Risk factors for major external structural birth defects among children in Kiambu County, Kenya: a case-control study [version 2; peer review: 2 approved, 1 approved with reservations]

George N. Agot, Marshal M. Mweu, Joseph K. Wang’ombe

School of Public Health, College of Health Sciences, University of Nairobi, Nairobi, Kenya

Abstract

Background: Although major external structural birth defects continue to occur globally, the greatest burden is shouldered by resource-constrained countries with no surveillance systems. To our knowledge, many studies have been published on risk factors for major external structural birth defects, however, limited studies have been published in developing countries. The objective of this study was to identify risk factors for major external structural birth defects among children in Kiambu County, Kenya.

Methods: A hospital-based case-control study was used to identify the risk factors for major external structural birth defects. A structured questionnaire was used to gather information retrospectively on maternal exposure to environmental teratogens, multifactorial inheritance, and sociodemographic-environmental factors during the study participants' last pregnancies. Descriptive analyses (means, standard deviations, medians, and ranges) were used to summarize continuous variables, whereas categorical variables were summarized as proportions and percentages in frequency tables. Afterward, logistic regression analyses were conducted to estimate the effects of the predictors on the odds of major external structural birth defects in the country.

Results: Women who conceived when residing in Ruiru sub-county (adjusted odds ratio [aOR]: 5.28; 95% CI; 1.68-16.58; P<0.01), and Kiambu sub-county (aOR: 0.27; 95% CI; 0.076-0.95; P=0.04), and preceding siblings with history of birth defects (aOR: 7.65; 95% CI; 1.46-40.01; P=0.02) were identified as the significant predictors of major external structural birth defects in the county.

Conclusions: These findings pointed to MESBDs of genetic, multifactorial inheritance, and sociodemographic-environmental etiology. Thus, we recommend regional defect-specific surveillance programs, public health preventive measures, and treatment
strategies to understand the epidemiology and economic burden of these defects in Kenya. We specifically recommend the integration of clinical genetic services with routine reproductive health services because of potential maternal genetic predisposition in the region.

**Keywords**
Major external structural birth defects, risk factors, case-control study, Kenya

**Corresponding author:** George N. Agot (nyadimogeorge@gmail.com)

**Author roles:** Agot GN: Conceptualization, Data Curation, Formal Analysis, Funding Acquisition, Investigation, Methodology, Project Administration, Resources, Software, Supervision, Validation, Visualization, Writing – Original Draft Preparation, Writing – Review & Editing; Mweu MM: Conceptualization, Supervision, Writing – Review & Editing; Wang'ombe JK: Conceptualization, Supervision

**Competing interests:** No competing interests were disclosed.

**Grant information:** The author(s) declared that no grants were involved in supporting this work.

**Copyright:** © 2021 Agot GN et al. This is an open access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

**How to cite this article:** Agot GN, Mweu MM and Wang’ombe JK. Risk factors for major external structural birth defects among children in Kiambu County, Kenya: a case-control study [version 2; peer review: 2 approved, 1 approved with reservations]
F1000Research 2021, 10:59 https://doi.org/10.12688/f1000research.50738.2
First published: 01 Feb 2021, 10:59 https://doi.org/10.12688/f1000research.50738.1
Amendments from Version 1

To address the reviewers’ comments, we have clarified that; the hypothesized odds ratio used in sample size calculation was set at 2.0 (universally accepted), different types of birth defects were collectively used in sample size calculation rather individual defects due to the rarity of the defects coupled with the unavailability of surveillance systems for birth defects in the country, and that data collection period spanned three months from May 31st 2019 to July 31st 2019. Additionally, some of these defects are potentially fatal and could possibly introduce survivor bias in the study, thus we clarified being cognizant of this epidemiological phenomenon, however we could not minimize it considering that pathological examinations to determine the causes of death in stillbirths and miscarriages is not a routine practice in the country. Maternal residence, at conception and paternal age were controlled for in the multivariable analyses; however maternal cigarette smoking was not controlled for due to negligible responses received from the study participants. We have also redesigned our conceptual framework, defined maternal exposure to pesticides and chronic illnesses in the context of this study. We also improved the texts in the introduction, discussion, and conclusion sections of the manuscript, including changes to Figure 1 and all Tables.

Any further responses from the reviewers can be found at the end of the article.

Introduction

Worldwide, an estimated 7.9 million children are born every year with a birth defect of which approximately 3.3 million die before age five and around 3.2 million could be physically disabled for life\(^1\). More than 94% of such defects occur in developing countries where about 95% of these children do not survive beyond childhood\(^1\). Birth defects are defined as abnormalities of body structures or functions that develop during the organogenesis period (first trimester of gestation) and detectable during pregnancy, at birth, or soon after\(^2\). These defects may be classified as major when associated with significant adverse health effects requiring medical/surgical care; otherwise, they are described as minor\(^3\). Alternatively, they can be classified as external when visible at birth or soon after; or internal when advanced medical imaging techniques are required for their detection\(^4\). Consequently, the phrase ‘major external structural birth defects’ (MESBDs) denotes congenital physical abnormalities that are clinically obvious at birth or soon after which call for medical and/or surgical interventions\(^1,2\). The causes of these defects can be classified into three categories: (i) identifiable environmental factors (teratogens/micronutrient deficiencies); (ii) identifiable genetic factors; and (iii) complex genetic and idiopathic environmental factors, described as multifactorial inheritance\(^1,4,5,10\). One-third of these causes are attributed to identifiable environmental and genetic factors, whereas the rest are believed to be of multifactorial aetiology\(^5,6,7,8,10\). Additionally, an environmental endowment of women of reproductive age is thought to operate through their socioeconomic and sociodemographic characteristics leading to causes of MESBDs, described as sociodemographic-environmental factors\(^1,4,8,10\).

Organogenesis occurs in the first eight weeks of gestation; however, approximately half of pregnancies are usually unplanned/unintended, thus not recognized until the end of the second trimester\(^1,3,7,13,15\). Completing more years of education could improve maternal health because educated women are more likely to make informed reproductive health choices than those with low levels of education to improve birth outcomes\(^16,17\). Some of the notable maternal decisions include planned pregnancy, preconception folic acid intake in anticipation of conception, and subsequently prompt prenatal care\(^18,19\). Supplemental vitamins with folic acid are dispensed during routine antenatal care (ANC) visits, as well as health education on adequate nutrition, avoidance of environmental teratogens, and maternal infections as public health preventive strategies for MESBDs\(^18,20\). These measures could be effective only when pregnant women promptly began antenatal care within eight weeks of gestation before the intrauterine formation of MESBDs\(^4\). Folic acid is essential for normal development of the brain and spinal cord during the first 4 weeks of conception, and have been found to reduce the occurrence of neural tube defects, orofacial clefts, limb reduction defects, urinary system defects, and omphalocele; some of the most prevalent defects in the county\(^21-23\). Thus, the recommended first ANC at the 12th week of pregnancy could be a sub-optimal preventive strategy for these defects, nevertheless it improves experiences of the women during pregnancy and childbirth\(^24\). Maternal occupation as a predictor of MESBDs could be dependent on educational levels, nonetheless occupations such as farming could expose women of reproductive age to teratogenic pesticides\(^25\).

Maternal residence at conception is similarly a significant predictor of MESBDs determined by environmental etiology attributed to widespread poverty, environmental pollution, inadequate health care services, and ineffective preventive strategies; factors largely found in developing countries\(^26,30\). Parental age is a multifaceted risk factor whose mechanisms of actions in the intrauterine formation of MESBDs are underpinned by human biology and socio-economic endowment among women of reproductive age. From the biologic standpoint, the female gametogenesis begins before birth with the initial meiotic division (prophase stage) expected to complete shortly before ovulation, however, this is not the case always because the process may delay up to 45 years to conclude\(^27\). Thus, the oocytes take exceedingly long in the prophase stage increasing the likelihood of meiotic errors due to exposure to the environmental teratogens\(^28\). Advancing maternal age beyond 35 years is similarly a risk factor for MESBDs of genetic etiology due to chromosomal abnormalities\(^24,31,32\). Similarly, from the genetic viewpoint, genetic mutations and accumulation of chromosomal aberrations during the maturation of male germ cells have been attributed to the formation of MESBDs in utero\(^33,34\). The amount of deoxyribonucleic acid damage in sperm of men aged 36–57 is three times that of men <35 years, increasing the likelihood of these defects in aging couples\(^34\). From the socio-economic perspective, parental age could be associated with MESBDs of multifactorial etiology ascribed to physiological interactions between complex genetic and idiopathic environmental attributes of women of reproductive age\(^29,30,36,37\).

