</td>
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<td>• Relatively inexpensive</td>
<td>• Current markers identify genetic variants related to marker metabolism but not filtration</td>
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<td>• Repeated measures often readily available to assess trajectory</td>
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<th>Kidney histology</th>
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<tr>
<td>• Allows for classification based on structural patterns of damage</td>
<td>• Often aggregates a diverse set of underlying disease-initiating events under a common histological damage pattern (e.g., FSGS), thereby potentially introducing functional and genetic heterogeneity</td>
</tr>
<tr>
<td>• Standardized classification scheme for most glomerular diseases</td>
<td>• Limited accessibility in resource-constrained settings</td>
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<tr>
<td>• Current reference standard for clinical management with established clinical workflow</td>
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<tr>
<td>• Histology classifications may reflect a more homogenous pathophysiology than kidney-function markers</td>
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<th>Nontraditional molecular markers (e.g., markers quantified with high-throughput omics technologies)</th>
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<tr>
<td>• Can segregate kidney-disease populations into more-homogenous subgroups and thereby facilitate the identification of underlying disease causes and drivers</td>
<td>• Emerging technologies with limited accessibility in resource-constrained settings need to establish cost-effective readouts readily attainable in low- and middle-income countries</td>
</tr>
<tr>
<td>• Enables systems genetics analysis of kidney disease</td>
<td>• Access to large biobanks required for disease subtyping</td>
</tr>
<tr>
<td>• Comprehensive multi-omics profiling possible (e.g., metabolomics, proteomics, exposomics)</td>
<td>• Some marker levels may vary by kidney function</td>
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eGFR, estimated glomerular filtration rate; FSGS, focal segmental glomerulosclerosis.

Considerations for genetic testing

A positive family history, early age of onset, and presence of extrarenal symptoms are associated with a higher probability of monogenic disease. In addition, the clinical diagnosis is highly predictive of diagnostic yield and will guide the choice of genetic tests, motivating a thorough clinical workup prior to genetic testing. For example, glomerular and tubulointerstitial disorders are associated with a higher diagnostic yield than diabetic kidney disease. In general, because of the genetic heterogeneity of most forms of nephropathy, genetic testing with phenotype-driven gene panels, or exome or genome sequencing, is more efficient than sequential single-gene analyses.

Genetic testing is usually performed subsequent to a clinical workup, but in some situations, early genetic testing may be advantageous. For example, prospective kidney donors related to a recipient with a known genetic condition should be tested early during the donor-evaluation process. Other situations in which early genetic testing should be considered are listed in Table 4. For healthy children and adults, currently, no data support predictive or presymptomatic genetic testing, even if a family history is present. Nevertheless, once a pathogenic variant is identified in a proband, cascade testing of family members and genetic counseling in mutation carriers are the standard practices in clinical genetics.

Most countries do not have guidelines to help determine which nephrology patients should be referred to genetic testing and counseling. Nephrology communities would therefore benefit from developing guidelines based on best evidence and practices in clinical genetics. Overall, guidance should take into account the potential benefit of a genetic diagnosis for specific patients and their families (e.g., treatment changes, family planning, ending a diagnostic odyssey) and balance the risk of false-positive results that could engender unnecessary clinical workup for patients and their families. A position paper by the European Renal Association–European Dialysis and Transplant Association (ERA-EDTA) Working Group for Inherited Kidney Diseases (WGIKD) and the Molecular Diagnostics Taskforce of the European Rare Kidney Disease Reference Network (ERKNet) has been recently issued to delineate indications for genetic testing in CKDs.86

