

## **Title**

**Association of linear growth velocities between 0 and 6 years with kidney function and size at 10 years: a birth cohort study in Ethiopia.**

## **Name of authors**

Beakal Zinab<sup>1,3</sup>, Rahma Ali<sup>2,3</sup>, Bikila S Megersa<sup>3</sup>, Tefera Belachew<sup>1</sup>, Elias Kedir<sup>4</sup>, Tsinuel Girma<sup>5</sup>, Alemseged Abdisa<sup>6</sup>, Melkamu Berhane<sup>5</sup>, Bitiya Admasu<sup>2</sup>, Henrik Friis<sup>3</sup>, Mubarek Abera<sup>7</sup>, Mette F Olsen<sup>3,8</sup>, Gregers S Andersen<sup>9</sup>, Jonathan CK Wells<sup>10</sup>, Suzanne Filteau<sup>11</sup>, Rasmus Wibæk<sup>9</sup>, Dorothea Nitsch<sup>11</sup> \* and Daniel Yilma<sup>12</sup> \*.

<sup>1</sup> Department of Nutrition and Dietetics; and <sup>2</sup>Department of Population and Family Health, Faculty of Public Health, Jimma University, Jimma, Ethiopia.

<sup>3</sup> Department of Nutrition, Exercise, and Sports, University of Copenhagen, Copenhagen, Denmark.

<sup>4</sup> Department of Radiology; and <sup>5</sup>Department of Pediatrics and Child Health Faculty of Medical Sciences, Jimma University, Jimma, Ethiopia.

<sup>6</sup> Armauer Hansen Research Institute, Addis Ababa, Ethiopia.

<sup>7</sup> Department of Psychiatry, Faculty of Medical Sciences, Jimma University, Jimma, Ethiopia.

<sup>8</sup> Department of Infectious Diseases, Rigshospitalet, Copenhagen, Denmark

<sup>9</sup> Steno Diabetes Center Copenhagen, Herlev, Denmark.

<sup>10</sup> Childhood Nutrition Research Center, UCL Great Ormond Street Institute of Child Health, London, United Kingdom.

<sup>11</sup> Faculty of Epidemiology and Population Health, London School of Hygiene & Tropical Medicine, London, United Kingdom.

<sup>12</sup> Department of Internal Medicine, Faculty of Medical Sciences, Jimma University, Jimma, Ethiopia.

\* These authors contributed equally to this work.

Corresponding author: Beakal Zinab ([beakalzinab1@gmail.com](mailto:beakalzinab1@gmail.com))

### **Conflict of Interest**

All authors declare no conflicting interests.

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### **Data sharing**

Data described in the manuscript, code book, and analytic code will be made available upon request pending application and approval.

### **Abbreviations**

iABC, infant Anthropometric and Body Composition; ADP, air displacement plethysmograph; DOHaD, Developmental Origins of Health and Disease; Glomerular Filtration Rate; LMICs, Low and Middle Income Countries; LBW, Low Birth Weight.

1 **Abstract**

2 **Background:** The risk of non-communicable diseases accrues from fetal life, with early childhood  
3 growth having an important role for the risk of adult disease. There is a need to understand how early  
4 life growth relates to kidney function and size.

5 **Objectives:** This study aimed to assess the association of linear growth velocities among children  
6 between 0 and 6 years with kidney function and size among children aged 10 years.

7 **Methods:** The Ethiopian Anthropometric and Body Composition (iABC) birth cohort recruited  
8 infants born at term to mothers living in Jimma, with a birth weight of  $\geq 1500$  grams, and without  
9 congenital malformations. Participants were followed up with 13 measurements between birth and 6  
10 years of age. The latest follow-up was at ages 7-12 years with measurement of serum cystatin C as a  
11 marker of kidney function, and ultrasound assessment of kidney dimensions. Kidney volume was  
12 computed using an ellipsoid formula. Linear spline multi-level modelling was used to compute linear  
13 growth velocities between 0-6 years. Multiple linear regression modelling was used to examine the  
14 associations of linear growth velocities in selected age periods with cystatin C and kidney size.

15 **Results:** Data were captured from 355 children, at a mean age of 10 (range 7-12) years. The linear  
16 growth velocity was high between 0-3 months and then decreased with age. There was no evidence  
17 of an association of growth velocity up to 24 months with cystatin C at 10 years. Between 24-48 and  
18 48-76 months, serum cystatin C was higher by 2.3% (95% CI 0.6, 4.2) and 2.1 % (95% CI 0.3, 4.0)  
19 for one SD higher linear growth velocity, respectively. We found a positive association between linear  
20 growth velocities at all intervals between 0-6 years and kidney volume.

21 **Conclusion:** Greater growth between 0-6 years of development was positively associated with kidney  
22 size and greater growth velocity after 2 years was associated with higher serum cystatin C level.

23 **Key words:** linear growth velocities, Kidney function, Kidney size, cohort study, Ethiopia.

## 24 **Introduction**

25 The developmental origins of health and disease (DOHaD) hypothesis states that adverse  
26 environmental factors acting early in life increase the risk of later-life disease vulnerability (1). For  
27 example, stressors in early life may result in structural and functional changes in the developing  
28 kidney, increasing individuals' vulnerability to kidney, and cardiovascular disease in later life (2,3).

29 In humans, kidney development begins during the ninth week of pregnancy and continues until the  
30 36th week (4). Except for extremely preterm neonates, there is no evidence of nephrogenesis in  
31 humans after birth (4). The normal human kidney has an average of 1 million nephrons, which  
32 consisting of a glomerulus (filter unit) and a tubule (controlling urinary composition). Multiple  
33 studies have shown that the total glomerular number may vary by thirteen -fold between individuals  
34 (5–7). The number of nephrons during adulthood reflects the difference between the number of  
35 nephrons at birth and the number of nephrons lost (8–10).

36 Trajectories of growth reflect the complex interplay of biological and environmental processes that  
37 influence life course health and development (11,12). Linear growth retardation in early life is a good  
38 indicator of a poor early-life environment and is associated with increased risk of morbidity later in  
39 life (13). Previous studies have shown that children with short stature exhibit reduced kidney size and  
40 a lower nephron number (14,15).

