

ВСЕМИРНЫЙ ДЕНЬ ПОЧКИ

11-й Всемирный день почки отмечается 10 марта 2016 года. Это ежегодное глобальное мероприятие, проводимое совместно Международным обществом нефрологов (International Society of Nephrology, ISN) и Международной федерацией нефрологических организаций (International Federation of Kidney Foundations, IFKF), стало эффективным средством информирования людей о серьезности заболеваний почек и их последствиях. В этом, 2016 году Всемирный день почки посвящен заболеваниям почек у детей и предпосылкам их развития у взрослых, которые могут начинаться в раннем детстве.

Пациенты, перенесшие острое повреждение почек (ОПП) в детском возрасте, могут иметь пролонгированные негативные последствия, вплоть до хронической болезни почек (ХБП). Кроме того, ХБП, перенесенная в детстве, особенно врожденная ХБП, а также осложнения от многих не почечных заболеваний могут привести не только к значительной заболеваемости и смертности в детском возрасте, но и к серьезным проблемам со здоровьем по истечении детского возраста. В действительности, детская смертность от длинного списка инфекционных заболеваний неразрывно связана с заболеваниями почек. Например, дети, которые заболевают холерой или другими желудочно-кишечными инфекционными заболеваниями, часто умирают не по причине данных инфекций, а от ОПП. Кроме того, множественные исследования доказывают, что артериальная гипертензия, протеинурия и ХБП в зрелом возрасте имеют «предшественников» в детском возрасте – еще с внутриутробного или перинатального периодов развития.

Всемирный день почки – 2016 имеет целью повысить общую осведомленность людей о том, что многие из взрослых заболеваний почек фактически начинаются в детстве. Понимание важности диагностирования заболеваний почек в детском возрасте позволяет выявлять и превентивно наблюдать за пациентами с повышенным риском ХБП на протяжении всей их жизни.

Статистические данные о заболеваемости ХБП и ОПП в детском возрасте в настоящее время ограничены во всех странах, хотя и наблюдается определенное улучшение ситуации. Распространенность ХБП в детском возрасте достаточно низка и составляет от 15 до 74,7 на 1 млн детей в зависимости от региона. Такой разброс скорее всего связан с тем, что данные о заболеваемости ХБП подвержены влиянию региональных и культурных факторов, а также используемой методологии. Всемирная организация здравоохранения (ВОЗ) не так давно добавила почечные и урологические заболевания в список мониторируемых по смертности во всем мире, поэтому с течением времени данные ВОЗ станут доступными. На сегодняшний день ВОЗ не предоставляет детализацию по возрастным группам (как это и должно быть в будущем, чтобы повысить ценность таких данных). Тем не менее, данные из таких источников, как Североамериканский педиатрический регистр почечных проб и сочетанных исследований (North American Pediatric Renal Trials and Collaborative Studies, NAPRTCS), Объединенная система данных о донорских почках США (U.S. Renal Data System,USRDS) и регистр Европейской ассоциации диализа и трансплантации (EDTA), включают информацию о заболеваемости хронической почечной недостаточностью (ХПН) у детей и (незначительно) по ХБП. А такие проекты, как «ItalKid» (перспективный популяционный регистр заболеваемости ХПН среди детей, Италия) и «СКiD» (перспективные когортные исследования заболеваемости ХПН среди детей, проводимые Национальными институтами здоровья, США), исследование «Global Burden of Disease Study 2013», а также ряд национальных регистров, которые стали появляться в последнее время во многих странах мира, предоставляют все более и более детальную и актуальную информацию по заболеваниям почек среди детского населения.

ОПП может привести к развитию ХБП (в соответствии с рядом «взрослых» исследований). Заболеваемость ХБП среди детей, поступающих в больницы, варьирует очень сильно – от 8 до 89%. Результат, как правило, зависит от имеющихся исследовательских ресурсов. Результаты глобальных исследовательских проектов, таких как «Всемирное исследование ОПП, почечной недостаточности и их эпидемиологии у тяжелобольных детей» (Assessment of Worldwide Acute Kidney Injury, Renal Angina and Epidemiology in critically ill children, «AWARE»; не путать с «AWAREness during REsuscitation», – иным, не связанным с заболеваниями почек медицинским исследованием) и еще одного исследования в пяти странах на предмет заболеваемости ХБП среди детей, еще только ожидаются. Моноинституциональные исследования, такие как мета-анализ по историям болезней пациентов, показывают, что и ОПП, и ХБП у детей составляют меньшинство из всего массива заболеваемости ХБП в мире. Тем не менее, становится все более очевидным, что заболевания почек в зрелом возрасте часто имеют «детское» наследие.

