What we do and do not know about women and kidney diseases; Questions unanswered and answers unquestioned: Reflection on World Kidney Day and International Woman’s Day.

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Abstract

Chronic Kidney Disease affects approximately 10% of the world’s adult population: it is within the top 20 causes of death worldwide, and its impact on patients and their families can be devastating. World Kidney Day and International Women’s Day in 2018 coincide, thus offering an opportunity to reflect on the importance of women’s health and specifically their kidney health, on the community, and the next generations, as well as to strive to be more curious about the unique aspects of kidney disease in women so that we may apply those learnings more broadly.

Girls and women, who make up approximately 50% of the world’s population, are important contributors to society and their families. Gender differences continue to exist around the world in access to education, medical care, and participation in clinical studies. Pregnancy is a unique state for women, offering an opportunity for diagnosis of kidney disease, but also a state where acute and chronic kidney diseases may manifest, and which may impact future generations with respect to kidney health. There are various autoimmune and other conditions that are more likely to impact women with profound consequences for child bearing, and on the fetus. Women have different complications on dialysis than men, and are more likely to be donors than recipients of kidney transplants.

In this editorial, we focus on what we do and do not know about women, kidney health, and kidney disease, and what we might learn in the future to improve outcomes worldwide.

Keywords: women, access to care, kidney health, acute and chronic kidney disease, inequities

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**Kidney Health and Women’s Health: a case for optimizing outcomes for present and future generations**

**Introduction**

Chronic Kidney Disease (CKD) affects approximately 10% of the world’s adult population: it is within the top 20 causes of death worldwide (1), and its impact on patients and their families can be devastating. World Kidney Day and International Women’s Day in 2018 coincide, thus offering an opportunity to reflect on the importance of women’s health and specifically their kidney health, on the community, and the next generations; as well as to strive to be more curious about the unique aspects of kidney disease in women, so that we may apply those learnings more broadly.

Girls and women, who make up approximately 50% of the world’s population, are important contributors to society and their families. Besides childbearing, women are essential in childrearing and contribute to sustaining family and community health. Women in the 21st century continue to strive for equity in business, commerce, and professional endeavours, while recognizing that in many situations, equity does not exist. In various locations around the world, access to education and medical care is not equitable amongst men and women; women remain under-represented in many clinical research studies, thus limiting the evidence base on which to make recommendations to ensure best outcomes (Figure 1).

In this editorial, we focus on what we do and do not know about women’s kidney health and kidney disease, and what we might learn in the future to improve outcomes for all.

**What we know and do not know**

Pregnancy is a unique challenge and is a major cause of acute kidney injury (AKI) in women of childbearing age; AKI and pre-eclampsia (PE) may lead to subsequent CKD, but the entity of the risk is not completely known (2-5). CKD has a negative effect on pregnancy even at very early stages (6-7). The risks increase with CKD progression thus posing potentially challenging ethical issues around conception and maintaining of pregnancies (6-8). We do know that PE increases the probability of hypertension and CKD in later years, but we have not evaluated a surveillance or reno-protective strategy to determine if progressive loss of kidney function can be attenuated (9-12).

Specific systemic conditions like Systemic Lupus Erythematosus (SLE), Rheumatoid Arthritis (RA), and Systemic Scleroderma (SS), are more likely to affect women than men. We do not know the relative contribution of these acute and chronic conditions on progression to end-stage renal disease (ESRD) in women.

In CKD cohorts, the prevalence in women is always less than in men, and they have slower progression to ESRD (13-15). We do not know why and how much of this is due to differences in identification of kidney impairment, different access to care, or true difference in disease severity and prevalence.

Women with CKD have a higher cardiovascular risk than women without CKD (16); but their risk is still lower than that of men with similar degrees of kidney impairment. In hemodialysis cohorts, there are differences in vascular access types in women versus men, which may be due to biological or systemic factors. In some locations there is differential use of peritoneal and hemodialysis in women and men.
Women are more likely to donate kidneys for transplantation than to receive them. We do not know if this is because of the differential incidence of CKD in men vs women, cultural factors, or other reasons.

There remain gender differences in access to care in different regions of the world, and we do not have data to directly evaluate the extent of these differences, in the poorest parts of the world in particular.

