The Kidney from Prenatal to Adult Life: Perinatal Programming and Reduction of Number of Nephrons during Development

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Introduction

The processes of development and maturation of organs occur continuously throughout the pre- and postnatal periods. Intrauterine growth is generally regulated by intrinsic growth potential, genetic endowment, and support of extrinsic growth provided by nutrients and oxygen from the materno-uteroplacental unit. However, during the postnatal period growth may be affected by environmental conditions and genetic background. Barker et al. [1, 2] and Brenner et al. [3] reported how modified embryonic-fetal development resulting in low birth weight (LBW) may lead to a reduced nephron endowment, hypertension and renal diseases in adulthood. Regarding the involvement of genetic factors, several environmental conditions may also contribute towards reducing the number of nephrons in the fetus and infant, subsequently constituting a health burden in later life.

Reduction in the Number of Nephrons and Disease

In humans the overall number of nephrons ranges between 617,000 and 1,075,000 (mean 850,000 nephrons). Variations in nephron number have been associated with different clinical conditions: hypertensive adults have a...
mean of ~702,000 nephrons, lower than that displayed by normotensive subjects with a mean of 1,429,000 [4].

The pathophysiological characteristics of diseases associated with a reduced number of glomeruli are linked to a cascade of events ultimately resulting in compensatory nephron hypertrophy. A low number of nephrons is associated with global renal reduced glomerular volume (although in a single nephron there is a compensatory increase in glomerular volume), reduced surface filtration area, impaired tubular function and changes in vascular permeability.

In the mouse model, the loss of one allele per glial cell line-derived neurotrophic factor (GDNF) resulted in an approximately 30% reduction in normal sized glomeruli. At a later stage of life the same animals developed high arterial blood pressure featuring glomerular hypertrophy and hyperfiltration of kidneys manifested as a compensatory mechanism for the reduced number of nephrons [5]. In spite of the finding in humans that small for gestational age (SGA) newborns displayed shorter renal length than appropriate for gestational age (AGA), on ageing they manifested either an accelerated renal maturation process or early compensatory kidney hypertrophy [6]. A low nephron endowment led to the onset of compensatory mechanisms in the residual nephrons consisting of glomerular hypertension, high nephron filtration rates, and lastly glomerular hypertrophy [7]. On the basis of the hyperfiltration hypothesis, the larger glomeruli predispose an individual to developing hyperfiltration injury and further deterioration of renal function from increased workload, proteinuria with glomerulosclerosis, tubulointerstitial inflammation and fibrosis [7, 8].

The risk factors underlying the onset of hypertension and renal disease are: family history of hypertension and renal disease, diabetes mellitus, obesity, dietary factors, insulin resistance, gestational diabetes, and at-risk ethnic groups. Oligonephronia, whether or not associated with LBW, should be added to a growing list of risk factors of progressive renal disease. Hypertension produces as a result of reduced glomerular number may also be responsible for microalbuminuria, a known early marker for renal disease and hypertension [9]. Moreover, the marked reduction in nephron number characterizing renal hypoplasia/dysplasia leads to a decline in renal function and development of chronic renal failure in childhood [10]. Nevertheless, a small reduction in nephron number in subjects featuring intrauterine growth retardation (IUGR) and LBW has also been associated with renal failure and hypertension in adult life [11].

Causes Underlying a Reduction in Nephron Number

Perinatal programming controls nephrogenesis during the developmental stage up to 34–36 weeks of gestational age. In infants born before 36 weeks of gestation, nephrogenesis is still ongoing following preterm delivery [12]. Intrauterine stress or prenatal or postnatal perturbation in preterm infants may result in a reduced nephron number. Any adverse event occurring prior to completion of nephrogenesis likely compromises renal growth and produces a longer-lasting effect on final renal potential. However, perinatal programming for hypertension and diabetes may elicit a synergistic impact with the reduced nephron number leading to the development of chronic kidney disease [12, 13].