Contributors (listed alphabetically):
Julie R. Ingelfinger, Kamyar Kalantar-Zadeh, Franz Schaefer

AVERTING THE LEGACY OF KIDNEY DISEASE – FOCUS ON CHILDHOOD

on behalf of the World Kidney Day Steering Committee*

ABSTRACT

World Kidney Day 2016 focuses on kidney disease in childhood and the antecedents of adult kidney disease that can begin in earliest childhood. Chronic kidney disease (CKD) in childhood differs from that in adults, as the largest diagnostic group among children includes congenital anomalies and inherited disorders, with glomerulopathies and kidney disease in the setting of diabetes being relatively uncommon. In addition, many children with acute kidney injury will ultimately develop sequelae that may lead to hypertension and CKD in later childhood or in adult life. Children born early or who are small-for date newborns have relatively increased risk for the development of CKD later in life. Persons with a high-risk birth and early childhood history should be watched closely in order to help detect early signs of kidney disease in time to provide effective prevention or treatment. Successful therapy is feasible for advanced CKD in childhood; there is evidence that children fare better than adults, if they receive kidney replacement therapy including dialysis and transplantation, while only a minority of children may require this ultimate intervention. Because there are disparities in access to care, effort is needed so that those children with kidney disease, wherever they live, may be treated effectively, irrespective of their geographic or economic circumstances. Our hope is that World Kidney Day will inform the general public, policy makers and caregivers about the needs and possibilities surrounding kidney disease in childhood.

«For in every adult there dwells the child that was, and in every child there lies the adult that will be».

– John Connolly, The Book of Lost Things

Introduction and Overview

The 11th World Kidney Day will be celebrated on March 10, 2016, around the globe. This annual event, sponsored jointly by the International Society of Nephrology (ISN) and the International Federation of Kidney Foundations (IFKF), has become a highly successful effort to inform the general public and policymakers about the importance and ramifications of kidney disease. In 2016, World Kidney Day will be dedicated to

kidney disease in childhood and the antecedents of adult kidney disease, which can begin in earliest childhood.

Children who endure acute kidney injury (AKI) from a wide variety of conditions may have long-term sequelae that can lead to chronic kidney disease (CKD) many years later.^{1–4} Further, CKD in childhood, much of it congenital, and complications from the many non-renal diseases that can affect the kidneys secondarily, not only lead to substantial morbidity and mortality during childhood but also result in medical issues beyond childhood. Indeed, childhood deaths from a long list of communicable diseases are inextricably linked to kidney involvement. For example, children who succumb to cholera and other diarrheal infections often die, not from the infection, but because of AKI induced by volume depletion and shock. In addition, a substantial body of data indicates that hypertension, proteinuria and CKD in adulthood have childhood antecedents – from as early as in utero and perinatal life (see Table 1 for definitions of childhood). World Kidney Day 2016 aims to heighten general awareness that much adult renal disease is actually initiated in childhood. Understanding high risk diagnoses and events that occur in childhood have the potential to identify and intervene preemptively in those people at higher risk for CKD during their lifetimes.

Worldwide epidemiologic data on the spectrum of both CKD and AKI in children are currently limited, though increasing in scope. The prevalence of CKD in childhood is rare—and has been variously reported at 15–74.7 per million children.³ Such variation is likely because data on CKD are influenced by regional and cultural factors, as well as by the methodology used to generate them. The World Health Organization (WHO) has recently added kidney and urologic disease to mortality information tracked worldwide, and should be a valuable source of such data over time – yet WHO does not post the information by age group.⁵ Databases such as the North American Pediatric Renal Trials and Collaborative Studies (NAPRTCS)⁶ the U.S. Renal Data System (USRDS)⁷ and the EDTA registry⁸ include data on pediatric ESRD, and some on CKD. Projects such as the ItalKid⁹ and Chronic Kidney Disease in Children (CKiD)¹⁰ studies, the Global Burden of Disease Study

*Members of the World Kidney Day Steering Committee are: Philip Kam Tao Li, Guillermo Garcia-Garcia, William G. Couser, Timur Erk, Julie R. Ingelfinger, Kamyar Kalantar-Zadeh, Charles Kernahan, Charlotte Osafo, Miguel C. Riella, Luca Segantini, Elena Zakharova.