**Pregnancy, preeclampsia, pregnancy-induced hypertensive disorders, and fetal health. The importance of women’s health to present and future kidney health**

**What we know**

PE is the principal cause of AKI and maternal death, particularly in developing countries (2,17). Pregnancy is the most common cause of AKI in women of childbearing age (10,18,19). Several diseases and conditions, besides PE, hypertensive disorders of pregnancy, and CKD, can lead to pregnancy-related AKI. Causes vary in different regions. Septic abortion after an illegal procedure is the leading cause of early AKI in countries where legal abortions are not available, while PE after assisted fertilization is becoming a leading cause in developed countries (12,20-22).

PE and hypertensive disorders of pregnancy occur in 3-10% of all pregnancies (2-3,18); in these disorders the kidney is the main target of an unbalanced pro-angiogenic and anti-angiogenic derangement, leading to hypertension, proteinuria, and widespread endothelial damage. The incidence of PE, higher in low-middle income countries (possibly reflecting undiagnosed predisposing diseases), peaks at the extremes of reproductive age for reasons mentioned above (12,20-22).

The relationship between kidney and placenta is biunivocal, and the presence of CKD is a risk factor for PE and hypertensive disorders of pregnancy (Figure 2). Besides CKD, other conditions cited as risk factors for PE (diabetes, immunologic diseases, baseline hypertension, obesity, and metabolic syndrome), are also risk factors for CKD. Given that even minor alterations of kidney function are present in many of these disorders, the importance of kidney function is indirectly recognized in the development of PE. Newer definitions of PE recognize differences between “placental” and “maternal” causes of PE, based on novel angiogenic-antiangiogenic markers (23,24), which may be important for management during and after pregnancy.

*There are long term effects of PE on both maternal and fetal health, but this remains an area of active research with many unknowns.*

PE is a risk factor for the future development of CKD and ESRD in the mother (3-5). The reasons are not fully understood; podocyte loss is a hallmark of PE, suggesting permanent glomerular damage (25). Endotheliosis, associated with PE, but also found in normal pregnancies, may herald glomerulosclerosis; tubular and vascular damage may co-exist (26,27).

Besides maternal risks, PE is associated with intrauterine and perinatal death, preterm delivery, and restricted intrauterine growth; the latter two are linked to “small babies” (2-3,5). Small babies and preterm babies have highly increased risks of neurological deficits and postnatal complications, especially sepsis (28-32). The risks may be higher in low-income countries, since survival and deficit-free survival depend on the provision of postnatal intensive care (20,21). In the long term, small babies are at risk for the development of diabetes, metabolic syndrome, cardiovascular diseases (CVDs), and CKD in adulthood.
Since kidney development is completed in the last phases of pregnancy, delayed, insufficient kidney growth, resulting in low nephron number is probably the basis of the increased risk of CKD and hypertension in small for gestational age, and preterm babies (33-37).

**Pregnancy in chronic kidney disease, dialysis, and transplantation**

**What we know**

**Chronic kidney disease**

CKD is a risk factor for adverse pregnancy outcomes from its early stages (Table 1) (6,38,39). The risks increase from CKD stage 1 to CKD stage 5, and may be higher in glomerular nephropathies, autoimmune diseases, and diabetic nephropathy (6,7,38-41). Results of pregnancy after kidney donation suggest that reduction of kidney parenchyma may be associated with a higher risk of PE and hypertensive disorders of pregnancy (42,43).

Hypertension and proteinuria at baseline are important modulators of pregnancy-related risks; among the risks, we know that malformations are not increased with respect to the overall population (out of the context of inherited diseases, such as reflux nephropathy, polycystic kidney disease, or congenital anomalies of the kidney and urinary tract), maternal death is unusual (in highly resourced countries), while the incidence of preterm delivery and of small for gestational age babies, intrinsically linked, is increased in stage 1 CKD patients, and rises with the worsening of kidney function. Likewise, the effect of pregnancy on CKD progression is not fully understood because of different study designs, obstetric policies, and duration of follow-up. Overall, short- and long-term decrease in kidney function is unusual in early CKD, but the risk increases as CKD severity increases (6,7,38-41,44-48).

Pregnancy is a potential occasion for the initial diagnosis of CKD. In poorly or unevenly resourced countries, advanced CKD may be discovered only during pregnancy. The implications of dialysis initiation may present important clinical and ethical issues; in highly resourced countries with established prenatal care, the diagnosis of earlier stages of CKD may lead to more intensive therapy and surveillance (49-51).