41 Linear growth failure manifested as stunting is a major public health problem in developing countries  
42 in general and in sub-Saharan Africa in particular (16)(17). Therefore, in this setting, there may be

43 subclinical differences in kidney function and volume that may already be detectable in childhood and  
44 may explain the susceptibility of individuals to kidney disease in adulthood (18). Understanding the  
45 natural history of kidney function, including subclinical differences and modifiable risk factors, is  
46 pivotal to designing and implementing efficient preventive strategies at the population level. Studies  
47 have described a high burden of acute and chronic pediatric kidney disease in low- and middle-  
48 income countries (19–21), but many of the existing studies on childhood predictors of adult kidney  
49 function were conducted in high-income countries (22–24). Evidence from high-income settings  
50 suggests that birth characteristics, fetal growth, and early childhood growth influence kidney function  
51 throughout one's life course (25). The current study was intended to fill current research gaps by  
52 investigating the association of early-life linear growth with a marker of kidney function and kidney  
53 size at 10 years using an Ethiopian birth cohort.

54

## 55 **Methods**

### 56 **Study setting and participants**

57 The study included children from the Ethiopian iABC birth cohort, which has been described earlier  
58 (26,27). Briefly, infants of mothers who lived in Jimma Town, born at term (gestational age at birth  
59  $\geq 37$  completed weeks) with a birth weight of  $\geq 1500$  gram and without congenital malformations were  
60 included in the cohort. The mother-child pairs were invited to attend a total of 13 study visits at birth,  
61 at 1.5, 2.5, 3.5, 4.5, 6 months, and 1, 1.5, 2, 3, 4, 5, and 6 years. A total of 644 mother-newborn dyads  
62 were recruited for the study between December 2008 and October 2012.

63 The current follow-up visits, hereafter referred to as 10-year follow-up, were conducted from June  
64 2019 to December 2020 and included 355 children aged 7-12 years. Families of all children were  
65 traced by previously provided phone number or residential location and invited to bring their children  
66 after receiving clear information about the study. In cases where phone numbers were not working, a  
67 study nurse visited the family's last known address.

### 68 **Data collection tools and procedures**

69 Experienced research nurses and laboratory technicians collected the data. For the 10-year follow-up,  
70 families were requested to bring their children fasting in early morning; samples were taken in the  
71 morning because of postprandial changes in plasma (28). Maternal and childhood characteristics  
72 were collected using questionnaires. Body dimensions, body composition, and renal size were  
73 measured using anthropometry, air displacement plethysmography, and ultrasound, respectively.  
74 Serum cystatin C was analyzed from blood samples as described below.

### 75 **Questionnaire data**

76 A pre-tested interviewer-administered structured questionnaire was used to collect information  
77 concerning socio-demographic and economic characteristics of the family. The tool includes

78 questions intended to capture family and child socio-demographic characteristics, status of the house  
79 they live, and ownership of properties. Additional relevant previous maternal and child characteristics  
80 were abstracted from iABC data.

### 81 **Anthropometric and body composition measurements**

82 Weight from birth to six months was measured to the nearest 1 gram using a PEA POD, an infant air  
83 displacement plethysmograph (ADP; COSMED, Rome, Italy); for the follow-up visits starting from  
84 12 months, weight was measured to the nearest 0.1 kg using an electronic UNICEF scale (SECA,  
85 Hamburg, Germany). Length was measured in a recumbent position for infants below 24 months to  
86 the nearest 0.1 cm using a SECA 416 Infantometer. In children 24 months and above, standing height  
87 was measured to the nearest 0.1 cm using a SECA 213 portable stadiometer (SECA, Hamburg,  
88 Germany). More detailed information on specific measurements at different follow-up visits is  
89 published elsewhere (29).

90 For the current visit at 10 years, the participants' weight was measured to the nearest 0.1 g by the  
91 BOD POD, a child/adult version of ADP, after removing heavy clothes. Height was measured to the  
92 nearest 0.1 cm using a stadiometer, according to the standard procedure without shoes. All  
93 anthropometric measurements were taken twice, and the average values were used.

### 94 **Body composition assessment**

95 Body composition was assessed at birth at 1.5, 2.5, 3.5, 4.5, and 6 months of age with the PEA POD,  
96 designed to measure infants between birth and 6 months of age. BOD POD, was used to assess body  
97 composition at the 4, 5, 6, and 10 years follow-ups. Children were fasting and wearing close-fitting  
98 underwear and a swimming cap during measurement. Both the PEA-POD and BOD-POD have high  
99 accuracy and precision and are feasible and safe for assessing body composition in infants and  
100 children, respectively (30,31)

### 101 **Ultrasound measurement**

102 Kidney size was measured by ultrasonography using a C1-5-D 2D convex probe (GE P6) (General  
103 electronics Co.Ltd Boston, USA). The kidney was identified in the sagittal plane along its longitudinal  
104 axis. Measures of maximal bipolar kidney length, width, and thickness were obtained for both  
105 kidneys. Renal width and thickness were measured at the level of the kidney hilum. All dimensions  
106 were measured to nearest 0.1 cm. All children were examined by the same certified radiologist.  
107 Kidney volume was calculated in cubic centimeters using the formula of an ellipsoid: length × width  
108 × depth × 0.523 (32,33). Total kidney volume was calculated as the sum of the right and left kidney  
109 volumes.

#### 110 **Blood sample collection and analysis**

111 Families were asked ahead of the visit to bring their children fasted overnight for 8 hours. Lab  
112 technicians collected blood samples (4 ml) after confirming that the child had fasted. The study nurses  
113 provided the children with a meal immediately after sample collection. In cases where children came  
114 non-fasting or where the ultrasound machine was not working, participants were given new  
115 appointments for outcome measurements.