The causes underlying a reduced number of nephrons in an individual are both genetic and environmental. The environmental impact on a genetic program yields the renal perinatal programming of each individual. Furthermore, an ongoing interaction between genes and the environment from prenatal to adult life will contribute towards forming the renal potential of an individual. Signal molecules and transcription factors have been implicated in determining segmental nephron identity and functional differentiation. While some of these genes (p53 gene family, hepatocyte nuclear factor-1β) promote the terminal epithelial differentiation fate, others (Notch, Brn-1, IRX, KLF4, and Foxi1) regulate the differentiation of specific nephron segments and cellular types [14, 15]. Congenital oligomeganephronia, clinically characterized by bilateral renal hypoplasia with a reduced number of enlarged nephrons, has been associated with PAX2 gene mutations [16]. Furthermore, the renin-angiotensin system plays a critical role in kidney development. Therefore, angiotensin-converting enzyme (ACE) gene insertion/deletion (I/D) polymorphism does not affect the risk of alterations in renal morphogenesis, but is involved in the physiopathology of disease progression from acquired glomerular disorders to renal hypoplasia, dysplasia, and uropathies [17]. Additionally, genetic issues underlie the development of small, undifferentiated kidneys, nephron hypoplasia, dysplasia and hypodysplasia, in which unknown factors may modify the severity of clinical presentation. Moreover epigenetic changes, characterized by alterations in chromatin structure through modification of histone by methylation, acetylation, phosphorylation and ubiquitylation, lead to stable and potentially hereditable changes in gene expression. In particular, DNA methylation has been strongly implicated in fetal renal development and disease [18, 19].

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Accordingly, congenital renal anomalies are more frequent in relatives of patients than in controls. Therefore, variability in the presence, type, unilateral or bilateral involvement of renal malformation may be explained by diverse gene-environmental interactions among individuals of affected families [20].

Several environmental stressors may act on specific genetic programming of low nephron number (fig. 1). Alterations to genetic pattern during fetal development may be produced by nature, timing, duration and severity of the renal insult. In particular, the time at which an adverse factor is involved during gestation before completion of nephrogenesis may affect kidney growth [21]. However, malnutrition caused by a low protein-calorie diet during intrauterine and neonatal life and bad eating habits in later life (excessive salt intake, unbalanced fat and protein in the diet) may have adverse implications on renal outcome [8, 9]. Moreover, additional stressors affecting renal structure and function are a history of LBW and IUGR, vitamin A deficiency, urinai tract malformations such as obstructive uropathy and infections, administration of nephrotoxic drugs (especially antibiotics and non-steroidal anti-inflammatory drugs, NSAIDs) to mothers and newborns [22, 23]. All factors may interact to increase potential nephron damage.

**LBW and Extreme Prematurity**

LBW represented an independent risk factor for chronic renal injury in experimental studies on rats. Animal kidneys showed reduced glomerular number and high glomerular size [24, 25].

Autopsy studies in newborns and children have reported a marked association between LBW and reduced nephron number [24, 26, 27]. At birth, LBW infants without renal failure displayed a compensatory higher glomerular volume [27–29]. Low glomerular number and high glomerular size have been associated with the development of hypertension, cardiovascular disease and increased susceptibility to renal disease in later life [21, 30, 31]. Moreover, a study performed in human twins suggested that the association between LBW and renal dysfunction is closely correlated with the involvement of individual fetoplacental factors, whereas genetic factors would appear to be of lesser importance [32]. Poor socioeconomic and nutritional status in the mother may imply a reduced supply of nutrients to the developing fetus. Similarly, factors that hinder the passage of nutrients through the placenta, such as smoking and hypertension, are associated with an increased risk of LBW. Factors affecting intrauterine space availability, such as primiparity, low maternal height, and mothers who were born SGA, are additional risk factors for LBW [33].

The overall prevalence of hypertension in pediatric age is <1%. Therefore, in the presence of a neonatal history of LBW or prematurity, blood pressure should be strictly monitored during early childhood in view of the risk of hypertension [34]. LBW implied an increased risk of disease in adult life, in particular coronary heart disease, reduced insulin sensitivity and increased blood pressure [2, 35]. However, very-LBW (VLBW) infants featured a high rate (10%) of hypertension after a follow-up of 6–12 years [36]. Furthermore, preterm children were characterized by a high rate of hypertension (6.8%), with no differences between AGA and SGA. After 5 years of follow-up, renal risk in children born preterm was similar between AGA and SGA, as well as between VLBW and LBW [34, 37].

LBW was associated with hypertension and renal disease in African Americans of southeastern USA [27, 38]. In Australian Aboriginals, LBW disposed to albuminuria and a high rate of early-onset chronic renal failure [39]. LBW and SGA in Aboriginals was more than twice that observed in the general Australian population, with hypertension, cardiovascular disease, and the frequency of albuminuria being twofold the normal rates. However, susceptibility to renal disease was significantly higher than in controls [40, 41]. Long-term renal follow-up of VLBW infants born without neonatal renal failure revealed a reduction in renal function following reduced nephron endowment [36]. Furthermore, VLBW infants with renal failure showed clear evidence of active glo-
eredulerosclerosis in addition to low glomerular supply [29]. Among VLBW children with neonatal acute renal failure, almost half developed chronic renal failure, a minority of which required kidney transplant [42]. Renal damage and hypertension may constitute a risk later in life due to the presence of glomerular stress following nephron hypertrophic changes in individuals born preterm or LBW with neonatal renal complications including renal failure and signs of glomerular and tubular damage.