**Dialysis and transplantation**

Fertility is reduced in ESRD; Australian and European data suggest a 1:10 ratio from general population to transplantation and from transplantation to dialysis (1:100 probability as compared to the general population) (52,53). The first sporadic cases of successful pregnancy on dialysis were described in the 70s, but in the new millennium this became an acknowledged real clinical possibility (8,54,55).

More than 1000 pregnancies have been reported in dialysis patients (55). The most important advance has been the demonstration of a strong relationship between the intensity (frequency and duration) of the dialysis sessions and positive pregnancy results: thus, intensifying dialysis up to daily, is the current standard of care (8,54). Changing attitudes towards counselling women with advanced CKD may be impacted, with the knowledge of positive outcomes on dialysis for women and their offspring.

Fertility is partly restored after kidney transplantation (56-60). However, even in an ideal situation (normal kidney function, no hypertension or proteinuria, at least 2 years after transplantation, without recent rejection episodes), the risk of complications is higher in women with transplanted kidneys than in the
general population. However, if teratogen drugs are avoided (mycophenolic acid and rapamycin), the outcomes of pregnancy after kidney transplantation shares the same risk factors as CKD (kidney function, hypertension, and proteinuria) (59).

Experience with pregnancy in patients with a reduced renal function or failing kidney graft is limited and counseling is still forcibly based on personal experience or indirect evidence (61,62). Assisted fertilization techniques are increasingly popular in some settings, but dedicated studies in CKD patients are few; multiple pregnancies may bear an added risk in CKD patients, with both native and transplanted kidneys.

Autoimmune diseases, women, and kidney disease

What we know

Autoimmune diseases such as SLE, RA, and SS preferentially affect women and are characterized by systemic inflammation leading to target organ dysfunction, including kidneys. Sex differences in the incidence and severity of these diseases result from a complex interaction of hormonal, genetic, and epigenetic factors (Table 2). The public health burden of autoimmune diseases, which collectively represent a leading cause of morbidity and mortality among women throughout adulthood, is substantial (63-65).

SLE is an autoimmune disease with multiple organ involvement, affecting approximately five million people worldwide; disproportionately predominant in women (9:1 female to male ratio) and individuals of non-European ancestry. The highest female predominance (up to 15:1) is in peak reproductive years. The biology of these differences has been explored: one explanation is the number of X chromosomes and genetic variants on the X chromosome (66-68); another important etiological explanation is the role of estrogen in SLE. Estrogen’s primary effects are mediated by transcription activity of the intracellular estrogen receptors, whose profile is altered in T-cells from female SLE patients (69,70). Cathepsin S protein has recently been identified as a potential cause of lupus, triggering the immune system to attack healthy cells, particularly in females (71). Numerous non-HLA genetic markers may predispose individuals of European, Hispanic, and Afro-American ancestry to lupus (72). Susceptibility to SLE during pregnancy is also multifactorial; one factor being upregulation of IFN-α. Elevated IFN-α, expressed by the placenta, plays a pathogenic role in SLE, contributing both to the success of placental reproduction and to increased susceptibility to SLE (73). Regulatory T-cells (which may be the key to cell modulating feto-maternal tolerance) have abnormalities of structure and function, and may contribute to pregnancy pathology in women with SLE and to challenges of managing them during pregnancy (74). SLE affects kidneys in about 50% of patients, including glomerular, interstitial, and vascular lesions. Lupus nephritis is a major risk factor for overall morbidity and mortality in SLE, and despite potent therapies still leads to significant impairment of kidney function for many patients (75). Kidney disease is a critical concern in counseling women with lupus considering pregnancy, with previous kidney involvement and lower C4 levels conferring high risk of active nephritis occurring in pregnancy (76). Socioeconomic disparities are also linked to the health of patients with lupus. Poverty is associated with an increased long-term level of accumulated disease-associated damage and a 1.67-times increased likelihood of experiencing a clinically meaningful increase in damage. Frequency of adverse pregnancy outcomes in women with lupus is twofold higher in black and Hispanic women than in white women. In blacks, socioeconomic status was a determinant of pregnancy outcomes and a key contributor to adverse pregnancy outcomes (77,78).

