Introduction

In children and adults chronic kidney disease (CKD) is associated with an increased rate of cardiovascular complications [1,2,3]. Traditional risk factors like hypertension, dyslipidemia, obesity and alterations in glucose metabolism as well as uremia-related factors including volume overload, anemia, abnormal calcium and phosphate metabolism, malnutrition, and inflammation contribute to the development of the various forms of the cardiorenal syndrome (CRS) [1,4]. Accelerated atherosclerosis and left-ventricular hypertrophy with diastolic and systolic dysfunction are present already in children with CKD [4,5].

In adult and pediatric cardiology n-terminal pro-B-type natriuretic peptide (nt-proBNP), the inactive cleavage product of brain natriuretic peptide (BNP) is an established marker of cardiovascular dysfunction [6,7]. Increased levels of circulating nt-proBNP are the result of re-expression of the cardiac embryonic gene program and induce cardiac remodeling and fibrosis [8,9]. During human development the cardiac natriuretic peptide system, comprising atrial natriuretic peptide (ANP) and BNP is functional by mid-gestation, and nt-proBNP concentrations have been found to be elevated in conditions associated with increased volume or pressure load [10–16].

The aim of our study was to analyze the association between renal and cardiovascular parameters in fetuses with severely reduced or absent renal function secondary to isolated urinary tract malformations. For that, nt-proBNP, cystatin C, β2-microglobulin, and hemoglobin were determined from fetal blood samples. Additionally, echocardiography and color Doppler examination of the fetal cardiovascular system was performed, and myocardial wall dimensions and cardiothoracic area were measured.

Cardiorenal Syndrome is Present in Human Fetuses with Severe, Isolated Urinary Tract Malformations

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Abstract

Objective: We analyzed the association between renal and cardiovascular parameters in fetuses with isolated severe urinary tract malformations.

Methods: 39 fetuses at a mean gestational age of 23.6 weeks with nephropathies or urinary tract malformations and markedly impaired or absent renal function were prospectively examined. Fetal echocardiography was performed, and thicknesses of the interventricular septum, and left and right ventricular wall were measured. Blood flow velocity waveforms of the umbilical artery, middle cerebral artery, and ductus venosus were obtained by color Doppler ultrasound. Concentrations of circulating n-terminal pro-B-type natriuretic peptide (nt-proBNP), cystatin C, β2-microglobulin, and hemoglobin were determined from fetal blood samples.

Results: Malformations included 21 cases of obstructive uropathy, 10 fetuses with bilateral nephropathy, and 8 cases of bilateral renal agenesis. Marked biventricular myocardial hypertrophy was present in all cases. The ratio between measured and gestational age-adjusted normal values was 2.01 (interventricular septum), 1.85, and 1.78 (right and left ventricular wall, respectively). Compared to controls, levels of circulating nt-proBNP were significantly increased (median (IQR) 5035 ng/L (5936 ng/L) vs. 1874 ng/L (1092 ng/L); p < 0.001). Cystatin C and β2-microglobulin concentrations were elevated as follows (mean ± SD) 1.85 ± 0.391 mg/L and 8.44 ± 2.423 mg/L, respectively (normal range 1.66 ± 0.202 mg/L and 4.25 ± 0.734 mg/L, respectively). No correlation was detected between cardiovascular parameters and urinary tract morphology and function. Despite increased levels of nt-proBNP cardiovascular function was preserved, with normal fetal Doppler indices in 90.2% of cases.

Conclusion: Urinary tract malformations resulting in severe renal impairment are associated with biventricular myocardial hypertrophy and elevated concentrations of circulating nt-proBNP during fetal life. Cardiovascular findings do not correlate with kidney function or morphology.

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Our hypothesis was that renal and cardiac function is interrelated during gestation, and that changes in cardiovascular function are present in fetuses with impaired or absent renal function.

Materials and Methods

Patients

Patients referred to our center between January 2007 and October 2011 with a diagnosis of bilateral nephropathy or urinary tract malformation who underwent fetocide or fetal blood sampling (FBS) were eligible. Fetocide was performed as part of termination of advanced pregnancy in accordance with national legislation. Indications for FBS included rapid karyotyping due to late referral or kidney function analysis. We excluded fetuses with any of the following conditions: unilateral nephropathy, unilateral urinary tract malformation, associated structural malformations, aneuploidy, moderate or severe anemia [17], monochorionic twin pregnancies, and cases with intrauterine growth restriction (estimated fetal birth weight <10. percentile). Gestational age (GA) was confirmed by first-trimester ultrasound. For nT-proBNP analysis, previously established reference values served as controls. Details of the control group have already been published [16].

Ethics Statement

The study was approved by the university of Bonn ethics committee. All participating pregnant women gave their written consent.