116 Samples were stored in the lab fridge for a maximum of 4 hours in K2-EDTA tubes. Blood samples  
117 underwent centrifugation for 10 minutes and were stored at -80°C until further lab analysis. Serum  
118 cystatin C was determined using an enhanced immune turbidimetric assay on a Cobas c 702 analyser  
119 (Roche Diagnostic, Germany). Cystatin C is a low molecular weight protein, produced at a relatively  
120 constant rate. The concentration of serum cystatin C is highly correlated with directly measured  
121 Glomerular Filtration Rate (GFR) values, and small reductions in GFR can be detected more readily  
122 with serum cystatin C (34,35). The estimated glomerular filtration rate was calculated using  
123 Zappitelli's formula  $eGFR_{Cyst} = 75.94 / [CysC^{1.17}]$  (36). Because this formula is not validated for  
124 the target population, the results are unlikely to reflect true eGFR in these children; further analyses  
125 were carried out only using cystatin C as the outcome.



126 **Statistical analyses**

127 Data were double-entered in Epi Data version 4.4.2.0 (Denmark). Descriptive data were presented as  
128 mean (standard deviation [SD]) for normally distributed data, median (interquartile range (IQR)) for  
129 continuous non-normally distributed data, and. count (proportion) was used for categorical variables.  
130 Since serum cystatin C was not normally distributed, it was log-transformed, before regression  
131 analyses. Estimates from these models were back-transformed and presented as a percentage change.  
132 The normality of the residuals was checked visually by histogram, pnorm, and qnorm plots. Residuals  
133 were plotted against the fitted values to check the homogeneity of variance of the residuals.

134 **Linear growth velocity 0 to 6 years**

135 The non-linear relationship of length/height as a function of age were modeled using a series of linear  
136 splines (37). Linear-spline multilevel (piecewise linear multilevel) models are increasingly used to  
137 model childhood growth since they address many of the challenges associated with analyzing  
138 longitudinal data (38,39) Knot points were placed at 3, 6, 24 and 48 months while taking into  
139 consideration data density, previous knowledge and model fit statistics. Linear growth velocity  
140 between 0 to 3 months is the difference between predicted length at 3 months and length at birth  
141 divided by 3 to get cm/month, and similarly for the other growth periods. These individual specific  
142 monthly linear length velocities over discrete time intervals from 0 to 6 years of age were generated  
143 using R version 4.2.0 (R Foundation for Statistical Computing).

144 **Association of linear growth velocity with kidney function and size**

145 Linear regression models were used to test associations of cystatin C and kidney size with estimates  
146 of each child's birth length, and length growth velocity from 0-3, 3-6, 6-24, 24-48, and 48-76 months.  
147 To obtain comparable estimates across the different growth periods, sex-based standardization of  
148 growth velocities was done by subtracting the mean from the individual's score and dividing by the  
149 standard deviation. These sex-based standardized growth velocities were used for subsequent

150 multiple linear regression analyses as exposure variables. Thus, the estimates indicate the change in  
151 cystatin C or kidney size per study population SD increase in length/height velocity.

152 Three models were fitted separately for birth length and each of the length/height velocity exposures.  
153 Model 1 was adjusted for sex and current age. Model 2 additionally adjusted for birth weight,  
154 gestational age, birth order and current fat mass. The adjustment for current fat mass was done to  
155 remove any effect of fat mass on cystatin C measurements (40). Model 3 was additionally adjusted  
156 for maternal education and height at birth. Stata version 14 (StataCorp LLC College Station, Texas,  
157 USA) was used to fit the multiple linear regression models.

### 158 **Sensitivity analyses**

159 We investigated whether there was a difference identified in serum cystatin C level between low birth  
160 weight (LBW) and normal birth weight children who attended the 10<sup>th</sup> follow-up. Cross-sectional  
161 analyses of associations of height at the latest follow-up with kidney parameters were carried to sense-  
162 check the results. Instead of total kidney volume, a separate regression model computed for kidney  
163 dimensions of each kidney. To investigate whether associations of growth with kidney size were  
164 driven by body surface area (BSA), sensitivity analyses investigated associations between linear  
165 growth velocities from 0 to 6 years with kidney volume divided by BSA (derived using the Boyd  
166 formula,  $BSA [m^2] = Weight [kg]^{0.4838} \times Height [cm]^{0.3} \times 0.017827$ )).

### 167 **Ethical considerations**

168 Ethical permission was obtained from Jimma University Ethical Review Board (Letter No.  
169 IHRPHD/333/18), and London School of Hygiene and Tropical Medicine ethics committee.  
170 Parents/guardian signed consent forms before entry into iABC and the current 10 year follow-up.  
171 Any abnormal findings detected during clinical and laboratory evaluations were communicated to  
172 families of children and they were linked to Jimma University Medical Center for further evaluation.

173

## 174 **Results**

### 175 **Characteristics of study participants and mothers**

176 A total of 644 mother-infant pairs attended the birth visit. We excluded 10 preterm and 63 children  
177 not living in Jimma. The mean maternal age ( $\pm$ SD) at the infant's birth was  $22.1 \pm 4.5$  years, and at  
178 the time, 61% of the mothers had attended primary school.

179 Of the remaining 571, 355 (62%) were recruited at age 7-12 for the 10-year follow-up visit for  
180 assessment of kidney function and size. Two participants were excluded because they had only one  
181 height measurement from birth to 6 years, leaving 353 for the 10-year analysis (Figure 1). Of these  
182 353 children, 51.8% were male, the mean ( $\pm$ SD) age was  $9.8 \pm 1.0$  years, and 48.7% were first born  
183 (Table 1).

184 Children who were lost to follow-up were similar with respect to most variables, but had lower birth  
185 weight and birth length, and were less likely to be second- or third-born children compared to those  
186 who attended the current visit (Supplementary Table 1). Reasons for loss to follow-up were migration  
187 out of Jimma, death, and refusal to participate in further follow-up assessments. In addition, we could  
188 not obtain serum cystatin and kidney size measurements for 6 and 3 children respectively, because  
189 participants came from far away which meant that a reappointment was not possible.