Defining actionable genes in kidney diseases. Actionable genes in kidney diseases refer to genes that, when significantly altered, confer a high risk of serious disease that could be prevented or mitigated if the risk were known.87 A set of 73 actionable genes have been proposed by the American College of Medical Genetics and Genomics (ACMG), many of which are associated with phenotypes relevant to nephrology (PALB2, GLA, HNF1A, MEN1, MAX, RET, SDHAf2, SDHB, SDHC, SDHD, VHL, TMEM127, TSC1, TSC2, WT1). Although these genes were selected based on the possibility that targeting them may prevent overall morbidity and/or mortality, one can conceive of additional, kidney-specific actionable genes, nominated based on availability of interventions, that could prevent renal morbidity (Figure 4).
Examples include the following: early initiation of general renoprotective therapies (e.g., renin-angiotensin blockade for carriers of pathogenic variants in type IV collagen genes); initiation of targeted therapies (e.g., enzyme therapy for Fabry disease or CoQ10 supplementation for nephrotic syndrome due to CoQ10 deficiency); avoidance of treatment that would be futile and perhaps even deleterious (e.g., prolonged immunosuppressive therapies for genetic podocytopathies); and surveillance for recurrence of disease after kidney transplantation (e.g., atypical hemolytic uremic syndrome/thrombotic microangiopathy [aHUS/TMA], primary hyperoxaluria). ClinGen, an international initiative to define robust disease-gene associations and curate pathogenic variants, now has a kidney expert work group that is developing a stable list of nephropathy-associated genes and variants. This group is also expected to provide guidance for actionability for kidney genes and nominate them for the ACMG list. Awareness of the ClinGen Initiative should be promoted in the kidney community, along with messaging regarding the importance of variant submission to public databases such as ClinVar and the value of creating interdisciplinary expert boards to discuss controversial

Figure 2 | Genome-wide association studies (GWAS), exome- or genome-sequencing studies of kidney function markers and kidney diseases.

**The largest study focused on urinary albumin-to-creatinine ratio. Several included serum albumin studies.

**Pediatric population. ***For case–control studies, the total sample sizes were plotted. CKD, chronic kidney disease; eGFRcr, estimated glomerular filtration rate from serum creatinine; FSGS, focal segmental glomerulosclerosis; IgA, IgA nephropathy; KF, kidney failure; LN, lupus nephritis; MCD, minimal change disease; MN, membranous nephropathy; Scr, serum creatinine; SRNS and SSNS, steroid-resistant and steroid-sensitive nephrotic syndrome; T1DN and T2DN, type 1 and type 2 diabetic nephropathy; WES, whole-exome sequencing; WGS, whole-genome sequencing.

Figure 3 | Associations of APOL1 high-risk genotype and various kidney diseases or their progression. APOL1 high-risk genotype: G1G1, G2G2, or G1G2. Studies were ordered by PubMed identifier (PMID), a proxy for publication date. *Compared with White patients. APOL1, apolipoprotein L1; CKD, chronic kidney disease; DRC, Democratic Republic of Congo; FSGS, focal segmental glomerulosclerosis; HIVAN, human immunodeficiency virus–associated nephropathy; HTN, hypertension; KF, kidney failure; LN, lupus nephritis.
variants of uncertain significance (VUS) and discussing the most complex cases. Additional efforts to harmonize gene and gene-panel curation, such as the Genomics England panel app (https://panelapp.genomicsengland.co.uk) are listed in Supplementary Table S1.

In addition to rare pathogenic variants, common genetic variants or polygenic scores may become appropriate for clinical reporting if they are shown to alter patient management, indicate need for surveillance for progression or associated comorbidities, or inform familial screening. In complex diseases, the current best candidates for reporting include APOL1 risk alleles, genetic risk score for membranous nephropathy based on PLA2R1, NFKB1, IRF4, and HLA risk alleles, extremes of a polygenic risk score for eGFR, and pharmacogenetic variants that are informative about risk of adverse events, pharmacokinetics, and pharmacodynamics for specific drugs, some of which may be especially relevant to CKD patients (for example, azathioprine, tacrolimus, warfarin, clopidogrel, simvastatin, voriconazole, and allopurinol). However, we currently lack evidence for actionability for polygenic scores—that is, evidence that reporting can improve clinical outcomes.