Correspondence:

World Kidney Day,
International Society of Nephrology, in collaboration with International Federation of Kidney Foundation
Rues de Fabriques 1B, 1000, Brussels, Belgium
E-mail: myriam@worldkidneyday.org

Table 1

Definitions of Stages of Early Life

Perinatal Period	22 completed weeks of gestation to Day 7 of postnatal life
Neonatal Period	Birth to Day 28 of postnatal life
Infancy	Birth to 1 year of age
Childhood	1 year of age to 10 years of age
Adolescence	10 years of age to 19 years of age

The data in this table are as defined by the World Health Organization. The perinatal period is defined as 22 completed weeks of gestation to Day 7 of life; the neonatal period, as up to 28 days of life; infancy as up to one year of age; childhood as year 1 to 10; and adolescence from 10 years to age 19. There is variation worldwide in how these stages of early life are defined. Some would define «young people» as those age 24 or less. In the United States, childhood is as a whole defined as going to age 21.

Table 2

Etiology of Chronic Kidney Disease in Children

CKD		ESRD	
Etiology	Percentage (Range)	Etiology	Percentage (Range)
CAKUT	48–59%	CAKUT	34–43%
GN	5–14%	GN	15–29%
HN	10–19%	HN	12–22%
HUS	2–6%	HUS	2–6%
Cystic	5–9%	Cystic	6–12%
Ischemic	2–4%	Ischemic	2%

Rare causes include congenital NS, metabolic diseases, cystinosis/Miscellaneous causes depend on how such entities are classified CAKUT: Congenital anomalies of the kidney and urinary tract; GN: Glomerulonephritis; HN: Hypertension; HUS: Hemolytic uremic syndrome *from Harambat et al. CKD data are from NAPRTCS, the Italian Registry and the Belgian Registry. ESRD data are from ANZDATA, ESPN/ERA-EDTA, UK Renal Registry and the Japanese Registry.

2013, as well as registries that now exist in many countries provide important information, and more is required.¹¹

AKI may lead to CKD, according to selected adult population studies.¹² The incidence of AKI among children admitted to an intensive care unit varies widely – from 8 to 89%.¹ The outcome depends on the available resources. The results from projects such as the AWARE study, a five-nation study of AKI in children are awaited.¹³ Single center studies, as well as meta-analyses indicate that both AKI and CKD in children account for a minority of CKD worldwide.^{2,3} However, it is increasingly evident that kidney disease in adulthood often springs from a childhood legacy.

Spectrum of Pediatric Kidney Diseases

The conditions that account for CKD in childhood, with a predominance of congenital and hereditary disorders, differ substantially from those in adults. To date, mutations in more than 150 genes have been found to alter kidney development or specific glomerular or tubular functions.¹⁴ Most of these genetic disorders present during childhood, and many lead to progressive CKD. Congenital anomalies of the kidney and urinary tract (CAKUT) account for the largest category of CKD in children (see Table 2) and include renal hypoplasia/dysplasia and obstructive uropathy. Important subgroups among the renal dysplasias are the cystic kidney diseases, which originate from genetic defects of the tubuloepithelial cells' primary cilia. Many pediatric glomerulopathies are caused by genetic or acquired defects

of the podocytes, the unique cell type lining the glomerular capillaries. Less common but important causes of childhood CKD are inherited metabolic disorders such as hyperoxaluria and cystinosis, and atypical hemolytic uremic syndrome, a thrombotic microangiopathy related to genetic abnormalities of complement, coagulation or metabolic pathways.

In various classifications it is not clear how to categorize children who have suffered AKI and apparently recovered, or how and whether to include those children who have had perinatal challenges, likely resulting in a relatively low nephron number.

Among children with childhood-onset end-stage renal disease (ESRD) glomerulopathies are slightly more and congenital anomalies less common (Table 2), due to the typically more rapid nephron loss in glomerular disease. However, recent evidence suggests that many patients with milder forms of CAKUT may progress to ESRD during adulthood, peaking in the fourth decade of life.¹⁵

There are national and regional differences in the types and course of both AKI and CKD during childhood and beyond. Death from kidney disease is higher in developing nations, and national and regional disparities in care and outcome must be addressed. Further, access to care is variable, depending on the region, the country and its infrastructure. By focusing on kidney disease in childhood, cost-effective solutions may be reached, as treating disease early and preemptively may prevent later, more advanced CKD. Expectations depend on the availability of care and management. Treating children,

even from infancy, who have AKI and CKD that requires renal replacement therapy can be effective in mitigating the burden of kidney disease in adulthood. Doing so requires resources that focus on the most expeditious and cost-effective ways to deliver acute RRT in childhood.