### 190 **Linear growth velocity between 0-6 years**

191 The velocity of linear growth was fastest in the first 3 months of life (4.1cm/month), and then  
192 decreased with age. Males had faster linear growth velocity between 0-3 months while it was faster  
193 in females between 6 to 24 months of age (Table 2).

### 194 **Kidney function and volume**

195 The median (IQR) serum cystatin C and estimated glomerular filtration rate were 0.93 (0.83; 1.01)  
196 mg/dl and 82.7 (75.1; 94.4) ml/min per 1.73m<sup>2</sup>, respectively. The median (IQR) combined kidney  
197 volume was 117.2 (103.0; 132.4) cm<sup>3</sup> (Table 3). We found no evidence of a difference in serum  
198 cystatin C level between children with LBW and normal birth weight who attended the 10th follow-  
199 up (Supplementary table 2).

#### 200 **Association of linear growth velocity with kidney function**

201 Associations of estimated standardized linear growth velocity between 0-3, 3-6, 6-24, 24-48 and 48-  
202 76 months with serum cystatin C are presented in Figure 2. In the fully adjusted models, there was  
203 no evidence that linear growth velocities at 0-3, 3-6, and 6-24 months were associated with log-serum  
204 cystatin C. However, between 24-48 and 48-76 months, a one SD higher linear growth velocity was  
205 associated with 2.3 % (95% CI 0.6, 4.2) and 2.1 % (95% CI 0.3, 4.0) higher serum cystatin C,  
206 respectively. Additionally, a positive and significant association was observed between serum  
207 cystatin C and observed height at the 10<sup>th</sup> year follow-up (Supplementary Table 3).

#### 208 **Association of linear growth velocity and kidney size (volume)**

209 Across all models, linear growth velocities between different knots from 0 to 6 years were positively  
210 associated with kidney volume at 10 years. The strongest association was seen for linear growth  
211 velocity from 48-76 months (Figure 3). In sensitivity analyses, linear growth at all intervals from 0-  
212 6 years was positively associated with both right and left kidney length (Supplementary Figures 1 &  
213 2). Conversely, only linear growth velocity after two years of age was associated with kidney anterior-  
214 posterior diameter (depth) (Supplementary Figures 3 & 4). There was a positive association found  
215 between observed height at the 10<sup>th</sup> year follow-up and kidney size (Supplementary Table 3). As  
216 depicted in Supplementary Figure 7, once kidney volume was divided by BSA, there was no evidence  
217 for an association with linear growth velocities.

218 **Discussion**

219 In this study, linear growth velocities at 24-48 and 48-76 months, but not at other age intervals, were  
220 positively associated with serum cystatin C level, indicating that greater growth in these periods is  
221 associated with comparatively lower kidney function when compared to peers. On the other hand, the  
222 observed positive association between faster linear growth velocity and cystatin C might partly  
223 explained by non-renal factors.

224 To the best of our knowledge, this study is the first to report the longitudinal relationship between  
225 early life linear growth velocities and cystatin C and kidney size in an African context. Although we  
226 cannot infer causality, our results suggest that faster linear growth beyond 2 years may be related to  
227 later life kidney function deficits. This finding is consistent with multiple studies of cardiometabolic  
228 markers, including blood pressure, which show a positive association with faster linear growth after  
229 2 years of age (41–44).

230 The underlying mechanisms for the associations of linear growth between 2 to 6 years, and kidney  
231 function are not well understood. One potential explanation of the findings is that faster growth in  
232 children and adolescents imposes a greater functional burden on kidneys, and that demands on renal  
233 capacity made by rapid childhood growth after 24 months of age may not be entirely met by renal  
234 development, resulting in compensatory increase in blood pressure (45). Although being taller as an  
235 adult appears healthier with lower non-communicable disease risk (46), the current study indicated  
236 that, having faster linear growth velocity in childhood after 2 years is not beneficial for kidney  
237 function. That would suggest that the more favorable pathways in terms of kidney function are for a  
238 child to realize its genetic growth potential before the age of 2 years.

239 In this study, faster growth velocities between 0-6 years were positively associated with kidney  
240 volume at 10 years. Our results are similar to other studies that performed radiological measurements

241 of renal size (47,48). Kidney size, though an imperfect proxy for nephron number, is positively  
242 associated with kidney function (49). It is well known that in the context of reduced nephron number,  
243 the remaining nephrons increase in size (50). We speculate that the persisting positive association of  
244 linear growth with kidney size, as observed in this study, may be related to this compensatory  
245 mechanism to meet the child's metabolic requirements to reach the required BSA, but at the expense  
246 of kidney function. This is in line with associations of linear growth velocities and kidney volume  
247 appearing to be mediated by BSA at the latest follow-up.

248 Regression models for each kidney dimension separately (data shown in Supplementary Figures 1-6)  
249 confirm that associations are robust for both kidneys. On the other hand, only linear growth velocity  
250 after two years of age was associated with kidney anterior-posterior diameter (depth). Combined with  
251 the higher serum cystatin C levels with greater linear growth after two years, the results are consistent  
252 with a previous study of 8-year-old children in Nepal, in that "thicker" kidneys appear to be less  
253 favorable for cardio-metabolic health when compared to longer kidneys (51).

#### 254 **Strengths and limitations**

255 A major strength of the study is that it used prospectively measured growth data. To date, most such  
256 research has been conducted in high-income countries and studies that associated linear growth with  
257 the later development of non-communicable diseases generally and kidney diseases, particularly, are  
258 scarce. The observed effect estimates in the present study are not in the range where one would  
259 consider these to explain overt kidney disease in children, i.e. are associations within the norm. These  
260 associations may be important from an etiological and developmental perspective because the  
261 subclinical variation of kidney function in childhood may well impact later life kidney function,  
262 similar to the tracking of pediatric blood pressure measurements with later cardiovascular risk.(52)  
263 (53)(54). Future follow-ups of this cohort will investigate this as the children reach adulthood.