Ultrasonography and Echocardiography

High-resolution ultrasound equipment (Philips IU22, Philips Hamburg, Germany; GE logiq 9, Voluson E8, Voluson730 expert, GE, Munich, Germany) was used in all cases with 5 to 9 and 2 to 6, 7, and 8 MHz respectively, convex transducers. The spatial peak temporal average power output was kept at <100 W/cm², applying only the fetal use adapted ultrasound machine settings. A detailed ultrasound of the fetal anatomy, echocardiography with measurement of myocardial wall dimensions, and color Doppler examination of the fetal and uterine vessels was performed. Thicknesses of the interventricular septum (IVS) and the right and left ventricular wall (RVW-GA and LVW-GA) were calculated for the interventricular septum (IVS-GA) and the right and left ventricular wall (RVW-GA and LVW-GA) using the following formulas: RVT-W GA = (0.00018 * GA2); RVW-GA = -0.1473+0.02045 * GA; LVW-GA = -0.136+0.01967 * GA; (0.00019 * GA2); (0.00019 * GA2); (0.00019 * GA2), respectively. Ratios were calculated by dividing measured anion values by normal values.

Statistical Analysis

To adjust for the effect of gestational age hemoglobin values were transformed into multiples of the median (MoM). Expected hemoglobin values were calculated with the following formula: e((2.84-5.56/GA) [17]. MoMs were then calculated by dividing the measured values by the expected values. Anemia was classified as mild (0.84–0.85 MoM), moderate (0.64–0.55 MoM), and severe (<0.55 MoM) [17].

Gestational age-adjusted normal values and standard deviations were calculated for the interventricular septum (IVS-GA) and the right and left ventricular wall (RVW-GA and LVW-GA) using the following formulas: IVS-GA = -0.1321+(0.01927 * GA)-0.0018 * GA2; RVW-GA = -0.1473+0.02045 * GA; LVW-GA = -0.136+0.01967 * GA; (0.00019 * GA2). Ratios were calculated by dividing measured anion values by normal values.

For normally distributed values, between group comparisons of continuous variables were performed by independent samples t-test or one-way ANOVA with post-hoc analysis by Tamhane-T2 Test. For nT-proBNP and IVS, Mann-Whitney-U or Kruskal-Wallis test was used. Categorical variables were compared by x² test. For correlation analysis Pearson’s coefficient was calculated. Unless indicated otherwise, results are reported as mean ± SD.

Results

41 analyses were performed in 39 fetuses. In two fetuses with obstructive uropathy measurements were taken twice (before and after shunting) with two weeks interval. Cystatin C and hemoglobin values could not be determined in three cases, β2-microglobulin in two. Mean gestational age was 23.6 (±3.57 weeks of gestation), 30 fetuses were male (76.9%). A normal karyotype was present in all examined cases (n = 36, 92.3%).

Malformations were classified in three categories:

(I) Obstructive uropathy, n = 21 (53.9%); lower (urethral) urinary tract obstruction with or without development of urinoma or urinouretic ascites.

(II) Bilateral nephropathy, n = 10 (25.6%); autosomal recessive polycystic kidney disease; bilateral multicystic dysplastic kidneys; unilateral multicystic dysplastic kidney with contralateral renal agenesis or dysplasia.

(III) Bilateral renal agenesis, n = 8 (20.5%).
Severe oligohydramnios (amniotic fluid index \(\leq 2\) cm) or anhydramnios was present in all cases. Pulmonary hypoplasia, defined as cardiothoracic area ratio (CTAR) >0.3 could be demonstrated in 36 fetuses (87.8%); CTAR was normal in 3 (7.3%), and measurement was not available for 2 cases (4.9%).

Myocardial wall dimensions were markedly altered, see Figure 1 and Table 1. No correlation was detected between myocardial wall thicknesses and any of the serological parameters under investigation, and no difference was present between the malformation groups. Qualitative echocardiography revealed no signs of cardiac failure in any of the fetuses. Pulsed wave Doppler sonography of fetal vessels was normal in 90.2% of cases (n = 37). Three fetuses had abnormal DV pulsatility indices (including one case with abnormal PSV-MCA), in one case PSV-MCA only was abnormal. 29 pregnancies (74.4%) were terminated, five (12.8%) ended in live births, and five were lost to follow-up.

Compared to controls, nt-proBNP concentrations were significantly increased in the study group (median [IQR] 5035 ng/L [5936 ng/L] vs. 1874 ng/L [1092 ng/L], p<0.001), see Figure 2; no difference was present between the three malformation categories. Cystatin C, \(B_2\)-microglobulin and GA-adapted hemoglobin concentrations are listed in Table 2. Compared to fetal reference values [19,20] cystatin C and \(B_2\)-microglobulin were found to be increased. Additionally, cystatin C and \(B_2\)-microglobulin concentrations were significantly higher in bilateral renal agenesis compared to obstructive uropathies (p = 0.005 and p<0.001, respectively).