To our knowledge, many studies on the risk factors have been published in developed countries, however, such publications are scanty in developing countries owing to the rarity of the defects, unplanned/unintended pregnancies, and difficulties in
identifying these women until the end of the second trimester when the defects have already formed\textsuperscript{3}. To address this gap, this study investigated maternal periconceptional exposure to environmental teratogens, sociodemographic-environmental, and multifactorial risk factors for MESBDs in Kiambu County, Kenya. The study assessed: maternal periconceptional exposure to farm-sprayed pesticides, and teratogenic therapeutic medicines proxied by maternal chronic illnesses (epilepsy and depression); multifactorial inheritance proxied by the history of siblings with birth defects, sex of the “last born” current child, nature of pregnancy, and parity; and sociodemographic-environmental factors consisting of maternal age, paternal age, residence, level of education, occupation, and adequate prenatal care proxied by gestational age at first ANC, and preconception folic acid intake. The findings of this study could provide great public health opportunities for the formulation of specific treatment strategies, preventive measures, risk-based surveillance systems, and clinical genetic services for the most prevalent MESBDs, regionally and nationally. Consequently, the objective of this study was to identify the risk factors for MESBDs among children in Kiambu County, Kenya.

Methods

Study design and settings

A hospital-based case-control study was conducted to identify the risk factors for MESBDs. The study participants were recruited as they presented to the child welfare clinics, neonatal/pediatric units, and occupational clinics for care during the data collection period from May 31\textsuperscript{st} 2019 to July 31\textsuperscript{st} 2019. A case-control design was the optimal design for this study considering its suitability for the investigation of rare outcomes, as is the case with MESBDs. Even though a population-based design would have been preferable, the ease of recruiting case and control subjects within the hospital settings disproportionately favored the hospital-based design. This was an observational study, therefore was reported as per the STROBE guidelines\textsuperscript{38}.

The study was conducted in 13 hospitals comprising three-county referral hospitals (Kiambu, Gatundu, and Thika), eight sub-county hospitals (Karuri, Kihara, Wangige, Nyathuna, Lari, Tigoni, Lussigetti, and Kigumo), and two faith-based hospitals (Presbyterian Church of East Africa Kikuyu Orthopedic and African Inland Church Cure International) situated within Kiambu County, Kenya. Notably, neither population-based nor hospital-based surveillance systems for MESBDs existed in the county nor the study hospitals. Nonetheless, cases detected by primary health providers during childbirth and neonatal care were recorded for the compilation of monthly hospital reports and subsequent entry into the District Health Information System (DHIS). The cases were drawn from Kiambu, Thika, Gatundu, Tigoni, Kikuyu, and Cure hospitals, which provided occupational and rehabilitative health services to children with MESBDs. The controls, on the other hand, were drawn from Kiambu, Gatundu, Thika, Karuri, Kihara, Wangige, Nyathuna, Lari-Rukuma, Tigoni, Lussigetti, and Kigumo hospitals which provided child welfare services to the under-fives. Kiambu is the second-most densely inhabited county with an estimated population of 2.4 million people out of an estimated national population of 47.5 million\textsuperscript{39}. Its economic mainstay is largely agriculture, comprising tea, coffee, and dairy farming\textsuperscript{39}. Of the county’s total estimated population, approximately 2.2% aged ≥5 years are living with lifelong disabilities\textsuperscript{37}. A study carried out in the county between 2014 and 2018 observed defects of the musculoskeletal system as the most prevalent single system defects followed by central nervous, orofacial clefts genital, ocular, and anal organ defects\textsuperscript{21}.

Study population and eligibility of participants

The study population consisted of children aged ≤5 years old seeking health services at the study hospitals during the study period spanning from May to July 2019. All children whose mothers consented to participate in the study were recruited.

Case definition and recruitment

Cases were defined as children aged ≤5 years born with at least one MESBD to resident women of Kiambu County and seeking health care services at the neonatal units, pediatric wards, child welfare clinics, and/or occupational therapist clinics of the study hospitals during the three-month study period. The Research Assistants (RAs) liaised with team leads of the departments listed above to identify cases of MESBDs. The team leads had been working in these departments, thus were conversant with the cases seeking services. The team leads invited the mothers of the children who met the case definition to comfortable private rooms within the departments where informed consent was sought and interviews conducted by the RAs. All cases that met this definition and whose caregivers consented to participate were prospectively recruited into the study until the required sample was attained (see Sample size determination).

Control definition and recruitment

Controls were defined as children aged ≤5 years born without any forms of birth defects to resident women of Kiambu County and attending routine child welfare clinics at the study hospitals during the same three-month study period. The Research Assistants liaised with team leads of the child welfare clinics to identify the children without any form of birth defects and were seeking routine immunization, and growth monitoring services. The team leads had been working in these clinics, hence were familiar with most of the under-fives seeking the services. These services are provided between 8.00 am and 5.00 pm from Monday to Friday; the team leads introduced the RAs who then briefed the potential participants on the study objectives. Because of the relatively large number of controls available, they were selected by simple randomization using sealed envelopes upon definition of the sample population and frequency-matched to the cases by the day of presentation. Informed consent was sought from the study participants who met the study eligibility criteria; those who consented to participate in the study were prospectively recruited and invited to secluded comfortable rooms within the clinics where face-to-face interviewer questionnaires were administered till the desired sample size was achieved (see sample size determination).

Sample size determination

The sample size was estimated as per the Kelsey JL et al.\textsuperscript{40} formula specified for case-control studies as follows: -
\[ n_1 = \frac{(Z_\alpha + Z_\beta)^2 \bar{pq} (r + 1)}{r(p_1 - p_2)^2} \]

\[ \bar{q} = 1 - \bar{p} \]

\[ n_2 = r n_1 \]

\[ p_1 = \frac{p_1 OR}{1 + p_2 (OR - 1)} \]

\[ \bar{p} = \frac{p_1 + r p_2}{r + 1} \]

Where: \( n_1 \) is the number of cases and \( n_2 \) is the number of controls; \( p_1 \) is the proportion of cases whose caregivers did not begin prenatal care in the first trimester (primary exposure), \( p_2 \) is the proportion of controls whose caregivers did not begin prenatal care in the first-trimester set at 57%\(^{1,12}\). Remarkably, \( Z_\alpha \) (1.96) and \( Z_\beta \) (-0.84) are the values specifying the desired two-tailed confidence level (95%) and statistical power (80%), respectively. The odds ratio (OR) for the effect of the primary exposure (cases whose caregivers did not begin prenatal care in the first trimester) was hypothesized to be 2.0 (universally accepted). The ratio \( r \) of unexposed to exposed individuals was set at 3.0, and given the estimates, a total sample size of 408 participants was derived (102 cases, and 306 controls).

**Data collection process and study variables**

Before data collection, four nursing graduate interns were recruited and trained as RAs on sound interviewing techniques, and information derivation/validation from antenatal care (ANC) booklets. This was to ensure the data collection process spanning three months (May 31\(^{st}\) 2019 to July 31\(^{st}\), 2019) was conducted in a standardized manner. The ANC booklet contains maternal profile, medical/surgical history, previous pregnancy history, clinical notes, and physical examination findings on ANC visits, among others. The maternal profile includes name, age, parity gravidity, height, weight, last menstrual period (LMP), expected date of delivery (EDD), and date of first ANC. Face-to-face structured questionnaires (see Extended data) were administered to the mothers of the study participants by RAs in comfortable secluded rooms within neonatal units and occupational therapy clinics for cases and child welfare clinics for the controls. Data were gathered retrospectively on exposures to environment-teratogens (farm-sprayed pesticides, and teratogenic medicines proxied by chronic illnesses), multifactorial inheritance (parity, nature of pregnancy, history of siblings with birth defects and sex of the “lastborn” (current) child), and sociodemographic-environmental factors (maternal age, paternal age, residence, education level, occupation, and adequate prenatal care proxied by gestational age at first ANC and preconception folic acid intake). The predictors were assessed as shown in Table 1.