Microalbuminuria was the most interesting urinary marker of renal outcome in preterm infants. Indeed, the latter showed variable results in line with different study protocols, methods and populations investigated. The prevalence of microalbuminuria measured by means of traditional methods was 11%, affording variable results ranging from 2.7 [43] to 12.5% [36]. The prevalence of microalbuminuria at young adult age of SGA subjects was 3.8%, 2.4 times higher than in AGA subjects [43]. On the contrary, no difference was reported for microalbuminuria between AGA and SGA, VLBW and LBW infants and in subjects with normal and high blood pressure [37]. The variability of the results observed, linked possibly to different preterm populations studied, the number of infants and methods of analysis, is somewhat questionable.

Although IUGR was associated with both a low nephron endowment and increased risk of hypertension, only circumstantial evidence obtained in animal studies has been advocated to support the hyperfiltration hypothesis after prenatal programming [44]. Recently, the findings of several human studies did not support the hypothesis that intrauterine growth restriction and prematurity contribute to the alteration of renal function in childhood and young adult life [45, 46].

**Nutritional Setting**

Maternal nutrition may have an important influence on renal programming [2]. The majority of studies investigating the relationship between nutrition and nephrogenesis have focused on animal models. In rats, a restricted supply of nutrients to the mother during the critical window in which nephrogenesis occurs led to a reduced number of glomeruli per kidney, activation of the renin-angiotensin system, glomerular enlargement, and hypertension in later life [47].

Different nutritional conditions (IUGR, overfeeding and IUGR plus overfeeding) were investigated in 4-month-old rats to evaluate the amount of protein urinary excretion as a marker of glomerular damage. Early postnatal overfeeding in rats improved postnatal nephron number. Therefore, enhanced nephron number was associated with elevated arterial pressure and glomerulosclerosis [48].

**Vitamin A Deficiency**

Vitamin A is a determinant in fetal renal programming of rats in view of its capacity to closely modulate nephron number and vascular supply. Vitamin A and its analogs (retinoids) are important regulators of cell proliferation, differentiation, immune function and apoptosis [49]. In 21-day-old rats, the number of nephrons was directly correlated with plasma vitamin A. Vitamin A deficiency led to a reduction in nephron number in rats. Under conditions of vitamin A deprivation, proto-oncogene c-ret expression was decreased in metanephron. The role of c-ret in renal formation is considered essential since null mice for these genes exhibited renal agenesis or rudimental kidneys [50]. On the contrary, vitamin A supply restored nephron endowment to normal in offspring of rat mothers exposed to protein restriction [51].

Furthermore, indirect evidence has emphasized the role of vitamin A in kidney development in humans. Vitamin A was seen to be lacking in cord and maternal blood in IUGR neonates. However, low circulating levels were common in women who were smokers, were known to abuse alcohol, or implemented inadequate dietary practices, all situations associated with IUGR delivery [49]. Finally, maternal vitamin A deficiency accounted for subtle renal hypoplasia in Indian newborns [52].

**Obstructive and Refluxing Nephropathy**

Obstructive uropathy during the perinatal period may be a leading cause of a reduction in nephron number. Mechanical stretch, implicated in the oncoming nephropathy, leads to tubular responses via the activation of ion channels, increases in intracellular calcium levels and cellular apoptosis [53].

Unilateral obstruction and reduction in the nephron number were investigated in neonatal rat models. Following relief of unilateral urinary obstruction, the ultimate growth of the post-obstructed kidney was impaired, the number of glomeruli reduced, and GFR decreased. The major renal damage observed following unilateral ureteral obstruction was renal tubular apoptosis and atrophy [54]. The opposite kidney showed significant compensatory growth. Moreover, glomerular sclerosis, tubular atrophy, macrophage infiltration, and interstitial fibrosis were significantly increased not only in the post-obstructed kidney, but also in the opposite kidney...
[55–57]. On the contrary, urinary obstruction during adulthood did not elicit a reduction in the number of nephrons. The kidney seems to be vulnerable to urinary tract obstruction both during and immediately following nephrogenesis [56].

In newborns, unilateral obstructed kidney was characterized by a poor outcome even when surgery was performed at an early stage, due to the fact that kidneys continued to develop tubular atrophy and interstitial fibrosis [58, 59]. The persistently low growth and volume of this unilateral obstructed kidney suggested a reduction in the total nephron number subsequent to apoptosis and/or inhibition of glomerulogenesis. The unobstructed kidney developed a higher volume to compensate for the persistently low function of the affected organ [60, 61].