APOL1 presents a special case in clinical nephrology because biallelic inheritance of 2 common variants in this gene, which present at high frequency in some populations of African ancestry, increases risk for several kidney disorders. Potential benefits for APOL1 screening include improved risk stratification and opportunities for education. However, only a minority of patients with APOL1 risk genotypes develop nephropathy, and currently no data support early intervention in asymptomatic individuals to reduce future risk of disease. Potential drawbacks to screening include potential for anxiety, stigma, or apathy, and the lack of evidence-based interventions. Combined, these drawbacks could lead to misunderstanding among patients, mistrust of the medical system, and perceived or real racial bias, given that APOL1 risk variants are found predominantly in those with African ancestry. On the other hand, the failure to offer a test that could be most informative in a specific ancestry group also could be perceived as bias. For transplant patients, APOL1 screening could prevent harm to living donors and satisfy recipients’ right to know, but screening could also reduce rates of living donation, waste deceased donor kidneys, and exacerbate organ shortage. The APOL1 Long-term Kidney Transplantation

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**Figure 4 | Actionable genes in kidney diseases.** Actionability refers to the potential for genetic test results to lead to specific clinical actions for prevention or treatment of a condition, supported by recommendations based on evidence. ADPKD, autosomal dominant polycystic kidney disease; aHUS, atypical hemolytic uremic syndrome; RAAS, renin–angiotensin–aldosterone system; SRNS, steroid-resistant nephrotic syndrome.
Outcomes Network (APOLLO) study, which is in progress and expected to end in 2023, is prospectively evaluating the impact of APOL1 risk alleles on donor and recipient outcomes.94 Moreover, the initiation of genotype-driven clinical trials may change the approach to diagnostic testing for APOL1 and other genetic disorders. These considerations emphasize the importance of further research into the usefulness of APOL1 testing.

**Reporting and terminology standards**

Differences in how diagnostic laboratories evaluate and report variants pose a significant challenge in molecular diagnosis. Therefore, standardization of evaluation and reporting among different laboratories and countries is an agreed-on key priority. The determination regarding pathogenicity is a semi-quantitative process that takes into account variant allele frequency, predicted impact on protein function, and prior reports of occurrence with disease. The ACMG and the Association for Molecular Pathology (AMP) published standards and guidelines for the interpretation of sequence variants.95 These guidelines are reviewed periodically and refined by the ClinGen Initiative to reduce discrepancies in variant interpretation among laboratories and clinicians.

The ACMG criteria classify variants into 1 of 5 tiers, with tiers 4 and 5 (i.e., likely pathogenic and pathogenic) classified as diagnostic variants.95 All variant classes can be later upgraded or downgraded based on novel information or interpretation, perhaps necessitating periodic review of clinical genetic reports. However, the abundance of class 3 VUS has created a particular challenge and urgency for improving evaluation and reporting. The definition and relevance of VUS may be unclear to physicians or patients, causing incorrect assignment of diagnoses and/or psychological distress to patients and families. This situation necessitates proper communication with patients to inform and educate them about the possibility of VUS, in which case familial segregation analysis might be recommended. Additionally, VUS should be reported only after interdisciplinary contact between the clinician and geneticist has occurred.96 Future reinterpretation of variants can be facilitated by diagnostic reports that provide detailed description of ACMG classification criteria that were applied at the time of reporting. Although no guidelines exist currently, incidental carrier status for autosomal recessive inheritance is not routinely reported in standard diagnostic reports. Guidelines for systematic reporting of these variants should be developed. Heterozygosity associated with a mild phenotype is increasingly recognized in human genetics, for example, for COL4A3/COL4A4 variants.97

**Unified disease terminology.** The consensus among conference participants was that establishing a unified disease terminology that takes into account genetic disease nomenclature is an important goal for the community. In support of unified, precise disease terminology, a suggested approach is 2-part (“dyadic”) naming, comprising both the clinical condition and the gene name (Figure 5), although some controversy regarding this approach remains.98,99 An important example is adoption of 2-part naming in autosomal dominant tubulointerstitial kidney disease (ADTKD), in which ADTKD is followed by reference to the underlying genetic defect, such as in ADTKD-UMOD and ADTKD-MUC1.100 Two-part names provide flexibility, in that some users (patients/clinicians) can use the first part only (ADTKD), and others (patients/clinicians/researchers) can use the whole name (ADTKD-UMOD). When clinical presentation is unspecified, or very heterogeneous, use of the gene name followed by “kidney disease” (e.g., PAX2-kidney disease) is encouraged. Potential limits to this approach include the possibility of classifying a patient with a benign prognosis as having a potentially progressive disorder, as well as the challenge of adding a second or gene name to conditions already described in International Classification of Disease codes. In this regard, participants of this KDIGO Controversies Conference did not reach consensus regarding renaming of traditional disease terms, such as Alport Syndrome.