**Table 1. Study variables and their assessments.**

<table>
<thead>
<tr>
<th>Variable (type)</th>
<th>Method of assessment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exposure to farm-sprayed pesticides (nominal)</td>
<td>Captured as &quot;yes&quot; for those who sprayed farms with pesticides and &quot;no&quot; for those who did not spray farms with pesticides</td>
</tr>
<tr>
<td>Teratogenic therapeutic medicines for chronic illnesses (nominal)</td>
<td>Captured as a nominal variable, categorized, and labelled; 1= &quot;medicines for hypertension&quot;, 2= &quot;no medicines for chronic illnesses&quot; and 3= &quot;medicines for other chronic illnesses&quot;</td>
</tr>
<tr>
<td>ANC began 8 weeks post-conception began (nominal)</td>
<td>Captured as yes/no</td>
</tr>
<tr>
<td>Gestational age (weeks) at first ANC (continuous)</td>
<td>Captured in weeks, categorized, and labelled; 1&lt;9 weeks, and 2≥ 9 weeks at first ANC visit.</td>
</tr>
<tr>
<td>Preconception folic acid intake (nominal)</td>
<td>Captured as yes/no</td>
</tr>
<tr>
<td>Sex of the “lastborn” current child (nominal)</td>
<td>Entered as male or female</td>
</tr>
<tr>
<td>History of siblings with birth defects (nominal)</td>
<td>This was captured as yes/no</td>
</tr>
<tr>
<td>Parity (continuous)</td>
<td>Abstracted from the ANC booklet as a continuous variable, categorized as and labelled; (1=) &quot;primiparous&quot;, and (&gt;1=) &quot;multiparous&quot;</td>
</tr>
<tr>
<td>Nature of pregnancy (nominal)</td>
<td>Entered as single or multiple</td>
</tr>
<tr>
<td>Maternal age (continuous)</td>
<td>Captured in years</td>
</tr>
<tr>
<td>Paternal age (continuous)</td>
<td>Captured in years</td>
</tr>
<tr>
<td>Level of education (ordinal)</td>
<td>Captured as no schooling, primary, secondary, college certificate, college diploma, and university degree, categorized and labelled; (1=) primary, (2=)secondary, and (3=)tertiary</td>
</tr>
<tr>
<td>Maternal occupation (nominal)</td>
<td>Captured as a nominal variable, categorized into three groups: 1=farming, 2=employed, and (3=)unemployed.</td>
</tr>
<tr>
<td>Residence (nominal)</td>
<td>Captured as a nominal variable, and categorized into five groups: 1=Thika, 2=Gatundu, 3=Kiambu, 4=Ruiru, and 5=other sub-counties</td>
</tr>
</tbody>
</table>

ANC, antenatal care; MESBDs major external structural birth defects.
The conceptual framework was organized based on the three causal categories of MESBDs (multifactorial inheritance, environmental teratogens, and sociodemographic-environmental factors). Nonetheless, because disentangling genetic etiology (identifiable, and complex) was a scientific limitation of observational studies as is the case in our study, analysis of such factors sufficed as a multifactorial inheritance in this conceptual framework to measure maternal genetic predispositions. A conceptual framework depicting the predictor-outcome relationship is displayed in Figure 1. The flow chart of the simple-random systematic sampling strategy is shown in Figure 2.

Ethical considerations
Ethical approval for this study was obtained from the Kenyatta National Hospital [KNH]-University of Nairobi [UoN] Ethics Review Committee [Ref. No: KNH-ERC/A/44]. The purpose of the study was explained to participants and written informed consent was obtained from the mothers of the study subjects before engaging in the study.

Minimizing bias
Considering potential biases inherent in case-control studies that were likely to invalidate the study results, deliberate attempts were made to minimize their occurrence. First and foremost, the research assistants were trained on sound interviewing techniques and information derivation/validation from ANC booklets to minimize interviewer and minimize information biases, respectively. In a bid to minimize recall bias, gestational age at the first ANC was estimated from the dates of the last menstrual period and dates of the first ANC obtained from the ANC booklets.

Data processing and statistical analysis
Following data collection, filled questionnaires were manually checked daily for accuracy and completeness and subsequently entered into a Microsoft Excel spreadsheet (Microsoft Office Professional Plus 2019) by two independent data managers to reduce potential errors. The excel dataset was validated and exported to Stata software version 14.0 (Stata Corporation, Texas, USA) for further cleaning, coding, and analyses. Descriptive analyses (means, medians, standard deviations, and ranges) were used to summarize continuous variables, whereas proportions and percentages for categorical variables were generated and presented in frequency tables. Afterward, the effect of each predictor on the odds of MESBDs was assessed using univariable logistic regression models at a liberal P-value ($P \leq 0.20$). Gestational age at first ANC as a continuous variable was categorized into groups (<9 weeks and ≥9 weeks) for

![Figure 1. Causal diagram of factors thought to influence major external structural birth defects (MESBDs) among children in Kiambu County, Kenya.](image-url)
evaluation in the univariable analyses. Additionally, parity as a continuous variable was grouped into two groups: 1=primiparous or >1=multiparous categories for assessment in the univariable analyses. However, maternal age as a continuous variable was insignificant in the univariable analyses, thus recategorized into two groups: <35 years, and ≥35 years and reassessed for statistical significance; women aged at least 35 years have previously been reported to have an increased likelihood of giving birth to children with MESBDs. Paternal age as a continuous variable was similarly insignificant in the univariable analyses, thus recategorized into seven groups and reassessed for statistical significance yet still insignificant; males aged at least 35 years have previously been associated with increased likelihood of defect-affected births in their female counterparts. Variables found statistically significant in the univariable analyses were fitted to a multivariable model where a backward stepwise approach was used to eliminate variables from the model at P-value >0.05. Nature of pregnancy was however collinear in the multivariable analyses thus dropped in the final multivariable analysis. To minimize the confounding effects, elimination of non-significant predictors was only considered when their exclusion from the model did not yield more than a 30% change in the effects of the remaining variable. Two-way interactions were fitted between the remaining variables of the final model and assessed for significance. A Hosmer-Lemeshow test was used to assess the goodness of fit of the logistic model, with a P-value of >0.05 being suggestive of a good fit.

Results

A total of 408 study respondents (102 cases and 306 controls) were enrolled in this study. The cases consisted of cleft lip with palate 1 (0.98%), cleft palate 3 (9.94%), clubbed hand 1 (0.98%), club foot 91 (89.22%), hydrocephalus 1 (0.98%), limb defects 4 (3.92%), and persistent cloacal 1 (0.98%).

Descriptive statistics

Sociodemographic-environmental factors: The median age of the study respondents was 26 years with a mean of 27.31 years (SD=5.73, R; 17-47) (Table 2). The median age of mothers in the case group was 28 years with a mean of 28.73 (SD=5.95, R; 19-47), whereas the median age of mothers in the control group was 26 years with a mean of 26.84 (SD=5.58, R; 17-42) (Table 2). The mean paternal age of the study respondents was 32.02 years with a standard deviation of 6.34 years, and a median age of 31 years ranging between 19 and 56 years (Table 2). Of the 408 study participants, 184 (45.10%) had attained a secondary level of education; 38 (37.25%) and 146 (47.71%) in the case and control groups, respectively (Table 2).

Environmental-teratogens: Of the 408 study respondents, 15 (3.68%) were exposed to farm-sprayed pesticides, of which four (3.92%) were in the case group and 11 (3.59%) were in the control group (Table 2).

Multifactorial inheritance: Of the 408 study respondents, 404 (98.77%) had single gestations for the current child, of which 99 (97.06%) and 304 (99.35%) were in the case and control groups, respectively (Table 2). Of the study participants,
Table 2. Descriptive statistics of the study respondents (N=408).