No correlation was present between nt-proBNP and markers of kidney function or hemoglobin, whereas cystatin C and \(B_2\)-microglobulin concentrations were closely correlated (\(r = 0.813, p<0.001\)).

A significant inverse correlation between MoM Hb and GA (\(r = -0.357, p = 0.028\)) was detected, see Figure 3. Mild anemia was present in 5 cases (13.2%); anemic fetuses did not show any difference in the variables under investigation.

In cases with obstructive uropathy, 14 samples were taken before and nine after vesico-amniotic shunting. There was no difference in any of the variables except for hemoglobin, which was significantly higher after shunting (MoM Hb before versus after shunting: p = 0.010).

**Discussion**

In our group of fetuses with severely impaired or absent renal function secondary to isolated nephropathies and urinary tract malformations biventricular myocardial hypertrophy with preserved myocardial function was present. Nt-proBNP concentrations were significantly elevated in the study group compared to controls; no difference was present between the various malformation categories.

**Nt-proBNP during Fetal Life**

During human development, natriuretic peptides (NPs) are involved in cardiac morphogenesis. By mid-gestation the cardiac natriuretic system (ANP and BNP) is assumed to be functional. Fetal and maternal compartments are separated and placental transfer of NPs does not occur [10,21]. In contrast to cardiac morphogenesis NPs are not contributing to renal development. Knock-out mice for natriuretic peptide receptor-A, the principal receptor for atrial natriuretic peptide (ANP) or ANP do not display renal anomalies [22]. The role of nt-proBNP as a marker of increased myocardial work load during fetal life has been investigated in intrauterine growth restriction, structural cardiac malformations, and anemia. Both, increased pressure load secondary to high resistance in the placental bed or outflow tract obstructions and increased volume load secondary to anemia are able to trigger nt-proBNP release [11–16]. The longer half-life and...
higher stability make nt-proBNP a more suitable candidate for analysis compared to BNP [23].

Myocardial Hypertrophy and Urinary Tract Anomalies during Fetal Life

Limited data are available on the association of myocardial hypertrophy and structural or functional abnormalities of the urinary tract during fetal life. In a series of 50 cases with cardiomyopathy 10% were associated with renal abnormalities [24]. Pedra et al. 2002 [25] in their analysis however did not find nephropathies or urinary tract malformations in association with fetal cardiomyopathy (55 cases). Although levels of circulating nt-proBNP were elevated in our study group no correlation was present between blood levels and myocardial wall thicknesses. Our results therefore do not support involvement of BNP in the development of ventricular wall hypertrophy. Cardiomyocyte hypertrophy was detected in transgenic rats expressing excessive amounts of prorenin [26], and transcription profiling of prorenin-stimulated cardiomyocytes revealed expression of genes involved in actin-filament modification [27]. Further studies are needed to clarify this issue.

Nt-proBNP and Cardiorenal Syndrome (CRS) in Postnatal Life

The role of nt-proBNP as marker of cardiovascular dysfunction in CKD has been a matter of debate [28]. Meanwhile, it has been proven that cardiac stimulators contribute independently to BNP release in CKD; despite impaired renal clearance nt-proBNP concentrations in CKD are predominantly related to cardiac pathology [29]. Furthermore, correlation between nt-proBNP and

Table 1. Cardiac dimensions in fetuses with urinary tract malformations (n = 37) and gestational age-adjusted normal values.

| Thickness (mm), mean ± SD | Adjusted normal value (mm), mean ± SD | Ratio*, mean ± SD |
|--------------------------|--------------------------------------|------------------|
| Interventricular septum  | 4.30±1.163                           | 2.14±0.643       | 2.01±0.372       |
| Right ventricular wall   | 3.97±1.055                           | 2.15±0.597       | 1.85±0.333       |
| Left ventricular wall    | 3.78±0.943                           | 2.14±0.570       | 1.78±0.342       |

*measured value/gestational age-adjusted normal value.
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Figure 2. Nt-proBNP concentration in fetuses with urinary tract malformations and controls [16]. Depicted values are log-transformed. *,** denotes outliers. Mann-Whitney-U test: p<0.001.
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myocardial dysfunction has been demonstrated in children with CKD [30]. Nt-proBNP, therefore, serves as a marker of cardiorenal burden in pediatric and adult medicine.

In contrast, the increase in circulating nt-proBNP observed in our study group was not associated with major changes in the fetal cardiovascular function. Therefore, our results do not support a role of nt-proBNP as marker of cardiovascular impairment in nephropathies or urinary tract malformations during fetal life.