**Genomic discovery and implications for chronic kidney diseases**

As demonstrated by the first GWAS for eGFR, common genetic variants that are associated with complex kidney traits usually have small effects and therefore require very large sample sizes for discovery.101 Accordingly, success has been limited in identifying common kidney disease susceptibility variants in individual observational studies of adult102–105 and pediatric106–108 CKD. Conference participants therefore recognized the importance of collaborative consortia, such as the Chronic Kidney Disease Genetics Consortium (CKDGen),11,109 Cohorts for Heart and Aging Research in Genomic Epidemiology (CHARGE) Consortium,66,110 International Genetics & Translational Research in Transplantation Network (iGeneTRAIN),111 and Continental Origins and Genetic Epidemiology Network (COGENT) Kidney Consortium,112,113 that aggregate and harmonize genetic and phenotypic data across multiple studies for combined genetic discovery. In addition to enlarging sample size and providing a platform for replication studies, expanding consortia to international sites can enable studies of more ancestrally and geographically diverse populations. For more specific but less...
frequent primary kidney disorders, such as IgA nephropathy, membranous nephropathy, and steroid-sensitive nephrotic syndrome, aggregating multiple international case–control cohorts is even more important to assure adequate power. Additionally, more diverse ancestral composition of analyzed cohorts facilitates fine-mapping of GWAS loci, enables discovery of ancestry-specific effects, and assures broader generalizability of genetic findings.

The identification of causal genes and variants underlying GWAS associations and definition of their pleiotropic effects are recognized as important challenges in the field. Examples such as *UMOD*, the locus with the strongest common variant association with CKD,66 support the existence of a spectrum of risk variants, from monogenic to complex. No examples currently exist of successful translation of insights from GWAS in CKD to new therapies, but the discovery of the *MYH9* locus,114,115 followed by the identification of *APOL1* as the causative gene,69 refinement of nephrototoxic mechanisms of *APOL1* risk variants,89 and an ongoing phase IIa study of a small molecule *APOL1* inhibitors (ClinicalTrials.gov identifier NCT04340362) represent promising steps toward that end.

Conference participants recognized the emerging importance of electronic health record (EHR)-based genetic research for linking genetic information with a wide range of laboratory parameters and medical conditions. EHR-linkage is possible in various settings, ranging from existing biobanks in research settings, hospitals, or healthcare systems to entire countries, such as Iceland, Estonia, and Finland. Examples of EHR-linked biobanks, institutions, healthcare systems, and country-wide efforts are UK Biobank,116 Million Veteran Program (MVP),117 HUNT Biobank,118 deCODE,119 FinnGen,90 BioBank Japan (BBJ),120 Vanderbilt University biobank (BioVU),121 Michigan Genomics Initiative (MGI),122 Electronic Medical Records and Genomics (eMERGE) Network,123 and the National Institutes of Health All of Us Research Program.124 The development of standardized, scalable, and portable computable phenotypes is time consuming and represents many challenges,125 but it can empower future genetic studies by automated identification of kidney-disease patients in large EHR databases.26,126 Notably, just as important as accurately defining those with a disease is defining those without to serve as healthy controls in genetic studies, which is often harder. We envision that computable phenotyping can be used to find patients with or without CKD, hypertension, kidney stones, and glomerular disease, as well as patients who have received a kidney biopsy or kidney transplant. In nephrology, computable phenotyping is underway.26,127–129 with CKD phenotyping perhaps best positioned for widespread implementation, given the availability of new algorithms based on International Classification of Diseases codes and laboratory values that are routinely measured in clinical practice.26,126

In addition to genomic discovery, EHR-linked genetic research may allow for recontacting of patients with a specific genotype for detailed clinical and molecular studies. Linking of EHR and genetic data also can be used to investigate pleiotropic associations of genetic variants originally discovered for a specific condition (e.g., *APOL1* or *UMOD*) with additional traits captured in medical records using phenome-wide association approaches.44,26,42,130 Such studies can be further complemented with Mendelian randomization methods to clarify associations between genetic variants, biomarkers, and phenotypes.131