RA also preferentially affects women (4:1 ratio to men) with the peak incidence at age 45-55, coinciding with the perimenopausal years. This suggests a possible association between estrogen deficiency and
disease onset. Female-to-male incidence ratio after age 60 years is approximately 1:1, potentially implicating changes in sex hormones in the development of RA, and a pattern of RA symptom improvement or even remission during pregnancy is well recognized (79-81). Renal involvement in RA is relatively common and multifactorial and is a predictor of mortality in RA patients. The risk of CKD is significantly higher in patients with RA than in the general population. The development of CKD may result from several ongoing processes, including specific renal involvement associated with RA (e.g., glomerulonephritis, interstitial nephritis), chronic inflammation, comorbidities, and nephrotoxic anti-rheumatic drugs. The strong association between RA activity and AA amyloidosis increases morbidity and is the main cause of ESRD with RA and nephropathy. Importantly, some of the life-long and combined RA pharmacotherapy can lead to various renal side effects (82-84).

SS predominantly affects women (female-to-male ratios ranging from 3:1 to 14:1), with the peak incidence in the fifth and sixth decades. Estrogen may play a role in scleroderma pathogenesis through its stimulatory effect on transforming growth factor-beta 1 receptor and platelet-derived growth factor receptor (85). Vasculopathy is an important disease-related manifestation in SS, and the low estrogenic state associated with menopause has been suggested to aggravate vascular manifestations in affected women (86). SS can also be complicated by a number of different forms of kidney disease, including scleroderma renal crisis, which represents a form of malignant hypertension with acute renal failure; or more commonly ischemic nephropathy leading to slowly progressive CKD, accompanied by hypertension and albuminuria (78). Nondiabetic acute renal failure in patients with SS may be caused by interstitial nephritis or ANCA vasculitis, a separate entity in scleroderma with poor outcome (87-89).

**Women, chronic kidney disease, and access to renal replacement therapies**

**What we know**

Although renal replacement therapy (RRT), including dialysis and transplantation is life-sustaining, not all patients receive RRT. The rate of ESRD treated by RRT differs greatly between countries and regions, and intricately depends on the economy of a country and health care system (90,91). Worldwide, only 50% of patients requiring RRT receive treatment (92), and in low and middle-income countries and regions, even less; in large parts of Sub-Saharan Africa, less than 2% of ESRD are treated by RRT (93). The equality of access to RRT for women and girls is of particular concern because, in many societies, they are disadvantaged by discrimination rooted in sociocultural factors (94,95).

**Sex differences in access to dialysis**

At least 2.284 million people may have died prematurely due to lack of access to RRT with treatment gaps being much larger in low-income countries, with conservative estimates in Asia and Africa of 1.907 million and 432,000 people not receiving RRT. By 2030, the estimated number of RRT should be more than double to 5.439 million (3.899–7.640 million), with the most growth in Asia (0.968 million to a projected 2.162 million [1.71–3.14 million]) (92). These numbers are derived from an extensive systematic review.

There are few data to compare the gender difference for the treatment gaps. Studies in Africa show that men were more likely to receive RRT than women (96,97). In Japan, the incidence of treated ESRD in females was less than half of that in males (3,287 in males vs. 1,764 women per million population treated) (91): no explanations are given for this finding. One US study reports women having significantly higher odds ratio of 1.70 for late initiation of dialysis compared to men (98). Awareness levels of previous kidney
disease in women were reported much lower than in men (2.9% ± 1.6% in women vs. 17.9% ± 5.9% in men), which may contribute to later initiation of RRT (99).

Mortality rates are similar in men and women on dialysis, but the incident rates of some dialysis-associated complications and morbidity are higher in women. A US report of hospitalizations in 111,653 patients undergoing maintenance hemodialysis describes higher hospitalization rates in women, and higher risk for 30-day readmissions (100).

In addition, the prevalent use of arteriovenous fistula, which is associated with reduced mortality, complication, and costs, is lower among female than male hemodialysis patients (101). This may be due to a number of different factors, including anatomical/surgical issues relating to vessel size, timing of referral, and attitudinal differences. This has not been systematically studied.

Dialysis dose, which is evaluated by Kt/V may result in under-dialysis in women who have an average smaller volume of urea distribution or total body water than men (102). Women receiving dialysis have also been reported to have worse clinical parameters including anemia, nutrition, and quality of life (103). Reasons are not certain.