Table 2. Umbilical vein concentrations of cardiac and renal function tests in fetuses with isolated urinary tract malformations.

|                     | Fetal reference values | Study group | Obstructive uropathya | Bilateral nephropathya | Bilateral renal agenesis |
|---------------------|------------------------|-------------|-----------------------|------------------------|--------------------------|
| nt-proBNP (ng/L), median (IQR) (n = 41) | 1874 (1092)b | 5035 (5936) | 4854 (5994) | 5686 (8279) | 4872 (4084) p = 0.001c |
| Cystatin C (mg/L), mean ± SD (n = 38) | 1.66±0.202 | 1.84±0.391 | 1.67±0.350 | 1.93±0.354 | 2.18±0.299 p = 0.005d |
| β2-Microglobulin (mg/L), mean ± SD (n = 39) | 4.25±0.734 | 8.44±2.423 | 7.55±2.127 | 8.45±2.640 | 10.79±1.133 p<0.001e |
| MoM Hb* (n = 38) | 0.84–1.16 | 0.95±0.101 | 0.96±0.115 | 0.95±0.084 | 0.93±0.079 |

*a see text for details; b78 cases; c study group vs. reference values (Mann-Whitney-U test); d bilateral renal agenesis vs. obstructive uropathy (Tamhane post-hoc test); e multiples of the median: 5.–95. percentile.

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Figure 3. Hemoglobin concentration in fetuses with urinary tract malformations. Multiples of the median hemoglobin (MoM Hb) during the course of pregnancy in fetuses with isolated malformations of the urinary tract (n = 38). Horizontal lines: Upper and lower limit, reference range [17]. R² = 0.127. Equation for the line: y = 1.183–0.010 x GA (gestational age).
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Nt-proBNP and Urinary Tract Anomalies during Fetal Life

Various factors may contribute to the increase in n-t-proBNP concentration in fetuses with nephropathies and urinary tract malformations.

Firstly, alterations in the vascular bed of the urinary tract may cause increased afterload.

Renal blood flow in the fetus amounts to 3% of the combined cardiac output and is characterized by high vascular resistance. Vascular resistance within the often grossly enlarged kidneys of severe nephropathy has not been measured. In fetuses with lower urinary tract obstruction increased downstream impedance in the common iliac arteries has been demonstrated and was interpreted to be caused by compression of the distended bladder [31]. Additionally, interstitial fibrosis and hydronephrosis accompanying obstructive uropathy may be associated with increased renovascular resistance.

Secondly, impaired or absent kidney function may result in a net increase in total body water in fetuses with reduced afterload. In fetal lambs, intra- and extracellular volume is reduced after second-trimester nephrectomy [32], the same, however, may not apply for nephropathies and urinary tract malformations during fetal life. Postnatally, end-stage renal disease is associated with volume retention which correlates with increasing BNP-levels [33]. During intrauterine life the placenta acts as clearing organ for fetal waste products and is involved in the fluid balance of fetal compartments and amniotic fluid. The distribution of total body water (TBW) in the fetus is subject to dynamic changes; parallel to the reduction in extracellular and increase in intracellular water the TBW to weight ratio decreases. Furthermore, transcapillary fluid filtration into the interstitial space is facilitated during fetal life. Despite placental function these factors may result in a net increase in total body water in fetuses with restricted or absent renal function.

Thirdly, even minor degrees of anemia are able to induce a rise in n-t-proBNP [34]. Although moderate or severe anemia was not present in our case series, we observed a significant reduction in hemoglobin with advancing gestational age; additionally, hemoglobin levels were higher after vesico-angiomatic shunting.

Limitations

The number of cases is small, thereby limiting the significance of our findings. Nevertheless, the study represents the largest such cohort in the literature to date. Bokemkamp et al. 2001 [20] and Berry et al. 1995 [35] reported on 20 and 15 fetuses, respectively. Confounding may have been introduced by selection bias. However, all major urinary tract malformation classes were included, and minor or unilateral anomalies and cases with associated malformations excluded. We are therefore confident that our findings are valid. We did not perform quantitative echocardiography and may have missed early signs of myocardial dysfunction. Since qualitative echocardiography and Doppler waveform analyses of fetal vessels including umbilical veins were within normal range, no case with major myocardial dysfunction was overlooked. For cardiac dimension calculations we would have preferred to utilize Z-scores [36]. However, nomograms for the parameters under investigation have not been established. Since the measurements in our study group are markedly abnormal we assume our finding to be valid.

Conclusion

Nephropathies and urinary tract malformations are associated with biventricular myocardial hypertrophy and elevated levels of circulating n-t-proBNP during fetal life. The cardiovascular findings do not correlate with kidney function or morphology, but may be caused by changes in prorenin concentrations and increased afterload.

Supporting Information

Dataset S1. (SAV)

Author Contributions

Conceived and designed the experiments: WMM UG. Analyzed the data: WMM RF KK. Contributed reagents/materials/analysis tools: BS-W. Wrote the paper: WMM UG.

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