Congenital Kidney Disease and Developmental Origins of Health and Disease, Renal Endowment and Implications

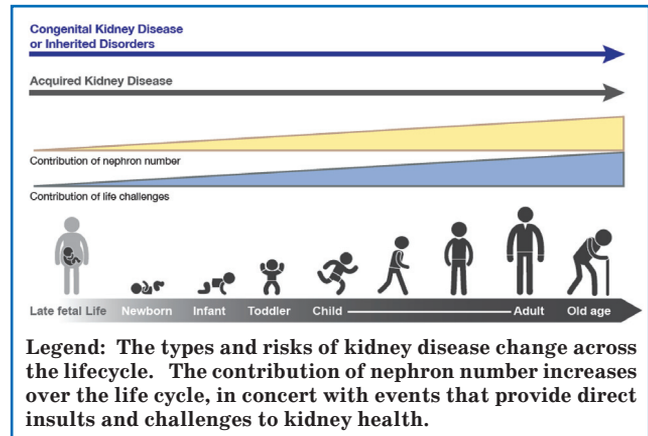
In regions where antenatal fetal ultrasounds are routine, many children with urologic abnormalities are identified antenatally, which permits early intervention. However, in much of the world, children with structural abnormalities are not identified until much later, when symptoms develop. While generalized screening for proteinuria, hematuria and urinary tract infections are carried out in some countries and regions, there is a lack of consensus as to its effectiveness. However, there is general agreement that children with antenatal ultrasound studies that indicate possible genitourinary anomalies, children with a family history of kidney disease, and children with signs such as failure to thrive or a history of urinary tract infection, voiding dysfunction or an abnormal appearing urine should be examined. Initial screening would include a focused physical examination and a urine dipstick, formal urinalysis and a basic chemistry panel, followed by a more focused evaluation if indicated.

Depending on the diagnosis, definitive therapy may be indicated. However, the evidence that therapy will slow progression of CKD in childhood remains limited. Angiotensin converting enzyme inhibitors, angiotensin receptor blockers, antioxidants and, possibly, dietary changes may be indicated, depending on the diagnosis. However, dietary changes need to permit adequate growth and development. The ESCAPE trial provided evidence that strict blood pressure control retards progression of CKD in children irrespective of the type of underlying kidney disease.¹⁶

Some very young children may require renal replacement therapy in early infancy. Recent data pooled from registries world wide indicate good survival, even when dialysis is required from neonatal age.^{2,17} Kidney transplantation, the preferred renal replacement therapy in children, is generally suitable after 12 months of age, with excellent patient and allograft survival, growth and development.

Evidence is accumulating that childhood-onset CKD leads to accelerated cardiovascular morbidity and shortened life expectancy. Ongoing large prospective studies such as the (Cardiovascular Comorbidity in Children with CKD (4C) Study are expected to inform about the causes and consequences of early cardiovascular disease in children with CKD.¹⁸

In addition to those children with congenital kidney disease, it is now known that perinatal events may affect future health in the absence of evident kidney disease



in early life.¹⁹ Premature infants appear to be particularly at risk for kidney disease long after they are born, based both on observational cohort studies, as well as on case reports. Increasingly premature infants survive, including many born well before nephrogenesis is complete.²⁰ The limited data available indicate that in the process of neonatal ICU care, such babies receive many nephrotoxins, and that those dying prior to discharge from the nursery have fewer and larger glomeruli.²¹ Additionally, those surviving have evidence of renal impairment that may be subtle.²² Even more concerning, abundant epidemiologic data indicate that persons born at term but with relatively low birth weights may be at high risk for hypertension, albuminuria and CKD in later life.²³ When direct measurements are pursued, such persons, as adults, may have fewer nephrons, thus a low cardiorenal endowment.

In focusing on children for World Kidney Day, we would note that it is key to follow kidney function and blood pressure throughout life in those persons born early or small-for-dates. By doing so, and avoiding nephrotoxic medications throughout life, it may be possible to avert CKD in many people.