Despite the large size of consortia and EHR-linked studies, certain groups of patients are still underrepresented in genetic research. For instance, the paucity of pediatric patients with genetic information has limited both longitudinal phenotype analyses from childhood to adulthood and the ability to identify genetic drivers of kidney diseases or traits of childhood. Also, there is urgent need to expand ancestral diversity of participants in genetic studies, specifically aiming to increase the representation of non-European populations.132 Additional challenges include harmonizing data for rare kidney conditions that necessitate aggregating cases from across several biobanks and EHRs; identifying ancestry-matched controls for case–control analyses; handling of missing data; and harmonizing genotypes in the presence of different types of available genetic data.133,134

Partnerships between academic labs and industry allow efficient exchange of ideas and resources to promote investigation of disease mechanisms, biomarkers, and therapeutic targets. Such partnerships can enable academic labs, biobanks, and institutions and healthcare systems to conduct large-scale multi-omic studies that would not be feasible with only support from internal funds or extramural grants, and facilitate follow-up studies to “functionalize” key genes or genetic variants. Successful partnerships must achieve a balance between companies’ incentive to invest and academics’ freedom in research and publishing. Key principles and processes, such as intellectual property, publications, and data sharing and access, also must be aligned. These partnerships have been particularly valuable for generating functional genomic data from primary kidney tissues and allow for rapid implementation of new methods.135–138 Generation of additional such data from primary kidney tissue and cell types should continue to be a research priority because the kidney is underrepresented in many existing public databases, including the Encyclopedia of DNA Elements (ENCODE),139 Roadmap Epigenomics,140 and the Genotype-Tissue Expression (GTEx) project.141 The Kidney Precision Medicine Project (KPMP)142 and similar new initiatives aim to address some of these important gaps by generating and harmonizing new multidimensional molecular data for human kidney tissue in health and disease.

**Polygenic scores.** Polygenic scores (PGSs) are based on the results of GWAS and aggregate the effects of trait- or disease-associated variants across the genome. PGSs capture a greater proportion of genetic variance compared to individual SNPs and potentially may be useful to risk-stratify populations, enhance screening, and ultimately inform
diagnosis, prognosis, and/or treatment. PGs have been shown to modify the penetrance of monogenic variants for hypercholesterolemia, hereditary breast and colon cancer, and obesity, although this effect has not yet been examined for kidney diseases. PGs for kidney disease can be constructed using a smaller set of genome-wide significant SNPs only, such as a 147-SNP score for eGFR (odds ratio of 2 for individuals in the highest 10% of the score) or a 5-SNP score for membranous nephropathy (odds ratio of >20 for those in the highest 10% of the score), or by using genome-wide scores with hundreds of thousands of variants, such as the UK Biobank score for CKD. Currently, most scores are derived from European populations and do not include rare or population-specific variation, potentially creating a new health disparity between individuals of European descent and others. Given that scores are constructed from GWAS for complex traits and diseases, they may reflect heterogeneous mechanisms and therefore may not necessarily point to targeted interventions.

Conference participants agreed that before applying PGs in clinical nephrology, more research is needed to derive the most accurate and cosmopolitan scores for kidney disease. Also necessary are the following: proof of clinical utility in surveillance, diagnosis, prognosis, or treatment of kidney disease; a better understanding of dependence on the clinical context, including disease stage, ancestry, sex, or demographics; and cost-effectiveness and added value beyond standard clinical risk factors. PGs computation needs to be robust, open-source, and able to be incorporated into points of care. Quality standards for PGs have recently been defined by ClinGen, providing a framework for evaluating clinical translation and utility.

Achieving implementation in clinical medicine

Clinical knowledge. Often, insufficient experience and knowledge are major barriers for implementing genetic evaluation in nephrology practice. To ensure equitable access to genetic testing, all nephrologists should have a sufficient knowledge base for discerning which patients would benefit from genetic testing, and at minimum, be able to collect personal and family histories. Although optimally all nephrologists would be able to recommend screening for at-risk family members if applicable, would conceptually understand types of genetic tests, including their risks and benefits, and would remain aware of local regulations around genetic testing, those who lack experience in these domains should collaborate with a clinical geneticist and/or a genetic counselor. In addition, reporting of positive genetic results to patients necessitates individual and family counseling and referrals. Hence, a multidisciplinary approach is key for successfully implementing use of genetics in the clinic.