**Sex differences in access to kidney transplantation**

Transplantation represents the best form of RRT in patients without contraindications. Worldwide data describes that women are less likely than men to be kidney transplant recipients, either from a cadaveric or living donor, but are more likely to serve as living donors for kidney transplantation (104). Data from different countries, including the US, France, China, and India, confirm differential kidney transplant rates (lower in women than men), less likelihood of women being registered on national transplant waiting lists, and longer time from dialysis initiation to listing. Mothers are more likely to be donors, as are female spouses (91,105-108). Sex inequality also exists in the pediatric population. A survey from 35 countries participating in the European Society for Pediatric Nephrology/European Renal Association-European Dialysis and Transplant Association Registry, reported girls had a lower access to renal transplantation than boys (109).

Socioeconomic factors undoubtedly play a role in the inequality of transplantation between sexes, especially in the low and middle-income countries and regions. Generally, men provide the major income for their family which may discourage them to donate kidneys. Different employment status and incomes between genders may contribute to sex differences in transplantation because employment and income status is usually associated with better healthcare insurance which cover the costs for transplantation. Psychosocial factors and education of women have been suggested as a contribution to sex disparity. US data found black women were less likely to want living donor kidney transplantation compared with men, despite being twice as likely as men to receive unsolicited offers for kidneys. They were also less likely to have been evaluated for a kidney transplant (110). Other reports describe disparities in age and sex in access to kidney transplantation which originate at the time of pre-referral discussions about kidney transplantation; irrespective of age, women were more likely not to have had discussions with medical professionals. This result may imply that there is a need for better clinical guidelines and education for women, their social network, and their providers (111).

**Present and future what we do not know**
Given the data presented above with respect to pregnancy, AKI, autoimmune diseases, CKD, dialysis and transplantation, there are many unanswered questions. In high income countries with increasing maternal age and assisted fertilization, there may be an increase in PE which may impact future generations if associated with adverse fetal outcomes. The increase in in-vitro fertilization techniques for those of advanced maternal age may lead to multiple pregnancies, which may predispose to PE, intrauterine growth restriction, or both. Will this lead to an increase in CKD and CVD for women in the future?

Due to the high heterogeneity of CKD, we do not know if and how pregnancy outcomes are modulated by the different nephropathies, as besides the most common ones such as IgA or lupus nephropathy, diabetic nephropathy, and reflux nephropathy, evidence is scant (44,45,112-114). How should we define preconception risks of pregnancy with respect to current proteinuria cut offs? Indications on when to start dialysis in pregnancy are not well established, nor is the specific role of frequency and duration. In those with kidney transplants, given the changing expanded donor policies, higher age at transplantation, and reduced fertility in older women, there may be changes in attitudes towards pregnancy with less than optimal kidney function (56,60). How this will impact short and long-term outcomes of mothers and their babies is not clear.

Teen pregnancies are very common in some parts of the world, and are often associated with low income and cultural levels. The uneven legal rules for assisted fertilization and the lack of systematic assessment of the kidney function point to the need for further research.

Despite elegant demonstrations for the role of sex hormones in vascular health and immunoregulation, the striking predominance in females of SLE, RA, and SS remains unexplained relative to other systemic diseases such as ANCA vasculitis and hemolytic-uremic syndrome. Note that thrombotic thrombocytopenic purpura has a higher incidence in women, though this is likely due to the association with other conditions more common in women. The incidence of kidney involvement in SLE during pregnancy and similarities/differences in those with PE have not been well studied. The role of different medications and responses to medications for autoimmune diseases relative to sex has also not been well studied.

More attention to similarities between conditions, the importance of sex hormones in inflammation, immune-modulation, and vascular health, may lead to important insights and clinical breakthroughs over time. If women are more likely to be living donors, at differential ages, does this impact both CVD risk, and risk for ESKD: have we studied this well enough, in the current era, with modern diagnostic criteria for CKD and sophisticated tools to understand renal reserve? Are the additional exposures that women have after living donation compounded by hormonal changes on vasculature as they age? And are the risks of CKD and PE increased in the younger female kidney living donor?

In the context of specific therapies for the treatment or delay of CKD progression, do we know if there are sex differences in therapeutic responses to ACEi/ARB? Should we look at dose finding/adjustments by sex? If vascular and immune biology is impacted by sex hormones as described earlier, do we know the impact of various therapies by level or ratio of sex hormones? In low-middle income countries how does changing economic and social cultures impact women’s health, and what is the nutritional impact on CKD of increasing predominance of obesity, diabetes, and hypertension?