Resources and Therapeutics for Children – Differences from Therapeutics in Adults

Disparities exist in the availability of resources to treat AKI in children and young people; consequently, too many children and young adults in developing nations succumb if AKI occurs. To address the problem the ISN has initiated the Saving Young Lives Project, which aims both to prevent AKI with prompt treatment of infection and/or delivery of appropriate fluid and electrolyte therapy, and to treat AKI when it occurs. This ongoing project in SubSaharan Africa and South East Asia, in which four kidney foundations participate equally (IPNA, ISN, ISPD and SKCF),* focuses on establishing and maintaining centers for the care of AKI, including the provision of acute peritoneal dialysis. It links with the ISN's 0 by 25 project, which calls on members to ensure by 2025 that nobody dies from preventable and acute kidney injury.

In view of the preponderance of congenital and hereditary disorders, therapeutic resources for children

*The four partners are (in alphabetical order): IPNA (International Pediatric Nephrology Association), ISN (International Society of Nephrology), ISPD (International Society for Peritoneal Dialysis), SKCF (Sustainable Kidney Care Foundation).

with CKD have historically been limited to a few immunological conditions. Very recently, progress in drug development in concert with advances in genetic knowledge and diagnostic capabilities has begun to overcome the long-standing «therapeutic nihilism» in pediatric kidney disease. Atypical HUS, long considered ominous, with a high likelihood of progression to ESRD and post-transplantation recurrence, has turned into a treatable condition – with the advent of a monoclonal antibody that specifically blocks C5 activation.²⁴ Another example is the use of vasopressin receptor antagonists to retard cyst growth and preserve kidney function in polycystic kidney disease.²⁵ First proven efficacious in adults with autosomal dominant polycystic kidney disease, therapy with vaptans holds promise also for the recessive form of the disease, which presents and often progresses to ESRD during childhood.

However, patient benefit from pharmacological research breakthroughs is jeopardized on a global scale by the enormous cost of some of the new therapeutic agents. The quest for affordable innovative therapies for rare diseases will be a key issue in pediatric nephrology in the years to come.

The identification of children likely to benefit from novel therapeutic approaches will be greatly facilitated by the development of clinical registries that inform about the natural disease course, including genotype-phenotype correlations. Apart from disease-specific databases, there is also a need for treatment-specific registries. These are particularly relevant in areas where clinical trials are difficult to perform due to small patient numbers and lacking industry interest, as well as for therapies in need of global development or improvement. For instance, there is currently a large international gradient in the penetration and performance of pediatric dialysis and transplantation. Whereas pediatric patient and technique survival rates are excellent and even superior to those of adults in many industrialized countries, it is estimated that almost half of the world's childhood population is not offered chronic renal replacement therapy (RRT) at all. Providing access to RRT for all children will be a tremendous future challenge. To obtain reliable information on the demographics and outcomes of pediatric RRT, the International Pediatric Nephrology Association (IPNA) is about to launch a global population-based registry. If successful, the IPNA RRT registry might become a role model for global data collection.

Transition from Pediatric to Adult Care

Transition of care for adolescents with kidney disease into an adult setting is critical both for patients and their caregivers. Non-adherence is a too-frequent hallmark of transition from pediatric to adult care for young patients with chronic disease states.^{26–28} Hence, considered steps combined with systematically defined procedures supported by validated pathways and credible guidelines must be in place to ensure successful outcomes.

In the process of change from pediatric to adult care «transition», which should occur gradually, must be distinguished from «transfer», which is often an abrupt

and mechanistic change in provider setting. Introducing the concept of transition should be preemptive, starting months to years prior to the targeted time, as children move into adolescence and adulthood. The ultimate goal is to foster a strong relationship and individualized plan in the new setting that allows the patient to feel comfortable enough to report non-adherence and other lapses in care.

A transition plan must recognize that the emotional maturity of children with kidney disease may differ widely. Assessment of the caregiver and the family structure as well as cultural, social, and financial factors at the time of transition are key, including a realistic assessment of caregiver burden.⁴ The appropriate timing and format of transition may vary widely among different patients and in different settings; therefore, a flexible process without a set date and even without a delineated format may be preferred.