264 This study also has some limitations that could affect the interpretation of the results. The observed  
265 association between linear growth velocity with cystatin C and size could be confounded by other  
266 potential prenatal factors such as maternal morbidity status and diet. At this age, we cannot ascertain  
267 whether the observed associations represent early subclinical kidney function deficits or normal  
268 growth-related phenomena. We cannot exclude the possibility of reverse causality in the absence of  
269 kidney phenotyping prior to the current assessment. We used cystatin C instead of estimating the  
270 glomerular filtration rate as the existing formulas are not validated for our study setting. We were  
271 unable to obtain pubertal status, which in high income settings, has been associated with cystatin C  
272 in healthy children (55) . Additionally, the loss to follow-up of substantial number of children with  
273 LBW may introduce bias. However, we found no association between serum cystatin C level and  
274 LBW status amongst children who attended the 10<sup>th</sup> follow-up, though this analyses may have been  
275 underpowered.

276 In conclusion, based on our findings, greater growth between 0 and 6 years favors kidney size to meet  
277 requirements of a given BSA, however greater growth after 2 years of age is associated with serum  
278 cystatin C level. Thus, existing programs that target the first 1000 days of life are still important and  
279 should be strengthened, but interventions to address linear growth in children over the age of two  
280 should also be in place. Additionally, we recommend other researchers carry out similar studies with  
281 a sizable sample size in contexts of LMICs.

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284 individuals involved in the data collection and supervision.

## 285 **Authors' contribution**

286 The authors' contributions were as follows HF, JCKW, TG, GSA, RW, DN, SF, DY, and BZ designed  
287 the study. BZ, RA, and BSM supervised the data collection; HF, JCKW, TG, GSA, RW, MFO, DY,  
288 SF, DN, EA, MA, AA, MB, TB and BA participated in methodology. Data analysis done by BZ, RW  
289 and GSA. BZ wrote the first draft of the manuscript and had responsibility for the whole work. BZ,  
290 DY, DN, and HF had primary responsibility for the final content. All authors contributed to the  
291 manuscript revisions and read the final manuscript and approved it for submission.



## References

1. Awazu K. Early life origins of human health and disease. *Ir Med J* [Internet]. 2010;103(8):2009–10. Available from: [https://www.researchgate.net/publication/260744098\\_Early\\_Life\\_Origins\\_of\\_Human\\_Health\\_and\\_Disease](https://www.researchgate.net/publication/260744098_Early_Life_Origins_of_Human_Health_and_Disease)
2. Tain YL, Hsu CN. Developmental origins of chronic kidney disease: Should we focus on early life? *Int J Mol Sci* [Internet]. 2017;18(2):1. Available from: <https://pubmed.ncbi.nlm.nih.gov/28208659>
3. Hsu CN, Tain YL. Developmental origins of kidney disease: Why oxidative stress matters? *Antioxidants* [Internet]. 2021;10(1):1–18. Available from: <https://www.mdpi.com/2076-3921/10/1/33>
4. Hoy WE, Hughson MD, Bertram JF, Douglas-Denton R, Amann K. Nephron number, hypertension, renal disease, and renal failure. *J Am Soc Nephrol* [Internet]. 2005;16(9):2559. Available from: <https://jasn.asnjournals.org/content/jnephrol/16/9/2557.full.pdf>
5. Luyckx VA, Brenner BM. The clinical importance of nephron mass. *J Am Soc Nephrol* [Internet]. 2010;21(6):1–3. Available from: <https://www.researchgate.net/publication/41424947>
6. Viviani S. Nephron Number in Patients with Primary Hypertension. *N Engl J Med* [Internet]. 2011;348(2):105. Available from: <https://www.nejm.org/doi/full/10.1056/NEJMoa020549>
7. Bertram JF, Douglas-Denton RN, Diouf B, Hughson MD, Hoy WE. Human nephron number: Implications for health and disease. *Pediatr Nephrol* [Internet]. 2011;26(9):1531–3. Available

from: <https://pubmed.ncbi.nlm.nih.gov/21604189/>

8. Luyckx VA, Bertram JF, Brenner BM, Fall C, Hoy WE, Ozanne SE, et al. Effect of fetal and child health on kidney development and long-term risk of hypertension and kidney disease. *Lancet* [Internet]. 2013;382(9888):275. Available from: [http://dx.doi.org/10.1016/S0140-6736\(13\)60311-6](http://dx.doi.org/10.1016/S0140-6736(13)60311-6)
9. Puelles VG, Hoy WE, Hughson MD, Diouf B, Douglas-Denton RN, Bertram JF. Glomerular number and size variability and risk for kidney disease. *Curr Opin Nephrol Hypertens* [Internet]. 2011;20(1):7–8. Available from: <https://pubmed.ncbi.nlm.nih.gov/21099687/>
10. Hoy WE, Hughson MD, Bertram JF, Douglas-denton R, Amann K. Nephron Number, Hypertension, Renal Disease, and Renal Failure. *J Am Soc Nephrol* [Internet]. 2005;16:2557–8. Available from: <https://europepmc.org/article/MED/16049069>
11. Batty GD, Shipley MJ, Gunnell D, Huxley R, Kivimaki M, Woodward M, et al. Height, wealth, and health: An overview with new data from three longitudinal studies. *Econ Hum Biol* [Internet]. 2009;7(2):138. Available from: <https://pubmed.ncbi.nlm.nih.gov/19628438/>
12. Neal Halfon, Christopher B. Forrest, Richard M. Lerner EMF. *Handbook of Life Course Health Development* [Internet]. Springer; 2018. 1–2 p. Available from: <https://www.ncbi.nlm.nih.gov/books/NBK543707/>
13. Bogin B, Varela-Silva MI. Leg length, body proportion, and health: A review with a note on beauty. *Int J Environ Res Public Health* [Internet]. 2010;7(3):1047–75. Available from: <https://www.ncbi.nlm.nih.gov/pmc/articles/PMC2872302/>