<table>
<thead>
<tr>
<th>Variables</th>
<th>Measurements</th>
<th>Observations (N=408), n (%)</th>
<th>Cases (N=102), n (%)</th>
<th>Controls (N=306), n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Residence</td>
<td>Thika</td>
<td>125 (30.64)</td>
<td>33 (32.35)</td>
<td>92 (30.07)</td>
</tr>
<tr>
<td></td>
<td>Gatundu</td>
<td>62 (15.20)</td>
<td>13 (12.75)</td>
<td>49 (16.01)</td>
</tr>
<tr>
<td></td>
<td>Kiambu</td>
<td>104 (25.49)</td>
<td>15 (14.71)</td>
<td>89 (29.08)</td>
</tr>
<tr>
<td></td>
<td>Ruiru</td>
<td>38 (9.31)</td>
<td>20 (19.61)</td>
<td>18 (5.88)</td>
</tr>
<tr>
<td></td>
<td>Others</td>
<td>79 (19.36)</td>
<td>21 (20.59)</td>
<td>58 (18.95)</td>
</tr>
<tr>
<td>Maternal age</td>
<td>&lt;35</td>
<td>356 (87.25)</td>
<td>82 (80.39)</td>
<td>274 (89.54)</td>
</tr>
<tr>
<td></td>
<td>≥35</td>
<td>52 (12.75)</td>
<td>20 (19.61)</td>
<td>32 (10.46)</td>
</tr>
<tr>
<td>Mean</td>
<td></td>
<td>27.31</td>
<td>28.73</td>
<td>26.84</td>
</tr>
<tr>
<td>Median</td>
<td></td>
<td>26</td>
<td>28</td>
<td>26</td>
</tr>
<tr>
<td>Standard deviation (SD)</td>
<td></td>
<td>5.73</td>
<td>5.95</td>
<td>5.58</td>
</tr>
<tr>
<td>Range (R)</td>
<td></td>
<td>17-47</td>
<td>19-47</td>
<td>17-42</td>
</tr>
<tr>
<td>Paternal age</td>
<td>&lt;35</td>
<td>251 (67.11)</td>
<td>64 (70.33)</td>
<td>187 (66.08)</td>
</tr>
<tr>
<td></td>
<td>≥35</td>
<td>123 (32.29)</td>
<td>27 (29.67)</td>
<td>96 (33.92)</td>
</tr>
<tr>
<td>Mean</td>
<td></td>
<td>32.02</td>
<td>31.3</td>
<td>32.25</td>
</tr>
<tr>
<td>Median</td>
<td></td>
<td>31</td>
<td>30</td>
<td>31</td>
</tr>
<tr>
<td>Standard deviation (SD)</td>
<td></td>
<td>6.34</td>
<td>5.47</td>
<td>6.59</td>
</tr>
<tr>
<td>Range (R)</td>
<td></td>
<td>19.56</td>
<td>21-54</td>
<td>19-56</td>
</tr>
<tr>
<td>Maternal education</td>
<td>≤Primary</td>
<td>94 (23.04)</td>
<td>27 (26.47)</td>
<td>67 (21.90)</td>
</tr>
<tr>
<td></td>
<td>Secondary</td>
<td>184 (45.10)</td>
<td>38 (37.25)</td>
<td>146 (47.71)</td>
</tr>
<tr>
<td></td>
<td>Tertiary</td>
<td>130 (31.86)</td>
<td>37 (36.27)</td>
<td>93 (30.39)</td>
</tr>
<tr>
<td>Maternal occupation</td>
<td>Farming</td>
<td>24 (5.88)</td>
<td>7 (6.86)</td>
<td>17 (5.56)</td>
</tr>
<tr>
<td></td>
<td>Unemployed</td>
<td>206 (50.49)</td>
<td>40 (39.22)</td>
<td>166 (54.25)</td>
</tr>
<tr>
<td></td>
<td>Employed</td>
<td>178 (43.63)</td>
<td>55 (53.92)</td>
<td>123 (40.20)</td>
</tr>
<tr>
<td>Parity</td>
<td>Primiparous</td>
<td>127 (37.35)</td>
<td>28 (35.00)</td>
<td>99 (38.08)</td>
</tr>
<tr>
<td></td>
<td>Multiparous</td>
<td>213 (62.65)</td>
<td>52 (65.00)</td>
<td>161 (61.92)</td>
</tr>
<tr>
<td>Mean</td>
<td></td>
<td>2.12</td>
<td>2.14</td>
<td>2.12</td>
</tr>
<tr>
<td>Median</td>
<td></td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Standard deviation (SD)</td>
<td></td>
<td>1.21</td>
<td>1.41</td>
<td>1.22</td>
</tr>
<tr>
<td>Range (R)</td>
<td></td>
<td>1–8</td>
<td>1–6</td>
<td>1–8</td>
</tr>
<tr>
<td>Nature of pregnancy</td>
<td>Multiple</td>
<td>5 (1.23)</td>
<td>3 (2.94)</td>
<td>2 (0.65)</td>
</tr>
<tr>
<td></td>
<td>Single</td>
<td>403 (98.77)</td>
<td>99 (97.06)</td>
<td>304 (99.35)</td>
</tr>
<tr>
<td>Sex of the “lastborn” current child</td>
<td>Female</td>
<td>199 (48.77)</td>
<td>45 (44.12)</td>
<td>154 (50.33)</td>
</tr>
<tr>
<td></td>
<td>Male</td>
<td>209 (51.23)</td>
<td>57 (55.88)</td>
<td>152 (49.67)</td>
</tr>
<tr>
<td>Sibling with a history of birth defects</td>
<td>No</td>
<td>393 (96.32)</td>
<td>93 (91.18)</td>
<td>300 (98.04)</td>
</tr>
<tr>
<td></td>
<td>Yes</td>
<td>15 (3.68)</td>
<td>9 (8.82)</td>
<td>6 (1.96)</td>
</tr>
<tr>
<td>Variables</td>
<td>Measurements</td>
<td>Observations (N=408), n (%)</td>
<td>Cases (N=102), n (%)</td>
<td>Controls (N=306), n (%)</td>
</tr>
<tr>
<td>-----------------------------------------------</td>
<td>--------------</td>
<td>-----------------------------</td>
<td>----------------------</td>
<td>-------------------------</td>
</tr>
<tr>
<td>Gestational age at first ANC</td>
<td>&lt;9 weeks</td>
<td>23 (9.09)</td>
<td>9 (18.75)</td>
<td>14 (6.83)</td>
</tr>
<tr>
<td></td>
<td>≥9 weeks</td>
<td>230 (90.91)</td>
<td>39 (81.25)</td>
<td>191 (93.17)</td>
</tr>
<tr>
<td>Mean</td>
<td></td>
<td>20.1</td>
<td>18.35</td>
<td>20.40</td>
</tr>
<tr>
<td>Median</td>
<td></td>
<td>20</td>
<td>18</td>
<td>21</td>
</tr>
<tr>
<td>Standard deviation (SD)</td>
<td></td>
<td>7.54</td>
<td>8.13</td>
<td>7.36</td>
</tr>
<tr>
<td>Range</td>
<td></td>
<td>4–40</td>
<td>4–35</td>
<td>4–40</td>
</tr>
<tr>
<td>Exposure to farm-sprayed pesticides</td>
<td>No</td>
<td>393 (96.32)</td>
<td>98 (96.08)</td>
<td>295 (96.41)</td>
</tr>
<tr>
<td></td>
<td>Yes</td>
<td>15 (3.68)</td>
<td>4 (3.92)</td>
<td>11 (3.59)</td>
</tr>
<tr>
<td>Teratogenic therapeutic medicines for chronic illnesses</td>
<td>Medicines for hypertension</td>
<td>17 (4.17)</td>
<td>4 (3.92)</td>
<td>13 (4.25)</td>
</tr>
<tr>
<td></td>
<td>No medicines for chronic illnesses</td>
<td>382 (93.63)</td>
<td>96 (94.12)</td>
<td>286 (93.46)</td>
</tr>
<tr>
<td></td>
<td>Medicines for others chronic illnesses</td>
<td>9 (2.21)</td>
<td>2 (1.96)</td>
<td>7 (2.29)</td>
</tr>
<tr>
<td>Preconception folic acid intake</td>
<td>No</td>
<td>230 (56.65)</td>
<td>59 (57.84)</td>
<td>171 (56.25)</td>
</tr>
<tr>
<td></td>
<td>Yes</td>
<td>176 (43.35)</td>
<td>43 (42.16)</td>
<td>133 (43.75)</td>
</tr>
<tr>
<td>ANC began eight weeks post-conception</td>
<td>No</td>
<td>330 (80.88)</td>
<td>77 (75.49)</td>
<td>253 (82.68)</td>
</tr>
<tr>
<td></td>
<td>Yes</td>
<td>78 (19.12)</td>
<td>25 (24.51)</td>
<td>53 (17.32)</td>
</tr>
</tbody>
</table>