Importantly, transition may need to be slowed, paused or even reversed temporarily during crises such as disease flares or progression, or if family or societal instability occurs. A recent joint consensus statement by the International Society of Nephrology (ISN) and International Pediatric Nephrology Association (IPNA) proposed steps consistent with the points just outlined, aiming to enhance the transition of care in kidney disease in clinical practice.^{29,30}

Call for Generating further Information and Action

Given vulnerabilities of children with kidney disease including impact on growth and development and future life as an adult, and given the much greater proportion of children in developing nations facing resource constraints educating everyone involved is imperative in order to realign communications and actions.^{31,32} These efforts should foster regional and international collaborations and exchange of ideas between local kidney foundations, professional societies, other not-for-profit organisations, and states and governments, so as to help empower all stakeholders to improve the health, well-being and quality of life of children with kidney diseases and to ensure their longevity into adulthood.

Until recently, however, the WHO consensus statement on non-communicable diseases (NCD) included cardiovascular disease, cancer, diabetes and chronic respiratory disease, but not kidney disease.^{33,34} Fortunately, due, in part, to a global campaign led by the ISN, the Political Declaration on NCDs from the United Nations Summit in 2011 mentioned kidney disease under Item.^{19,35}

Increasing education and awareness about renal diseases in general and kidney disease in childhood in particular is consistent with the objectives of the WHO to reduce mortality from NCD with a 10 year target population level initiatives focusing on changes in life style (including tobacco use reduction, salt intake control, dietary energy control, and alcohol intake reduction) and effective interventions (including blood pressure, cholesterol and glycemic control). Heightened efforts are needed to realign and expand these multidisciplinary collaborations with more effective focus on early detec-

tion and management of kidney disease in children. Whereas the issues related to kidney disease may be overshadowed by other NCDs with apparently larger public health implications such as diabetes, cancer, and cardiovascular diseases, our efforts should also increase education and awareness on such overlapping conditions

as cardiorenal connections, the global nature of the CKD and ESRD as major NCDs, and the role of kidney disease as the multiplier disease and confounder for other NCDs. White papers including consensus articles and blueprint reviews by world class experts can serve to enhance these goals.³⁶

Comprehensive Style of the References (with PMID and PMCID)