Similar to congenital obstruction, high-grade vesico-ureteral reflux leads to poor renal outcome. Vesico-ureteral reflux diagnosed at birth on prenatal ultrasonography was associated with early congenital damage [62]. In particular, refluxing kidneys feature smaller dimensions compared to contralateral non-refluxing organs. Prenatal diagnosis and early treatment did not modify renal outcome, thereby suggesting early fetal damage during nephrogenesis. Renal growth failure seems to correlate with the severity of vesico-ureteral reflux since it worsens in patients with an increased grade of severity [63]. Altered renal programming leading to a reduction in glomerular number or apoptosis should also be taken into account in high-grade vesico-ureteral reflux.

Nephrotoxic Drugs

Several drugs have been shown to produce an adverse affect on kidneys when administered during pregnancy or in preterm infants when the exposure occurs during active nephrogenesis. In prematurity, shorter nephrogenesis and a reduced number of glomerular layers have been associated with a peculiar magnitude of damage following exposure to nephrotoxic medication [44]. However, mothers of preterm infants developing acute renal failure had invariably taken larger quantities of drugs during pregnancy and delivery, mainly antibiotics and NSAIDs. Furthermore, preterm infants who developed renal failure had unfailingly received more drugs, mainly antibiotics, NSAIDs and diuretics [64].

Several observations on drug-related nephrotoxicity were derived from rat models. β-Lactam antibiotics (amoxicillin and ampicillin) administered to the rat mother may cross the placenta and reach the fetus. Penicillin given at therapeutic doses produced mild oligonephronia and cystic tubule dilation. Indeed, ceftriaxone at high doses led to impairment of nephrogenesis and blocked kidney development leading to permanent severe renal defects [65]. Aminoglycoside antibiotics given to pregnant females across the placenta and accumulate in the fetal kidney. Exposure to gentamycin at early stages of gestation lowered the final number of nephrons leading to an increased rate of protein urinary excretion [66, 67]. Following administration of gentamycin in rats, renal changes detected by light and electron microscopy were consistent with significant changes in glomerular and tubular structure [68].

Preterm SGA infants administered aminoglycosides showed impaired glomerular and tubular function to a greater extent than those not receiving the drug [69]. Moreover, following the administration of drugs to the mother or postnatal exposure to exposure, some degree of nephrotoxic action was frequently observed in term newborns. The onset of nephrotoxicity is particularly frequent with specific classes of antibacterial drugs (aminoglycosides and vancomycin), although generally being reversible on discontinuation of the drug [70, 71].

The risk of development of acute renal failure in the neonate is high following maternal exposure to NSAIDs. During pregnancy, when nephrogenesis occurs, exposure to NSAIDs may lead to hypoperfusion of the kidneys and long-term renal dysfunction. After long-term in utero exposure, NSAIDs induced nephrotoxicity in the fetus by reducing PGE2 leading to vasomotor nephropathy from renal vasoconstrictions [72]. Indomethacin, in exposed fetuses, led to cystic changes in the developing nephrons [73], and acute/chronic renal failure in newborns or infants born at or before 30 weeks of gestation [74, 75]. Conversely, the risk of renal impairment is lower, although not null, in newborns exposed in utero to NSAIDs once nephrogenesis has been completed [76]. The nephrotoxic actions of NSAIDs at postnatal ages are produced by ibuprofen leading to acute renal failure in term newborns [75]. Moreover, indomethacin exhibited potent suppressive effects on renal COX-2 and vasodilator prostanoids, important regulators of renal development and function, when administered during early postnatal life [77]. Furthermore, similar to effects described with indomethacin, nimesulide has been associated with maternal oligohydramnios, acute or chronic renal failure in the newborns of treated mothers following in utero exposure lasting from a few days to months. Indeed, some of the above patients manifested transient acute renal failure, prolonged acute renal failure and required peritoneal dialysis [78–81].
Adverse effects related to ACE inhibitors have been well documented and include hypotension, oliguria, acute renal failure and hyperkalemia. The major adverse effect is a reduction in glomerular filtration. Particularly bilateral ACE inhibitors should not be administered to patients during pregnancy because human fetopathies have been seen when these drugs were given after the first trimester of pregnancy, the period of fetal kidney development; in particular oligohydramnios, renal tubular dysgenesis and neonatal anuria were observed [82].

Other authors reported adverse fetal and neonatal renal effects after intrauterine exposure to lisinopril, enalapril or other ACE inhibitors. In addition, the ACE inhibitor fetopathy syndrome has been associated with a reduction in amniotic fluid volume as a consequence of the reduced production of fetal urine [83–91].