SD, standard deviation; R, range; Gatundu North and South sub-counties categorized as Gatundu sub-county, whereas Thika East and West sub-counties categorized as Thika sub-county.

approximately 3.68% (15) of the study participants reported a history of siblings with birth defects consisting of 9 (8.82) in the case group and 6 (1.96) in the control group (Table 2). Of the 15 study participants, 12 stated the name or described the nature of the defects in their previous pregnancies/births, however, 3 participants were unable to do so. Of the 12 study respondents, 7 of the case subjects with congenital talipes equinovarus reported a history of birth defects in their previous births of which 4 subjects reported a recurrence of congenital talipes equinovarus, whereas 3 reported foot aversion, internally rotated shorthand (phocomelia), and congenital scoliosis. On the other hand, 5 control subjects reported a history of siblings with birth defects in their preceding births comprising 3 cases of congenital talipes equinovarus, 1 case of autism, and 1 case of deafness (Table 3).

**Logistic regression analyses**

Notably, the factors assessed for statistical significance in the univariable analyses and found associated with MESBDs at P≤0.20 included maternal age, residence, education, occupation, ANC visits beginning eight weeks post-conception, gestational (age) at first ANC visits, nature of pregnancy, and history of siblings with birth defects (Table 4). Subsequently, these variables were fitted to the multivariable model for the final analysis, except education being distal relative to occupation, gestational age at first ANC visits, and ANC beginning eight weeks post-conception. (Figure 1).

In the multivariable analysis, only maternal residence at conception, and history of siblings with birth defects were shown as the significant predictors MESBDs at a 5% significance level (Table 5). Compared to women who conceived while residing in other sub-counties, women who conceived when residing in Ruiru sub-county were 5.28 times likely to give birth to children with MESBDs (aOR: 5.28; 95% CI: 1.68-16.58; P<0.01); whereas women who conceived when residing in Kiambu sub-county were 27% less likely give birth to children with MESBDs (aOR: 0.27; 95% CI: 0.076-0.95; P =0.04) holding all factors constant. Additionally, compared to siblings without a history of birth defects, siblings with a history of birth defects were 7.65 times likely to be born with MESBDs (aOR: 7.65; 95% CI; 1.46-40.01; P =0.02) holding all factors constant (Table 5).

**Discussion**

To our knowledge, this was the first case-control study conducted to identify the risk factors for MESBDs in the entire county. Our study results mimicked other findings across the world that maternal residence at conception and history of siblings with birth defects are strongly associated with the intrauterine formation of MESBDs\(^1\)\(^2\)\(^3\)\(^4\)\(^6\). Our study observed orofacial clefts comprising 1 (0.98%) cleft lip with the palate, and 3 (9.94%) cleft palates; limb reduction defects comprising 1 (0.98%) clubbed hand, and 4 (3.92%) limb defects; defects of the musculoskeletal system consisting of 91 (89.22%) clubfeet; and neural tube defects comprising 1 (0.98%) hydrocephalus and 1 (0.98%)
### Table 3. History of siblings with birth defects among case and control subjects.

<table>
<thead>
<tr>
<th>Types of MESBDs</th>
<th>Cases (n=102)</th>
<th>Controls (n=306)</th>
<th>Total (n=408)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Congenital talipes equinovarus</td>
<td>4</td>
<td>3</td>
<td>7</td>
</tr>
<tr>
<td>Autism</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Deafness</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Foot aversion</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Internally rotated shorthand</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Congenital scoliosis</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>7</strong></td>
<td><strong>5</strong></td>
<td><strong>12</strong></td>
</tr>
</tbody>
</table>

### Table 4. Univariable analysis of factors associated with MESBDs among children in Kiambu County, Kenya.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Value</th>
<th>Odds ratio</th>
<th>95% CI</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Residence</td>
<td>Others Reference</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Thika</td>
<td>0.99</td>
<td>0.52-1.86</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Gatundu</td>
<td>0.73</td>
<td>0.33-1.61</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>Kiambu</td>
<td>0.47</td>
<td>0.22-0.98</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Ruiru</td>
<td>3.07</td>
<td>1.37-6.89</td>
<td></td>
</tr>
<tr>
<td>Maternal age</td>
<td>&lt;35 Reference</td>
<td>2.09</td>
<td>1.13-3.85</td>
<td>0.02</td>
</tr>
<tr>
<td></td>
<td>≥35</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Paternal age</td>
<td>≥35 Reference</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>&lt;35</td>
<td>1.22</td>
<td>0.73-2.03</td>
<td>0.45</td>
</tr>
<tr>
<td>Maternal education</td>
<td>Tertiary Reference</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Secondary</td>
<td>0.65</td>
<td>0.39-1.10</td>
<td>0.18</td>
</tr>
<tr>
<td></td>
<td>≤Primary</td>
<td>1.01</td>
<td>0.56-1.82</td>
<td></td>
</tr>
<tr>
<td>Maternal occupation</td>
<td>Farming Reference</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Employed</td>
<td>1.09</td>
<td>0.43-2.77</td>
<td>0.003</td>
</tr>
<tr>
<td></td>
<td>Unemployed</td>
<td>0.59</td>
<td>0.23-1.51</td>
<td></td>
</tr>
<tr>
<td>Preconception folic acid intake</td>
<td>No Reference</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Yes</td>
<td>0.94</td>
<td>0.60-1.47</td>
<td>0.78</td>
</tr>
<tr>
<td>ANC began eight weeks post gestation</td>
<td>No Reference</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Yes</td>
<td>1.55</td>
<td>0.90-2.66</td>
<td>0.11</td>
</tr>
<tr>
<td>Gestational age at first ANC</td>
<td>&lt;9 weeks Reference</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>≥9 weeks</td>
<td>0.32</td>
<td>0.13-0.79</td>
<td>0.01</td>
</tr>
<tr>
<td>Parity</td>
<td>Primiparous Reference</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Multiparous</td>
<td>1.14</td>
<td>0.68-1.93</td>
<td>0.62</td>
</tr>
<tr>
<td>Nature of pregnancy</td>
<td>Multiple Reference</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Single</td>
<td>0.22</td>
<td>0.04-1.32</td>
<td>0.10</td>
</tr>
</tbody>
</table>
Table 5. Multivariable analysis of factors associated with MESBDs among children in Kiambu County, Kenya.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Value</th>
<th>aOR</th>
<th>95% CI</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maternal residence</td>
<td>Other sub-counties</td>
<td>0.27</td>
<td>0.076-0.95</td>
<td>0.04</td>
</tr>
<tr>
<td></td>
<td>Kiambu</td>
<td>0.07</td>
<td>0.63-0.95</td>
<td>0.004</td>
</tr>
<tr>
<td></td>
<td>Ruiru</td>
<td>5.36</td>
<td>1.99-14.58</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Siblings with a history of birth defects</td>
<td>No</td>
<td>7.65</td>
<td>1.46-40.01</td>
<td>0.02</td>
</tr>
<tr>
<td></td>
<td>Yes</td>
<td>0.09</td>
<td>0.34-3.52</td>
<td>0.88</td>
</tr>
</tbody>
</table>

aOR, adjusted odds ratio; CI, confidence interval; MESBD, a major external structural birth defect.