1. Goldstein, S.L. Acute kidney injury in children and its potential consequences in adulthood. *Blood Purif*33, 131–137 (2012).
2. Harambat, J., van Stralen, K.J., Kim, J.J. & Tizard, E.J. Epidemiology of chronic kidney disease in children. *Pediatr. Nephrol*27, 363–373 (2012).
3. Warady, B.A. & Chadha, V. Chronic kidney disease in children: the global perspective. *Pediatr. Nephrol*22, 1999–2009 (2007).
4. Furth, S.L., et al. Design and methods of the Chronic Kidney Disease in Children (CKiD) prospective cohort study. *Clin. J. Am. Soc. Nephrol*11, 1006–1015 (2006).
5. Health statistics and information systems: Estimates for 2000–2012. http://www.who.int/healthinfo/global_burden_disease/estimates/en/index1.html.
6. NAPRTCS Annual Reports <https://web.emmes.com/study/ped/annlrept/annlrept.html>.
7. Saran, R., et al. US Renal Data System 2014 Annual Data Report: Epidemiology of Kidney Disease in the United States. *Am. J. Kidney Dis*65, A7 (2015).
8. ESPN/ERA-EDTA Registry, European Registry for Children on Renal Replacement Therapy <http://www.espn-reg.org/index.jsp>.
9. Ardissino, G., et al. Epidemiology of chronic renal failure in children: data from the ItalKid project. *Pediatrics*111, e382–387 (2003).
10. Wong, C.J., Moxey-Mims, M., Jerry-Fluker, J., Warady, B.A. & Furth, S.L. CKiD (CKD in children) prospective cohort study: a review of current findings. *Am. J. Kidney Dis*60, 1002–1011 (2012).
11. Global Burden of Disease Study 2013 Collaborators. Global, regional, and national incidence, prevalence, and years lived with disability for 301 acute and chronic diseases and injuries in 188 countries, 1990–2013: a systematic analysis for the Global Burden of Disease Study 2013. *Lancet*. 386, 743–800 (2013).
12. Coca, S.G., Singanamala, S. & Parikh, C.R. Chronic kidney disease after acute kidney injury: a systematic review and meta-analysis. *Kidney Int*81, 442–448 (2012).
13. Basu, R.K., et al. Assessment of Worldwide Acute Kidney Injury, Renal Angina and Epidemiology in critically ill children (AWARE): study protocol for a prospective observational study. *BMC Nephrol*16, 24 (2015).
14. Eckardt, K.U., et al. Evolving importance of kidney disease: from subspecialty to global health burden. *Lancet*382, 158–169 (2013).
15. Wuhl, E., et al. Timing and outcome of renal replacement therapy in patients with congenital malformations of the kidney and urinary tract. *Clin. J. Am. Soc. Nephrol*8, 67–74 (2013).
16. Group, E.T., et al. Strict blood-pressure control and progression of renal failure in children. *N. Engl. J. Med*361, 1639–1650 (2009).
17. van Stralen, K.J., Borzych-Duzalka, D., Hataya, H. & et al. Survival and clinical outcomes of children starting renal replacement therapy in the neonatal period. *Kidney Int*86 (2014).
18. Querfeld, U., et al. The Cardiovascular Comorbidity in Children with Chronic Kidney Disease (4C) study: objectives, design, and methodology. *Clin. J. Am. Soc. Nephrol*8, 1642–1648 (2010).
19. Hoy, W.E., et al. The early development of the kidney and implications for future health. *Journal of developmental origins of health and disease*1, 216–233 (2010).
20. Flynn, J.T., et al. The effect of abnormal birth history on ambulatory blood pressure and disease progression in children with chronic kidney disease. *J. Pediatr*165, 154–162 e151 (2014).
21. Rodriguez, M.M., et al. Histomorphometric analysis of postnatal glomerulogenesis in extremely preterm infants. *Pediatric and developmental pathology : the official journal of the Society for Pediatric Pathology and the Paediatric Pathology Society*7, 17–25 (2004).
22. Abitbol, C.L., et al. Long-term follow-up of extremely low birth weight infants with neonatal renal failure. *Pediatr. Nephrol*18, 887–893 (2003).
23. Hodgin, J.B., Rasoulpour, M., Markowitz, G.S. & D'Agati, V.D. Very low birth weight is a risk factor for secondary focal segmental glomerulosclerosis. *Clin. J. Am. Soc. Nephrol*4, 71–76 (2009).
24. Verhave, J.C., Wetzels, J.F. & van de Kar, N.C. Novel aspects of atypical haemolytic uraemic syndrome and the role of eculizumab. *Nephrol. Dial Transplant*29 Suppl. 4, iv131–141 (2014).
25. Torres, V.E. Vasopressin receptor antagonists, heart failure, and polycystic kidney disease. *Annu Rev. Med*66, 195–210 (2015).
26. Jarzembowski, T., et al. Impact of non-compliance on outcome after pediatric kidney transplantation: an analysis in racial subgroups. *Pediatr. Transplant*8, 367–371 (2004).
27. Watson, A.R. Non-compliance and transfer from paediatric to adult transplant unit. *Pediatr. Nephrol*14, 469–472 (2000).
28. Aujoulat, J., et al. Non-adherence in adolescent transplant recipients: the role of uncertainty in health care providers. *Pediatr. Transplant*15, 148–156 (2011).
29. Watson, A.R., et al. Transition from pediatric to adult renal services: a consensus statement by the International Society of Nephrology (ISN) and the International Pediatric Nephrology Association (IPNA). *Kidney Int*80, 704–707 (2011).
30. Watson, A.R., et al. Transition from pediatric to adult renal services: a consensus statement by the International Society of Nephrology (ISN) and the International Pediatric Nephrology Association (IPNA). *Pediatr. Nephrol*26, 1753–1757 (2011).
31. Gallieni, M., et al. The burden of hypertension and kidney disease in Northeast India: the Institute for Indian Mother and Child noncommunicable diseases project. *The Scientific World Journal*2014, 320869 (2014).
32. White, A., Wong, W., Sureshkumar, P. & Singh, G. The burden of kidney disease in indigenous children of Australia and New Zealand, epidemiology, antecedent factors and progression to chronic kidney disease. *Journal of paediatrics and child health*46, 504–509 (2010).
33. Zarocostas, J. Need to increase focus on non-communicable diseases in global health, says WHO. *BMJ*341, c7065 (2010).
34. Gulland, A. WHO agrees to set up body to act on non-communicable diseases. *BMJ*346, f3483 (2013).
35. Feehally, J. Chronic kidney disease: Health burden of kidney disease recognized by UN. *Nat Rev Nephrol*8, 12–13 (2011).
36. Couser, W.G., Remuzzi, G., Mendis, S. & Tonelli, M. The contribution of chronic kidney disease to the global burden of major noncommunicable diseases. *Kidney Int*80, 1258–1270 (2011).