Conference participants recognized workforce education as a critical need. Genetics is currently not part of the nephrology fellowship curriculum in the US, and indeed, fellows report lacking competency in genetic renal disease. Similarly, in Australia, less than half of nephrologists feel confident in using results of genomic testing in clinical practice. No genetics core competency guidelines are in place for nephrologists, nor are guidelines for evaluating competencies for clinical genetic consent and return of results. Based on published data and information, a compiled list of core competencies expected from nephrologists at different levels of expertise can be found in Supplementary Table S2. These gaps can be remedied by including more robust genetics curricula in medical school, residency, and fellowship training. Education for current practicing nephrologists can be achieved via workshops at national and international societies, continuing medical education, review papers in nephrology journals, and introduction of clinical genetic questions to relicensing tests. One can also envision an advanced training or subspecialty track in genetic nephrology, similar to transplant, oncology, or glomerular diseases subspecialization. Supplementary Table S1 provides an overview of clinical genetics web resources to aid nephrologists.

Clinical practice. Centers of expertise are sites where patients can receive comprehensive, coordinated care from a multidisciplinary team that includes a relatively small number of nephrologists with a high skill set for genetic diagnosis (Figure 6). These centers also play an important role in training and research. Centers of expertise, or reference, are concentrated in Europe, with ERKNet constituting a consortium of more than 30 centers in 12 countries, supported by the European Union. In most regions of the globe, including the US, no centralized accreditation mechanisms are in place for developing centers of expertise or reference. The establishment of such centers can facilitate standardized variant interpretation, identify “actionable” genes associated with kidney diseases, train the future generation of physicians with dual expertise in genetics and nephrology, develop guidelines for referral and testing of patients with kidney diseases, disseminate implementation knowledge, and develop collaborative research projects and clinical trials for rare disorders.

Cost and access. Often, genetic testing is not affordable for either patients or healthcare systems. In regions with cost coverage or reimbursement, access can still be unequal, as genetic testing is based on clinical presentation, and obtaining coverage is often easier in children than in adults. Many countries do have genetic protection acts, laws, or regulations to ensure equitable access to genetic testing without fear of discrimination. However, legislation alone is not always sufficient for allaying patient concerns about the potential for prejudice.

Logistically, remote sample collection and telemedicine have potential for increasing access to genetic counseling. However, adequate physical evaluation and identification of
extrarenal manifestations can be more complicated or impossible with telemedicine. In addition, although the severe acute respiratory syndrome coronavirus-2 (SARS-CoV-2) pandemic has accelerated the deployment of telemedicine across many health systems, not all patients and physicians are comfortable with remote, video-based communications.

For most genetic conditions, we lack large-scale cost-effectiveness analyses to demonstrate the benefits of genetic testing. Recent data suggest that genetic testing has a high diagnostic yield in patients with CKD of unknown etiology and may reduce costly diagnostic workups, hopefully increasing the coverage of genetic testing for those patients.\(^3\),\(^8\) Also important is the demonstration of the clinical value of genetic testing beyond diagnosis, such as its impact on long-term outcomes and health economics. A comparison of the cost-effectiveness of genetic testing in nephrology across different healthcare coverage systems could provide key insights and an evidence base for expanding testing.

**Patient voice.** Patient engagement is vital for successful treatment and advances in research. To advocate for their own genetic testing, patients need to have an awareness of and education regarding genetics and kidney disease and the relative benefits and risks of genetic testing.\(^1\),\(^2\) The complex ethical, psychosocial, and familial implications for genetic testing, including presymptomatic testing, can make decision-making challenging and require an understanding of patient values, goals, and priorities.\(^3\),\(^5\) To engage and activate patients and patient communities, educational content needs to be

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**Table 5 | Recommended practices for value-based measures of implementation and quality assurance of clinical genetics in nephrology**