A study reported by Cooper et al. [92] has shown that almost 9% of children whose mothers were prescribed ACE inhibitors during the first trimester (but not later) had major congenital anomalies (cardiovascular, central nervous system and renal malformations), a rate 2.7 times that among unexposed infants [92].

Many papers have reported fetal anomalies that are very similar to those produced by maternal treatment with ACE inhibitors, in association with angiotensin II receptor antagonists. Thus, maternal treatment with angiotensin II receptor antagonists should be avoided [93, 94].

**Detection of Reduced Nephron Reserve**

Early detection of potential indicators of hyperfiltration, such as impaired renal reserve and blunted solute clearance, may provide subtle clues to the presence of reduced nephron number, increased obsolete glomeruli, thus providing early objective evidence of hypertension, microalbuminurias and renal risks [9]. Since serum Cys-C levels are frequently superior to serum creatinine levels as an index of renal function [95] with some limitations, it might be introduced as a marker of renal function and follow-up in these patients. Moreover, low renal volume detected by ultrasound measurement of renal length [96, 97] and 3-dimensional ultrasound volume [98, 99] might represent suitable parameters of reduced renal reserve in newborns and children. Scintigraphic measurement may provide a comparison of the renal reserve between kidneys and their relative variation by age in congenital unilateral reduced renal reserve [63]. Renal biopsy should only be performed in selected cases and it is not ethically acceptable for wide application.

**Treatment**

Several intrauterine risk factors are associated with low nephron number. The early identification of children at higher risk of reduced renal reserve allows consequent monitoring and treatment [22]. However, when possible, the administration of potentially nephrotoxic agents should be carefully evaluated when performed prior to completion of nephrogenesis in the presence of risk factors of reduced renal reserve.

Children with low renal reserve from congenital or acquired renal damage, showing elevated blood pressure or albuminuria, may benefit from prolonged treatment with renal protective agents [23, 100]. Renoprotective advantages afforded by ACE inhibitors have been demonstrated in animal models, in which angiotensin II played a role in unilateral partial ureteral obstruction regulating glomerular cell apoptosis. Moreover, subsequent to completion of nephrogenesis, angiotensin receptor blockades (ARBs) prevented interstitial and glomerular cell apoptosis in newborn pigs with unilateral ureteral obstruction [101]. Finally, ACE inhibitors decreased interstitial renal fibrosis and preserved renal tubules in newborn dogs with partial urethral obstruction [102].

Having demonstrated effective antiproteinuric and renoprotective actions, today both ACE inhibitors and ARBs are widely used in children to treat numerous chronic renal alterations. Despite different mechanisms of action of these two drugs, they seem to have synergic and adjunctive properties. Accordingly, more structured studies should be implemented to investigate the role of ACE inhibitors and ARBs in managing proteinurias and glomerulosclerosis in children with renal conditions characterized by reduced nephron number and glomerular hypertrophic changes [79].

However, adverse effects related to ACE inhibitors have been well documented and must be considered in the costs/benefits balance. They include hypotension, oliguria, acute renal failure, and hyperkalemia. The major adverse effect is a reduction in glomerular filtration, particularly in patients with bilateral renal artery stenosis [103].

**Conclusion**

Both intrauterine and perinatal events may affect proper long-term renal function, particularly in children at risk of reduced nephron number, such as those born preterm or with LBW. The involvement of stressors dur-
ing fetal life and active nephrogenesis may lead to reduced nephron generation. However, overall nephron availability constituting the renal potential of an individual is linked to specific genetic background, velocity of nephron loss following a specific insult, age at insult, and duration of the same. Furthermore, the latter lead to reduced renal function altering glomerular structure, eliciting fibrosis and cell apoptosis, and decreasing renal reserve. By responding to the higher requirement of electrolytes and fluid balance per glomerulus, the remaining nephrons will be subjected to hypertrophy and hyperfiltration. Similar conditions are manifested by nephrons in conditions of stress, with modification of local vasoactive conditions resulting in hypertension and proteinuria and, in the long term, cardiovascular diseases and irreversible renal damage. Unfortunately, to date no investigation method for the early detection of reduced nephron reserve is available. Future advances in radiological techniques and biochemical indicators of underlying structural changes in the kidneys may provide important findings. However, in view of the current lack of more specific methods, close monitoring of children and young adults at risk of reduced renal reserve should be carried out to enhance the early detection of potential changes in renal function, paying particular attention to subtle modifications in renal size and volume, and challenges to renal function.

References