Persistent cloacal. These are some types of MESBDs associated with genetic, partially genetic, and multifactorial etiology. The prevalence of such defects have been observed to vary by regions attributed to ethnical, and socioeconomic differences globally. Siblings with a positive history of MESBDs among their preceding siblings are at most risks of being born with MESBDs, have a recurrence of similar defects among the siblings, and/or among their offspring. This was indeed evident in this study where 4 of the case subjects with clubfoot similarly reported clubfoot in their preceding siblings, whereas 3 of the case subjects with clubfoot reported foot aversion, internally rotated shorthand (phocomelia), and congenital scoliosis each in their preceding siblings. Our study similarly made remarkable observations where case subjects with clubfoot reported concurrence of congenital pes planus, and arthrogryposis each, whereas a case subject with hydrocephalus reported concurrence of congenital pes planus, and two case subjects of limb defects reported concurrence with Down syndrome each. On the other hand, 5 control subjects reported a history of siblings with birth defects in the preceding births comprising 3 cases of clubfoot, 1 case of autism, and 1 case of deafness.

Positive siblings and familial history of specific types of MESBDs have been associated with increased risks of recurrence in subsequent pregnancies. Worldwide, the recurrence rate of NTD and Down syndrome have been approximated at 2-5% and 1%, respectively. Thus, accurate knowledge of birth defects by families when given to the clinicians is similarly of public health significance to improve risk assessments and reproductive health planning for couples susceptible to birth
defects of genetic, and multifactorial origin\textsuperscript{30}. Even though our study did not show a significant statistical association between MESBDs with parental age, advanced age has been strongly associated with defects of chromosomal etiology (Down syndrome), and non-syndromic etiology (neural tube defects and orofacial clefts)\textsuperscript{1,30,34,46}. Nonetheless, our study alluded to an increased risk of chromosomal abnormalities thus suggestive of the prevalence of MESBDs of genetic origin in the county. High prevalence of Down syndrome has been observed in developing countries attributed to many older women becoming pregnant, limited family planning services, unavailability of prenatal genetic screening, diagnosis, and related services\textsuperscript{1,30}. MESBDs are considered defects of public health importance, however the presence of certain defects; rare or common, minor or major, internal or external, functional or structural sometimes act as pointers to latent defects of similar significance because of the multiple genetic epidemiology, thus diagnosable later using advanced medical imaging techniques\textsuperscript{3,46}.

Our study similarly observed maternal residence at conception as a predictor of the intrauterine formation of MESBDs. The study showed that women who got pregnant when residing in Ruiru sub-county were 5.28 times likely to give birth to children with MESBDs compared to those who got pregnant residing in other sub-counties within Kiambu County. Conversely, the study showed that women who got pregnant when residing in Kiambu sub-county were 27% less likely to give birth to children with MESBDs compared to those who got pregnant residing in other sub-counties within the county. The study showed that Kiambu sub-county was protective implying it was relatively safe for women of reproductive age to become pregnant while residing in the sub-county. Maternal residence at the time of conception as a risk factor for MESBDs could be ascribed to variations in maternal genetic, multifactorial, sociodemographic-environmental attributes. From the genetic perspective, increased frequency of single-gene defects in developing countries has been associated with increased frequency of common recessive disorders such as hemoglobin disorders, sickle cell anemia, thalassemia, ocoulocutaneous albinism, and cystic fibrosis because of the discerning advantage for carriers to the mortal effects of malaria, as well as recessive conditions associated with high rates of consanguineous (couple) marriages\textsuperscript{1,30}. Additionally, high prevalence of defects of chromosomal etiology in developing countries has been ascribed to women delaying childbearing beyond 35 years, limited maternal access to family planning services, and absence of clinical genetic services\textsuperscript{1,24,30,48}. Sociodemographic-environmental characteristics, and physiological interactions between complex genetic disorders, and idopathic environmental factors could also lead to the occurrence of MESBDs associated with ethnic and geographic differences\textsuperscript{1,30}. Thus, the epidemiology of MESBDs in the county underscore an underlying genetic, multifactorial, sociodemographic-environmental etiology contributing to the global debate on the burden of a “silent” public health problem in developing countries\textsuperscript{1,30}.

Although our study did not show an association between MESBDs with known environmental factors (teratogens and micronutrient deficiencies), pregnancies in developing countries are at increased risk of potential teratogens because of high prevalence of intrauterine infections, maternal malnutrition, low socioeconomic levels, low levels of education, deficient environmental protection policies, and insufficiently regulated access to medicines\textsuperscript{1,30}. This could imply the county is performing relatively well in controlling potential environmental causes of MESBDs. The teratogens consist of; (i) congenital infections; (ii) maternal and altered metabolism; and (iii) recreational and therapeutic drugs\textsuperscript{30}. Congenital infections comprise toxoplasmosis, other infections (syphilis, varicella-zoster, human parvovirus B19), rubella, cytomegalovirus, and herpes, denoted by an acronym “TORCH”\textsuperscript{1,30}. Epilepsy and insulin-dependent diabetes are the examples of maternal illnesses and altered metabolism, whereas statins and alcohol are the examples of therapeutic and recreational drugs, respectively\textsuperscript{1,30}. Our study also did not show significant associations between MESBDs with maternal occupation, gestational age at first ANC, and ANC beginning 8 weeks post-conception; factors thought to influence maternal iron-folic acid supplementation\textsuperscript{1,16,21}. Folic acid is crucial for the biosynthesis, and methylation of deoxyribonucleic acid (DNA) and ribonucleic acid (RNA) which are important for cell division, differentiation, and regulation of gene expression, during rapid cell division such as embryogenesis, thus is necessary for the growth and smooth functions of human cells\textsuperscript{24,49}.

Nevertheless, some limitations were inherent in this study; there was a likelihood of differential recall bias among the study respondents; cases were more likely to remember their pre-conception period owing to the experience of MESBDs in the last birth than the controls, thus recall bias could affect estimates of the odds ratios. The study participants with a history of siblings with birth defects either stated or described the nature of the defects however the researchers could not ascertain accuracy of the diagnoses/descriptions, while others did not know the names of the defects. Survivor bias was also an inherent limitation in this study because some defects such as neural tube defects are potentially fatal, however the study could not establish the causes of deaths among stillbirths, and miscarriages in the study hospitals because it was not a pathological standard operating procedure in the entire Kenya. Additionally, due to the extreme rarity and stochasticity of MESBDs because of the absence of public health surveillance systems, the researchers lumped all types of MESBDs in calculating the sample size, yet births defects are largely heterogenous in their etiology, thus could also lead to underestimation of the effects of the predictors on the odds of MESBDs.

Conclusions
These findings were indeed suggestive of genetic, multifactorial, and sociodemographic-environmental etiology of MESBDs in Kiambu County, Kenya. Thus, these findings could provide the greatest public health opportunities for health planners in the region to establish defect-specific surveillance programs, implement proven public health preventive strategies, and provide appropriate treatment interventions for the most prevalent
MESBDs. Therefore, we would like to provide the following priority public health policy recommendations: establishment hospital-based surveillance systems for the most common MESBDs, and integration of clinical genetic services with routine reproductive health services, nationally. The genetic services should consist of counseling, screening, diagnosis, and associated treatments including elective termination of pregnancies for anomalies in jurisdictions with favorable legislative frameworks. Additionally, we would recommend further epidemiological, and economic evaluation studies to understand the epidemiology and economic burden of these defects in Kenya.

Data availability

Underlying data


Extended data


This project contains the following extended data:
- Questionnaire_mesbds_kmbu_ke.pdf (copy of questionnaire)
- Dofile_mesbds_kmbu_ke.do (syntax used for analysis)

Data are available under the terms of the Creative Commons Zero “No rights reserved” data waiver (CC0 1.0 Public domain dedication).