14. Millward DJ. Nutrition, infection and stunting: The roles of deficiencies of individual nutrients and foods, and of inflammation, as determinants of reduced linear growth of children. *Nutr Res Rev* [Internet]. 2017;30(1):50–72. Available from:  
<https://www.cambridge.org/core/journals/nutrition-research-reviews/article/nutrition-infection-and-stunting-the-roles-of-deficiencies-of-individual-nutrients-and-foods-and-of-inflammation-as-determinants-of-reduced-linear-growth-of-children/195A1401085226>
15. Hoy WE, Hughson MD, Singh GR, Douglas-Denton R, Bertram JF. Reduced nephron number and glomerulomegaly in Australian Aborigines: A group at high risk for renal disease and hypertension. *Kidney Int* [Internet]. 2006;70(1):104–10. Available from:  
<https://www.sciencedirect.com/science/article/pii/S0085253815517419>
16. Atukunda P, Ngari M, Chen X, Westerberg AC, Iversen PO, Muhoozi G. Longitudinal assessments of child growth: A six-year follow-up of a cluster-randomized maternal education trial. *Clin Nutr* [Internet]. 2021;40(9):5106–13. Available from:  
<https://doi.org/10.1016/j.clnu.2021.08.007>
17. Takele BA, Gezie LD, Alamneh TS. Pooled prevalence of stunting and associated factors among children aged 6-59 months in Sub-Saharan Africa countries: A Bayesian multilevel approach. *PLoS One* [Internet]. 2022;17(10 October):1–19. Available from:  
<http://dx.doi.org/10.1371/journal.pone.0275889>
18. Brophy PD, Charlton JR, Carmody JB, Reidy KJ, Harshman L, Segar J, et al. Chronic Kidney Disease : A Life Course Health Development Perspective. 2018; Available from:  
<https://www.ncbi.nlm.nih.gov/books/NBK543705/>

19. Kayange NM, Smart LR, Tallman JE, Chu EY, Fitzgerald DW, Pain KJ, et al. Kidney disease among children in sub-Saharan Africa: Systematic review. *Pediatr Res* [Internet]. 2015;77(2):272–81. Available from: <https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4426498/>
20. Jha V, Parameswaran S. Community-acquired acute kidney injury in tropical countries. *Nat Rev Nephrol* [Internet]. 2013;9(5):278–90. Available from: <http://dx.doi.org/10.1038/nrneph.2013.36>
21. Stanifer JW, Muiro A, Jafar TH, Patel UD. Chronic kidney disease in low- and middle-income countries. *Nephrol Dial Transplant* [Internet]. 2016;31(6):868–74. Available from: <https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4876969/>
22. Schmidt IM, Mølgaard C, Main KM, Michaelsen KF. Effect of gender and lean body mass on kidney size in healthy 10-year-old children. *Pediatr Nephrol* [Internet]. 2001;16(4):366–70. Available from: <https://link.springer.com/content/pdf/10.1007/s004670100568.pdf>
23. Bakker H, Gaillard R, Franco OH, Hofman A, Der AJ Van, Steegers EAP, et al. Fetal and Infant Growth Patterns and Kidney Function at School Age. *J Am Soc Nephrol* [Internet]. 2014;25:2607–15. Available from: <https://jasn.asnjournals.org/content/jnephrol/25/11/2607.full.pdf>
24. Barker DJP, Osmond C, Forsén TJ, Kajantie E, Eriksson JG. Trajectories of Growth among Children Who Have Coronary Events as Adults. *N Engl J Med* [Internet]. 2005;353(17):1802–9. Available from: [https://www.nejm.org/doi/10.1056/NEJMoa044160?url\\_ver=Z39.88-2003&rfr\\_id=ori:rid:crossref.org&rfr\\_dat=cr\\_pub\\_0www.ncbi.nlm.nih.gov](https://www.nejm.org/doi/10.1056/NEJMoa044160?url_ver=Z39.88-2003&rfr_id=ori:rid:crossref.org&rfr_dat=cr_pub_0www.ncbi.nlm.nih.gov)
25. Bakker H, Gaillard R, Franco OH, Hofman A, Van Der Heijden AJ, Steegers EAP, et al. Fetal

and infant growth patterns and kidney function at school age. *J Am Soc Nephrol.*

2014;25(11):2607–15.

26. Andersen GS, Girma T, Wells JCK, Kæstel P, Leventi M, Hother AL, et al. Body composition from birth to 6 mo of age in Ethiopian infants: Reference data obtained by air-displacement plethysmography. *Am J Clin Nutr.* 2013;98(4):885–94.
27. Wibaek R, Vistisen D, Girma T, Admassu B, Abera M, Abdissa A, et al. Body mass index trajectories in early childhood in relation to cardiometabolic risk profile and body composition at 5 years of age. *Am J Clin Nutr.* 2019;110(5):1175–85.
28. Payne RB. Tests of Kidney Function [Internet]. Second Edi. Scientific Foundations of Biochemistry in Clinical Practice. D. L. Williams and V. Marks; 1994. 325–337 p. Available from: <http://dx.doi.org/10.1016/B978-0-7506-0167-2.50024-8>
29. Wibaek R, Vistisen D, Girma T, Admassu B, Abera M, Abdissa A, et al. Associations of fat mass and fat-free mass accretion in infancy with body composition and cardiometabolic risk markers at 5 years : The Ethiopian iABC birth cohort study. *PLoS Med* [Internet]. 2019;16(8):1–22. Available from: <https://journals.plos.org/plosmedicine/article?id=10.1371/journal.pmed.1002888>
30. Ellis KJ, Yao M, Shypailo RJ, Urlando A, Wong WW, Heird WC. Body-composition assessment in infancy: Air-displacement plethysmography compared with a reference 4-compartment model. *Am J Clin Nutr* [Internet]. 2007;85(1):90–5. Available from: <https://pubmed.ncbi.nlm.nih.gov/17209182/>