Summary
Women have unique risks for kidney diseases: kidney diseases, as well as issues related to access to care, have a profound impact on both the current and next generations. Advocating for improved access to care for women is critical to maintain the health of families, communities, and populations.

Focused studies on the unique contribution of sex hormones, or the interaction of sex hormones and other physiology, is important to improve our understanding of the progression of kidney diseases. Immunological conditions such as pregnancy (viewed as a state of tolerance to non-self) as well as SLE and other autoimmune and systemic conditions common in women, better studied may also lead to breakthroughs in understanding and care paradigms.

There is a clear need for higher awareness, timely diagnosis, and proper follow up of CKD in pregnancy. In turn, pregnancy may also be a valuable occasion for early diagnosis of CKD, allowing planning of therapeutic interventions.

On this occasion, World Kidney Day and the International Women’s Day 2018 are commemorated on the same day, offering us the opportunity to highlight the importance of women’s health and particularly their kidney health. On its 13th anniversary, World Kidney Day promotes affordable and equitable access to health education, healthcare, and prevention for all women and girls in the world.

The coinciding of World Kidney Day and International Women’s Day offers an opportunity to develop and define best practices and future research agendas, and ultimately, to optimize the outcomes of all people living with or at risk for kidney disease.

Disclosure

All authors have contributed to the manuscript equally. None of the authors have direct conflicts of interest with this paper and material contained herein; full disclosures are listed in the individual authors’ Conflict of Interest forms.
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Table 1. Adverse pregnancy outcomes in patients with chronic kidney disease and in their offspring

<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
<th>Main Issues</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maternal death</td>
<td>Death in pregnancy or within 1 week-1 month postpartum</td>
<td>Too rare to be quantified, at least in highly resourced settings, where cases are in the setting of severe flares of immunologic diseases (SLE in primis). Still an issue in AKI; and in low resourced countries; not quantified in low-resourced countries, where it merges with dialysis need.</td>
</tr>
<tr>
<td>CKD progression</td>
<td>Decrease in GFR, rise in sCr, shift to a higher CKD stage</td>
<td>Differently assessed and estimated; may be linked to obstetric policy (anticipating delivery in the case of worsening of the kidney function); between 20% and 80% in advanced CKD. Probably not increased in early CKD stages.</td>
</tr>
<tr>
<td>Immunologic flares and neonatal SLE</td>
<td>Flares of immunologic diseases in pregnancy</td>
<td>Once thought to be increased in pregnancy, in particular in SLE, are probably a risk in patients who start pregnancy with an active disease, or with a recent flare-up. Definition of a “safe” zone is not uniformly agreed; in quiescent, well controlled diseases do not appear to be increased with respect to non-pregnant, carefully-matched controls.</td>
</tr>
<tr>
<td>Transplant rejection</td>
<td>Acute rejection in pregnancy</td>
<td>Similar to SLE, rejection episodes are not increased with respect to matched controls; may be an issue in unplanned pregnancies, in unstable patients.</td>
</tr>
<tr>
<td>Abortion</td>
<td>Fetal loss, before 21-24 gestational weeks</td>
<td>May be increased in CKD, but data are scant. An issue in immunologic diseases (eventually, but not exclusively linked to the presence of LLAC) and in diabetic nephropathy.</td>
</tr>
<tr>
<td>Stillbirth</td>
<td>Delivery of a nonviable infant, after 21-24 gestational weeks</td>
<td>Probably not increased in early CKD, maybe an issue in dialysis patients; when not linked to extreme prematurity, may specifically linked to SLE, immunologic diseases and diabetic nephropathy.</td>
</tr>
<tr>
<td>Perinatal death</td>
<td>Death within 1 week – 1 month form delivery</td>
<td>Usually a result of extreme prematurity, which bears a risk of respiratory distress, neonatal sepsis, cerebral hemorrhage.</td>
</tr>
<tr>
<td>Small, very small baby</td>
<td>A baby weighting &lt; 2500-1500 g at birth</td>
<td>Has to be analyzed with respect to gestational age.</td>
</tr>
<tr>
<td>Preterm, early extremely preterm</td>
<td>Delivery before 37 – 34 or 28 completed gestational weeks</td>
<td>Increase in risk of preterm and early preterm delivery across CKD stages; extremely preterm may be an important issue in undiagnosed or late referred CKD and PE-AKI.</td>
</tr>
<tr>
<td>SGA (IUGR)</td>
<td>&lt; 5th or &lt; 10th centile for gestational age</td>
<td>Strictly and inversely related to pre-term delivery; SGA and IUGR are probably related to risk for hypertension, metabolic syndrome and CKD in adulthood.</td>
</tr>
<tr>
<td>Malformations</td>
<td>Any kind of malformations</td>
<td>Malformations are not increased in CKD patients not treated by teratogen drugs (MMF, mTor inhibitors, ACEI, ARBS); exception: diabetic nephropathy (attributed to diabetes); hereditary diseases, such as PKD, reflux nephropathy, CAKUT may be evident at birth.</td>
</tr>
<tr>
<td>Hereditary kidney diseases</td>
<td>Any kind of CKD</td>
<td>Several forms of CKD recognize a hereditary pattern or predisposition; besides PKD, reflux and CAKUT, Alport’s disease, IgA, kidney tubular disorders and mitochondrial diseases have a genetic background, usually evident in adulthood and not always clearly elucidated.</td>
</tr>
<tr>
<td>CKD - hypertension</td>
<td>Higher risk of hypertension and CKD in adulthood</td>
<td>Late maturation of nephrons results in a lower nephron number in preterm babies; the risks are probably higher in SGA-IUGR babies than in pre-term babies adequate for gestational age.</td>
</tr>
<tr>
<td>Other long-term issues</td>
<td>Developmental disorders</td>
<td>Mainly due to prematurity, cerebral hemorrhage or neonatal sepsis, are not specific of CKD, but are a threat in all preterm babies.</td>
</tr>
</tbody>
</table>