Acknowledgments

We would like to acknowledge the National Commission for Science, Technology, and Innovation (Ref. No: NACOSTI/P/19/75586/28325), Kiambu County Commissioner (Ref. No: ED.12 (A)/1/VOL.11/107), and Kiambu County Director of Health. (Ref. No: KIAMBU/HRDU/AUTHO/2019/03/06/AgotGN), for permitting us to carry out this study in the county. We would also like to acknowledge the Medical Superintendents and Directors of the thirteen hospitals for granting us permission and support during data collection. Lastly, we would like to acknowledge the efforts of all data collectors in making this exercise a reality.

References


Open Peer Review

Current Peer Review Status: ![Green check mark] ![Green check mark] ![Question mark]

Version 2

Reviewer Report 25 May 2021

https://doi.org/10.5256/f1000research.56198.r84270

© 2021 Kishimba R. This is an open access peer review report distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Rogath Kishimba
1 Tanzania Field Epidemiology and Laboratory Training Program, Dar es Salaam, Tanzania
2 Tanzania Ministry of Health, Dodoma, Tanzania

The manuscript has improved a lot however the below issues need to be addressed before indexed.

Title: When you look on studied birth defects you realize they are all non fatal may be untreated hydrocephalus if any can be fatal. I advice they change the title to read, "Risk factors for non fatal major external structural birth defects among children in Kiambu County, Kenya: a case-control study".

Abstract: Well written

Methods:
- There is no a definition of "resident women". This is very important as it may link environmental factors and the observed outcome.
- Recall bias is not only subjected to gestation age at first ANC. There are a lot of questions which you asked referring to the index pregnancy such as teratogenic medicines, folic acid intake etc. and when exactly they used. How did you mitigate this? It needs clarification.
- What is your sampling unit? A mother? A Newborn? How did you take care of multi pregnancy? Suppose both or all offspring had a birth defect.

Results: Analysis of newborn age is important to be included under the results section.

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Maternal and Child Health Epidemiologist

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.
Title:
The title of this article is correct in my opinion considering that persistent cloacal and hydrocephalus are potentially fatal depending on the severity of the cases.

Methods:
Residence: Residence was defined as a maternal residence at conception whereas resident women were described as women who got pregnant while living in the respective counties of residence within Kiambu County.

Recall bias: The mothers of cases were asked if they used teratogenic drugs during 12 weeks before conception and 8 weeks after conception; the period when women of reproductive age are highly susceptible to teratogenicity. Additionally, inclusion criteria considered under-fives, however, the oldest newborn children whose mothers were interviewed were aged under four years old, thus helped in minimizing potential recall bias.

Sampling unit: Newborn child was the sampling unit, however, questions were administered to the caregivers because being exposed to teratogenicity predisposed unborn children to the intrauterine formation of major external structural birth defects, detectable prenatally, at birth, or soon after birth.

Multiple pregnancies: Data were gathered on multiple pregnancies however, no positive responses on the defects were recorded among the twins, thus not reported in the results.

Analysis:
Newborn age: This was included in the definition of the inclusion criteria, however, the age of the newborn child is not a risk factor for major external structural birth defects in my opinion, thus were not considered for analysis in this study.

Competing Interests: No competing interests.
for the reference group which is appropriate. However, they also deleted p values for exposures that were not the reference group but were or were not statistically significant.

**Competing Interests:** No competing interests were disclosed.

**I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.**

---

**Reviewer Report 09 April 2021**

https://doi.org/10.5256/f1000research.53822.r81820

© 2021 Kishimba R. This is an open access peer review report distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

**Rogath Kishimba**

1 Tanzania Field Epidemiology and Laboratory Training Program, Dar es Salaam, Tanzania
2 Tanzania Ministry of Health, Dodoma, Tanzania

This is a good and important research area for newborn health particularly now when a lot has been done on infectious unlike non infectious diseases. We have observed a great decrease of infant mortality given the available maternal and newborn interventions on infectious diseases. A higher contribution of non infectious disease particularly birth defects may be observed on neonatal and infant mortality with time. Below are my inputs and comments regarding this study;

- The title, abstract and introduction are well written.

- Current citations were used.

- The study design is appropriate however selection of cases was not appropriate given the study title and objectives. It can be admitted as one of study limitation.

- Cases were sampled from child welfare clinics, neonatal/pediatric units, occupational and rehabilitation clinics. All these data sources represent survivors of MESBDs and most probably non fatal MESBDs. It is difficult to get fatal MESBDs like neural tube defects (NTDs) cases from this subpopulation as majority will not survive to meet them in rehabilitation clinics.

- The ascertainment period from the case definition is too high (5 years and below). This may lead to potential recall bias as it will be very difficult for a mother to remember what happened in her pregnancy in the 3-4 years ago. Again may lead to recruitment of survivors and non fatal MESBDs cases. This could be mitigated for at least to consider/restrict enrolment into the study for children below 1 or 2 years only.
- I understand well that the data sources were the above mentioned clinics which are complimented by the ANC booklets. However the methodology section again mentioned about DHIS and I was wondering whether it was also another data source which was used. It needs clarity for the reader to well understand sources of data for this study.

- The methodology section need more clarity on maternal age. Is it the age of the mother during conception of the referred case? or the age of the mother during the data collection? It is also very important to define "residence" as it has implication on maternal exposures. The residence is important during conception and antenatal period. This is the period when environmental exposures can have impact on the unborn child. There is no any significance of considering residence post delivery.

- Sample size calculation is Ok. However you can not estimate proportion of controls ($p_2$) using a study with a different objectives from your intended study.

- The hypothesized odds ratio for the effect of the primary exposure is too high. This is the risk which you allow to be detected in your study. At least you can allow a minimal risk of odds ratio between 1.5 and 2.

- Results were well written however there is a need to your interpretation and conclusion to reflect your exact results. If the maternal age $\leq 34$ years was found to be protective does not mean the maternal age $\geq 35$ years is a risk. Remember this age category was your reference. If you want to refer the age category $\geq 35$ years then make the other category " $\leq 34$ years" a reference in your logistic regression analysis. Otherwise I advise to interpret and make conclusion exactly as what you found in your result section.

**Is the work clearly and accurately presented and does it cite the current literature?**
Partly

**Is the study design appropriate and is the work technically sound?**
Partly

**Are sufficient details of methods and analysis provided to allow replication by others?**
Partly

**If applicable, is the statistical analysis and its interpretation appropriate?**
Yes

**Are all the source data underlying the results available to ensure full reproducibility?**
Partly

**Are the conclusions drawn adequately supported by the results?**
Partly

**Competing Interests:** No competing interests were disclosed.
Reviewer Expertise: Maternal and Child Health Epidemiologist

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Author Response 11 Apr 2021

George Agot, University of Nairobi, Nairobi, Kenya

Sampling of cases: Survivor bias has since been included as a limitation of this study because some of these defects for example neural tube defects are potentially fatal, yet such data could not be gathered because we adopted a retrospective approach to gather the information.

The ascertain period for cases: It is true this period could potentially attract recall bias. This period was preferred considering the defects are extremely rare coupled with unavailability of surveillance systems in the county. Even though we preferred <5 years, we encountered children aged <4 years and approximately 78% were aged <2 years, thus, this limitation was somehow reduced by default.

DHIS: This was not used as a source of data in this study, however was intended to inform the readers on how data on birth defects are routinely gathered in the county in the absence of surveillance systems.

Maternal age: Maternal age was defined as maternal age at conception.

Residence: Residence was defined as maternal place of residence at conception, thus introduced in the model and controlled for in multivariable logistic analysis.

Sample size calculation: The choice of literature for p2 was informed by maternal peri-conceptional period, especially 8 weeks of gestation when defects are expected to have formed yet most women do not plan their pregnancies, thus making it difficult for health care workers to identify these women in good time to effectively implement public health preventive measures before the defects form. Majority of such women attend first ANC at the end of second trimester of gestation when the defects have already formed.

Hypothesized odds ratio: Hypothesized odds ratio for the effects of the primary exposure is actually 2.0 (universally accepted) and not 3.0 as presented in the manuscript. This was a typographical error and has been corrected.

Results: The reference category for maternal age was changed to <35 years, but showed no association in the univariable analysis.