<table>
<thead>
<tr>
<th>Measure</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Measure nephrologist adoption of genetic testing and appropriate referral to genetic testing</td>
<td>Define disease-specific outcomes that can be measured</td>
</tr>
<tr>
<td>Measure nephrologist utilization of genetic results (to determine if appropriate changes in diagnosis and care have occurred)</td>
<td>Rate of kidney disease progression</td>
</tr>
<tr>
<td>Define disease-specific outcomes that can be measured</td>
<td>Change in treatment</td>
</tr>
<tr>
<td></td>
<td>Access to genetically stratified clinical trials</td>
</tr>
<tr>
<td></td>
<td>Donor risk evaluation, or deceased-donor organ evaluation</td>
</tr>
<tr>
<td></td>
<td>Recipient risk evaluation (e.g., improved matching, customized immunosuppression, etc.)</td>
</tr>
<tr>
<td></td>
<td>Utilization of information for family planning</td>
</tr>
<tr>
<td></td>
<td>Patient-reported outcome (quality of life, etc.)</td>
</tr>
<tr>
<td></td>
<td>Hospitalization, cardiovascular outcomes, mortality</td>
</tr>
<tr>
<td></td>
<td>Diagnosis of at-risk family members</td>
</tr>
<tr>
<td></td>
<td>Define and measure potential harmful impacts of genetic testing (e.g., wrongful impact on change of treatment)</td>
</tr>
<tr>
<td>Define audits/assessments for centers that offer genetic testing in nephrology as quality-assurance activity</td>
<td>Potentially apply USPSTF and EGAPP methods to analyze the implementation of genetic testing for kidney diseases</td>
</tr>
</tbody>
</table>

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USPSTF, United States Preventive Services Task Force; EGAPP, Evaluation of Genomic Applications in Practice and Prevention.
accessable and sensitive to patients in terms of culture, language, and literacy, as well as be shared across multiple platforms. The topics of race and ancestry have been widely debated in genetics, as well as nephrology. In specific terms, race is a social, categorical construct, whereas ancestry is based on inherited genetic variants without categorization. In principle, genetics research is agnostic to race, and identifying disease-causing variants could obviate reliance on race or ancestry as a proxy for probability of carrying a risk allele.

Within nephrology, patient-reported outcome measures can provide doctors, investigators, and policymakers with important insights into patient symptoms and experiences that cannot be identified through laboratory or imaging studies alone. Research communities that engage with patients and include the patient voice can better advocate for more research and development in rare kidney diseases.

**Research in implementation.** Evidence-based frameworks for evaluating quality of care in genetic testing have been put forth by ACMG, ERKNet, and others. These frameworks cover different methods for evaluating the analytic and clinical validity, as well as the clinical utility, of genetic tests. Nephrology outcomes used in clinical trials have included those that are disease-specific or represent more general longer-term outcomes, such as kidney failure, cardiovascular death, or mortality, which require large datasets. Yet this space is evolving, as demonstrated by development of novel trial designs using Bayesian methodology, inclusion of patient-reported outcomes, and additional economic evaluation of genetic risk. Steps for expanding measures to best inform value-based implementation and quality assurance of clinical genetics in nephrology are listed in Table 5. This large and critical space underpins clinical translation and mainstreaming, with much research and work anticipated in the coming years.

**Conclusions**

This KDIGO Controversies Conference on Genetics in Chronic Kidney Disease discussed many technical, logistical, ethical, and/or research questions related to the definition and epidemiology of monogenic and complex kidney diseases, applications of genetic findings in clinical medicine, and utilization of genomics for defining and stratifying CKD. Identified areas of consensus and future research priorities provide a roadmap toward realizing the promises of genomic medicine for nephrology.

**APPENDIX**

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All Steering Committee Members contributed equally. The conference planning and the drafting and critical revision of this manuscript were performed by the Steering Committee Members, the Conference Co-Chairs, and Jennifer King, with important intellectual content contributions provided by the remaining authors.

SUPPLEMENTARY MATERIAL

Supplementary File (PDF)

Table S1. Online clinical genetics resources to aid nephrologists. Adapted with permission from Annals of Internal Medicine, Milo Rasouly H, Aggarwal V, Bier L, Goldstein DB, Gharavi AG. Aggarwal V, Bier L, Goldstein DB, Gharavi AG. Cases in kidney genetics in the 21st century. Kidney Int. 2018;94:363–371. under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/). © The Author(s) 2019.

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