SLE = Systemic Lupus Erythematosus; AKI = acute kidney injury; GFR = glomerular filtration rate; sCr = serum creatinine; CKD = chronic kidney disease; LLAC = Lupus-like anticoagulant; PE-AKI = preeclampsia acute kidney injury; SGA = small for gestational age; IUGR = intrauterine growth restriction; MMF = mycophenolate mofetil; mTor = mechanistic target of rapamycin; ACEI = angiotensin-converting-enzyme inhibitor; ARBS = angiotensin II receptor blockers; PKD = polycystic kidney disease; CAKUT = congenital anomalies of the kidney and urinary tract; IgA = immunoglobulin A
Table 2. Sex differences in the incidence and severity of autoimmune diseases

<table>
<thead>
<tr>
<th></th>
<th>SLE</th>
<th>RA</th>
<th>SS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peak Incidence</td>
<td>Reproductive age</td>
<td>Perimenopausal</td>
<td>After 50-60 years</td>
</tr>
<tr>
<td>Female/Male Ratio</td>
<td>Peak 15:1</td>
<td>Peak 4:1</td>
<td>Peak 14:1</td>
</tr>
<tr>
<td></td>
<td>Total 9:1</td>
<td>After 60 years</td>
<td>Total 3:1</td>
</tr>
<tr>
<td>Influence of Estrogen</td>
<td>High levels</td>
<td>Negative</td>
<td>Positive</td>
</tr>
<tr>
<td></td>
<td>Low levels</td>
<td>Unknown</td>
<td>Negative</td>
</tr>
</tbody>
</table>

SLE = Systemic Lupus Erythematosus; RA = Rheumatoid Arthritis; SS = Systemic Scleroderma

Figure 1. Sex differences throughout the continuum of CKD care

SLE = Systemic Lupus Erythematosus; RA = Rheumatoid Arthritis; SS = Systemic Scleroderma; AKI = acute kidney injury; CKD = chronic kidney disease; AI = autoimmune; AVF = arteriovenous fistula; HD = hemodialysis; KT = kidney transplant
Figure 2. Pregnancy and kidney function: complex interactions between 2 organs, the kidney and placenta

From the placenta to the kidney
PE may induce permanent kidney damage, via AKI, tubular damage, podocyte loss

Placenta and kidney are highly vascularized; filter blood; divide compartments; are sophisticated metabolic machines
Pregnancy is a precious occasion to diagnose CKD.

From the kidney to the placenta
CKD may induce placental dysfunction with an increased risk of pre-term delivery, hypertensive disorders of pregnancy and PE

PE = preeclampsia; AKI = acute kidney injury; CKD = chronic kidney disease