31. Ma G, Yao M, Lin Y, Lin A, Zou H, Urlando A, et al. Validation of a new pediatric air-displacement plethysmograph for assessing body composition in infants. *Am J Clin Nutr* [Internet]. 2004;79(4):653–60. Available from:  
<https://academic.oup.com/ajcn/article/79/4/653/4690163>
32. Mohamed M, Behery E, Ibrahiem MA, Siam S, Seksaka MA. Fetal Renal Volume and Fetal Doppler in Normal and Growth Restricted Fetuses : Is there a Correlation ? *Gynecology & Obstetrics*. 2012;2(2):2–6. Available from: <https://www.longdom.org/open-access/fetal-renal-volume-and-fetal-doppler-in-normal-and-growth-restricted-fetuses-is-there-a-correlation-37756.html>
33. Sampaio FJB. Theoretical kidney volume versus real kidney volume: comparative evaluation in fetuses. *J Clin Anat* [Internet]. 1995;71–5. Available from:  
<https://link.springer.com/content/pdf/10.1007/BF01629504.pdf>
34. Ferguson TW, Komenda P, Tangri N. Cystatin C as a biomarker for estimating glomerular filtration rate. *Curr Opin Nephrol Hypertens*. 2015;24(3):295–300.
35. Dönmez O, Korkmaz HA, Yldz N, Ediz B. Comparison of serum cystatin C and creatinine levels in determining glomerular filtration rate in children with stage i to III chronic renal disease. *Ren Fail*. 2015;37(5):784–90.
36. Zappitelli M, Parvex P, Joseph L, Paradis G, Grey V, Lau S, et al. Derivation and Validation of Cystatin C – Based Prediction Equations for GFR in Children. *Am J Kidney Dis* [Internet]. 2006;48(2):221–30. Available from:  
<https://www.sciencedirect.com/science/article/pii/S0272638606008286?via%3Dihub>

37. Pan H, Goldstein H. Multi-level repeated measures growth modelling using extended spline functions. *Stat Med*. 1998;17(23):2755–70.
38. Anderson EL, Tilling K, Fraser A, Macdonald-Wallis C, Emmett P, Cribb V, et al. Estimating trajectories of energy intake through childhood and adolescence using linear-spline multilevel models. *Epidemiology* [Internet]. 2013;24(4):507–15. Available from: [https://journals.lww.com/epidem/Fulltext/2013/07000/Estimating\\_Trajectories\\_of\\_Energy\\_Intake\\_Through.5.aspx](https://journals.lww.com/epidem/Fulltext/2013/07000/Estimating_Trajectories_of_Energy_Intake_Through.5.aspx)
39. Howe LD, Tilling K, Matijasevich A, Petherick ES, Santos AC, Fairley L, et al. Linear spline multilevel models for summarising childhood growth trajectories: A guide to their application using examples from five birth cohorts. *Stat Methods Med Res* [Internet]. 2016;25(5):1854–74. Available from: [https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4074455/pdf/10.1177\\_0962280213503925.pdf](https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4074455/pdf/10.1177_0962280213503925.pdf)
40. Wang Y, Levey AS, Inker LA, Jessani S, Bux R, Samad Z, et al. Performance and Determinants of Serum Creatinine and Cystatin C–Based GFR Estimating Equations in South Asians. *Kidney Int Reports* [Internet]. 2021;6(4):962–75. Available from: <https://doi.org/10.1016/j.ekir.2021.01.005>
41. Zhang X, Martin RM, Oken E, Aris IM, Yang S, Kramer MS. Growth during Infancy and Early Childhood and Its Association with Metabolic Risk Biomarkers at 11.5 Years of Age. *Am J Epidemiol* [Internet]. 2020;189(4):286–93. Available from: <https://www.ncbi.nlm.nih.gov/pmc/articles/PMC7305788/pdf/kwz234.pdf>
42. Jones A, Charakida M, Falaschetti E, Hingorani AD, Finer N, Masi S, et al. Adipose and height

growth through childhood and blood pressure status in a large prospective cohort study.

Hypertension [Internet]. 2012;59(5):919–25. Available from:

<https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3428923/pdf/ukmss-47724.pdf>

43. Regnault N, Kleinman KP, Rifas-Shiman SL, Langenberg C, Lipshultz SE, Gillman MW. Components of height and blood pressure in childhood. *Int J Epidemiol* [Internet]. 2014;43(1):149–59. Available from:  
<https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3937979/pdf/dyt248.pdf>
44. Adair LS, Fall CHD, Osmond C, Stein AD, Martorell R, Ramirez-Zea M, et al. Associations of linear growth and relative weight gain during early life with adult health and human capital in countries of low and middle income: Findings from five birth cohort studies. *Lancet* [Internet]. 2013;382(9891):525–34. Available from:  
<https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3744751/>
45. Weder AB, Schork NJ. Hypothesis Adaptation, Allometry, and Hypertension. *Hypertensio*. 1994;24:145–56.
46. Nüesch E, Dale C, Palmer TM, White J, Keating BJ, van Iperen EPA, et al. Adult height, coronary heart disease and stroke: A multi-locus Mendelian randomization meta-analysis. *Int J Epidemiol*. 2016;45(6):1927–37.
47. Dinkel E, Ertel M, Dittrich M, Peters H, Berres M, Wissermann HS. Pediatric Radiology Sonographical growth charts for kidney length and volume. *Pediatr Radiol* [Internet]. 1985;2:38–43. Available from: <https://link.springer.com/content/pdf/10.1007/BF02387851.pdf>