Conclusions: The conclusions have been aligned to the study findings and interpretations.

Competing Interests: None
Marcia L. Feldkamp  
Department of Pediatrics, University of Utah, Salt Lake City, UT, USA

The investigators present a hospital-based case-control study conducted in Kiambu County, Kenya. The paper is well-written and the methodology easy to follow that was used to investigate risk factors for major external structural malformations. The investigators are to be commended for evaluating risk factors for structural malformations in a developing country. This is an important step toward understanding potential risk factors for the ultimate goal of primary prevention. I have a few suggestions for the investigators to consider that may strengthen the paper.

Methods:
1. There are inconsistencies with the specified time period of data collection/enrollment of subjects: “May 31, 2018 to and July 31, 2019”; “May to July 2019”; “three-month study period”.
2. Exposure time period should be clarified for pesticides and how the exposure determined. The investigators should also consider investigating maternal tobacco use, pre-gestational diabetes (specifically), and periconception infections.
3. The investigators conducted a sample size calculation based on all cases. The challenge with this idea is that birth defects are a very heterogeneous group and lumping them altogether suggests that their risk factors are similar. Unfortunately, this is not the case.

Results:
1. Table 3: no need to have a p value listed for the reference group.
2. Since the investigators report an increased risk for maternal age >35 years, did any of the cases have a chromosomal etiology?
3. Did affected siblings have the same type of birth defect?

Discussion - Limitations: small sample size, based on combining several different types of birth defects.

Is the work clearly and accurately presented and does it cite the current literature?  
Yes

Is the study design appropriate and is the work technically sound?  
Yes

Are sufficient details of methods and analysis provided to allow replication by others?  
Yes

If applicable, is the statistical analysis and its interpretation appropriate?  
Partly

Are all the source data underlying the results available to ensure full reproducibility?  
Yes

Are the conclusions drawn adequately supported by the results?
Partly

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Epidemiology, birth defects

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

---

Author Response 11 Apr 2021

George Agot, University of Nairobi, Nairobi, Kenya

**Methods**

1. **Data collection period:** This has been corrected to imply a three-month data collection period from May 31, 2019 to July 31, 2019. It was a typing error, but has since been corrected.

2. **Exposure time period for pesticides:** This variable has been defined as maternal exposure to farm-sprayed pesticides at conception. Increased likelihood of maternal exposure to pesticides was considered because agriculture is the economic mainstay in the county. **Maternal tobacco use:** This variable was assessed as a variable of interest, however was dropped because only 2 out of 408 respondents reported its use. **Pre-gestational diabetes:** This was not assessed as a variable of interest, however it was alluded to and assessed among other chronic illnesses as a proxy for measuring maternal use of teratogenic therapeutic agents for conditions such as epilepsy, depression, hypertension and diabetes mellitus. **Infections:** Infection is indeed an important predictor for birth defects, however, it was beyond the scope of this study since data were gathered retrospectively.

3. **Sample size calculation:** Birth defects are largely heterogeneous in their etiology, however the defects were lumped together for sample size calculation because of the extreme rarity of these defects coupled with unavailability of hospital-based/population-based surveillance programs in the county. Nevertheless, this has been cited as a limitation of the study.

**Results**

1. **Table 3:** P-values for binomial variables were deleted, whereas likelihood ratio test (LRT) was performed for nominal variables/variables with more than two categories to estimate the associated P-values for each variable.

2. **Maternal age:** The reference category for maternal age was changed to <35 years, but showed no association in the univariable analysis. Nonetheless, the study results were suggestive of chromosomal etiology because some cases were reported to occur with down syndrome, whereas autism was also reported by control subjects as a defect in the previous births.

3. **Siblings with same types of birth defects:** Yes this was observed in the study. A recurrence of clubfoot was reported by some case subjects, whereas other types of birth defects were reported by case subjects to have occurred with clubfoot

**Discussion:** Small sample sample based on combined several different types of birth defects has been cited as a limitation of this study.
It is interesting, technically sound and intelligibly written manuscript.

There are minor points to be improved.

In the abstract, the results and the conclusions part should show consistent interpretation and conclusion. The conclusion should base on age < 35 years. The reference is Age > 35 years (would have been better to take the <35 years as a reference). While presenting the classes, since it is dichotomized, it is better to show a common number as a margin of the classes. E.g. <35 and >35 OR <9 and >9.

The Conceptual Framework should reflect the classifications of the risk factors presented in the Introduction. Some of the variables need to be regrouped in themes and the Framework should be redesigned accordingly.

Some of the variables need to be defined. E.g. Pesticide exposure, chronic illness,…

Check the Sample Size Determination part - the Epi Info calculation does not show the same number.

In the univariable analyses, why p-values for the reference categories are included?

The discussion is a bit shallow. Comparison with more literatures, more in-depth look in to the implications and significances of the findings, and addressing also key relevant factors without significant association in the current study can improve the Discussion part. The paternal age was also mentioned as key factor in previous studies but not assessed in the current study. Why? There are also other possible limitations not mentioned. E.g. survivor bias and not controlling for some relevant variables in the multivariable analysis like the paternal age.

The conclusion is a bit beyond the scope of the study. E.g. awareness level of couples or the community is not assessed. Detailed and in-depth discussion by citing other relevant literatures can help readers to better understand the situation and to deduce more appropriate conclusions.

Is the work clearly and accurately presented and does it cite the current literature?
Yes

**Is the study design appropriate and is the work technically sound?**
Yes

**Are sufficient details of methods and analysis provided to allow replication by others?**
Yes

**If applicable, is the statistical analysis and its interpretation appropriate?**
Yes

**Are all the source data underlying the results available to ensure full reproducibility?**
Yes

**Are the conclusions drawn adequately supported by the results?**
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Public Health and Epidemiology

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Author Response 11 Apr 2021

**George Agot,** University of Nairobi, Nairobi, Kenya

**Abstract:** Conclusions have been aligned to the study results and interpretations. Maternal age <35 years has been used as the reference category, and presented as <35 years, and >=35 years, however it was not included in the inclusion because it was no longer associated with dependent variable in the univariable analyses.

**Gestational age:** Gestational age has also been presented as <9 weeks, and >=9 weeks.

**Pesticides exposure:** Exposure redefined as maternal exposure to farm-sprayed pesticides before conception.

**Chronic illnesses:** Chronic illnesses were used as a proxy for measuring maternal use of teratogenic therapeutic agents for chronic conditions such as epilepsy, depression, hypertension and diabetes mellitus. In this respect, it was not used as a measure for particular chronic illnesses.

**Conceptual framework:** Redesigned to reflect the three classes of major external structural birth defects described in the study introduction.

**Univariable analyses:** P-values for binomial variables were deleted, whereas likelihood ratio test (LRT) was performed for nominal variables/variables with more than two categories to estimate the associated P-values for each variable.

**Discussion:** Discussion of the significant variables was improved to include their implications, and significance. Variables that showed no associations were also explained with reference to other studies.

**Paternal age:** Paternal age was introduced in the model, however it showed no association
with the defects in the univariable analyses. Nevertheless, it was controlled for in the multivariable, but still showed no association. Further, because of potential collinearity with materteral age, paternal age was controlled for without maternal age in the multivariable analyses, however, still showed no association with birth defects.

**Limitations of the study:** Survivor bias was included as a limitation of this study because some of these defects for example neural tube defects are potentially fatal.

**Conclusions:** Conclusions were aligned to the study findings and interpretations.

**Competing Interests:** None

Author Response 11 Apr 2021

**George Agot**, University of Nairobi, Nairobi, Kenya

**Sample size determination:** The sample computation was reconfirmed and found consistent with formula provided, however the hypothesized odds ratio is 2.0 (universally accepted, and not 3.0 as presented. This typing error has since been corrected.

**Competing Interests:** None

The benefits of publishing with F1000Research:

- Your article is published within days, with no editorial bias
- You can publish traditional articles, null/negative results, case reports, data notes and more
- The peer review process is transparent and collaborative
- Your article is indexed in PubMed after passing peer review
- Dedicated customer support at every stage

For pre-submission enquiries, contact research@f1000.com