48. Whitehall J. Anthropometry and renal size of children suffering under sustained conflict in Sri Lanka. *J Paediatr Child Health*. 2008;44(11):656–60.
49. Bakker H, Kooijman MN, van der Heijden AJ, Hofman A, Franco OH, Taal HR, et al. Kidney size and function in a multi-ethnic population-based cohort of school-age children. *Pediatr Nephrol* [Internet]. 2014;29(9):1589–98. Available from: <https://link.springer.com/content/pdf/10.1007/s00467-014-2793-8.pdf>
50. Fong D, Denton KM, Moritz KM, Evans R, Singh RR. Compensatory responses to nephron deficiency: Adaptive or maladaptive? *Nephrology*. 2014;19(3):119–28.
51. Wells JCK, Devakumar D, Grijalva-eternod CS, Manandhar DS. Blood Pressure and the Capacity-Load Model in 8-Year-Old Children from Nepal : Testing the Contributions of Kidney Size and Intergenerational Effects. *Am J of Human Biol*. 2016;565:555–65.
52. Yang L, Magnussen CG, Yang L, Bovet P, Xi B. Elevated blood pressure in childhood or adolescence and cardiovascular outcomes in adulthood: A systematic review. *Hypertension*. 2020;948–55.
53. Juhola J, Magnussen CG, Viikari JSA, Kähönen M, Hutri-Kähönen N, Jula A, et al. Tracking of serum lipid levels, blood pressure, and body mass index from childhood to adulthood: The cardiovascular risk in young Finns study. *J Pediatr*. 2011;159(4):584–90.
54. Luyckx VA, Bertram JF, Brenner BM, Fall C, Hoy WE, Ozanne SE, et al. Effect of fetal and child health on kidney development and long-term risk of hypertension and kidney disease. *Lancet* [Internet]. 2013;382(9888):273–83. Available from: <http://dx.doi.org/10.1016/S0140->

6736(13)60311-6

55. Nitsch D, Sandling JK, Byberg L, Larsson A, Tuvemo T, Syvnen AC, et al. Fetal, developmental, and parental influences on cystatin C in childhood: The uppsala family study. *Am J Kidney Dis* [Internet]. 2011;57(6):863–72. Available from: <http://dx.doi.org/10.1053/j.ajkd.2010.12.025>

Table1: Description of child and maternal characteristics.

<b>Characteristics</b>	<b>N</b>	<b>Mean (<math>\pm</math>SD) or n (%)</b>
<b>Maternal characteristics at birth</b>		
Age at delivery (years)	343	22.1 $\pm$ 4.5
Height (cm)	342	157.6 $\pm$ 5.9
Education	353	
No school		21 (6.0)
Primary school		214 (60.6)
Secondary school		67 (19.0)
Higher education		51 (14.4)
<b>Child characteristics at birth</b>		
Gestational age (weeks)	353	39.0 $\pm$ 1.0
Fat mass (kg)	349	0.22 $\pm$ 0.2
Fat free mass (kg)	349	2.84 $\pm$ 0.3
Birth weight (kg)	351	
$\leq$ 2.5		30 (8.5)
$>$ 2.5 $\leq$ 3.0		116 (33.1)
$>$ 3.0 $\leq$ 3.5		152 (43.3)
$>$ 3.5		53 (15.1)
Birth order	351	
First born		171(48.7)
Second born		94 (26.8)

>= Third born 86 (24.5)

**Current child characteristics**

Sex (male)	353	183 (51.8)
Age (years)	353	9.79 ± 1.0
Weight (kg)	353	27.3 ± 6.0
Height (cm)	353	132.2 ± 7.7
BMI (kg/m <sup>2</sup> )	353	15.5 ± 2.2
Fat mass (kg)	351	5.6 ± 3.5
Fat free mass (kg)	351	21.7 ± 3.4

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<sup>1</sup> Abbreviations: BMI, body mass index. <sup>2</sup> Data are mean (±SD) for continuous and count (%) for categorical variables

Table 2. Length at birth and non-standardized linear growth at birth and growth velocity 0-6 years.

	Male (N=183)	Female (N=170)	P-value
Length at birth (cm)	49.61 ± 1.6	49.0 ± 1.5	<0.001
Growth velocity 0-3 months (cm/month)	4.09 ± 0.3	3.93 ± 0.22	<0.001
Growth velocity 3-6 months (cm/month)	1.72 ± 0.2	1.74 ± 0.19	0.63
Growth velocity 6-24 months (cm/month)	0.91 ± 0.1	0.94 ± 0.11	0.03
Growth velocity 24-48 months (cm/month)	0.60 ± 0.2	0.60 ± 0.06	0.46
Growth velocity 48-76 months (cm/month)	0.56 ± 0.02	0.56 ± 0.02	0.85

<sup>1</sup>Data presented by mean (±SD). <sup>2</sup>Independent samples t-test.

Table 3: Markers of kidney function and kidney volume at the age of 7-12 years

Kidney outcomes	Median (IQR)	
	Male	Female
Serum cystatin C (mg/dl )	0.93 (0.83; 1.00)	0.94 (0.83; 1.01)
Estimated GFR <sub>cystatin C</sub> (ml/min per 1.73m <sup>2</sup> )	83 (76; 94)	82 (75; 94)
Right kidney volume (cm <sup>3</sup> )	54 (47; 63)	53 (45; 61)
Left kidney volume (cm <sup>3</sup> )	64 (56; 74)	63 (57; 72)
Combined kidney volume (cm <sup>3</sup> )	117 (104; 136)	117. (103; 132)

<sup>1</sup>Abbreviations: GFR, Glomerular Filtration Rate. <sup>2</sup>Data are presented on median (Interquartile range)

## Figure legends

**Fig 1:** Flow diagram showing number of participants followed up at different time points. Out of 355 recruited at 10-year follow-up, 2 were excluded analysis they only had 1 height measurement from birth to 6 years, which makes the final sample for the 10-year analysis 353.

**Fig 2:** Association of standardized linear growth velocities 0-76 months with Log serum cystatin C at age of 10 years. The Y-axis represents the estimate with 95% CI. Model 1: Adjusted for sex and Current age. Model 2: as model 1, further adjusted for birth weight, gestational age, birth order and fat mass at 10<sup>th</sup> follow-up. Model 3: as model 2, further adjusted for maternal education and maternal post-partum height.

**Fig 3:** Association of standardized linear growth velocities 0-6 years with kidney volume at 10 years at age of 10 years. The Y-axis represents the estimate with 95% CI. Model 1: Adjusted for sex and current age. Model 2: as model 1, further adjusted for birth weight, gestational age, birth order, and fat mass at 10<sup>th</sup> follow-up. Model 3: as model 2, further adjusted for maternal education and maternal post-partum height.